Particle-Associated Differ from Free-Living Bacteria in Surface Waters of the Baltic Sea

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Many studies on bacterial community composition (BCC) do not distinguish between particle-associated (PA) and free-living (FL) bacteria or neglect the PA fraction by pre-filtration removing most particles. Although temporal and spatial gradients in environmental variables are known to shape BCC, it remains unclear how and to what extent PA and FL bacterial diversity responds to such environmental changes. To elucidate the BCC of both bacterial fractions related to different environmental settings, we studied surface samples of three Baltic Sea stations (marine, mesohaline, and oligohaline) in two different seasons (summer and fall/winter). Amplicon sequencing of the 16S rRNA gene revealed significant differences in BCC of both bacterial fractions among stations and seasons, with a particularly high number of PA operational taxonomic units (OTUs at genus-level) at the marine station in both seasons. "Shannon and Simpson indices" showed a higher diversity of PA than FL bacteria at the marine station in both seasons and at the oligohaline station in fall/winter. In general, a high fraction of bacterial OTUs was found exclusively in the PA fraction (52% of total OTUs). These findings indicate that PA bacteria significantly contribute to overall bacterial richness and that they differ from FL bacteria. Therefore, to gain a deeper understanding on diversity and dynamics of aquatic bacteria, PA and FL bacteria should be generally studied independently.

Keywords: microbial communities, microbial diversity, particle-associated and free-living bacteria, Baltic Sea, salinity gradient, seasons, 454-pyrosequencing

INTRODUCTION

Nutrient availability, pH, temperature, and salinity have been shown to be major environmental drivers for community structure and activities of aquatic organisms. Temporal and spatial gradients in environmental factors shape community composition and metabolic activities of aquatic organisms, and hence biogeochemistry of aquatic ecosystems. Although macroscopic organic aggregates (>100 µm) play a crucial role in the ocean's carbon cycle (Fowler and Knauer, 1986; Alldredge and Silver, 1988; Simon et al., 2002), and particle-associated (PA) bacteria colonize nearly all types of particulate organic matter (POM), studies in microbial ecology traditionally focus on total or exclusively free-living (FL) bacteria (Grossart et al., 2010). This is in contrast to findings indicating that macroscopic organic aggregates produced by aggregation of various organic material, i.e., marine snow (Simon et al., 2002) or lake snow (Grossart and Simon, 1993) are densely colonized by bacteria. They represent nutrient rich “hotspots” of microbial activity even in oligotrophic environments (e.g., Simon et al., 2002; Grossart, 2010).
Differences in hydrodynamic conditions and primary production, however, may lead to changes in organic matter quantity and quality resulting in different aggregation dynamics and consequently of microbial particle colonization. Ortega-Retuerta et al. (2013) suggested that bacterial community composition (BCC) of the PA fraction in the ocean is rather determined by particle quality than by their quantity, since differences between PA and FL bacteria are not always correlated to POM concentrations, whereas the ratio between PA and FL bacteria relates well to suspended particulate matter (SPM) quality (Doxaran et al., 2012). Such differences in BCC and dynamics of both bacterial fractions indicate that PA and FL bacteria should be regarded as independent components of a bacterial assemblage to better understand the bacterial response to a changing environment. Yet, many studies in aquatic microbial ecology do not distinguish between PA and FL bacteria or remove most PA bacteria with particles from the sample when using pre-filtration steps.

