Planktonic community production and respiration and the impact of bacteria on carbon cycling in the photic zone of Lake Kinneret

T. Berman*, A. Parparov, Y. Z. Yacobi

Yigal Alon Kinneret Limnological Laboratory, Israel Oceanographic and Limnological Research, PO Box 447, Migdal 14 950, Israel

ABSTRACT: The following parameters were determined in the photic zone of Lake Kinneret from January 2001 through December 2002: primary production (PP), community respiration (CR), bacterial biomass production (BBP), bacterial numbers (BN) and biological oxygen demand (BOD₅). Average values over the 2 yr period for these parameters in the photic zone (0 to 15 m) were PP, 1539 mg C m⁻² d⁻¹; CR, 1653 mg C m⁻² d⁻¹; BBP, 887 mg C m⁻² d⁻¹; BN, 4.6 × 10⁶ cells ml⁻¹; and BOD₅, 6366 mg C m⁻². We used these data together with literature-based assumptions about the ratio of net to gross primary production, bacterial and zooplankton respiration to estimate bacterial growth efficiencies (BGE) ranging from 44 to 64%. Bacterial specific growth rates averaged 0.33 d⁻¹, ranging from 0.13 to 0.93 d⁻¹. CR was significantly correlated with both PP and BBP. A significant correlation was found between BBP and BOD₅, suggesting that the indigenous bacterial populations were strongly dependent on substrates measured by BOD₅. The potentially labile fractions of TOC, as quantified by BOD₅, were rapidly cycled by heterotrophic bacteria, within ~4 d on average. With the exception of 6 to 8 wk in early summer, the photic zone of this warm mesotrophic-eutrophic lake, with low inputs of allochthonous organic carbon, was net autotrophic. BBP and BGE values were high in comparison to those reported from even more eutrophic aquatic systems. Our results indicate that bacteria are by far the major biological agents of organic carbon cycling in Lake Kinneret, and contradict the general perception that the microbial loop plays a less important role in carbon cycling in eutrophic than in oligotrophic aquatic systems.

KEY WORDS: Bacterial production · Growth efficiency · Community respiration · Primary production · BOD₅ · Lake Kinneret

INTRODUCTION

The processes of phytoplankton photosynthetic carbon fixation (primary production, PP) and community respiration (CR) are responsible for major flows of carbon in the upper waters of most aquatic systems. Over the past 2 decades, it has become increasingly evident that heterotrophic bacterial uptake of dissolved organic carbon and heterotrophic respiration can also account for a further significant portion of total carbon flux within the photic zone of oceans and lakes (del Giorgio et al. 1997, Cole 1999, Biddanda & Cotner 2002). The precise quantification of these processes, which largely determine ecosystem functions, has been and still remains a goal for marine and freshwater ecologists.

More than 70 yr of sustained effort in measuring PP (Williams et al. 2002) has provided generally accepted estimates of global aquatic PP, mainly based on results achieved through variations of the ¹⁴C method introduced by Steemann-Nielsen (1952). Although the importance of concomitant CR measurements has long been recognised, there were relatively few studies of CR in either marine or freshwater systems until the mid-1980s (but see Winberg 1971 for a summary of work in the USSR). Since then, improvements of the classical Winkler oxygen assay have permitted many measurements of planktonic community respiration.
with satisfactorily high sensitivity and precision, even in oligotrophic waters. Several other techniques for measuring CR have been applied less frequently. Some studies have also attempted to include phytoplankton respiration explicitly together with measurements of PP (e.g. Cole et al. 1992, Luz et al. 2002).

Heterotrophic bacterial productivity (bacterial biomass production, BBP) in natural waters has most often been measured by the uptake of either radioactive thymidine (Fuhrman & Azam 1982) or leucine (Kirchman et al. 1985, Simon & Azam 1989). Less frequently, BBP has been determined by following the outgrowth of bacteria in dilution cultures (Biddanda et al. 2001), or by counting the frequency of dividing cells (Hagström et al. 1979, Tuomi 1997). There is now a considerable amount of data on bacterial productivity and growth rates for many different aquatic systems. In contrast, reliable determinations of bacterial respiration (BR) in aquatic environments have proved to be more difficult (Jahnke & Craven 1995).

Despite uncertainties inherent in all of the above methods, the application of existing techniques has led to an impressive characterization and quantification of these major carbon fluxes in many marine and freshwater systems. Such measurements have led to the perception of various aquatic environments as being net autotrophic or net heterotrophic (e.g. del Giorgio et al. 1997, Williams 1998, del Giorgio & Duarte 2002), or by counting the frequency of dividing cells (Hagstrom et al. 1979, Tuomi 1997). There is now a considerable amount of data on bacterial productivity and growth rates for many different aquatic systems. In contrast, reliable determinations of bacterial respiration (BR) in aquatic environments have proved to be more difficult (Jahnke & Craven 1995).

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Here we report the results of a series of concomitant measurements of PP, CR and BBP made over a 2 yr period in Lake Kinneret, and examine the relationships between these parameters. We show that the photic zone of this lake is net autotrophic for most of the year, with a brief period of net heterotrophy in early summer. From these data we use a novel approach to estimate BBG for the lake’s photic zone and suggest that determination of biological oxygen demand, BOD₅, made in the same water samples, provided a quantitative measure of the labile fraction of the total organic carbon (TOC) pool.

