INTRODUCTION

Bacterial mineralization and uptake of organic matter (OM) represent fundamental steps in carbon (C) cycling in aquatic ecosystems. OM quantity and quality can both affect bacterial community composition and substrate utilization (Sinsabaugh & Findlay 2003), implying that different bacterial species vary in their capacity to exploit stored C, and in turn that bacterial community composition may itself influence the efficiency of OM processing (del Giorgio et al. 2011). While labile compounds such as amino acids are readily available for most bacteria, the decomposition of polymeric OM, such as chitin and cellulose, requires specialized pathways, usually controlled by complex regulation processes and interactions (del Giorgio & Newell 2012). Thus, it is assumed that only a specific fraction of natural bacterial communities can access the vast pool of organic polymers. Consistent with this, in many freshwater ecosystems, estuaries and coastal areas (representing the most productive systems and up to 20% of the surface waters on earth, and where most of the OM is directly accessible to only a very limited number of highly specialized bacteria) there appears to be a direct link between OM quality and bacterial community composition (Hutalle-Schmelzer et al. 2010). Bacteria in such communities frequently form syntrophic associations and/or co-metabolism with other species (McInerney et al. 2008), indicating that inter-
specific interactions impact the functioning of the entire ecosystem (Keller & Surette 2006). Consequently, changing community composition due to, for example, invasion by allochthonous organisms following climate change or flooding may cause important changes in system efficiency (Venail et al. 2008, Horrák & Corno 2012). Indeed, recent studies (e.g., Litchman 2010, Hahn et al. 2012) have suggested that microbial interactions are central to improving our understanding of polymeric organic matter degradation by natural microbial communities.

For example, predation of bacteria by protozoa can cause huge changes in bacterial community structure in waters, both by direct selection of the prey and by indirect release of dissolved labile C as exudate during the process of digestion of the captured prey. The concomitant direct and indirect effects of grazing on bacterial communities have been observed in large temperate lakes (Corno et al. 2008), in high mountains lakes (Callieri et al. 2006), and tested in laboratory-based experimental systems (Corno & Jürgens 2008). Indeed, even without protozoan grazing of bacteria, energy transfer and OM cycling will depend on interspecific interactions (Jiao et al. 2010, Nobu et al. 2015). For example, more diverse communities tend to lead to higher C-utilization efficiencies (Miki et al. 2014). Therefore, interspecific interactions that influence community composition, such as invasion by a single competitively dominant species, may influence community-level C-transfer efficiency.

The present study investigates the ecological interactions between microorganisms in simplified multispecies communities (Jessup et al. 2005), testing whether interactions among bacterial species and between bacteria and protozoan grazers influence community assembly and degradation of complex C polymers. We designed a bacterial model community of 4 freshwater species with different physiological behaviors and growth rates when grown on chitin or cellulose as the sole C-source. We selected chitin and cellulose as they represent two of the most abundant biopolymers on earth, particularly in aquatic ecosystems, with huge production rates at a global level (Gooday 1990). Furthermore, their decomposition is not a simple process, but there is no evidence of long-term accumulation of those substrates in aquatic ecosystems (Gooday 1990). This suggests complex and efficient degradation pathways involving a number of interactions (Mayor et al. 2014), as demonstrated in decomposing wood (Borsodi et al. 2005), sediments (Cardenas et al. 2008), wetlands (Ibekwe et al. 2003), and bioreactors (Hiibel et al. 2008).

In order to raise the number of potential interactions, we tested for the effect of a protist (Poterioochromonas, known to graze bacterial cells under a wide range of conditions; Callieri et al. 2006, Corno 2006, Blom & Pernthaler 2010), on bacterial community composition and degradation of biopolymers (chitin and cellulose). We also tested community-level responses to the introduction of an additional bacterial species that we expected to be competitively dominant under these conditions: Burkholderia, common in conditions with abundant polymeric OM (Hutalle-Schmelzer et al. 2010) and potentially able to express chitino- (Kong et al. 2001) and cellulolytic activity (Liang et al. 2014). We found that community-level responses to predation differed from those of individual species, and that introduction of a protozoan grazer can enhance bacterial degradation of both chitin and cellulose by inducing bacterial co-aggregation.

