



Potential impacts of black carbon on the marine microbial community

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ABSTRACT: Black carbon (BC) is the carbonaceous residue of the incomplete combustion of fossil fuels and biomass and encompasses a range of chemically heterogeneous substances from partly charred plant material to highly condensed soot aerosols. We addressed the potential role of BC aerosol deposition on marine microbial processes in the ocean by investigating the effects of BC reference material (and its exposure to simulated solar radiation) on viral and bacterial activity in batch cultures with aged seawater. Viruses and bacteria were rapidly adsorbed to BC. No difference between the effect of irradiated and non-irradiated BC on free viral parameters was observed. Bacterial leucine incorporation was higher in the BC treatments than in the BC-free controls. The stimulated bacterial production in the dark BC treatments might be caused by the reduction of viral infection due to adsorption of organic material or by direct use of BC material. Viral production was significantly lower in BC-amended treatments than in BC-free controls, and the estimated fraction of infected cells decreased with increasing BC concentration. Moreover, bacterial activity in the solar-radiation-exposed BC treatments was higher than in the dark BC treatments, indicating that radiation made BC more accessible to bacteria. Our data reveal that BC has the potential to stimulate bacterial activity in the water column, particularly after exposure to solar radiation. Rising BC levels in the atmosphere due to increasing anthropogenic emissions could have far-reaching effects, including potential stimulation of seawater heterotrophy and CO₂ production, through its effects on bacteria and viruses.

KEY WORDS: Black carbon aerosols · Light exposure · Viral lysis · Bacterial production

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INTRODUCTION

Black carbon (BC) refers to the carbonaceous products of incomplete combustion of vegetation and fossil fuels, and is defined as a continuum of compounds from partly charred plant material to highly condensed soot aerosols (Goldberg 1985, Schmidt &

Noack 2000, Mannino & Harvey 2004). These thermogenic compounds are chemically heterogeneous and are all characterized by high carbon content and aromatic structures, which are particularly resistant to biodegradation (Goldberg 1985, Schmidt & Noack 2000). BC particles constitute aggregates of small carbon spheres (Cattaneo et al. 2010), whose fractal

morphology, i.e. their high porosity, offers an important surface area for the adsorption of organic matter (Cornelissen et al. 2005, Koelmans et al. 2006). BC particles occur in a wide size range, and small soot particles (<1 μm) easily become airborne and can remain in the atmosphere for weeks and spread over remote areas including the open ocean (Masiello 2004, Hadley et al. 2007). During atmospheric transport, BC aerosols efficiently absorb solar radiation and are the second strongest contributor to current global warming (Ramanathan & Carmichael 2008).

BC is introduced into the ocean by atmospheric deposition of land-derived aerosols (Jurado et al. 2008), via river estuarine systems (Mitra et al. 2002, Kim et al. 2004, Elmquist et al. 2008, Stubbins et al. 2010), and by marine diesel engine exhausts (Lack et al. 2008). Dry and wet deposition of BC to the global ocean is estimated to be 2 and 10 Tg C yr^{-1} , respectively, with higher fluxes over the northern hemisphere (Jurado et al. 2008). A portion of BC becomes soluble over time and enters the dissolved organic matter (DOM) pool (Kim et al. 2004). Dissolved BC has been observed throughout different domains of the ocean accounting for 1 to 9% of the DOM pool with a strong coastal to open ocean concentration gradient (Mannino & Harvey 2004, Dittmar & Koch 2006, Dittmar & Paeng 2009, Ziolkowski & Druffel 2010, Dittmar et al. 2012). In open ocean sediments, BC is up to 14 000 yr older than the co-deposited bulk organic carbon, suggesting that the small and light BC particles are not used but are retained in the dissolved organic carbon (DOC) pool before being deposited in marine sediments (Masiello & Druffel 1998). However, isotopic analysis revealed the petrogenic origin of BC in marine sediments and suggested a significant overestimation of combustion-derived BC burial in sediments and underestimation of degradation processes in the water column (Dickens et al. 2004). Additionally, the low contribution of BC to marine DOC in the open ocean, i.e. less than 3.5% (Dittmar & Paeng 2009, Ziolkowski & Druffel 2010), suggests that BC is more labile than formerly believed. Recent studies on the photochemical (Stubbins et al. 2012) and microbial degradation of BC (Zimmerman 2010) indeed challenge the previous assumption on the refractory character of BC.

Despite the ubiquitous presence of BC in the oceanic particulate organic matter (POM) (Flores-Cervantes et al. 2009) and DOM pools, its effect on marine systems and its role in the marine carbon cycle remain largely enigmatic. It has been demonstrated that BC reference material rapidly adsorbs viruses and bacteria (Cattaneo et al. 2010) and stim-

ulates the aggregation of organic particles and bacterial production (Mari et al. 2014). Also, there is evidence that BC can have an influence on viral and bacterial diversity (Cattaneo et al. 2010, Weinbauer et al. 2012).

Viruses are important mortality agents in the ocean, structuring microbial communities and influencing biogeochemical cycles (Fuhrman 1999, Suttle 2007) by converting cells into DOM and increasing the recycling and retention of nutrients in the photic zone (Wilhelm & Suttle 1999). Organic particles, in turn, may provide shelter to bacteria from viral infection, although experimental results are controversial (Suttle & Chen 1992, Noble & Fuhrman 1997, Riemann & Grossart 2008, Sheik et al. 2014). As BC particles are a relevant type of organic particle, knowledge on the potential impact of BC-rich aerosols on production and viral-mediated loss of microorganisms and the consequences for biogeochemical cycles in the sea is needed.

To explore the potential mechanisms playing a role in the response of prokaryotes and associated viruses to BC, we set up 2 experiments in which we added various concentrations of standard BC to a standardized equilibrated community from aged seawater from the North Sea. Exposure of BC to solar radiation, a phenomenon that commonly occurs during atmospheric transport, was simulated, and its effect on BC bioavailability to bacteria was also investigated.

Our experiments provide insight into the potential effects of BC on the microbial community, such as: (1) a stimulating effect of BC on bacterial heterotrophic production, as BC is ultimately a carbon source, (2) an increased bioavailability of BC due to previous solar radiation mimicking atmospheric transport, (3) attachment of bacteria and viruses, as BC are organic particles with a high surface area, and (4) a deleterious effect on viral infectivity due to attachment and loss of activity.

MATERIALS AND METHODS

Experimental set-up

Two experiments were performed in May 2005 with aged surface seawater from the open North Sea collected in August 2004 from onboard the RV 'Pelagia' at the North Sea Oyster Grounds (54° 30' N, 04° 30' E) and kept in the dark at 20°C.

Aged seawater was chosen because it is not limited in inorganic nutrients, but instead in organic carbon.

During aging, more degradable organic matter is eliminated which otherwise would obstruct the use of a more refractory carbon source such as BC. Also, other organic particles present in un-aged seawater might adsorb viruses and bacteria and bias the interaction between BC particles and viruses or bacteria. Additionally, aged seawater was used to ensure that the microbial community was in a steady state. Steady-state conditions were assessed prior to the experiments by measuring selected bacterial parameters for 3 consecutive days; these parameters were sampled every 13 ± 3 h and exhibited variations of $\leq 4\%$ (data not shown). The rationale of the present study was to explore the potential mechanisms playing a role in the response of the microbial food web to BC rather than studying the effect of BC on any specific planktonic community. Therefore we used a standardized equilibrated community.

The first experiment was designed to assess the effect of a defined concentration of BC and consisted of duplicate incubations of seawater with and without receiving BC (experiment *BC-fix*). In the second experiment, a gradient of BC concentrations was tested in non-replicated incubations (experiment *BC-grad*). In both experiments, BC reference material (SRM 2975, Diesel Particulate Matter from the US Department of Commerce, National Institute of Standards and Technology [NIST], Gaithersburg, MD) was used. Soot carbon represents 95% of NIST diesel particulate matter (versus 5% organic carbon, Gustafsson et al. 1997), and the trace metal content of SRM 2975 is low (0.22 wt.%, Jensen 2006). The majority of BC reference material particles are within the size range of 1.5 to 4 μm equivalent spherical diameter (ESD). Details on the size distribution of SRM 2975 are given by Cattaneo et al. (2010).