**MATERIALS AND METHODS**

**Study site.** Lake Kinneret, the Biblical Sea of Galilee, is a warm monomictic lake with a surface area of 170 km², and mean and maximum depths of 24 and 43 m, respectively. Homothermy occurs between late December and early March, with minimum water temperatures usually >14°C. The lake is strongly stratified from about April to December, with maximum epilimnetic temperatures reaching 29 to 30°C. With the onset of stratification, the hypolimnion rapidly becomes anoxic, with high concentrations of sulphide (5.0 to 9.0 mg l⁻¹) and ammonia (~0.5 to 1.3 mg l⁻¹ N-NH₄⁺). The average annual PP over the period 1972 to 1993 was 610 g C m⁻² yr⁻¹ (Berman et al. 1995). Allochthonous inputs of organic carbon from riverine, littoral and benthic inflows and from dust deposition are about 5000 t C yr⁻¹ or approximately 30 g C m⁻² yr⁻¹ (A. Nishri pers. comm.), i.e. not more than about 5% of annual PP.

**Methods.** Samples were taken in the photic zone (0 to 15 m) at a central, pelagic site, Stn A. Chlorophyll and PP were determined biweekly at depths of 0, 1, 2, 3, 5, 7, 10 and 15 m; CR biweekly at 1, 2, 5, 10 and 15 m; BOD₅ as for CR, but not at 5 m; BBP and bacterial numbers (BN) monthly at 1, 5, 10 and 15 m. Linear integration was used to derive areal values for the photic water column. Note that not all parameters were measured on all occasions.

Chlorophyll concentrations were determined by acetone extraction and fluorometry (Holm-Hansen et al. 1965) and BN by DAPI staining and epifluorescence microscopy (Porter & Feig 1980).

For PP we used the ¹⁴C method (Steemann-Nielsen et al. 1952 as modified by Berman & Pollinger 1974), with a 3 h in situ incubation at Stn A.

Planktonic community respiration (CR) was measured in triplicate lake water samples in dark BOD bottles that were incubated in situ for 24 h. Potentiometric titrations (azide modification of the Winkler method) with a high precision (±2.0 µl) 719S Metrohm Titrisol titrator were made to determine oxygen concentrations. We used a factor of 0.3 to convert mg O₂ l⁻¹ to mg C l⁻¹ (i.e. respiration coefficient of 0.8).

All lake water samples for BOD₅ determinations were initially brought to oxygen saturation by intense stirring, prior to incubation for 5 d at 20°C in the dark. Initial and residual oxygen concentrations were measured by potentiometric titration as for CR. BOD₅ was determined as mg O₂ l⁻¹ and transformed to carbon (mg C l⁻¹) using a respiratory coefficient of 0.8.

We determined BBP using the leucine uptake method (Kirchman et al. 1985, Simon & Azam 1989) as modified by Smith & Azam (1993). Zero time controls were run for all samples. Leucine uptake was converted to carbon uptake using the conversion factors of Simon & Azam (1989) with an isotope dilution factor of 2.

**RESULTS**

In this paper we focus on data collected in the photic water layer (0 to 15 m) of the lake from January 2001 until December 2002. During this period, average water temperatures in this layer varied from 15.9 to 30.2°C (Fig. 1). The general patterns of phytoplankton biomass (as chlorophyll, averaged for the photic layer)
are also shown in Fig. 1. The lake stratified and the hypolimnion became anoxic in early and mid-May in 2001 and 2002, respectively.

A summary of the parameters directly measured in our study: chlorophyll, PP, CR, BBP, BOD$_5$ and BN, averaged for the entire period from January 2001 through December 2002 and separately for the winter-spring (January through June) and summer-fall (July through December) seasons, is given in Table 1. Some ratios between the above parameters are shown in Table 2.

### Primary production

The annual pattern of PP during the 2 yr of this study (Fig. 2) did not show the late winter-spring peak associated with the dinoflagellate bloom that had occurred regularly for many years in Lake Kinneret (Berman et al. 1995). Other unusual features in the pattern of PP were the appearance of a secondary summer peak in August-September 2001 and again in the summer of 2002. Average PP for the period 2001 and 2002 was $1539 \pm 517 \text{ mg C m}^{-2} \text{ d}^{-1}$. There was no significant relationship between primary production and water temperatures.

### Community respiration

CR averaged $1653 \pm 762 \text{ mg C m}^{-2} \text{ d}^{-1}$, ranging from 343 (January 2002) to 3399 mg C m$^{-2}$ d$^{-1}$ (May 2001). There were higher levels of CR in the early summer months from May through June that were not accompanied by a similar increase in PP (Fig. 2). There was less variation in the seasonal rates of CR than in chlorophyll or PP (Table 1). CR was significantly correlated with PP as a power function and, less strongly, with water temperature (Table 3).

### Bacterial abundance

Bacterial cell counts in the epilimnion ranged from $1.9 \times 10^6 \text{ ml}^{-1}$ to $7.9 \times 10^6 \text{ ml}^{-1}$.
Averaging $4.6 \times 10^6$ ml$^{-1}$. Bacterial abundance was low at the beginning of this study period but increased >2-fold during spring 2001 and thereafter remained relatively high throughout 2002; comparable high BN has not been observed in Lake Kinneret since 1998 (Fig. 3). During the years 1988 to 1994 the average concentrations of bacteria were similar ($4.0 \times 10^6$ ml$^{-1}$ for the 0 to 10 m water column) but with greater variability, ranging from $5 \times 10^5$ ml$^{-1}$ to $2.3 \times 10^7$ ml$^{-1}$ (T. Berman unpubl.).