The Baltic Sea is an ideal model system to address general differences in BCC and dynamics of both bacterial fractions at different salinities. It is one of the largest brackish (mesohaline) ecosystems in the world where salinity declines along a ca. 2000 km long horizontal gradient from salinities of around 32 in the Kattegat to 2 in the northern Bothnian Bay. The narrow and shallow Danish straits limits water exchange with the North Sea resulting in a water residence time of the surface waters of ca. 25–30 years (Kautsky and Kautsky, 2000) and relatively stable horizontal and vertical salinity gradients. Salinity has been shown to shape the distribution pattern of macroorganisms, with higher organisms being either adapted to freshwater or marine systems resulting in a relatively lower diversity in the freshwater-marine transition zone (Remane, 1934; Telesh and Khlébovich, 2010). Protozoa, however, did not show a clear salinity-dependent distribution pattern (Telesh and Khlébovich, 2010) and bacterial communities in the Baltic Sea revealed only little differences in α-diversity along the salinity gradient (Herlemann et al., 2011). In contrast, BCC greatly varies along the horizontal and vertical salinity gradients in the Baltic Sea (Edwards and John Pujeta, 1997; Holmfeldt et al., 2009; Herlemann et al., 2011). None of these studies, however, distinguished between PA and FL bacteria communities, and at present we do not know whether the diversity of PA bacteria is shaped by the salinity gradient and to what extent they differ from FL communities.

The majority of studies differentiating between PA and FL bacteria has been either conducted in marine or in freshwater habitats (Simon et al., 2002; Ghiglione et al., 2007, 2009; Fontanez et al., 2015). Only a few studies compared BCC of both bacterial fractions at different salinities in marine, brackish and freshwater ecosystems (e.g., Crump et al., 1999, 2004; Ortega-Retuerta et al., 2013; Bižić-Ionescu et al., 2014; Simon et al., 2014). Nevertheless, all of these studies found general differences in BCC between habitats and bacterial fractions. Whereas some studies indicated a high specialization of bacteria on particles (Crump et al., 1999; Rösel and Grossart, 2012) others point to a high exchange between FL and PA bacteria fractions (Hollibaugh et al., 2000; Ghiglione et al., 2007). Since enzymatic capabilities greatly differ among various bacterial phyla and life-styles (Martinez et al., 1996; Lyons and Dobbs, 2012), knowledge about BCC of both PA and FL bacteria fractions provides hints on metabolic differences. For example, Ganesh et al. (2015) focusing on dissimilatory processes of the nitrogen cycle in the oxygen minimum zone of the Eastern Tropical North Pacific revealed 8- to 15-fold higher bacterial gene counts in the FL fraction, which was mainly related to anammox-associated transcripts.

In particular, little is known about PA bacteria from the marine-freshwater transition zone. Interestingly, Crump et al. (1998) found different PA bacteria at the transition from the river to the estuary and coastal sites, whereas FL bacteria remained similar. Garneau et al. (2009) found the strongest differences in bacterial fractions at the estuarine station with the highest POM content. Moreover, Ortega-Retuerta et al. (2013) have analyzed various samples from a river, the coastal and open ocean and found significant differences in BCC of total bacteria between samples at different salinities. However, BCC of PA and FL bacteria differed only in the open ocean, although particle concentration was much higher in the riverine and coastal samples. The authors also found a generally higher diversity for PA bacteria (Shannon, Simpson, and Chao indices) than FL. We hypothesized that (1) PA bacteria significantly contribute to overall bacterial richness, and (2) that salinity has a strong impact on the composition of PA bacteria. During two seasons, we assessed PA and FL BCC at three stations in the Baltic Sea with different salinities using 454-pyrosequencing of partial 16S rRNA genes (spanning variable regions V3 and V4). In summer, we simultaneously measured the cell-specific bacterial protein production (csBPP) by 14C-leucine incorporation. Our results indicate that PA bacteria substantially contribute to overall bacterial α- and β-diversity at all stations in both sampling seasons (summer and fall/winter). BCC substantially differed among both bacterial fractions, whereby α- and β-diversity of PA and FL bacteria varied with season and salinity. The highest bacterial α-diversity occurred in the PA fraction of the marine station in fall/winter. Although PA bacteria contributed rather little to total bacterial abundance, csBPP rates of PA were significantly higher than FL bacteria. This suggests that PA bacteria represent an important component of overall bacterial α- and β-diversity and presumably activity in the Baltic Sea.