Before being employed in the experiments, BC stock solutions (BC added to ultrapure [MilliQ, Millipore] water at concentrations of 20 mg l^{-1} in experiment *BC-fix* and 100 mg l^{-1} in experiment *BC-grad*) were either kept in the dark or exposed to artificial solar radiation in quartz tubes (2.8 cm inner diameter) for 15 h and 23 h, in *BC-fix* and *BC-grad*, respectively. Artificial solar radiation was supplied by 3 different types of light sources. Two HQI-T Powerstar (Osram) lamps provided photosynthetically active radiation (400–700 nm wavelength range), 2 TL 100W/10R fluorescent light tubes (Philips) were used to provide UV-A (320–400 nm), and 3 UVA-340 fluorescent light tubes (Q-Panel) supplied UV-A and UV-B (300–320 nm). The solar simulator was adjusted to 30–60% of the local maximum radiation intensity in late spring measured on a cloudless day (Pausz &

Herndl 2002). Thus, the dose rate received was similar to that expected in the top surface layer of the North Sea water column. For the dark control (unexposed) treatment, the BC stock was kept in the dark in glass tubes wrapped in aluminum foil in the same incubator. During the exposure to artificial solar radiation, BC stocks were kept in a flow-through water bath connected to a temperature control unit (LAUDA RCS/RC-6).

In experiment *BC-fix*, 10 ml of the light-exposed (BC-light treatment) and unexposed BC stock (BC-dark treatment) were added to 1 l of unfiltered aged seawater in duplicate acid-rinsed borosilicate bottles to obtain a final BC concentration of 200 $\mu\text{g l}^{-1}$. For experiment *BC-grad*, the BC stock solution was added to 1 l of aged seawater to obtain the following concentration gradient: 200, 500, 1000, and 2000 $\mu\text{g l}^{-1}$ light-exposed BC l^{-1} (BC-light treatments) and 500 and 2000 $\mu\text{g l}^{-1}$ unexposed BC l^{-1} (BC-dark treatments). The highest BC concentration measured to date in coastal seawater is 162 $\mu\text{g l}^{-1}$ (Mannino & Harvey 2004), i.e. similar to the concentrations added in experiment *BC-fix* (200 $\mu\text{g l}^{-1}$). Experiment *BC-grad* with up to 10-fold higher additions of BC than has been found in marine environments was designed to unveil the basic effects of BC on microbes, rather than simulating scenarios at ecologically relevant concentrations. Elevated concentrations of BC should also amplify the sorption dynamics and potentiate the interaction between particles and cells or viruses, respectively.

In both experiments, 10 ml of ultrapure water were added to the BC-free controls, and all bottles were incubated in the dark at 20°C for 4 d and sampled every 13 ± 6 h. We started sampling (t_0) approximately 10 min after adding the BC material. The incubation bottles were thoroughly mixed prior to sampling.

Determination of DOC and inorganic nutrients

Samples for the measurement of dissolved inorganic nutrient concentrations (NH_4^+ , NO_3^- , NO_2^- , PO_4^{3-}) were pre-filtered through 0.2 μm polycarbonate filters (GTTP, Millipore), and subsequently analyzed in a TRAACS 800 autoanalyzer system. NH_4^+ was detected with the indo-phenol blue method (pH 10.5) at 630 nm (Helder & De Vries 1979). NO_2^- , NO_3^- , and PO_4^{3-} were determined following the Joint Global Ocean Fluxes Study recommendations (Gordon et al. 1993). NO_2^- was detected after diazotation with sulfanilamide and N-(1-naphthyl)-ethylene

diammonium-dichloride as the reddish-purple dye complex at 540 nm. NO_3^- was reduced in a copper cadmium coil to nitrite and then measured as nitrite. PO_4^{3-} was determined via the molybdenum blue complex at 880 nm.

Samples for DOC measurements were filtered through Whatman GF/F filters using a glass filtration system. The Whatman GF/F filters and all of the glassware were combusted (450°C for 4 h) prior to the filtration. Eight ml of filtered samples were transferred to combusted glass ampoules, and 50 μl of 40% phosphoric acid were added to each sample. Afterwards, the ampoules were sealed and stored at 4°C in the dark and analyzed within 1 mo. DOC concentrations were determined in triplicate using a Shimadzu TOC-5000 analyzer (Benner & Strom 1993).

Determination of total bacterial abundance, respiring bacteria, and 'live' versus 'dead' bacteria

Water samples for total bacterial abundance (1.8 ml) were fixed with paraformaldehyde (1%) and glutaraldehyde (0.05%), kept at room temperature for ca. 10 min, and then flash-frozen in liquid nitrogen. Within a few days, the samples were thawed, stained with SYBR Green I (Molecular Probes, Invitrogen) at $10\times$ dilution for 10 min, amended with Fluoresbrite[®] Yellow Green (YG) beads of 0.98 μm diameter (Polysciences) as an internal standard, and analyzed in a Becton Dickinson FACSCalibur flow cytometer as described previously (Gasol & del Giorgio 2000).

Bacteria were detected in plots of 90° light scatter (SSC) and green DNA fluorescence (Fig. 1). Differences in the green fluorescence and SSC signature in the cytometric plot allowed us to separate different bacterial populations according to their nucleic acid content, i.e. low (LNA) and high nucleic acid (HNA) content, as previously described (Gasol et al. 1999).

In the samples with BC additions, we detected an additional population with a SYBR Green derived fluorescence comparable to HNA bacteria, yet with a much higher SSC (Fig. 1). The bacteria in that gate were operationally considered to be particle-associated bacteria (PAB) assuming that the increase in SSC without parallel increase in nucleic acid-derived fluorescence had to be due to the association of bacteria with particles.

The numbers of actively respiring bacteria were determined by the fluorogenic tetrazolium dye CTC (5-cyano-2,3-ditolyl tetrazolium chloride) labeling of

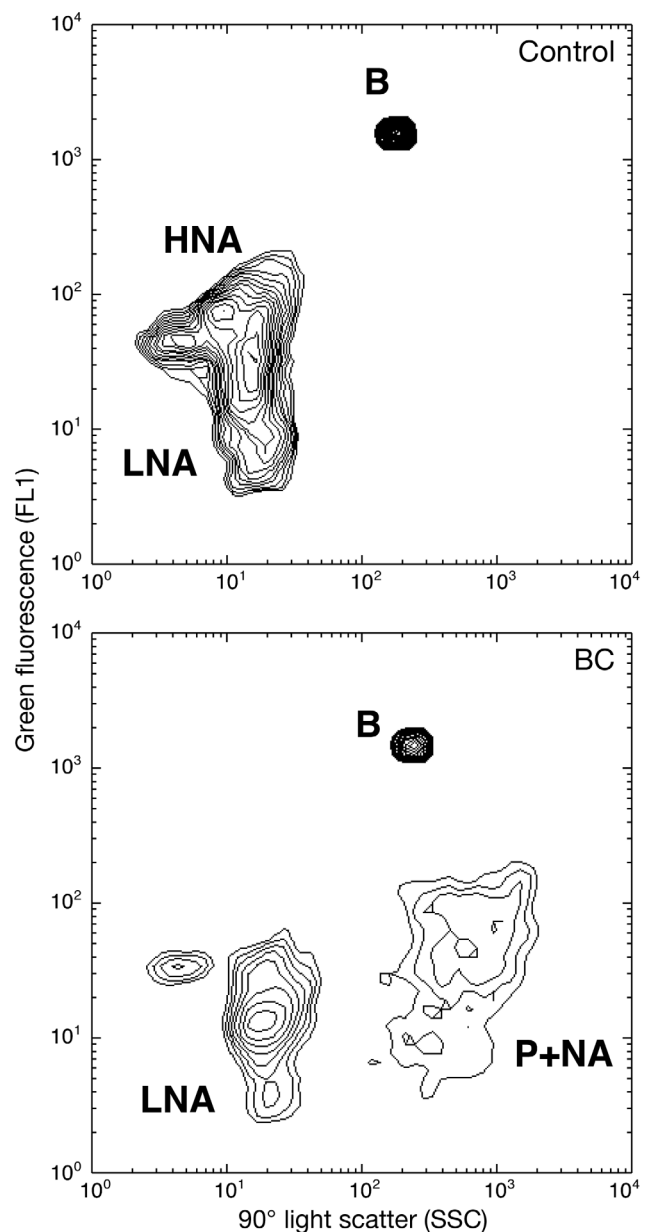


Fig. 1. Flow cytometric plot of side scatter (SSC) versus green fluorescence (FL1) of SYBR-Green-stained bacteria from experiment *BC-grad*. Examples are given of bacteria that had received black carbon (BC; lower panel) and bacteria from the BC-free control (upper panel). A secondary population of particle-associated bacteria (P+NA) with higher SSC and similar fluorescence to high nucleic acid (HNA) bacteria is evident in the lower panel. B: beads; LNA: low nucleic acid bacteria

highly active cells (del Giorgio et al. 1997, Sherr et al. 1999, Sieracki et al. 1999). Five mM CTC of a daily prepared batch was added to water samples and incubated at 20°C for 3 h. After incubation, the samples were analyzed with a Becton Dickinson FAC-

S-Calibur flow cytometer as described in detail elsewhere (Gasol & Arístegui 2007).