**Bacterial biomass production**

Bacterial biomass production (BBP), integrated to 15 m depth, averaged 887 ± 469 mg C m$^{-2}$ d$^{-1}$, ranging from 300 to 2148 mg C m$^{-2}$ d$^{-1}$ over the 2 yr period, with markedly higher levels during most of 2001 than in 2002 (Fig. 4). For all samples, BBP ranged from 16 to 151 mg C m$^{-3}$ d$^{-1}$ and tended to be evenly distributed within the epilimnion at all seasons. On 2 occasions (16 June and 13 August 2002), exceptionally high rates of BBP were measured but we believe that these results were due to experimental error since all other parameters (chlorophyll, PP, CR, BOD) determined in the same water samples were not exceptional. Therefore we have excluded these results from our considerations here.

No significant relationships were observed between BBP and water temperatures, phytoplankton biomass (as chlorophyll) or PP. However, BBP was significantly correlated with CR (Table 3). Also, significant correlations were found between BBP and BOD$_5$ either with data from all samples taken at 1 and 10 m depths or with data integrated for the photic water column (Table 3). From January 2001 to December 2002, bacterial cell-specific productivity in the epilimnion, calculated as BBP divided by BN (DAPI counts), averaged 14.7 fg C cell$^{-1}$ d$^{-1}$, varying from 4.4 to 38.4 fg C cell$^{-1}$ d$^{-1}$, with highest levels during spring and summer 2001.

Assuming a relatively high average carbon content of 30 fg cell$^{-1}$ for bacterial cells in a mesotrophic-eutrophic lake (Lee & Fuhrman 1987, Fukuda et al. 1998) we calculated that, in the epilimnion of Lake Kinneret, the average carbon turnover time for a bacterial cell (defined as the time required to replace bacterial carbon biomass, or doubling time) fluctuated from 0.7 to 5.3 d (Fig. 4) with an overall average of 2.9 ± 1.3 d. Cell-specific growth rates averaged 0.33 ± 0.22 d$^{-1}$ and ranged from 0.93 to 0.13 d$^{-1}$.

**Biological oxygen demand**

We measured BOD$_5$ as a proxy for the readily labile fractions of organic carbon in the TOC pool (Osta-
penya 1971). BOD$_5$ integrated for the upper 15 m water column averaged 6366 ± 2473 mg C m$^{-2}$ over the period of this study, varying from 2723 to 12 908 mg C m$^{-2}$ (Fig. 5). Generally higher BOD$_5$ was found during the winter-spring period, when phytoplankton biomass was greatest (Table 1). Maximum BOD$_5$ levels were observed during March–May 2001, probably reflecting the higher standing stocks of phytoplankton in that season.

**Relationships between parameters**

Within the photic zone, CR was on average 106% of measured PP, ranging from 36 to 230%, with 2 periods of elevated values in May–June 2001 and 2002 (Fig. 2). We postulate that PP measured by the $^{14}$C method as used by us was close to net primary production (Cole et al. 1992, Marra 2002). If net primary production in Lake Kinneret was generally between 70 and 80% of gross primary production (GPP) (Berman & Gerber 1980, Berman & Kaplan 1984), then, on average, CR would have been between 86 and 75% of GPP (Table 2), ranging from 25 to 184% over the 2 yr period. Overall, for most of the year, the photic zone of the lake was net autotrophic, with a short period in May–June 2001 and 2002 of net heterotrophy (i.e. when CR > GPP).

For data integrated over 15 m, BBP averaged 60% of PP (Table 2) and fluctuated from 27 to 141% of PP for all sampling dates. BBP averaged 55% of CR, ranging from 21 to 97% (Table 2). These percentages were markedly higher in 2001 than in 2002 because of the drop in BBP in the latter year (Table 1, Fig. 4).

**Bacterial growth efficiencies**

BGE, defined as BBP divided by total bacterial carbon uptake, is a critical ecosystem parameter that is extremely difficult to measure directly in natural aquatic systems (del Giorgio & Cole 1998, 2000). Total carbon uptake, or gross BBP, is the sum of net BBP plus bacterial respiration, BR.

The range of BGE in Lake Kinneret was estimated from the set of measurements and assumptions in Table 4. We posited that our measurements of BBP and PP were close to net BBP (Biddanda & Cotner 2002, M. Simon pers. comm.) and net PP (Cole et al. 1992, Marra 2002), respectively. In the example in Table 4, we used previously reported values of 20% daily carbon loss from GPP due to phytoplankton respiration, PR (Berman & Kaplan 1984) together with minor amounts of exudation and photorespiration (Berman 1976). Since our measurement of CR reflected the total of bacterial, zooplankton and phytoplankton respiration (CR = BR + ZR + PR), it follows that CR – PR = BR + ZR.

**Table 4. Bacterial growth efficiency (BGE) estimate from Lake Kinneret data. All values in mg C m$^{-2}$ d$^{-1}$. GPP, PP: gross and net primary production, respectively. CR, PR, BR, ZR: community, phytoplankton, bacterial and zooplankton respiration, respectively**

<table>
<thead>
<tr>
<th>Conditions</th>
<th>PR = 20 % GPP</th>
<th>BR = 2 ZR</th>
<th>PR = 20 % GPP</th>
<th>BR = ZR</th>
</tr>
</thead>
<tbody>
<tr>
<td>GPP</td>
<td>1539</td>
<td>PP$^a$</td>
<td>1924</td>
<td>1924</td>
</tr>
<tr>
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<td>385</td>
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<td>CR$^a$</td>
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<tr>
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<td>1268</td>
<td>CR-PR = BR + ZR</td>
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<td></td>
</tr>
<tr>
<td>BR (if BR = 2 ZR)</td>
<td>850</td>
<td>BR (if BR = ZR)</td>
<td>634</td>
<td></td>
</tr>
<tr>
<td>BBP$^a$</td>
<td>887</td>
<td>BBP$^a$</td>
<td>887</td>
<td></td>
</tr>
<tr>
<td>BGE = BBP/(BBP + BR)</td>
<td>51.1%</td>
<td>BGE = BBP/(BBP + BR)</td>
<td>58.3%</td>
<td></td>
</tr>
</tbody>
</table>

$^a$Experimentally measured parameter, averaged from January 2001 to December 2002.
In Table 4, we used ratios of BR:ZR of either 1 or 2. Combining Lines 6 and 7 in Table 4 then gave a direct estimate of BGE, in this case, 51.1 or 58.3%, depending on the ratio BR:ZR.