**MATERIALS AND METHODS**

**Site Description and Field Work**

The sampling was performed in the scope of the ATKiM (“Abbaumarkt von artischem, terrigenem Kohlenstoff im Meer”) project during two cruises (M86-1, M87-3) of the RV Meteor in the Baltic Sea: (i) November/December 2011 (“fall/winter”) and (ii) May/June 2012 (“summer”). Sampling locations (Figure 1) were characterized by differences in salinity and defined as marine (Skagerrak, salinity ≈ 30-34; N 58°07′/59.88′′, E010°0′/0′′), mesohaline (Gotland Basin, salinity ≈ 7, N57°18′/20.52′′; E020°04′/41.5′′) and oligohaline (Bothnian Bay, salinity ≈ 2.7; N65°26′/42′′, E023°17′/53.88′′). Surface water of the three different sampling stations was collected with a 400 dm³
stainless steel water sampler (Hydrobios, Kiel, Germany) from which triplicate samples were taken (representing “technical replicates”). The samples were pre-filtered through a 100 µm mesh filter to exclude large zooplankton and then directly transferred into 25L polyethylene canisters from which subsamples were taken at 10°C for further analyses. The bacteria collected on the 5-µm filter were considered to represent the PA fraction and the bacteria collected on the 0.2–µm filters the FL fraction.

**Chemical Parameters**

Dissolved inorganic phosphate, nitrate, nitrite, silicate, and ammonium were analyzed following standard methods (Rohde and Nehring, 1979; Grasshoff et al., 1983). All samples were filtered through pre-combusted GF/F filters (pore size 0.65 µm; Whatman, Dassel, Germany) and hydrolyzed at 220°C, except for ammonia analysis. Phosphate, nitrate, nitrite, and silicate were analyzed by a continuous-flow analyzer “FLOWSYS” (Alliance Instruments GmbH, Airing, Germany). Ammonium concentration was determined by a UV mini 1240 photometer (Shimadzu).

For particulate organic carbon (POC), five-hundred ml samples were filtered through pre-combusted (500°C, 3 h) GF/F filters (Whatman, Dassel, Germany). Filters were dried for 4 h at 50°C in prewashed petri dishes and stored in a desiccator until analysis. The samples were analyzed by using an organic elemental analyzer “Flash EA 1112” coupled to an isotope ratio mass spectrometer “Finnigan MAT V” (Thermo Fisher Scientific Inc., Waltham, USA) in the stable isotope laboratory of the “Museum für Naturkunde”, Berlin, Germany.

Dissolved organic carbon (DOC) content was determined by filtering 30 ml per sample through pre-combusted (500°C, 3 h) GF/F filters (Whatman, Dassel, Germany) into high-density polyethylene [HDPE] vials which were kept frozen until analysis. For analysis, the “fall/winter” samples were acidified with 2 M HCl until the pH was below 4 in order to remove all inorganic carbon. DOC samples were then measured in duplicates on a Multi N/C 3100 Analyzer (Jena Analytics, Germany) by applying non-dispersive infrared detection (NDIR) after combustion. The “summer” samples were acidified to pH 2 with hydrochloric acid and measured in duplicates per sample and in triplicates per station by an organic carbon analyzer “TOC-VPCH and TNM-1” (Shimadzu Deutschland GmbH, Duisburg, Germany). For both analyses, an internal standard of potassium hydrogen phthalate (5 mg L⁻¹) was used.