**MATERIALS AND METHODS**

**Organisms**

_Flectobacillus_ sp. strain GC009 (Bacteroidetes, hereafter referred to as _Flectobacillus_) was isolated from an enrichment culture of mesotrophic Lake Schoehsee, Germany. _Arthrobacter agilis_ strain GC027, _Brevundimonas_ sp. strain GC044, and _Aeromonas hydrophila_ strain GC015 were isolated from an enrichment culture of Lake Zürich, Switzerland. Partial 16S rRNA sequences of our _A. agilis_, _Brevundimonas_, and _A. hydrophila_ strains were deposited in GenBank under accession numbers JN009621, JN009622, and KJ409640, respectively. _Burkholderia_ sp. strain AH62 was introduced into some of the microcosms after 3 d of preconditioning the resident bacterial community to the respective substrates (see below). We selected _Poterioochromonas_ sp. strain DS (hereafter referred to as _Poterioochromonas_ or ‘Protist’) as a bacterial grazer because it is common in freshwater, where it is one of the most efficient interception feeding predators (Rothhaupt 1997). Further details of the organisms used are given in the Supplement at www.int-res.com/articles/suppl/a076p027_supp.pdf.

**Experimental setup**

The experimental design consisted of replicate microcosms, each containing a community of 4 bacterial species grown in the dark at 20°C in inorganic artifi-
cial lake water (ALW) medium (Zotina et al. 2003) enriched with chitin or cellulose (27 mg l⁻¹) as the sole C source (final concentration of C = 9 mg l⁻¹). Prior to inoculation, bacteria were preconditioned for 3 d under the same conditions as in the main experiment. We grew such communities in 4 treatments: ‘Control’, with these 4 species only; ‘Burkholderia’, where we introduced an additional bacterial species expected to be competitively dominant under these conditions and therefore to act as an invasive species; ‘Protist’, where we introduced a protozoan grazer expected to act as a predator of bacteria; and ‘Burkholderia+Protist’, with both together. Prior to inoculation, *Burkholderia* was cultivated under the same conditions as all other bacteria, but in DEV Lactose Peptone Broth. In each treatment, we measured productivity, diversity and C transfer efficiency of the community.

Additionally, we grew the protist and *Burkholderia* both alone and in the presence of each individual bacterial species, to determine whether they could exploit those biopolymers in the absence of the mixed bacterial community. We also grew each bacterial species on its own in all 4 treatments to determine whether responses to Burkholderia+Protist, with both together. Prior to inoculation, Burkholderia was cultivated under the same conditions as all other bacteria, but in DEV Lactose Peptone Broth. In each treatment, we measured productivity, diversity and C transfer efficiency of the community.

For each microcosm, the 4 preconditioned bacterial species were inoculated into 500 ml ALW in a 1000 ml Erlenmeyer flask, resulting in an initial total cell abundance of ~4.5 × 10⁴ cells ml⁻¹. The same concentration was used to inoculate single-species controls, which were grown in triplicate on both C sources separately for 11 d in the dark at 20°C. Mixed communities were grown in triplicate on both C sources at 20°C in the dark. On Day 3, each replicate community was divided into two 250 ml sub-treatments, to one of which we added Burkholderia (at 1:100 Burkholderia:resident bacteria). On Day 5, each community incubation was again divided into two 125 ml sub-treatments, one receiving the protist *Petroioochromonas* (at 1:1000 protist:resident bacteria). At each division step the size of the Erlenmeyer flask was reduced accordingly, in order to maintain a constant volume:surface ratio (between 2.2 and 2.4) in the different treatments for the whole experiment. Samples for total cell counts and morphological analysis of resident bacteria in mixed control communities were taken on Days 0, 3, 5, 6, 8, 9, 10, and 11, whereas Burkholderia and Protist treatments were sampled from their creation onwards (Days 3 and 5, respectively). Subsamples (10 ml) for catalyzed reporter deposition-fluorescence in situ hybridization (CARD-FISH) were taken on Days 3, 5, and 11, to estimate the relative abundances of different bacterial species.