Table 5 shows BGE values, averaged for 2001–2002, obtained for a range of ratios of PP:GPP and BR:ZR, with PP varying from 70 to 90% of GPP and BR varying from BR = ZR to BR = 3ZR. Rather surprisingly, the estimated BGE varied only from 44 to 64%, over this fairly wide span for PP:GPP and BR:ZR.

### DISCUSSION

During this study, the seasonal development of phytoplankton in Lake Kinneret differed from the fairly regular annual cycle that prevailed from 1972 to 1994 (Berman et al. 1998). Phytoplankton maxima in February 2001 and May 2002 (Fig. 1) were due to *Aulacoseira granulata* and *Peridinium gatunense*, respectively. In 2002, after a lapse of 2 yr there was a moderate winter-spring dinoflagellate bloom similar to the ‘normal’ pattern (T. Zohary pers. comm.). A clear division between a *P. gatunense* bloom season and a chlorophyte-dominated summer season was a basic feature in attempts to quantify carbon flux in the Lake Kinneret system (Serruya et al. 1980, Stone et al. 1993, Hart et al. 2000). Unlike previous years, during the summers of 2001 and 2002 the phytoplankton was characterized by a mixture of cyanobacteria, chlorophytes and diatoms. The average values of PP for the years 2001 and 2002 (Table 1) were about 20% lower than the multi-year average PP from 1972 to 2002 (Y. Yacobi pers. comm.).

The pattern of bacterial abundance (Fig. 3) may also reflect the changing state of this ecosystem, with relatively higher levels in 1998 and again from the summer of 2001, without any clear annual cycles such as were observed in measurements made from 1989 to 1994 (not shown). We have no data to indicate whether there were any changes in the phylogenetic composition of the lake bacterial community during this period.

#### CR and GPP; net autotrophy and net heterotrophy in the phototrophic zone

Similarly to observations in other aquatic systems (del Giorgio & Peters 1994, Iriarte et al. 1996, Carignan et al. 2000) we found a significant power regression for CR with PP in Lake Kinneret (Table 3). CR also showed a significant power correlation to water temperatures, as has been reported elsewhere (Iriarte et al. 1996), although only about 20% of the variability in CR could be explained by changes in temperature.

Based on the average for the period from January 2001 to December 2002, CR in the photic zone accounted for 75 to 86% of GPP, when total carbon losses by phytoplankton respiration and exudation from GPP were between 20 and 30% (Berman & Gerber 1980, Berman & Kaplan 1984). This level of CR would imply that between 14 to 25% of annual GPP sedimanted out as particulate organic carbon (POC) from the phototrophic water layer, i.e. ~100 to 150 g C m⁻² yr⁻¹ or ~17 000 to 25 000 t organic C yr⁻¹ for a lake surface area of 170 km². These amounts of sedimented organic carbon are compatible with estimates of TOC exported from the trophogenic layer, based on calculations of the amounts of NH₄⁺ and H₂S generated within the hypolimnion by mineralization of organic matter (Nishri et al. 1998, A. Nishri pers. comm.).

In both 2001 and 2002, CR exceeded PP and GPP for only about 2 mo in early summer, coincident with the most rapid warming of the water mass, and towards the end of the late winter-spring phytoplankton bloom (Figs. 1 & 2). This appears to be the only period when the photic zone of Lake Kinneret was net heterotrophic. As noted above, the total amounts of organic carbon entering the lake from the River Jordan and minor river inflows, from littoral vegetation, benthos and atmospheric deposition are very small compared to the total amounts of these pools in the lake water (A. Nishri pers. comm.). Thus, >95% of TOC in this lake derives from autochthonous sources, in contrast to many lakes in which allochthonous organic carbon inputs support considerable heterotrophic activity (Cole et al. 2000).

A general pattern of increasing CR-GPP with increasing oligotrophy was suggested by del Giorgio & Peters (1994) and by del Giorgio et al. (1997), who posited that lakes tend to become net heterotrophic when chlorophyll levels fall to <15 µg l⁻¹ or to PP.
<120 µg C l⁻¹ d⁻¹. Note, however, that the average, photic zone chlorophyll concentrations and PP in Lake Kinneret during our study were 12.4 (±7.5) µg l⁻¹ and 103 ± 34 µg C l⁻¹ d⁻¹, respectively, with lengthy periods of chlorophyll concentrations <15 µg l⁻¹ (Fig. 1) and rates of PP below 100 µg C l⁻¹ d⁻¹. Nevertheless, as noted above, the phototrophic zone of the lake was net autotrophic during most of the year.

Our present observations on Lake Kinneret do not fit the general pattern of net autotrophic and net heterotrophic aquatic systems proposed by del Giorgio et al. (1997). Another exception to the trend suggested by del Giorgio et al. (1997) was reported by Carignan et al. (2000), who found a predominance of net autotrophy in 12 oligotrophic to mesotrophic Canadian shield lakes, with a median CR:GPP ratio of 0.59.