**Bacterial Enumeration**

For enumeration, bacterial samples were fixed with formaldehyde (2% v/v) and stored at 4°C. The following filtration was done within 4 weeks (Buesing, 2005). All samples (30 ml) were sequentially filtered through a 5.0 µm polycarbonate membrane (0 25 mm; Whatman, Dassel, Germany) for PA bacteria, and 5 ml of the filtrate were sieved through a 0.2 µm filter for FL bacteria. All filters were air-dried and stored at −20°C until microscopy analysis (Grossart et al., 2005). For counting FL bacteria, the bacteria were stained with a mixture of 1 µg ml⁻¹ DAPI (4′,6-diamidino-2-phenylindole) (AppliChem GmbH, Darmstadt, Germany) and the PA bacteria with 1:1000 SybrGold (LifeTechnologies, Carlsbad, USA). Both dyes were diluted in Citifluor AF1 mounting medium (Citifluor Ltd, London, UK) which was added directly on the filter before covering with the cover slide. All samples were analyzed with an epifluorescence microscope “Axio Imager.Z1” (Zeiss, Jena, Germany) at 630 x magnification with an automated cell counting approach by preparing pictures of 25–40 stacks at intervals of 0.3 µm (Zeder et al., 2011). The image analysis was performed with ImageJ (Abramoff et al., 2004) after Massana et al. (1997). Percentages of particle area were analyzed at 200x and 630x magnification which were used to relate PA bacteria counts to sample volume.

**Bacterial Protein Production (BPP) and Respiration**

BPP was measured for the summer samples by incorporation of L¹⁴C-leucine (Hartmann Analytic, Braunschweig, Germany) according to Simon and Azam (1989). For each station, six samples and two negative controls (2% v/v formaldehyde) were inoculated with a final concentration of 50 µCi to ensure the uptake of L¹⁴C-leucine of FL as well as PA bacteria. The incubation was performed in the dark at 10°C and stopped after 1 h by adding formaldehyde (2% v/v). After fixation, samples were filtered sequentially onto 5.0 and 0.2 µm cellulose nitrate filter (Whatman, Dassel, Germany) for separating PA and FL bacteria, respectively. Proteins were extracted for 5 min in 5% trichloroacetic acid (TCA). Thereafter, the filters were washed twice with ultrapure water and then with ethanol (50%), before they were transferred into 20 ml scintillation vials and dissolved with 500 µl of ethyl acetate (15 min). Before counting, 10 ml of the scintillation cocktail “Ultima Gold” (Perkin-Elmer, Downers Grove, USA) was added and incubated overnight. The counting was determined as disintegrations per minute (DPM) on a liquid scintillation analyzer “TriCarb 2810 TR” (PerkinElmer, Downers Grove, USA). The produced protein amount was converted by a
factor of 0.86 into carbon (Simon and Azam, 1989). Cell-specific BPP (csBPP) was calculated for both bacterial fractions based on their cell numbers.

Total bacterial respiration (BR) of the samples was measured by oxygen consumption of subsamples with a Sensor Dish Reader SDR 2 (PreSens, Regensburg, Germany) in air-tight glass vials using an optode system (Köster et al., 2012). For each sample, oxygen consumption rates were determined over 24 h twice per sample in the dark at 10°C. After a temperature correction by calculating the oxygen solubility at a given temperature using the Bunsen absorption coefficient α (T), the amount of consumed oxygen was converted into carbon by using 0.88 as a conversion factor (Fogg and Gerrard, 1990; Robinson et al., 1999). Bacterial growth efficiency (BGE) was calculated as BGE = BPP/ (BPP+BR).

Bacterial Community Composition

For DNA extraction, a total of 36 samples were analyzed. Each sample of 0.5 L was sequentially filtered over a 5.0 µm Nucleopore TrackEtch polycarbonate membrane (Whatman, Dassel, Germany) for the PA-fraction and the total 5.0 µm filtrate was sieved through a 0.22 µm Durapore membrane (Merck KGaA, Darmstadt, Germany) for the FL-fraction. Filters were shock-frozen in liquid nitrogen and stored at −80°C until use. DNA extraction was performed after Necesrian et al. (2005), using a Phenol-Chloroform protocol with cetyltrimethyl ammonium bromide (CTAB) as a complexing agent for polymeric substances.