We also discriminated 'live' or membrane-intact from 'dead' or membrane-damaged bacteria using the nucleic-acid double staining protocol (Gregori et al. 2001) as explained in detail elsewhere (Falcioni et al. 2008). In brief, samples were dually stained with propidium iodine ($10 \mu\text{g ml}^{-1}$), a membrane-impermeable red dye, and SYBR Green I (Molecular Probes, Invitrogen), a membrane-permeable green dye, for 15 min, and then analyzed in a Becton Dickinson FACSCalibur flow cytometer at a low speed (ca. $15 \mu\text{l min}^{-1}$). A dot plot of red (FL3 cytometric channel) versus green fluorescence (FL1 cytometric channel) allowed distinction of the 'live' cells from the 'dead' cells. 'Live' cells were considered those having more green than red staining, while 'dead' cells had more red than green.

Bacterial heterotrophic production

Subsamples for the measurement of bulk leucine incorporation by bacteria were taken from the different treatments and treated using the method as outlined by Simon & Azam (1989). Two 5 ml samples and one 5 ml formaldehyde-killed blank per treatment were inoculated with ^3H -leucine (20 nM final concentration, Amersham, specific activity 160 Ci mmol^{-1}) and incubated in the dark at 20°C for 2 h. Subsequently, the samples were fixed with formaldehyde (2% final concentration), filtered onto $0.2 \mu\text{m}$ Millipore GTTP polycarbonate filters, and rinsed 3 times with 10 ml of 5% ice-cold trichloroacetic acid. The filters were then transferred to scintillation vials and dried at room temperature. One ml of ethyl acetate and 8 ml of scintillation cocktail (Packard Gold Insta Gel) were added to each vial. The vials were counted in an LKB liquid scintillation counter after 18 h. The obtained disintegrations per minute (DPMs) were converted to leucine incorporation rates. Leucine incorporated into bacterial biomass was converted to bacterial carbon production using the empirical conversion factor $0.07 \times 10^{18} \text{ cells mol}^{-1} \text{ Leu}$ (Riemann et al. 1990) and assuming a C content of bacteria of $20 \text{ fg C cell}^{-1}$ (Lee & Fuhrman 1987). The application of this conversion factor resulted in similar bacterial heterotrophic production (BHP) estimates as with the theoretical factor of $1.55 \text{ kg C mol}^{-1} \text{ Leu}$ that assumes no isotope dilution (Simon & Azam 1989). BHP divided by bacterial biomass provided estimates of the specific growth rate of bacteria (d^{-1}).

Viral abundance

Viral abundance was enumerated by flow cytometry (FC), which currently cannot be combined with virus-aggregate disruption agents such as methanol, since it interferes with the staining dye (Weinbauer et al. 2009). Thus, bulk measurements represent likely free viral abundance (FVA). In specific samples, viruses attached to BC particles were enumerated by epifluorescence microscopy (see below).

For viral counts by FC, we followed the optimized protocol by Brussaard (2004). Subsamples (2 ml) were fixed with glutaraldehyde (0.5% final concentration), incubated at 4°C for 15 to 30 min, and subsequently frozen in liquid nitrogen and stored at -80°C . Upon thawing, viruses were stained with SYBR Green I (Molecular Probes, Invitrogen) at a final concentration of 0.5×10^{-4} of the commercial stock at 80°C for 10 min and quantified using a FACSCalibur (Becton and Dickinson) flow cytometer after dilution with TE buffer (10 mM Tris, 1 mM EDTA, pH 8).

Viral production

Viral production (VP) was estimated for selected samples using the virus reduction technique (Weinbauer et al. 2010). The rationale behind the virus reduction approach is to reduce virus abundance and thereby essentially prevent new viral infection. Thus, the viruses produced originate from already infected cells. Bacteria from 200 ml of raw seawater were concentrated using a $0.2 \mu\text{m}$ pore size tangential flow system (VIVAFLOW 50). To obtain virus-free seawater, the $0.2 \mu\text{m}$ pore-size ultrafiltrate was passed through a 100 kDa cartridge (VIVAFLOW 50). The bacterial concentrates were brought up to the original volume with virus-free seawater and incubated in duplicate 50 ml Falcon tubes (BD Biosciences) at $20 \pm 2^\circ\text{C}$ for 24 h. At t_0 of the experiments, 2 additional tubes were amended with mitomycin C (Sigma) at a final concentration of $1 \mu\text{g ml}^{-1}$ in order to induce the lytic cycle in prophages. Subsamples (2 ml) for viral abundance were taken every 3 to 4 h from each incubation, fixed with glutaraldehyde (0.5% final concentration), incubated at 4°C for 15 to 30 min, subsequently frozen in liquid nitrogen and stored at -80°C until counted by flow cytometry. Lytic VP was calculated as the increase in viral abundance over short time intervals ($\sim 4 \text{ h}$). An increase in viral abundance in the mitomycin C treatments represents lytic + lysogenic production (Paul & Weinbauer 2010, Weinbauer et al. 2010). VP was corrected for the changes in bacterial

abundance in the VP assays. Dividing the number of produced phages by an assumed burst size (BS) of 50 (Parada et al. 2006) yields the number of lysed cells and gives an estimate of the fraction of infected cells (FIC) when divided by the bacterial abundance at the start of the experiment (Weinbauer et al. 2002). Lysis rates were calculated by dividing VP by BS and were used to calculate virus-mediated mortality (VMM) as a percentage of bacterial standing stock (BSS) per day or as a fraction of BHP. Alternatively, VMM was related to FIC using the model by Binder (1999).

BC particle size and attachment of viruses and bacteria

Associations between BC particles, bacteria, and viruses were investigated by epifluorescence and confocal laser scanning microscopy (CLSM), respectively, in experiment *BC-grad* following the protocol of Cattaneo et al. (2010). The size and number of colonized particles were high enough to allow enumeration only in the treatment with the addition of BC at the highest concentration ($2000 \mu\text{g l}^{-1}$). Slides were prepared using a slightly modified version of the Noble and Fuhrman procedure (Noble & Fuhrman 1998). Subsamples of 2 ml were fixed with glutaraldehyde (0.5% final concentration) at 4°C for 15 to 30 min, frozen in liquid nitrogen, and stored at -80°C until analysis. Thawed samples were filtered onto $0.02 \mu\text{m}$ pore size AlO_3 filters (25 mm diameter, Anodisc, Whatman) by low-pressure filtration and stained with SYBR Gold (Molecular Probes, Invitrogen) diluted 1000-fold in autoclaved and $0.2 \mu\text{m}$ filtered MilliQ water. The filters were transferred onto slides with a mounting solution (0.1% *p*-phenylenediamine; freshly made from a frozen 10% aqueous stock Sigma-Aldrich, P-1519) in 50% glycerol-50% phosphate-buffered saline (PBS, 0.05 M Na_2PO_4 , 0.85% NaCl, pH 7.5) and Vectashield (1:6 v:v; Vector, Burlingame). The slides were stored at -20°C until examined under the CLSM.