We used a respiration coefficient of 0.8 to convert CR from oxygen to carbon, and thereby obtained lower carbon respiration rates than would have been obtained with the frequently used respiration coefficient of 1.0 (del Giorgio et al. 1997, Biddanda et al. 2001). As pointed out by Geider (1997), a respiration coefficient of 0.8 is probably more accurate for natural aquatic environments where the respiration of organic matter includes proteins and lipids in addition to carbohydrates. Recently, Robinson et al. (2002) experimentally determined a respiratory coefficient of 0.8 for 11 samples in the eastern Atlantic Ocean.

Our estimates for GPP are considerably lower than those reported for Lake Kinneret by Luz et al. (2002) using an ¹⁸O oxygen isotope method. At present we have no explanation for this discrepancy, although the question of the value assigned to the photosynthetic quotient in order to transform O₂ to C in this method is critical.

**BBP and PP**

White et al. (1991) listed bacterial growth rates ranging from 0.017 to 8.7 d⁻¹ for 10 eutrophic and 14 mesotrophic lakes, and Cole & Pace (1995) reported bacterial specific growth rates from 0.02 to 4.6 d⁻¹ for a variety of freshwater and marine locations. The average bacterial growth rates for photic zone bacteria in Lake Kinneret (0.46 d⁻¹ in 2001; 0.23 d⁻¹ in 2002; 0.33 d⁻¹ for all samples) were well within these ranges and similar to growth rates measured in lake water cultures by Pinhassi & Berman (2003). Nevertheless, the levels of both volumetric BBP and areal BBP measured in Lake Kinneret were at the high end of rates reported for a wide variety of aquatic environments (Table 6). This seems reasonable, considering that Lake Kinneret is mesotrophic-eutrophic, with higher phytoplankton standing stocks and warmer water than many (but not all) of the aquatic systems listed in Table 6. The average BBP values observed during this study were remarkably similar to those measured in Lake Kinneret during the period 1998 to 1999 using the same ¹⁴C leucine uptake method (average: 913 ± 619 mg C m⁻² d⁻¹) and to BBP determined by ³H thymidine uptake during the years 1988 to mid-1991 (average: 994 ± 1247 mg C m⁻² d⁻¹; T. Berman unpubl.).

We calculated bacterial cell-specific growth rates and turnover times (Fig. 4) on the basis of total BN as given by DAPI staining. However, as has been observed in Lake Kinneret (Berman et al. 2001) and elsewhere (Choi et al. 1999, Bernard et al. 2000, Søndergaard & Danielsen 2001), only a small proportion of the total bacterial population (perhaps ~20% as a conservative estimate) appear to be strongly metabolically active. If this is so, then the average cellular uptake, turnover time and cell-specific growth rate for ‘active bacteria’ in this lake would be 73.6 fg C cell⁻¹ d⁻¹, 0.57 d and 1.22 d⁻¹, respectively.

A partial explanation for the very rapid growth rates attributed to ‘active’ bacterial cells may be the observation that ‘no more than ~20% of bacterial cells are active at any given time’ (Berman et al. 2001). It is feasible that bacterial cells may be strongly active only intermittently. Thus, some of the inactive (or weakly active) cells observed at one point in time may subsequently become active (Choi et al. 1999, Berman et al. 2001), while previously active cells may cease activity. In this manner, all the DAPI-stained cells in the community may indeed be active but, on average, for only 20% of the day. In this case, estimates of the daily growth rates for each cell in the bacterial community become more reasonable. Of course, it is unlikely that all DAPI-counted bacteria are metabolically active or that ‘activity’ is an all-or-none phenomenon.

An important consideration is the question of whether BBP determinations using either radioactive thymidine or leucine give gross or net measurements (or something between). In the present work, similarly to that of Biddanda & Cotner (2002), Ducklow (2000) and M. Simon (pers. comm.), we have taken the radioactive ¹⁴C-leucine method with a 1 h incubation time as used here to give a close approximation of net bacterial production.

For the period of this study, the average BBP:PP measured for the photic water column was 60% (Table 2) and BBP was 42 or 48% of GPP, depending on the assumed daily phytoplankton respiration and exudation losses of either 30 or 20% (Berman & Gerber 1980, Berman & Kaplan 1984). A pioneering study by Winberg (1971), based on dark ¹⁴C-uptake and O₂ measurements of BBP and GPP, respectively, indicated a wide variation of BBP to GPP, from 16 to 78%, for a series of Russian lakes ranging from oligotrophic to...
eutrophic. The levels of BBP relative to GPP in Lake Kinneret are considerably higher than those recorded by Simon et al. (1998) in mesotrophic Lake Constance, but similar to those measured by Inkina (1985) for Lake Miastro. Both these lakes have levels of primary production roughly similar to those in Lake Kinneret. Scavia & Laird (1987) observed that annual bacterial carbon demand, i.e. gross BBP, was 65% of PP in Lake Michigan. Schwaerter et al. (1988) listed estimates of bacterial gross production ranging from 20 to 80% of phytoplankton gross production in lakes and from 24 to 60% in marine environments. Cole et al. (1988), in an extensive cross-system review of bacterial productivity, found that BBP to PP was generally between 20 and 40% in most aquatic environments. In a recent study of the Antofagasta coastal and oceanic regions within the Humbold Current off Chile, Troncoso et al. (2003) reported extremely high BBP:PP, ranging from 63 to 478%. Most marine studies have found much lower values; for instance, Ducklow et al. (2002) brought evidence to show that levels of BBP were previously overestimated in the North Atlantic Ocean and that BBP:PP could not be >20% in this oligotrophic marine system.