**Cell counts and determination of dissolved organic carbon/total organic carbon**

Overall bacterial and protist cell numbers were counted using a FACSaria II flow cytometer (Becton Dickinson) according to the protocol of Gasol et al. (1999). To confirm flow-cytometric counts, bacterial and protist numbers were additionally assessed by epifluorescence microscopy using a Zeiss Axio Imager 1 and 1000× magnification for every fourth sample (results correspondence always >95%, data not shown). Bacterial community composition analyzed by CARD-FISH and total (TOC) as well as dissolved organic carbon (DOC) were determined at Days 3, 5, and 11 (full protocols are provided in the Supplement).

**Aggregate features and C transfer**

Aggregate size was estimated by determining the maximal Feret dimension (the largest possible diameter fitting within the aggregate, Fe_max) of each aggregate detected on DAPI and CARD-FISH stained filters (specific probes for each strain were used, as explained in detail in the Supplement). Aggregates were grouped into size classes using a 10 μm step width. Cell clusters composed of a single strain and with Fe_max <10 μm were termed microcolonies (Corno et al. 2013). Volumes and organic C content of bacteria and flagellate cells were estimated according to Loferer-Krössbacher et al. (1998) and Menden-Deuer et al. (2001); the amount of C utilized by predators after ingestion of the bacterial prey was considered as 43% of the overall prey C content for *Petroioochromonas* (ingested prey biomass = estimated protist biomass gain × 0.43). The overall amount of C for each group of organisms was obtained by multiplying the amount of C estimated per cell for the number of cells of the group. The amount of C excreted by predators was estimated as 35% of the ingested C and considered as labile DOC (Pelegri et al. 1999).

**Statistical analyses**

We first tested whether bacterial growth differed between cellulose and chitin media by taking the average total bacterial abundance over time in each replicate, then using Welch’s *t*-test for an average difference between chitin and cellulose in the Control treatments. By using mixed linear models for
comparing different treatments (Protist, Burkholderia, Burkholderia+Protist or Control), we accounted for the non-independence within replicates in the same as well as in different treatments, by averaging over time and including replicate within chitin or cellulose media as a random effect (‘lme4’ package in R, R Development Core Team 2013). We fitted models by maximum likelihood and tested the significance of fixed effects (treatment and growth medium) by likelihood ratio tests comparing full and reduced models (Crawley 2007). In some cases we compared each treatment pairwise to the Control; in such cases we accounted for multiple testing by only accepting statistical significance when $p < \alpha$, with $\alpha$ adjusted by sequential Bonferroni correction. Bacterial abundances were log-transformed to improve the linear fit; this did not qualitatively affect results. We used the same approach to analyze the fraction of bacterial cells in aggregates at the end of the experiment, arcsine-transforming proportion data prior to analysis.

**RESULTS**

**Bacterial growth on chitin and cellulose with and without Protist and Burkholderia**

Growth of bacteria in pure cultures varied among species and depending on C source (cellulose or chitin; Fig. 1). After 3 d, *Aeromonas hydrophila* grew to ~0.5 × 10⁶ cells ml⁻¹ on chitin, but then rapidly decreased to ~0.15 × 10⁶ cells ml⁻¹, comparable to the same species on cellulose. *Brevundimonas* grew well on chitin, reaching ~10⁶ cells ml⁻¹ after 3 d before slowly declining until Day 11. This species was less successful on cellulose, but still formed distinct populations, with ~0.3 × 10⁶ cells ml⁻¹ after 3 d and decreasing thereafter. *Arthrobacter agilis* and especially *Flectobacillus* did not grow well on either C source (Fig. 1). By contrast, *Burkholderia*, predicted to be well-adapted to these conditions, grew steadily in pure culture, reaching its highest abundances at the final time point on both chitin (0.39 × 10⁶ cells ml⁻¹) and cellulose (0.17 × 10⁶ cells ml⁻¹).

The 4-species bacterial community grown in the absence of *Burkholderia* or the protist reached higher average cell numbers on chitin than on cellulose (Welch’s $t_{3.11} = 15.44$, $p < 0.001$; ‘C’ in Fig. 2). *A. hydrophila* and *Brevundimonas* were the most abundant species in control communities grown on both carbon sources (Fig. 3). By comparing bacterial abundances in pure and mixed cultures, we inferred that interspecific competitive interactions were relatively strong in chitin cultures, as shown by relatively low population densities in mixed cultures compared to a null model based on growth in pure cultures (Fig. S1 in the Supplement at www.int-res.com/articles/suppl/a076p027_supp.pdf).