Initially, abundances of free and BC-attached viruses and bacteria were assessed at 1200-fold magnification with an epifluorescence microscope (Axio-phot, Carl Zeiss). At least 25 microscopic fields were inspected. Colonized BC particles were then investigated with a CLSM (Leica SP2) equipped with an argon neon laser (excitation: 488 nm; emission spectrum: 530–550 nm). For each particle, stacks of images were acquired and 3-dimensional surfaces were measured for each particle as previously described (Cattaneo et al. 2010). The surface area of

particles was converted into ESD (i.e. the diameter of spheres with equivalent volume to nonspherical-shaped particles; Peduzzi & Weinbauer 1993). We quantified the abundance of attached viruses and bacteria, and manual counting was preferentially used over automated counting to ensure the required accuracy of the measurements (Luef et al. 2009, Cattaneo et al. 2010).

Statistics

All statistical analyses were performed with JMP 7.0 (SAS). The Shapiro-Wilk *W*-test was used to check for normal distribution of data. Analysis of covariance (ANCOVA) and 1-way ANOVA for normal distributions and Kruskal-Wallis tests for non-normal distributions were used to evaluate the differences between treatments. Spearman rank correlation for nonparametric data was performed to determine the relationships between the various parameters measured.

RESULTS

Effects of BC on nutrients and DOC

The addition of BC resulted in differences in inorganic nutrients and DOC concentrations between treatments already at t_0 : light-exposed BC additions produced a significant increase in DOC concentration in experiment *BC-fix* (ANCOVA, $p < 0.05$) compared to the BC-free control. Phosphate concentrations were higher in BC incubations than in the BC-free control, although the differences were only significant in experiment *BC-grad*. In both experiments, average ammonium concentrations were 5- to 6-fold higher with BC additions than in the controls (ANCOVA, $p < 0.05$, Table 1). In most cases, nutrient concentrations did not change significantly over the course of the experiments, but ammonium and nitrate concentrations in experiment *BC-fix* and nitrite concentration in experiment *BC-grad* increased (ANCOVA, Table 1).

Total BA and the number of active bacteria

In both experiments, BA did not vary significantly with time in the BC-free control nor in the treatment with the addition of $200 \mu\text{g BC l}^{-1}$ (ANCOVA, Table 2, Fig. 2). However, BA increased with time and BC con-

Table 1. Mean \pm SD values of the nutrient and dissolved organic carbon (DOC) concentrations (μM) analyzed in each of the different treatments as well as results of analyses of covariance with time (*BC-fix*) and with time and black carbon (BC) concentration (*BC-grad*) as covariates. Values are averaged over the experimental time ($n = 24$). Significantly different values are highlighted in **bold**: ** $p < 0.001$, * $p < 0.05$; ns: not significant. Treatments are as follows: BC-free: BC-free control; BC-dark: with 200 (*BC-fix*) or 500 and 2000 (*BC-grad*) $\mu\text{g BC l}^{-1}$ maintained in the dark; BC-light: with 200 (*BC-fix*) or 200, 500, 1000, and 2000 (*BC-grad*) $\mu\text{g BC l}^{-1}$ exposed to light

Experiment	Treatment	PO_4^{3-}	NH_4^+	NO_2^-	NO_3^-	DOC
<i>BC-fix</i>	BC-free	0.11 ± 0.02	$0.14 \pm 0.03^{**}$	0.01 ± 0.005	0.08 ± 0.02	81.3 ± 3.4
	BC-dark	0.16 ± 0.03	0.87 ± 0.23	0.02 ± 0.003	0.09 ± 0.03	80.9 ± 4.7
	BC-light	0.15 ± 0.03	0.93 ± 0.42	0.02 ± 0.01	0.11 ± 0.04	$91.1 \pm 13.6^*$
Covariate	Time	ns	*	ns	*	ns
<i>BC-grad</i>	BC-free	$0.10 \pm 0.00^*$	$0.17 \pm 0.10^*$	0.02 ± 0.02	0.07 ± 0.02	80.3 ± 3.8
	BC-dark	0.14 ± 0.02	0.76 ± 0.50	0.04 ± 0.01	0.12 ± 0.05	83.0 ± 6.1
	BC-light	0.16 ± 0.03	0.90 ± 0.59	0.04 ± 0.01	0.10 ± 0.03	83.6 ± 9.0
Covariate	Time	ns	ns	*	ns	ns
	BC concentration	ns	ns	ns	ns	ns

Table 2. Mean \pm SD values of organism abundance and activity analyzed in each of the different treatments as well as results of analyses of covariance with time (*BC-fix*) and time and black carbon (BC) concentration (*BC-grad*) as covariates. Values are averaged over experimental time. Significantly different values are highlighted in **bold**: *** $p < 0.0001$, ** $p < 0.001$, * $p < 0.05$. Treatments as in Table 1. BHP: bacterial heterotrophic production; BA: total bacterial abundance; PAB: particle-attached bacteria; CTC: 5-cyano-2,3-ditolyl tetrazolium chloride reducing cells; LIVE: membrane-intact cells using the nucleic acid double staining protocol (see Materials and Methods); FVA: free viral abundance

Experiment	Treatment	BHP $\mu\text{g C l}^{-1} \text{d}^{-1}$	BA $\times 10^5 \text{ ml}^{-1}$	PAB %	CTC %	LIVE %	FVA $\times 10^6 \text{ ml}^{-1}$
<i>BC-fix</i>	BC-free	1.7 ± 0.2	2.0 ± 0.2	-	62 ± 11	$83 \pm 4^{***}$	$12.7 \pm 0.5^*$
	BC-dark	1.9 ± 0.1	2.2 ± 0.2	-	64 ± 13	78 ± 6	11.3 ± 0.8
	BC-light	$3.0 \pm 0.8^{***}$	2.2 ± 0.2	-	$56 \pm 13^*$	75 ± 6	11.5 ± 0.7
Covariate	Time	*	ns	-	*	ns	ns
<i>BC-grad</i>	BC-free	1.7 ± 0.4	1.5 ± 0.1	$6 \pm 1^{***}$	49 ± 23	72 ± 4	$12.7 \pm 0.7^{**}$
	BC-dark	2.4 ± 1.2	1.8 ± 0.4	49 ± 10	54 ± 11	76 ± 6	7.3 ± 1.7
	BC-light	2.6 ± 1.5	1.8 ± 0.3	47 ± 10	54 ± 16	75 ± 5	8.2 ± 2.2
Covariate	Time	*	**	*	***	***	ns
	BC concentration	***	***	***	ns	*	***

centration in the other treatments (ANCOVA, $p < 0.0001$, Table 2, Fig. 2B), reaching 78% higher BA in the light-exposed and 68% higher BA in the dark-exposed BC treatment at 2000 $\mu\text{g BC l}^{-1}$ with respect to initial values. The percentage of actively respiring cells that reduced 5-cyano-2,3-ditolyl tetrazolium chloride (CTC) was significantly lower in the BC-light treatment than in all other treatments in experiment *BC-fix* (ANCOVA, $p < 0.05$, Table 2). The percentage of membrane-intact cells (LIVE) was significantly lower in the BC treatments than in the control in *BC-fix*, but increased slightly with BC concentration and time in *BC-grad* (ANCOVA, $p < 0.0001$, Table 2).

Abundance of free viruses

Initial free viral abundance (FVA) as measured by flow cytometry was 1.26×10^7 and 1.16×10^7

viruses ml^{-1} in experiments *BC-fix* and *BC-grad*, respectively. In the BC-free controls, these values remained roughly stable from t_0 throughout the experiment *BC-fix* (Fig. 3A) while in *BC-grad* they increased by 8% until t_{8h} and then stayed approximately unchanged until the end (Fig. 3B). Upon BC amendment, FVA decreased from t_0 in both experiments and remained roughly constant after the first sampling 8 h later (Fig. 3). In *BC-fix*, this decrease amounted to 13% with respect to the BC-free controls. After 80 h, FVA increased in all treatments and finally leveled off at ca. 1.34 ± 0.09 (range) $\times 10^7$ viruses ml^{-1} in the controls, and at $1.27 \pm 0.03 \times 10^7$ and $1.29 \pm 0.03 \times 10^7$ viruses ml^{-1} in the light and dark BC treatments, respectively. Average FVA was significantly lower in the BC treatments than in the BC-free controls and was not influenced by BC light exposure (Table 1, Fig. 3A, ANCOVA, $p < 0.05$).