All 36 samples were PCR amplified with 30 cycles using a 20 ng aliquot of each DNA sample for a 25 µl PCR reaction, the primer pair Bakt_341F (CCTACGGGNGGCWGCAG) and Bakt_805R (ACHVGGGTATCTAATCC) and the Fusion DNA Polymerase Hercule II (Agilent Technologies, Santa Clara, USA) according to Herlemann et al. (2011). Primers were tagged with sample-specific 5pb barcodes a 454-adptor region, spanning variable regions V3 and V4 of the 16S rRNA gene. PCR products were purified with Agencourt AMPure XP magnetic beads (Beckman Coulter GmbH, Krefeld, Germany), quantified with a Picogreen assay (LifeTechnologies, Carlsbad, USA), diluted and equally pooled. The bidirectional sequencing was performed on a 454 platform with Titanium Flex chemistry (Roche etc.). Our samples represent subsamples, from a run with 61 samples in total, which were sequenced at the Berlin Center for Genomics in Biodiversity Research (BeGenDiv, Berlin) with an average of 115,000 reads per run and accordingly on average 1885 reads per sample. After a brief quality check with Mothur (Schloss et al., 2009) (maximum number of N: 0, minimum sequence length: 150, and minimum exponential Q-score: 20) SILVAngs data analysis service (Yilmaz et al., 2013) was used to align the resulting sequences with the SILVA Incremental Aligner (SINA) and to remove contaminations of the dataset with non-rDNA sequences. SILVAngs performs an additional quality check by a minimal length cut-off (50 bases) as well as ambiguity and homopolymer check (max. 2%). After the quality control, identical reads were identified (dereplication), unique reads were clustered (OTUs), on a per sample basis, and the reference read of each OTU was classified. Dereplication and clustering was done using cd-hit-est version 3.1.2 (Li and Godzik, 2006). The clustering was performed with a minimum of 97% sequence identity to each other (pairwise distance and single linkage clustering). For each OTU clustering, the longest read was then used as a reference of this cluster for taxonomic classification. BLAST (version 2.2.28+) in combination with the SILVA SSURef dataset (release 119) was used to classify the sequences. The resulting classification of the reference sequence of a cluster was mapped to all members of the respective cluster as well as their replicates. Sequences having an average BLAST alignment coverage and alignment identity of less than 93% would be considered as unclassified. This method was first used by Klindworth et al. (2013) and Ionescu et al. (2012). PAST was used to generate rarefaction curves with an algorithm from Krebs (1989). To normalize the number of sequence reads between samples, a random subsampling according to the smallest n (430 sequences) was performed in R with the vegan package (Oksanen et al., 2011) and the rarefy function. For taxonomic analysis the resulting reads were normalized based on OTUs representing the genus-level diversity (97% similarity). All analyses of α and β-diversity were performed with the rarefied dataset.

Sequences

All sequences are deposited in the European Nucleotide Archive under the study accession number: PRIEB9483 and the sample accession numbers: ERS742024 to ERS742056.

Statistical Analysis

All statistics was performed using the software R (R Core Team, 2013) and PAST (Hammer et al., 2012).

For α-diversity calculation of species richness CHAO1 index, Buzas and Gibson’s Evenness [eH/H], Simpson Evenness [1-D] as well as Shannon [H] diversity indices were calculated with PAST. For β-diversity (BCC) Bray-Curtis dissimilarity matrixes were generated, visualized by non-metric multidimensional scaling (NMDS), unweighted-pair group method with arithmetic averages (UPMGA), and statistically verified by analysis of similarity (ANOSIM) with the R package vegan and the functions metaMDS, hclust (average) and anosim. To assess what taxa are primarily responsible for the observed differences between the sample groups a similarity percentage algorithm (SIMPER) was used with the function simper. For estimation of significant differences between PA and FL bacterial activities and richness a non-parametric Wilcoxon rank sum test for independent samples was performed in R.