Total bacterial abundance at the end of the experiment varied among our experimental treatments ($\chi^2 = 53.77$, df = 3, $p < 0.0001$; Fig. 2). Addition of *Poterioochromonas* (‘P’ in Fig. 2) caused significant changes in bacterial growth, with abundance of the resident bacteria rapidly increasing after the protist
was introduced on Day 5, reaching $4.68 \times 10^6$ cells ml$^{-1}$ on chitin and $3.58 \times 10^6$ cells ml$^{-1}$ on cellulose by Day 9. Thus, bacterial abundance after Day 5 was on average higher in grazed (Protist) than non-grazed (Control) communities ($\chi^2 = 32.32$, df = 1, $p < 0.0001$), although bacterial density decreased from Days 9 to 11 in the Protist treatment. The increased bacterial abundance in the Protist treatment was specific to mixed communities: it was not observed in pure cultures of the component species (Fig. S2 in the Supplement). Despite this, addition of the protist did not induce qualitative changes in bacterial community composition: *A. hydrophila* and *Brevundimonas* consistently dominated, with a combined proportion always above 80% (Fig. 3). Addition of *Burkholderia* (‘B’ in Fig. 2), which reached frequencies of up to 16% of the total bacterial population, also increased the average total bacterial abundance after it was added on Day 3 (compared to Control treatment: $\chi^2 = 11.07$, df = 1, $p = 0.001$). This effect was much weaker than that of the protist and was stronger in cellulose than chitin (carbon source × *Burkholderia* interaction: $\chi^2 = 27.09$, df = 1, $p < 0.0001$). Changes in bacterial abundance in the *Burkholderia*+Protist treatment (‘B+P’ in Fig. 2) were similar to those in the Protist treatment, being on average significantly higher than in the Control treatment after Day 5 ($\chi^2 = 27.92$, df = 1, $p < 0.0001$).

To determine whether the positive effects of the protist on total bacterial growth were due to exploitation of cellulose or chitin by the protist itself (which could increase the availability of metabolic by-products for bacterial growth), we measured growth of the protist in the absence of bacteria in chitin and cel-
These experiments showed that *Poterioochromonas* does not grow in the absence of bacteria (Fig. S3). Therefore, it very probably grazes planktonic bacterial cells (Rothhaupt 1997), even though its presence had a net effect of increasing total bacterial abundance in mixed communities.

**Induced morphotype formation**

We detected 3 main bacterial morphotypes: (1) single free-living cells, (2) microcolonies (generally composed of a single strain) of 3 to 10 cells, and (3) large aggregates (generally composed of more than 1 species) of up to a few hundred cells. On both substrates, the proportion of cells in microcolonies accounted for ~15% without and 20 to 25% with the protist (Fig. 4). The fraction of bacteria in aggregates at the end of the experiment varied depending on addition of the protist and *Burkholderia* (effect of treatment: $\chi^2 = 79.64$, df = 3, $p < 0.0001$). Specifically, without the protist large aggregates were absent in cellulose, whereas in chitin 4% of the total bacterial population were present in aggregates. Addition of the protist significantly increased the incidence of aggregates, which accounted for almost 20% of the total bacterial population in both chitin and cellulose (Fig. 4), such that the fraction of aggregated cells was higher in both Protist and *Burkholderia*+Protist treatments.

![Fig. 3. Bacterial community composition at the end of the experiment (Day 11) in the different treatments (C: Control; B: *Burkholderia*; P: Protists; B+P: *Burkholderia*+Protists). *Flectobacillus* is not visible as their proportion was very low and they quickly ran to extinction. Error bars: SD](image1)

![Fig. 4. Proportion of bacteria in each morphotype class with and without protists. Values are means of Control and *Burkholderia* treatment triplicates (without protists), and of Protist and *Burkholderia*+Protist treatments (with protists). Error bars: SD](image2)
compared to the Control (p < 0.0001 in pairwise comparisons; Fig. 4). The proportion of cells in each morphotype class also varied among bacterial species (Fig. S4), with *A. hydrophila*, *A. agilis* and *Burkholderia* being particularly prone to aggregation.