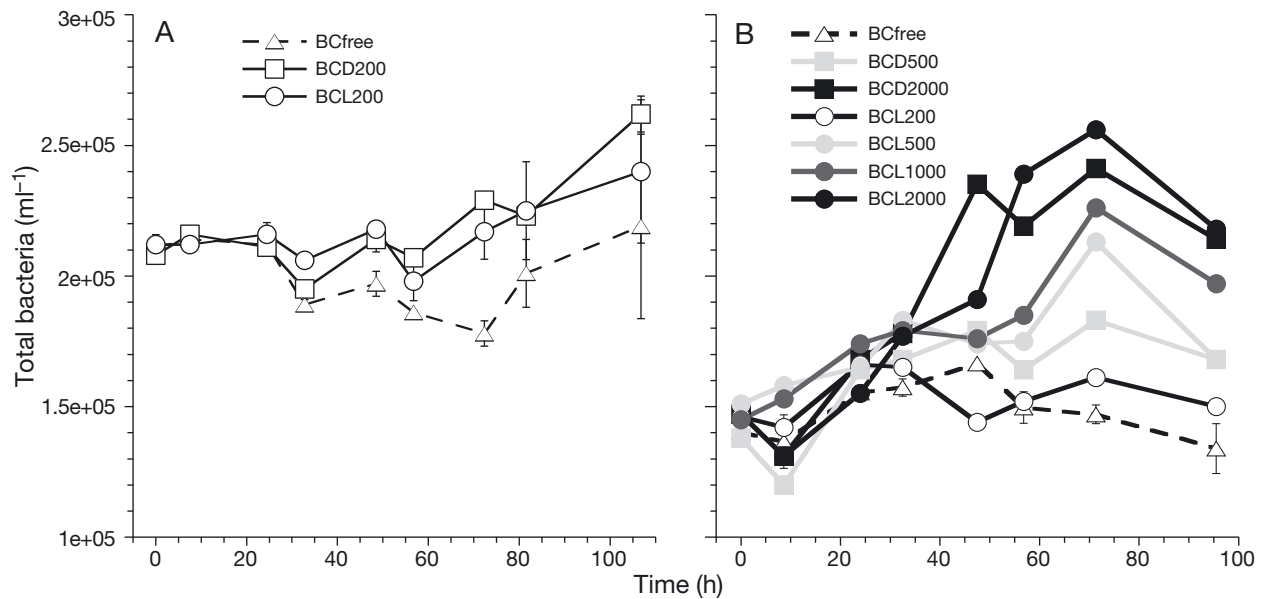


Fig. 2. Bacterial abundance in experiments (A) *BC-fix* and (B) *BC-grad* with addition of light-exposed (BCL) and non-exposed (BCD) black carbon (BC) and under control treatments (BCfree). Values are averages from duplicate treatments and error bars are the ranges in experiment *BC-fix* and in the BC controls of experiment *BC-grad*. BC concentrations are indicated ($\mu\text{g l}^{-1}$)

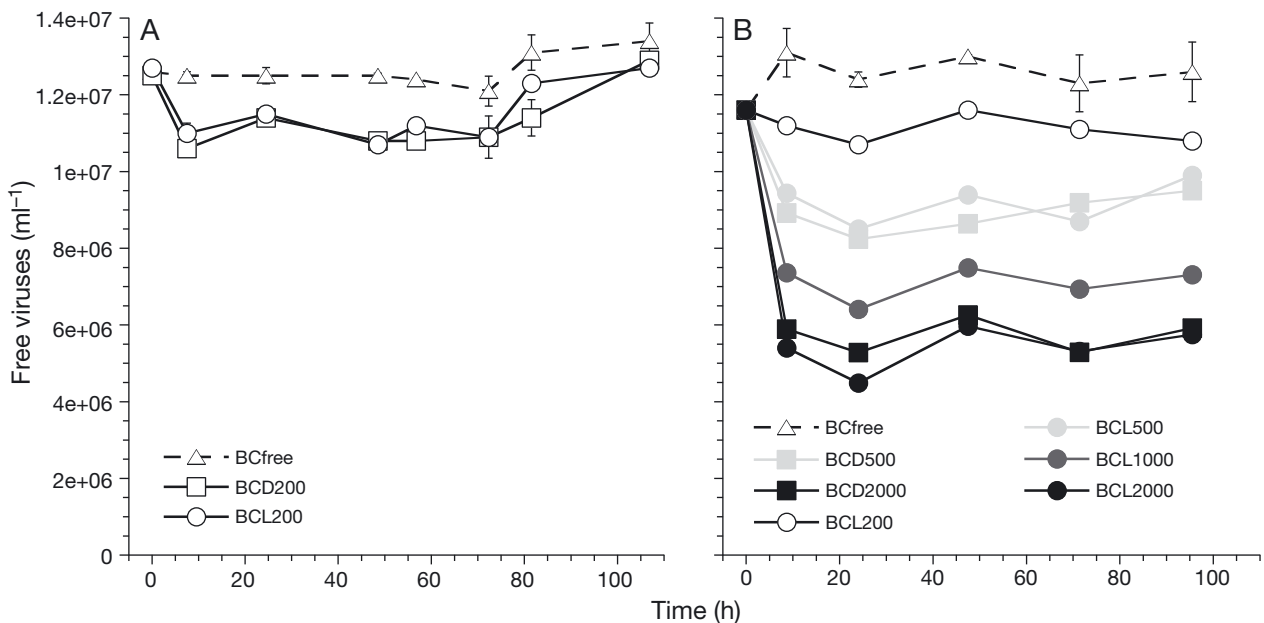


Fig. 3. As in Fig. 2, but for the abundance of free viruses in experiments (A) *BC-fix* and (B) *BC-grad*. Where not visible, error bars are contained within the symbol

In experiment *BC-grad*, FVA averaged over the experimental time decreased with increasing BC concentrations (ANCOVA, $p < 0.0001$) but was not influenced by BC light exposure (Table 2, Fig. 4B).

The virus-to-bacteria ratio (VBR) ranged from 21 to 109 and averaged 56 ± 16 without differences between experiments. VBR was significantly lower with BC addition only in the experiment *BC-grad* (47 vs. 88, ANCOVA, $p < 0.05$,