RESULTS

Site Characteristics

All three sampling stations (Table 1) were characterized by similar water temperatures between the “summer” and “fall/winter” season, mainly because of exceptionally high temperatures in late autumn 2011. The oligohaline station is an exception, with an “unproductive” phase at our summer sampling, due to its late ice-coverage (until the end of spring).
The oligohaline station was characterized by the lowest salinity (~2.7) and temperature (4.33 and 6.26°C in fall/winter and summer, respectively), but by the highest DOC (>5 mg L\(^{-1}\)), silicate (39 and 43 µM in fall/winter and summer, respectively), and oxygen concentrations (8.18 and 9.42 mg L\(^{-1}\) in fall/winter and summer, respectively) (Table 1). POC concentrations did not differ much between summer (0.13 mg L\(^{-1}\)) and fall/winter (0.19 mg L\(^{-1}\)). In summer, POC at the oligohaline station was lower than at the mesohaline station. Chla concentration, however, increased from 0.44 µg L\(^{-1}\) in fall/winter to 1.76 µg L\(^{-1}\) in summer - similar to the mesohaline station. Dissolved nitrogen concentration was highest at the oligohaline station (4.0 µM in fall/winter and 5.0 µM in summer).

**Bacterial Numbers and Activity**

At both sampling time points, the total number of bacterial cells (Table 1) was lowest at the marine station, and no trend in abundance was obvious between both seasons. Cell number was lower in summer at the mesohaline station, particularly for FL cells. The opposite trend occurred at the oligohaline station. Numbers of PA bacteria at the marine and mesohaline stations were higher in summer whereas those of the oligohaline station remained at the same high level in both seasons. Marine and

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**TABLE 1 | Physical, chemical and biological parameters at different stations and seasons in the Baltic Sea.**