**Carbon flux**

At the end of the experiment, the total amount of C stored in bacterial biomass was higher in the *Burkholderia*, Protist and *Burkholderia*+Protist treatments compared to the Control (Fig. 5). Note that in chitin cultures containing the protist, a significant fraction of additional C was stored in protist biomass. In cellulose cultures, even more C was stored in protists than in bacterial biomass. A schematic description of the C-flow on chitin and cellulose is given in Fig. 6.

**DISCUSSION**

Our results show that interactions among microbial species from different trophic levels play a fundamental role in bacterial exploitation of hard-to-degrade, complex biopolymers. Addition of a protist grazer to a simplified community of 4 bacterial species caused a marked increase in total bacterial abundance (Fig. 2) that was associated with formation of multi-species aggregates and an increase in total C-transfer of the entire community. These findings are particularly relevant to microbial metabolism of C in nature for 2 reasons. First, the C-sources in our experiments (chitin and cellulose) both constitute a large fraction of available C in natural aquatic environments (Gooday 1990, Hopkins & Vallino 2005). Second, aggregation of bacterial species in response to protist grazing has been observed in several other bacteria–protist combinations and experimental conditions (Grossart et al. 2006); anti-predation

![Fig. 5. Estimated proportions of particulate organic carbon bound in bacteria and flagellates at the end of the experiment (Day 11), on either chitin or cellulose. Diamonds: amount of organic carbon respired during the entire experiment.](image1)

![Fig. 6. Graphical and empirical modelling of the carbon fluxes. Fluxes (in mg l⁻¹) are considered within the 4 treatments when grown on chitin (values in red) or cellulose (values in green). Data are directly measured (total dissolved [DOC] and particulate organic carbon [POC]), extrapolated (respired C) or estimated by the mathematical/empirical models (see 'Materials and methods').](image2)
strategies such as formation of biofilms or microcolonies minimize the impact of flagellate predation (Matz & Kjelleberg 2005), although they can incur energetic costs (‘defence cost’ paradigm; Coustau et al. 2000), limiting the ecological success of grazing-resistant strains, and can be complicated by interactions with other organisms and viruses (Våge et al. 2014).

A number of ecological and physiological factors, as well as emerging interspecific interactions, may promote DOC-uptake efficiency and hence productivity in the presence of microcolonies and aggregates (Jousset et al. 2011, Lindström & Östman 2011). Furthermore, indirect effects of protozoan grazing including the release of chemical cues (e.g. allelochemicals; Corno & Jürgens 2006, Blom & Pernthaler 2010), and the accumulation of leftovers from digested bacterial cells (Pernthaler et al. 1997) could potentially promote growth of bacterial cells of the same species that are not preyed upon—especially in systems with limited labile C (as in our experimental system).

The presence of protists has been shown to selectively promote the bacterial degradation of organic detritus with low mineral content (Sherr et al. 1982), again, supporting the hypothesis of a positive effect of grazing on total bacterial productivity. The increased C-transfer to microbial biomass in our Protist treatment, where bacterial aggregation was more common, is consistent with evidence that bacteria in microcolonies or aggregates are more efficient in DOC-uptake and express a higher cell-specific production than free-living cells (Griffith et al. 1994, Grossart et al. 2003, 2007). This may reflect a general difference between diffusion-limitation of access to C-sources in free-living and aggregated cells (Guasto et al. 2012). However, in scenarios where aggregation results from interaction with a protist grazer, as in our experiments, increased productivity may also reflect the growth-promoting effects of the metabolic by-products of protist growth (flagellates such as *Poterioochromonas* excrete ca. 30 to 40% of ingested C as labile DOC; Pelegri et al. 1999), as well as C-transfer to the protist in the form of grazed cells. This is particularly relevant for spatially structured microenvironments such as microcolonies and aggregates (Hagström et al. 1988) where bacteria reach high densities and physiological coupling between microorganisms is likely, e.g. by releasing oxygen radicals (Diaz et al. 2013). Therefore, predation might have a strong impact not only on the composition of bacterial communities but also on their C-transfer efficiency.