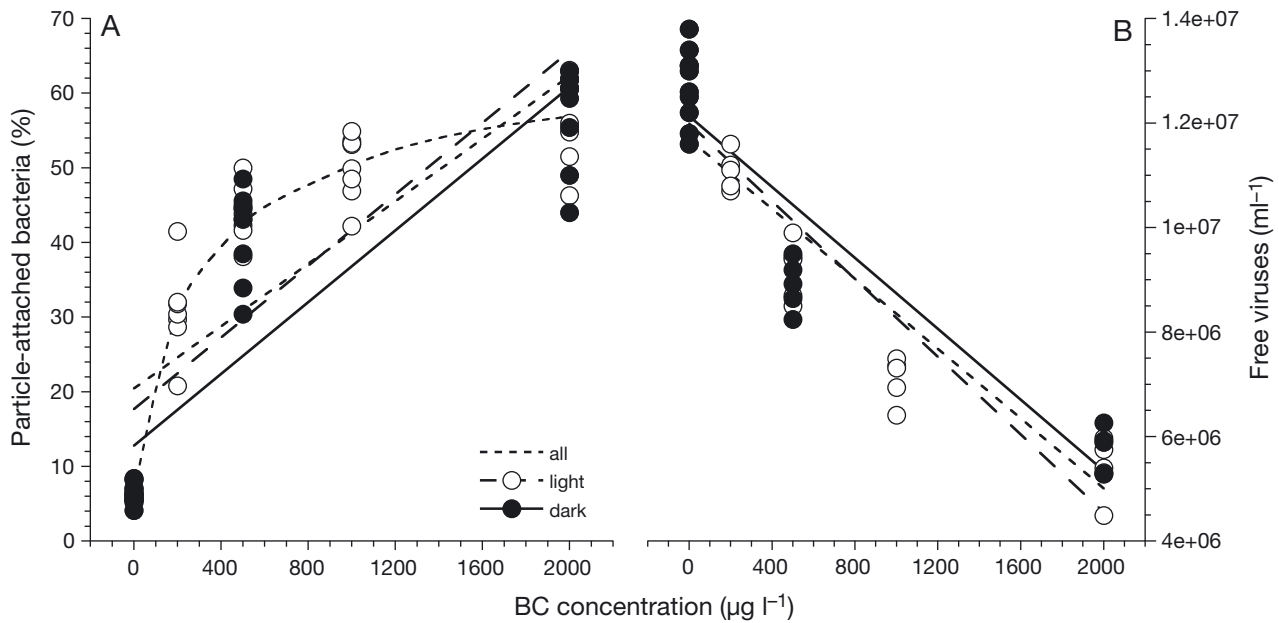


Fig. 4. (A) Percentage of particle-attached bacteria (PAB) and (B) abundance of free viruses (FVA) in relation to black carbon (BC) concentration in experiment *BC-grad*. When interpolating all data, it is apparent that BC particles were becoming saturated at a BC concentration of $1000 \mu\text{g BC l}^{-1}$, and at most about half of the bacteria were becoming attached. PAB increased with BC concentration, while FVA decreased with BC concentration: all PAB = $20\% + 0.02 \text{ BC concentration}$ ($R^2 = 0.67$); light BC treatment: PAB = $18\% + 0.02 \text{ BC concentration}$ ($R^2 = 0.69$); dark BC treatment: PAB = $13\% + 0.02 \text{ BC concentration}$ ($R^2 = 0.77$); Virus: all FVA = $1.17 \times 10^7 - 3344 \text{ BC concentration}$ ($R^2 = 0.87$); light BC treatment: FVA = $1.20 \times 10^7 - 3721 \text{ BC concentration}$ ($R^2 = 0.88$); dark BC treatment: FVA = $1.21 \times 10^7 - 3395 \text{ BC concentration}$ ($R^2 = 0.88$)

covariates: BC concentration: $p < 0.0001$, time: non-significant).

BC-attached viruses and bacteria and size of the BC particles

In *BC-grad*, the size of BC particles with attached bacteria and viruses as analyzed by CLSM ranged from 4.6 to $22.4 \mu\text{m}$ ESD (average: $9.5 \pm 5.4 \mu\text{m}$), but 58% of the particles were smaller than $10 \mu\text{m}$ ESD throughout the experiment except t_0 (details not shown). Fifty-six percent of the particles had fewer than 10 attached bacteria. The abundance of attached bacteria was positively related to BC particle size ($BA = -0.03 + 0.94 \text{ ESD}$, $n = 16$, $R^2 = 0.38$, Table 3), and the average number of bacteria per ESD was $0.9 \pm 0.7 \mu\text{m}^{-1}$. The number of BC-attached viruses ranged between 4 and 64 viruses per particle. A positive correlation was observed between the abundance of BC-attached viruses and BC particle size, while the number of BC-attached viruses per μm^2 was negatively correlated with BC particle size (Table 3).

At the end of the incubation, in the treatment with the addition of BC at the highest concentration

Table 3. Spearman rank correlation analysis of microbial abundances with black carbon (BC) particle size from the treatments with the addition of $2000 \mu\text{g BC l}^{-1}$ kept in the dark in experiment *BC-grad*. VBR: virus-to-bacteria ratio; ns: not significant

	ρ	n	p
Abundance of attached:			
viruses particle ⁻¹	0.640	18	<0.05
viruses μm^{-2}	-0.612	18	<0.05
bacteria particle ⁻¹	0.715	16	<0.05
bacteria μm^{-2}	-0.441	16	ns
VBR μm^{-2}	-0.738	16	<0.05

($2000 \mu\text{g l}^{-1}$), BC-attached viruses assessed by epifluorescence microscopy accounted for 40% of total viral abundance, and attached bacteria accounted for 50% of the total bacterial abundance. The percentage of particle-attached bacteria measured using the FC approach was only recorded in *BC-grad* and ranged from 4–8% in the BC-free controls (average $6 \pm 1\%$, considered to be the background counts of the technique), to 44–63% (average $57 \pm 6\%$) at $2000 \mu\text{g BC l}^{-1}$. The percentage of particle-attached bacteria increased significantly with increasing BC concentration (Fig. 4A, Table 2). Furthermore, per-

centage of particle-attached bacteria correlated positively and significantly with bacterial production ($\rho = 0.335$, $n = 55$, $p < 0.05$) and with the % HNA ($\rho = 0.310$, $n = 55$, $p < 0.05$). No effect of exposure to artificial solar radiation of BC on the percentage of particle-attached bacteria was detected.

Bacterial heterotrophic production

In *BC-fix*, the addition of irradiated BC led to a significant increase in BHP with respect to the other treatments after 24 h (ANCOVA, $p < 0.0001$), and this level of activity was maintained throughout the experiment (Fig. 5A). In contrast, BHP in the BC-dark treatment was not significantly different from the BC-free controls and increased by only 32% (vs. 148% in treatment BC-light, Fig. 5A).

In *BC-grad*, BHP increased significantly with increasing BC concentrations in all BC treatments (Table 2, Fig. 5B). BHP amended with light-exposed BC was up to 12 times higher than the initial BHP of $0.7 \mu\text{g C l}^{-1} \text{d}^{-1}$, whereas BHP in the dark BC treatment was only higher than the BC-free control at the highest BC concentration ($2000 \mu\text{g BC l}^{-1}$), reaching $5 \mu\text{g C l}^{-1} \text{d}^{-1}$. Time-averaged BHP for the time interval of 2 to 3 d was significantly higher with the addition of light-exposed BC than in the other treatments

(ANCOVA, $p < 0.05$). The specific growth rate of bacteria showed the same trend as BHP in both experiments (details not shown).

Viral production and infection

VP was measured at t_0 and after 2 d in all treatments of experiment *BC-fix*. In *BC-grad*, VP was measured only in some of the treatments (Table 4). Tangential flow filtration through a 100 kDa cartridge removed viruses efficiently, resulting in $<1\%$ of the original viral abundance. At t_0 , viral abundance in the VP incubations ranged between 9 and 40% of the original viral abundance. These viruses were introduced along with the bacterial concentrate into the incubations for estimating VP.

In *BC-fix*, lytic VP increased from $1.3 \pm 0.3 \times 10^6$ viruses $\text{ml}^{-1} \text{d}^{-1}$ at t_0 to $2.7\text{--}24.3 \times 10^6$ viruses $\text{ml}^{-1} \text{d}^{-1}$ after 2 d of incubation (Table 4). VP was significantly lower in the dark BC treatments compared to the BC control and the light-exposed BC treatments (Table 4, Kruskal-Wallis test, $p < 0.05$). At t_0 , $12 \pm 2\%$ bacteria were infected, corresponding to an estimated viral lysis rate of $2.5 \pm 0.6 \times 10^4$ bacteria $\text{ml}^{-1} \text{d}^{-1}$. At t_{2d} , significantly fewer bacteria were infected in the dark BC treatment than in the BC control and in the light BC treatment, and estimated viral lysis rates were signifi-