The relatively high percentage of BBP to PP that we observed implies that total bacterial carbon demand (gross BBP) in the trophogenic zone of Lake Kinneret was close to, or at times even in excess of GPP. If we take average annual values for BGE of 51 or 57% (Table 5, with BR = 2 ZR) and a ratio of BBP:GPP of 48 or 42% (depending on whether PR was taken to be 20 or 30%), then the proportion of GPP taken up by bacterial carbon demand was between 73 and 94% averaged over the 2 yr study. These are high but by no means impossible levels for gross BBP:GPP. The total bacterial carbon uptake by bacteria and by secondary and tertiary consumers can exceed GPP because of recycling within the system (Strayer 1988), as was shown to be the case for Lake Michigan (Scavia 1988). Our data confirm previous indications that bacteria in

<table>
<thead>
<tr>
<th>Location</th>
<th>BBP</th>
<th>BBP</th>
<th>Method</th>
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<tr>
<td></td>
<td>mg C m⁻² d⁻¹</td>
<td>mg C m⁻³ d⁻¹</td>
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<td>Aquatic systems</td>
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<td>T, L</td>
<td>Cole et al. (1988)</td>
</tr>
<tr>
<td>Freshwater</td>
<td>–</td>
<td>45.3 (0.5–302)</td>
<td>T</td>
<td>White et al. (1991)</td>
</tr>
<tr>
<td>Marine</td>
<td>–</td>
<td>18.0 (&lt;0.1–336)</td>
<td>T</td>
<td>White et al. (1991)</td>
</tr>
<tr>
<td>Estuarine and coastal waters</td>
<td>–</td>
<td>356 (0.03–23,767)</td>
<td>T</td>
<td>White et al. (1991)</td>
</tr>
<tr>
<td>Chesapeake Bay</td>
<td>–</td>
<td>20.0–280</td>
<td>T</td>
<td>Jonas &amp; Tuttle (1990)</td>
</tr>
<tr>
<td>Mississippi River Plume</td>
<td>–</td>
<td>5.0–90</td>
<td>T, L</td>
<td>Chin-Leo &amp; Benner (1992)</td>
</tr>
<tr>
<td>Gulf of Mexico (shelf, slope)</td>
<td>–</td>
<td>23.5 (19.0</td>
<td>T, L</td>
<td>Biddanda et al. (1994)</td>
</tr>
<tr>
<td>SubArctic, NE Pacific</td>
<td>–</td>
<td>0.5–4.5</td>
<td>T, L</td>
<td>Sherry et al. (1999)</td>
</tr>
<tr>
<td>Weddell, Scotia Seas</td>
<td>–</td>
<td>0.12 (0.02–0.34)</td>
<td>L</td>
<td>Moran et al. (2001)</td>
</tr>
<tr>
<td>Tuamota Atoll Lagoon</td>
<td>–</td>
<td>1.0–43.6</td>
<td>T</td>
<td>Torrton et al. (2002)</td>
</tr>
<tr>
<td>L. Hylke, Denmark</td>
<td>1520</td>
<td>–</td>
<td>T</td>
<td>Riemann &amp; Sandergaard (1986)</td>
</tr>
<tr>
<td>Frederiksberg Slotso</td>
<td>6640</td>
<td>–</td>
<td>T</td>
<td>Riemann &amp; Sandergaard (1986)</td>
</tr>
<tr>
<td>Lake Michigan</td>
<td>652</td>
<td>28.9 (0.2–90)</td>
<td>T</td>
<td>Scavia &amp; Laird (1987)</td>
</tr>
<tr>
<td>Lake Plußsee (photic zone, Apr–Oct)</td>
<td>–</td>
<td>9.6 (0.9–40.6)</td>
<td>T</td>
<td>Chrost &amp; Rai (1994)</td>
</tr>
<tr>
<td>Lake Constance</td>
<td>78.0</td>
<td>0.24–96</td>
<td>T, L</td>
<td>Gude (1990)</td>
</tr>
<tr>
<td>Lake Constance</td>
<td>(199–954)</td>
<td>–</td>
<td>L</td>
<td>Simon et al. (1998)</td>
</tr>
<tr>
<td>Lake Paul</td>
<td>50.0</td>
<td>–</td>
<td>L</td>
<td>Pace &amp; Cole (1994)</td>
</tr>
<tr>
<td>Crystal Lake</td>
<td>51.0</td>
<td>–</td>
<td>L</td>
<td>Pace &amp; Cole (1994)</td>
</tr>
<tr>
<td>Lake Trout</td>
<td>48.0</td>
<td>–</td>
<td>L</td>
<td>Pace &amp; Cole (1994)</td>
</tr>
<tr>
<td>9 Minnesota lakes (oxic layer)</td>
<td>–</td>
<td>12.6 (2.6–17.0)</td>
<td>L</td>
<td>Cole &amp; Pace (1995)</td>
</tr>
<tr>
<td>Lake Superior</td>
<td>–</td>
<td>20.6 (1.1–47.0)</td>
<td>L</td>
<td>Cole &amp; Pace (1995)</td>
</tr>
<tr>
<td>East Long Lake</td>
<td>–</td>
<td>4.3–19.6</td>
<td>DC</td>
<td>Biddanda et al. (2001)</td>
</tr>
<tr>
<td>Lake Michigan</td>
<td>–</td>
<td>0.58–1.15</td>
<td>L</td>
<td>Biddanda &amp; Cotner (2002)</td>
</tr>
<tr>
<td>Lake Kinneret (1988–1991)</td>
<td>0–20 m</td>
<td>994 ± 1247</td>
<td>50.0</td>
<td>T. Berman (unpubl.)</td>
</tr>
<tr>
<td>Lake Kinneret (2001–2002)</td>
<td>0–15 m</td>
<td>913 ± 619</td>
<td>61.0</td>
<td>T. Berman (unpubl.)</td>
</tr>
<tr>
<td>Lake Kinneret (1998–1999)</td>
<td>0–15 m</td>
<td>887 ± 469</td>
<td>56.4 (16.0–151)</td>
<td>This paper</td>
</tr>
</tbody>
</table>

*aTo 100 m depth; *b euphotic zone; *c to 20 m depth, maximum rates; *d maximum value

Table 6. Bacterial biomass production (BPP) in a variety of aquatic environments. L: leucine method; T: thymidine method; DC: dilution culture method
Lake Kinneret are by far the major agents of organic carbon cycling in this ecosystem (Stone et al. 1993, Hadas & Berman 1998, Hart et al. 2000).