Although the chemical structures of chitin and cellulose are homologous, their degradation rates in aquatic bacterial communities are different. Our data showed faster and more efficient degradation of chitin, confirming the results in similar systems observed on a different bacterial community by Peter et al. (2011). As in that study, we cannot exclude the fact that the observed better growth on chitin was promoted by the specific ability for chitin degradation of some of the strains in the community. At the same time, however, we suggest a possibly easier access of simplified bacterial communities to chitin than to other structurally heterogeneous polymers (e.g. cellulose).

Our experimental community included at least 2 strains belonging to genera that are recognized for their potential to degrade chitin and cellulose: chitinolytic activity has recently been attributed to a number of *Burkholderia* strains (Liang et al. 2014), and *Burkholderia gladioli* and *Aeromonas* sp. Strain 10S-24 show chitinolytic activity (Kong et al. 2001, Ueda et al. 2003). The evaluation and comparison of growth curves, as well as of the community compositions between treatments, does not support the hypothesis of a direct competitive advantage for *Burkholderia*. It never exceeded 10 to 16% of the total number of bacteria, while *Aeromonas* was the most abundant strain in every treatment except for the chitin control.

Microbial metabolism of chitin and cellulose is inherently important because (1) they are both highly abundant in nature (Kaplan & Newbold 2003, Beier & Bertilsson 2011, Eckert et al. 2012), (2) relatively few bacteria (1 to 5% of the total bacterial communities in coastal and freshwaters; Cottrell et al. 1999, Beier & Bertilsson 2011) are capable of exploiting such complex biopolymers, but (3) many bacteria utilize substrates that result from chitin or cellulose degradation (up to 40% of the total bacterial communities; Nedoma et al. 1994, Riemann & Azam 2002). Overall, the estimated amount of chitin degraded per day in coastal, estuarine or freshwater systems varies between 2 and 30% of the total pool, while in open seas this proportion is <1% (Gooday et al. 1991, Boyer 1994). Based on this finding, we propose that bacteria in microcolonies and on aggregates (with a relatively lower proportion in the open ocean than in coastal, estuary or freshwater ecosystems) form closely coupled meta-communities, enabling a potentially more efficient community metabolism (Fig. 6) than the suspended, free-living cells. In coastal and inland waters, ca. 20 to 30% of all bacteria occur on biofilms attached to particles and on aggregates (as estimated from data in Maranger & Bird 1995, Whitman et al.
1998, Amaral-Zettler et al. 2010), and although not necessarily chitinolytic or cellulolytic (as in our study), their potential to degrade cellulose and chitin increases by 5- and 10-fold in aggregates. Bacteria occurring on aggregates could thus account for almost 60 and 75% of cellulose and chitin degradation by bacteria in coastal and inland waters, respectively.

In conclusion, protist grazing of bacteria resulted in aggregate formation, and this in turn strongly influenced total bacterial population density and increased community-level metabolism of chitin and cellulose. This suggests that interspecific microbial interactions (Nobu et al. 2015) are of fundamental importance to understanding C-cycling in aquatic environments at a global scale. Thus, altering microbial interactions at the microscale (i.e. in microcolonies and on aggregates) has far-reaching implications for organic matter and C-cycling on earth. Yet it remains unknown how anthropogenic activities (e.g. input of nutrients, toxins, and microplastics) affect microbial interactions in the microscale, and what the ultimate consequences for earth’s biochemistry may be.

Acknowledgements. We thank F. Bulleri, D. Fontaneto, and E. Eckert for suggestions on the experimental modelling, E. Eckert for design and production of graphs, and S. Pinnow for operation of the flow cytometer. Poterioochromonas was provided by J. Blom (University of Zürich). This study was partially funded by the Short-Term Mobility programme 2014 of the National Research Council of Italy. The authors declare no conflict of interest.

LITERATURE CITED


Rothhaupt KO (1997) Nutrient turnover by freshwater bacterivorous flagellates: differences between a heterotro-


Editorial responsibility: Paul del Giorgio, Montreal, Canada

Submitted: February 9, 2015; June 29, 2015
Proofs received from author(s): August 11, 2015