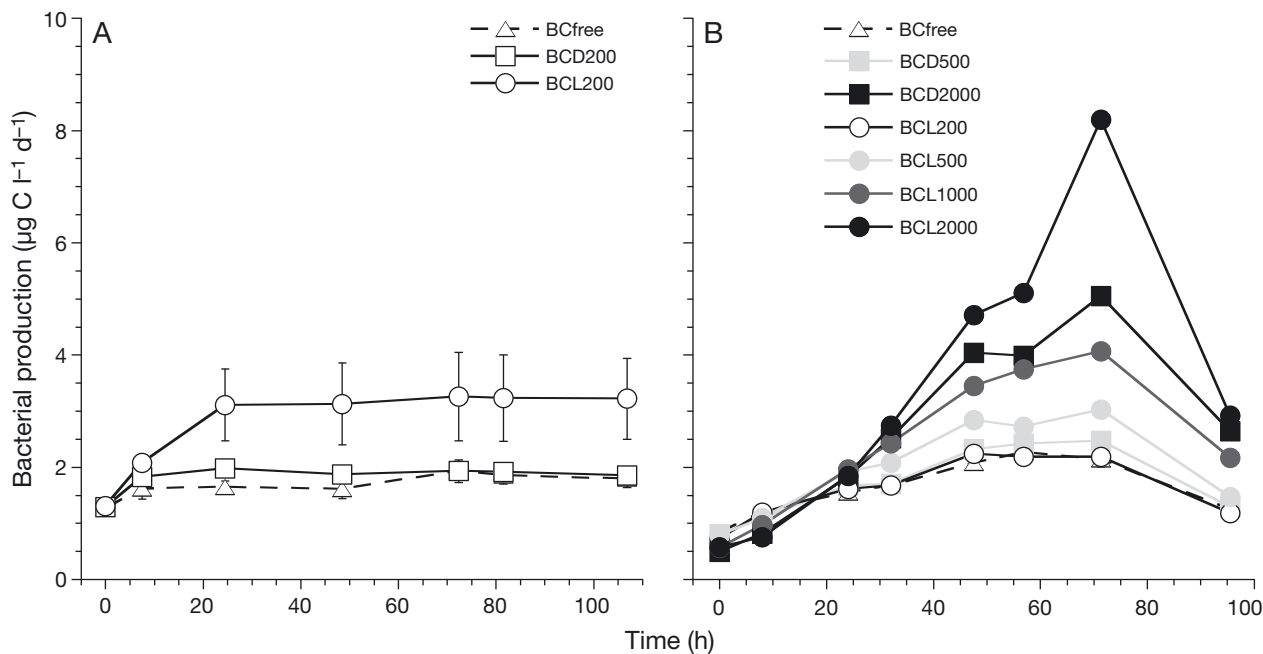


Fig. 5. Bacterial heterotrophic production in experiments (A) *BC-fix* and (B) *BC-grad* with addition of light-exposed (BCL) and non-exposed (BCD) black carbon (BC) and under control treatments (BCfree). Values are averages from duplicate treatments and error bars are the ranges in experiment *BC-fix* and in the BC controls of experiment *BC-grad*. BC concentrations are indicated ($\mu\text{g l}^{-1}$)

Table 4. Viral production (VP, viruses ml⁻¹ d⁻¹), frequency of infected cells (FIC, %), bacterial lysis rates and virally mediated loss of bacterial standing stock (VMM_{SS}, % d⁻¹) and viral mediated mortality related to FIC (VMM, %) by a model (Binder 1999). Data are given as means ± range of duplicate measurements. Significantly different values for treatments are highlighted in **bold** (p < 0.05, Kruskal-Wallis test). Treatments as in Table 1

Experiment	Time (d)	Treatment	VP × 10 ⁶ ml ⁻¹ d ⁻¹	FIC %	Lysis rate × 10 ⁴ ml ⁻¹ d ⁻¹	VMM _{SS} % d ⁻¹	VMM %
<i>BC-fix</i>	0		1.3 ± 0.2	12 ± 2	2.5 ± 0.4	12 ± 2	25 ± 4
	2	BC-free	9.9 ± 2.9	20 ± 5	19.7 ± 5.8	101 ± 26	46 ± 17
		BC-dark	3.6 ± 0.9	10 ± 3	7.2 ± 1.8	34 ± 9	21 ± 8
		BC-light	15.4 ± 9.5	27 ± 16	30.8 ± 19.1	140 ± 85	88 ± 68
<i>BC-grad</i>	0		2.0 ± 0.9	8 ± 3	4.0 ± 1.8	28 ± 12	16 ± 7
	2	BC-free	15.2 ± 1.8	25 ± 2	30.5 ± 3.5	183 ± 21	63 ± 8
		BC-dark 500	2.2 ± 2.2	3 ± 3	4.4 ± 4.4	24 ± 24	6 ± 6
		BC-light 500	6.2 ± 0.6	9 ± 1	12.3 ± 1.2	71 ± 7	19 ± 2
		BC-light 2000	1.3 ± 1.3	2 ± 2	2.7 ± 2.7	14 ± 14	3 ± 3

cantly lower in BC-dark ($7.2 \pm 1.8 \times 10^4$ bacteria ml⁻¹ d⁻¹) than in the other treatments ($26.0 \pm 15.1 \times 10^4$ bacteria ml⁻¹ d⁻¹, Table 4, Kruskal-Wallis test, p < 0.05). Virus-mediated mortality of bacterial standing stock per day (VMM_{SS}) varied between 12% d⁻¹ at t₀ and >100% d⁻¹ in the light-exposed BC treatment at t_{2d} and was significantly lower in the dark BC treatment than in the light-exposed BC treatment and the BC control (Table 4, Kruskal-Wallis test, p < 0.05). The estimated virally mediated loss of BHP ranged from 33% at t₀ to >100% in the light-exposed BC treatment and BC-free control without significant differences between treatments. Following the model by Binder (1999), VMM ranged from 7% in the dark-exposed BC treatment to >100% in the light-exposed BC treatment without differences between treatments.

In *BC-grad*, lytic VP increased from $2.0 \pm 0.9 \times 10^6$ at t₀ to $15.2 \pm 1.8 \times 10^6$ viruses ml⁻¹ d⁻¹ at t_{2d} in the BC-free control and was significantly higher than in the BC-amended treatments ($3.2 \pm 2.8 \times 10^6$ viruses ml⁻¹ d⁻¹, Table 4, Fig. 6, Kruskal-Wallis test, p < 0.05). At t₀, 8 ± 3% of bacteria were infected. At t_{2d}, FIC was significantly higher in the BC-free control than in the BC-amended treatments (Table 4, Kruskal-Wallis test, p < 0.05). Consequently, estimated viral lysis rates were, on average, lower with BC addition ($6.5 \pm 5.7 \times 10^7$ bacteria ml⁻¹ d⁻¹) than in the BC-free control (Table 4, Kruskal-Wallis test, p < 0.05). The resulting VMM_{SS} values were significantly lower in the presence of BC (on average $36 \pm 33\%$ d⁻¹) compared to the control (>100% d⁻¹, Table 4, Kruskal-Wallis test, p < 0.05). According to the model by Binder (1999) VMM was significantly lower with BC addition (9 ± 9%) than in the BC-free control (63 ± 8% Table 4, Kruskal-Wallis test, p < 0.05). No lysogens could be induced by mitomycin C in any of the experiments.

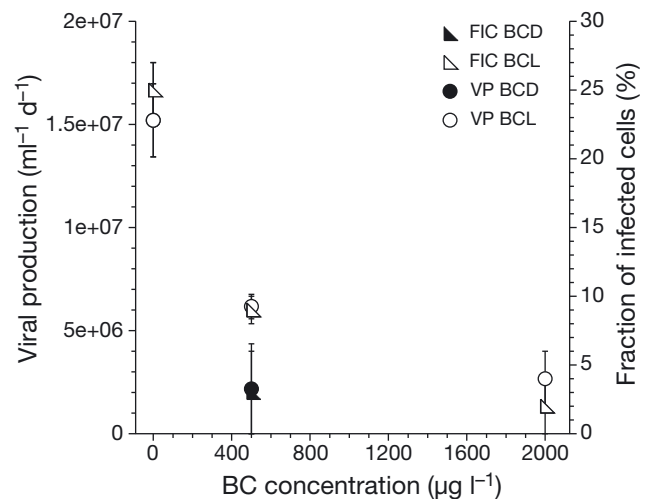


Fig. 6. Viral production (VP, viruses ml⁻¹ d⁻¹) and the fraction of infected cells (FIC, %) in experiment *BC-grad* from Day 2 as a function of black carbon (BC) concentration. Values are averages of duplicates from the virus reduction approach to measure VP and FIC, and error bars are the ranges. BCL: light-exposed BC; BCD: non-exposed BC

DISCUSSION

The effects of BC on bacterial abundance, production, and viral-mediated loss of bacterial production were assessed in this study using BC reference material and aged marine microbial communities. The main findings were that (1) BC addition increased bacterial biomass accumulation and reduced VP and infection and (2) exposure of BC to UV light increased bacterial production. These results suggest a stimulating effect of BC on aggregation processes and bacterial production and a suppressing effect on the viral shunt and associated nutrient regeneration with the potential to increase system heterotrophy.