Our finding that BBP was significantly correlated with CR (Table 3) would also imply that bacterial respiration is an important component of planktonic CR in this lake (as indicated in Table 4) similar to many (Hopkinson et al. 1989, Jahnke & Craven 1995, Sherry et al. 1999), but not all (Williams 1981, Sherry et al.1999), aquatic systems.

Although many investigators have observed a strong relationship between BBP and temperature in aquatic environments (e.g. Iriberri et al. 1985, White et al. 1991, Shah & Ducklow 1994, Simon et al. 1998, Sherry et al. 1999), no such relationship above 10°C was found by Scavia & Laird (1987) in Lake Michigan or by Cole & Pace (1995) in 9 North American lakes, and none was evident in Lake Kinneret.

Unlike previous observations in Lake Kinneret (Berman et al. 2001) and elsewhere (e.g. Cole et al. 1988, White et al. 1991), no correlation was found between BBP and primary production, or between BBP and chlorophyll. Pace & Cole (1994) also reported no significant correlation between BBP and PP in 3 Wisconsin lakes and suggested that other factors such as nutrient recycling and phytoplankton loss rates could be more significant than levels of PP in determining BBP.

**Estimating BGE**

For a direct evaluation of BGE, both BBP and BR need to be known. These parameters can be determined relatively easily in experiments with bacterial cultures, but in natural aquatic systems reliable measurement of BR is problematic (Jahnke & Craven 1995). Experimental measurements of BGE in batch and chemostat cultures have shown that this parameter can fluctuate widely, depending on specific organism and growth conditions (see references in del Giorgio & Cole 1998, 2000). Early estimates of BR in marine and freshwater (Hobbie & Crawford 1969) that were based on the addition of trace quantities of radioactive substrates (usually 14C-glucose or 14C-amino acids) tended to give relatively low BR and therefore relatively high BGE. For example, Williams (1970) reported a BGE of 67 and 78% for marine bacteria metabolizing glucose or amino acids, respectively, and Berman et al. (1979) found BGEs ranging from 39 to 75% in Lake Kinneret based on 14C-glucose uptake experiments. The results of such experiments are now thought to have generally underestimated BR and overestimated BGE, because they were based on bacterial utilization of a single or, at most, a few readily available substrates, whereas bacteria in the real life environment presumably exploit a broad spectrum of substrates of variable availability (del Giorgio & Cole 1998, 2000).

Because of the difficulties of direct measurement of BR, in more recent studies of bacterially mediated carbon flux in natural aquatic environments, some investigators have used assumed BGE values based on the literature (e.g. Scavia & Laird 1987, Strayer 1988, Baines & Pace 1991, Ducklow & Carlson 1992, Stone et al. 1993, Coveney & Wetzel 1995, Hart et al. 2000). Others have quantified BR directly by following the increase in ambient dissolved inorganic carbon (Cole et al. 1989) or, more frequently, by determining O2 uptake in water samples that were pre-filtered in order to exclude phytoplankton and protozoans (e.g. Hopkinson et al. 1989, Biddanda et al. 1994, 2001, Roland & Cole 1999, Sherry et al. 1999, Rivkin & Legendre 2001, Smith & Kemp 2003). Unfortunately, the method of estimating BR using filtration followed by a lengthy (usually ~24 h) incubation step radically changes the environment in which BR determinations are made by altering the patterns of organic carbon cycling in these samples. Additionally, there is the problem of ‘mixed’ timescales, whereby BBP is measured ‘instantaneously’ at the beginning of the experiment, while BR requires a prolonged incubation (see Roland & Cole 1999). Thus, the validity of BR estimates made in pre-filtered samples may be compromised.

Recently, Toolan (2001) successfully applied coulo-metric respiration measurement on filtered and unfiltered samples to directly estimate BR and CR with only ~10 h incubation. Measurements of electron transport system (ETS) activity (Packard 1971) have also been used to quantify planktonic respiration. In Lake Kinneret, ETS activity in GF/C-filtered samples was found to correlate reasonably well with ΔO2 and hydrolytic enzyme activity but not with BBP (Berman et al. 2001).

For logistic reasons (i.e. lack of time and money), we were unable to use any of the above methods to estimate BR, and in the present study we applied a different approach to derive BGE. By using the relatively robust determinations of PP, BBP and CR, and reasonable assumptions based on the literature for the possible ranges of PR (from 20 to 30% of GPP) and the ratio of BR:ZR (0.33 to 1), we calculated a range of potential average values for BGE in the photic zone of Lake Kinneret (Tables 4 & 5).

In their very extensive review on BGE in natural waters, del Giorgio & Cole (1998) showed that reported values for this parameter tended to be lowest in rivers, increased somewhat in oceans, then in lakes, and were highest in estuaries. The median BGE for their sample of about 20 lakes was ~28%, the median BGE for 6 estuaries was ~34%. Kristiansen et al. (1992) deter-
mined a mean BGE of 31% in a small, eutrophic lake, Frederiksborg Slotse. In a recent study in Lake Superior and 10 small Minnesota lakes with chlorophyll concentrations ranging from 0.6 to 52.7 µg l−1, Biddanda et al. (2001) reported BGE from 5 to 39%, increasing from oligotrophic to eutrophic systems. In Chesapeake Bay, Smith & Kemp (2003) measured a range of BGE from 20.4 to 41.3%, with spatial variations in efficiencies apparently increasing with increasing nutrient concentrations.