Interaction between microorganisms and BC particles

With the help of flow cytometry, we detected an additional bacterial population characterized by high SSC (Fig. 1). Since this population with high SSC but comparable SYBR green derived fluorescence to HNA bacteria was not observed in the BC-free controls, it is likely that it consisted of bacteria associated with particles. The strong correlation between this bacterial FC population and BC concentration (Fig. 4A) supports the notion that these were BC-associated bacteria. Also, the similarity of the data on the fraction of particle-associated bacteria data obtained by FC and by epifluorescence microscopy supports this idea. The values of the side scatter in the FC cytograms for this type of particle increased during the course of the experiment (data not shown). Assuming that light scatter is proportional to the size of the particles (Koch et al. 1996), the data indicate that particles were increasing in size. Hence, a progressive aggregation of BC including colloids and bacterial cells into larger particles might have taken place. Different stages of association between bacteria and BC material, which range from small BC colloids attached to bacteria to bacteria completely entrapped in BC have been shown previously (Cattaneo et al. 2010), suggesting BC-driven aggregation and particle formation. This is supported by findings that BC reference material stimulated the formation of a specific class of organic aggregates, the transparent exopolymeric particles (Weinbauer et al. 2012, Mari et al. 2014). Also, *in situ* evidence suggests that soot deposition can increase coagulation and aggregation of organic matter (Mari et al. 2014).

Viral abundance per unit surface of BC decreased with BC particle size similarly to patterns previously observed in BC reference material (Cattaneo et al. 2010) and in other suspended material (Simon et al. 2002, Luef et al. 2007, Mari et al. 2007). The decreasing concentration of attached viruses with increasing particle size can be assigned to the fractal structure of BC particles (Slowik et al. 2007) which translates into increasing porosity with increasing particle size. Increasing porosity implies a higher content of pore water with potentially lower microbial and viral densities.

Effect of BC on viral production and infection

Free viral abundance was significantly lower in the BC treatments than in the controls (Table 2) and de-

creased with BC concentrations (Fig. 4B). Viruses were found attached to BC particles as in experiments performed with water from the Mediterranean Sea (Cattaneo et al. 2010). VP and the fraction of infected cells were also significantly lower in the BC-amended treatments (Table 4, Fig. 6) except for the BC-light treatment in *BC-fix*. In this treatment, the negative effects of BC on VP as experienced in the dark BC treatment (see discussion below) could have been compensated by an increased viral infection due to the stimulated bacterial production in BC-light (see below) with the net outcome of no detectable effect of BC at a concentration of 200 $\mu\text{g l}^{-1}$.

Our data suggest that adsorption onto BC particles inactivated viruses and prevented infection of attached cells, or reduced infection by reducing the abundance of free viruses and thus encounter rates. Similarly, in a study with water from the Danube River, free viruses were scavenged on particles (Kernegger et al. 2009). Such a mechanism could have resulted in the increased BHP in the BC treatments as also observed for suspended material in the Danube River. Interestingly, particle quality was a determining factor for microbial attachment in these freshwater experiments, and significantly fewer viruses attached to mineral than to organic particles (Kernegger et al. 2009). In accordance, the addition of mineral particles, such as Saharan dust (Weinbauer et al. 2009, Pulido-Villena et al. 2014) or clay particles (Salter et al. 2011) to marine coastal waters led to a negligible sorption of viruses to particles and increased VP rates in contrast to the response of the microbial community to BC addition in the present study.

VP in solar radiation-exposed BC treatments was significantly stimulated with respect to the non-exposed BC (Table 4). This could be traced to the stimulation of bacterial production in BC-light treatments, since viral infection and production are related to bacterial activity and production (Wommack & Colwell 2000). Also, viral adsorption or inactivation by BC could have been reduced due to UV-exposure, thus allowing for higher infection rates.

Effect of BC on bacterial production

The addition of BC stimulated bacterial production in both experiments (Table 2). The positive correlation found between particle-associated bacteria and heterotrophic bacterial production as well as HNA bacteria, which are often the more active bacterial fraction (e.g. Gasol et al. 1999), suggests that

attached bacteria were particularly active and contributed significantly to the measured bacterial production. Such an increased metabolic activity of attached bacteria has been found following the up-regulation of enzyme activity upon colonization of organic aggregates (Riemann et al. 2000, Simon et al. 2002, Grossart et al. 2007). Historically, BC was assumed to be highly refractory (Schmidt & Noack 2000), but an increasing body of literature shows abiotic (Decesari et al. 2002, Lehmann et al. 2005, Cheng et al. 2006, Stubbins et al. 2010, 2012) and biotic oxidation of BC in soils (Potter 1908, Shneour 1966, Cheng et al. 2006, 2008, Zimmerman 2010) and sediments (Middelburg et al. 1999). For example, oxidation of turbidite in the Madeira Abyssal Plain removed about 77% and 64% of organic and soot carbon, respectively, thus challenging the assumption that BC is biologically and chemically recalcitrant (Middelburg et al. 1999). However, since the quantification of BC was beyond the scope of this study, there is no direct evidence of BC decomposition or its use as a carbon source during our experiments. The addition of ammonium along with the BC material (Table 1) could have stimulated bacterial production, since the simultaneous addition of ammonium and organic carbon can stimulate bacterial production (Kirchman & Rich 1997). However, the use of aged seawater and the finding that ammonium concentrations did not decrease with incubation time argues against such a mechanism. Another mechanism explaining increased bacterial activity could be the adsorption of DOC to BC particles, which could create hot spots of microbial activity similar to those observed in marine snow particles (Azam & Malfatti 2007). It is well known that soot particles can interact with organic compounds (Rockne et al. 2000, Slowik et al. 2007) and redistribute them in the environment (Ahrens & Morrisey 2005). There is additional evidence (Cattaneo et al. 2010, this study) that viruses, which are by definition part of the DOC pool, are rapidly adsorbed to BC. Overall, it is well known that organic compounds adsorb very efficiently to materials such as BC, generally exceeding adsorption for typical amorphous organic matter by a factor of 10 to 100 (Cornelissen & Gustafsson 2005, Cornelissen et al. 2005).

In both experiments, heterotrophic bacterial production was significantly higher after exposing BC to solar radiation. Photo-oxidation can induce chemical changes in the highly aromatic structure of BC-like molecules (Stubbins et al. 2010), thus offering sites for degradation and increasing BC bioavailability. Stubbins et al. (2012) reported a 20-fold decrease in marine dissolved BC concentration after exposure

of North Atlantic Deep Water to simulated solar radiation confirming the high photo-lability of dissolved BC. Maki et al. (2001) found that the exposure of biodegraded crude oil to sunlight significantly decreased the aromatic fraction and that this material favored the growth of seawater microbes as compared to non-irradiated material (Maki et al. 2001). Our data suggest that light exposure of BC in surface waters can alter BHP. As aerosols are normally exposed to solar radiation during atmospheric transport with radiation levels potentially exceeding those observed in surface waters, our results also suggest that the distance from the BC source and its residence time in the atmosphere prior to the deposition on the ocean surface could be an important parameter modulating the effect of BC on the marine ecosystem.

CONCLUSIONS

Our study indicates that BC deposition in the ocean can stimulate heterotrophic bacterial production either by reducing the impact of viral lysis, by serving as a carbon source to bacteria, and/or by serving as hotspots due to organic matter adsorption. Our experiments also indicate that atmospheric transport of BC, and the residence time in surface waters, will likely influence the effect on the microbial community, since exposure to solar radiation increased BC availability. Fossil-fuel and biomass burning will probably further increase in the future (Novakov et al. 2003), and the impact of BC on the functioning of the marine microbial food web is likely to become more important, particularly in coastal oceans. One of the potential consequences could be a shift in the metabolic balance of the planktonic ecosystem towards a more heterotrophic ocean.

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