Our estimated BGE values (Table 5) were distinctly higher than the <10 to 25% (del Giorgio et al. 1997) or <30% (Cole 1999) reported for a series of freshwater and marine systems, but more similar to those cited for productive estuarine systems (see also Table 1 in del Giorgio & Cole 2000). Applying the relationship BGE = 0.02 × PP0.41 suggested by del Giorgio et al. (1997) to the Lake Kinneret data, we obtained a BGE of 40%.

Rivkin & Legendre (2001) analyzed the results of studies from a range of polar, temperate and tropical marine systems in which BGE was computed from BBP and BR (measured by the uptake of DOC or of O2 in filtered samples), and noted a significant inverse relationship between BGE and water temperature. We did not find any relationship between BGE and temperature in Lake Kinneret. If the results shown by Rivkin & Legendre (2001) in their Fig. 1 apply also to freshwater, then they would imply that our BGE determinations are greatly overestimated for this warm lake.

**BOD5 as a measure of labile organic carbon**

The measurement of BOD is a commonplace and ‘classical’ indicator of water quality. However, with the exception of work in the former Soviet Union and Eastern European countries (e.g. Ostapenya 1971), BOD measurements have rarely been applied to ecological studies of aquatic environments.

Although some of the O2 uptake in the BOD bottles was due to dark respiration of phytoplankton and did not reflect direct extracellular hydrolysis of organic substrates by bacteria, nevertheless we found a highly significant correlation between BBP and BOD5 (Table 3). This result suggests that the indigenous bacterial populations were strongly dependent on the labile organic fractions measured by BOD5. Close to 40% of the variability of BBP could be attributed to variability in BOD5. In addition, CR was significantly correlated with BOD5 (Table 3). These relationships strengthen the premise that BOD5 can indeed serve as a useful and quantitative measure of the labile organic carbon pool in natural water bodies.

During the years 2001 and 2002, DOC concentrations in the epilimnion of Lake Kinneret varied from about 3.5 to 5 g C m−2, with an average of ~50 g C m−2 for the trophogenic water column. The DOC pool comprised ~75% of total organic carbon (TOC) (A. Parparov pers. comm.). The average integrated value for BOD5 down to 15 m was 6.4 ± 2.5 g C m−2. Thus, the labile fraction of organic carbon as inferred by our BOD5 measurements was only about 13 and 10% of the DOC and TOC pools, respectively.

The idea that BBP is dependent on readily available DOC is not new (e.g. Søndergaard et al. 1995). Søndergaard & Middleboe (1995) documented the levels of the labile DOC fraction of the total DOC pool in a variety of aquatic systems. In a recent study, Moran et al. (2001) found that BBP was strongly correlated to ambient DOC in Antarctic waters. We suggest, however, that BBP is dependent not only on the dissolved but also on some of the particulate organic carbon fractions that are readily hydrolyzed by extracellular enzymes.

The concentration of labile organic carbon given by BOD5 divided by the total bacterial carbon demand (i.e. BBP + BR) gives an estimate of the turnover time of the labile pool of TOC taken up by heterotrophic bacteria. Using our measured values for BBP and an estimated BGE of 50%, we calculated a turnover time of ~4 d for the labile fractions of TOC utilized by heterotrophic bacteria, with fastest rates occurring in summer-fall 2001. Using respiration measurements, Parparov et al. (1998) estimated that the average turnover time for POC in Lake Kinneret was about 14 d. Both these rates are much faster than the turnover times of 75 to 120 d for biodegradable DOC measured in batch cultures and in a bio-reactor in samples from Lake Esröm, Denmark, by Søndergaard et al. (2000). At least part of the discrepancy between these estimates may lie in differences in biological availability of the TOC pool components in the 2 systems.

**Conclusions**

The values that we measured for PP, CR, BBP, and our derived BGE give a consistent picture of the major carbon flows in the epilimnion of Lake Kinneret, including sedimentation of POC to the hypolimnion. In contrast to many previously described freshwater systems, the phototrophic zone of this monomictic, large, warm lake, with negligible amounts of allochthonous TOC input, was net autotrophic for most of the year. An independent confirmation of the patterns of net autotrophy/net heterotrophy reported above for this lake could be obtained by means of a gas-flux model (e.g. Cole et al. 2000), but this approach has yet to be applied to this lake (A. Nishri pers. comm.). Measured rates of BBP were high relative to those reported for more oligotrophic and cooler aquatic environments,
Acknowledgements. We are grateful to 2 reviewers who helped us to improve the original manuscript considerably. We thank Bina Kaplan and Sara Chava for devoted and excellent technical help and James Easton and Meir Hatab for skillful manuscript. We also thank Ami Nishri and Tamar Zohary for comments, enlightenment and use of unpublished data. This study was supported in part by Grant No. 95-002 from the United States-Israel Binational Foundation, Jerusalem, Israel, and by the Office of the Israel Water Commissioner (Lake Kinneret Ecosystem Modelling Project. Principal Investigators: T. Zohary, J. Imberger). This is a contribution of Israel Oceanographic and Limnological Research.

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Submitted: June 10, 2003, Accepted: September 18, 2003
Proofs received from author(s): January 7, 2004

Editorial responsibility: Karel Šimek, České Budějovice, Czech Republic