| Sample name abbreviations: T, temperature; O\(_2\), oxygen content; Chla, Chlorophyll a; DOC, dissolved organic carbon; POC, particulate organic carbon; NO\(_3\), nitrate concentration; NO\(_2\), nitrite concentration; PO\(_4\), phosphate concentration; SiO\(_4\), silicate concentration; NH\(_4\), ammonium concentration; PA, particle-associated; FL, free-living; Tot, total; bac, bacteria; BPP, bacterial protein production; csBP, cell specific bacterial production; BGE, bacterial growth efficiency. |
| :-----------------|-----------------|-----------------|-----------------|
| Marine | Mesohaline | Oligohaline | Marine | Mesohaline | Oligohaline |
| Salinity (°C) | 30.40 | 7.15 | 2.65 | 34.62 | 6.97 | 2.84 |
| T [°C] | 12.29 | 9.07 | 4.33 | 12.15 | 10.54 | 6.26 |
| O\(_2\) [µM] | 6.64 | 8.51 | 9.42 | 5.93 | 7.24 | 8.18 |
| Chla [µg L\(^{-1}\)] | 0.13 | 1.63 | 1.76 | 0.30 | 0.47 | 0.44 |
| DOC [mg L\(^{-1}\)] | 1.97 (± 0.12) | 4.48 (± 0.57) | 5.55 (± 0.30) | 1.20 (± 0.11) | 3.63 (± 0.10) | 5.20 (± 0.12) |
| POC [mg L\(^{-1}\)] | 0.09 | 0.39 | 0.13 | 0.08 | 0.13 | 0.19 |
| NO\(_3\) [µM] | 0.17 | 0.20 | 0.70 | 0.06 | 0.26 | 0.25 |
| NO\(_2\) [µM] | 0.00 | 0.00 | 5.04 | 0.36 | 0.14 | 4.09 |
| PO\(_4\) [µM] | −0.02 | 0.09 | 0.00 | 0.29 | 0.16 | 0.02 |
| SiO\(_4\) [µM] | 0.05 | 9.64 | 43.14 | 0.70 | 4.95 | 39.00 |
| NH\(_4\) [µM] | 0.44 | 0.11 | 0.22 | 0.48 | 0.81 | 0.64 |
| cells \(_{PA}\) [10\(^5\) cells ml\(^{-1}\)] | 0.26 (± 0.06) | 0.57 (± 0.13) | 0.76 (± 0.21) | 0.12 (± 0.069) | 0.13 (± 0.08) | 0.74 (± 0.03) |
| cells \(_{FL}\) [10\(^5\) cells ml\(^{-1}\)] | 5.16 (± 0.20) | 7.37 (± 1.06) | 16.55 (± 0.27) | 5.19 (± 0.17) | 11.30 (± 0.28) | 11.70 (± 0.08) |
| cells \(_{Tot}\) [10\(^5\) cells ml\(^{-1}\)] | 5.42 (± 0.24) | 7.94 (± 1.18) | 17.31 (± 0.29) | 5.31 (± 0.23) | 11.40 (± 0.35) | 12.50 (± 0.11) |
| PA, bac [%] | 4.85 | 7.17 | 4.38 | 3.28 | 0.93 | 5.84 |
| BPP, PA [µg C L\(^{-1}\) h\(^{-1}\)] | 0.18 (± 0.02) | 0.33 (± 0.07) | 0.17 (± 0.07) | NA | NA | NA |
| BPP, FL [µg C L\(^{-1}\) h\(^{-1}\)] | 0.12 (± 0.09) | 0.32 (± 0.06) | 0.25 (± 0.09) | NA | NA | NA |
| csBPP, PA [fg C cell\(^{-1}\) h\(^{-1}\)] | 0.30 (± 0.07) | 0.65 (± 0.05) | 0.34 (± 0.15) | NA | NA | NA |
| csBPP, FL [fg C cell\(^{-1}\) h\(^{-1}\)] | 0.23 | 0.43 | 0.15 | NA | NA | NA |
| csBPP, Tot [fg C cell\(^{-1}\) h\(^{-1}\)] | 0.55 | 0.81 | 0.24 | NA | NA | NA |
| Respiration [µg C L\(^{-1}\) h\(^{-1}\)] | 1.97 | 1.49 | 1.09 | NA | NA | NA |
| BGE [µB h\(^{-1}\)] | 0.11 | 0.30 | 0.28 | NA | NA | NA |

Beside its high salinity (>30), the marine station showed the highest temperature (~12°C) as well as the lowest DOC (~1–2 mg L\(^{-1}\)), POC (~0.08 mg L\(^{-1}\)), oxygen (~6–7 mg L\(^{-1}\)) and SiO\(_4\) concentrations (3.7 µM and 0.05 µM in fall/winter and summer, respectively) (Table 1). Nitrate content was higher in fall/winter than in summer (0.36 µM and below detection limit, respectively). The same pattern occurred for dissolved phosphate (0.29 µM vs. below the detection limit). Consequently, Chla concentration was highest in fall/winter than in summer (0.3 vs. 0.13 µg L\(^{-1}\)), whereby fall/winter values were in the same range as for the other sampling stations.

The mesohaline station showed large differences in POC and Chla concentrations between both seasons and was characterized by a salinity of around 7 (Table 1). DOC and SiO\(_4\) concentrations were close to the average of what can be expected in the Baltic Sea (DOC: 3.6 and 4.48 mg L\(^{-1}\) as well as 4.95 and 9.64 mg L\(^{-1}\) SiO\(_4\) in fall/winter and summer, respectively). Oxygen varied in both seasons between 7.24 and 8.51 mg L\(^{-1}\). In summer, POC and Chla values (0.39 mg L\(^{-1}\) and 1.63 µg L\(^{-1}\)) were clearly higher than in the fall/winter season (0.13 mg L\(^{-1}\) and 0.47 µg L\(^{-1}\)). In contrast, NO\(_3\), NH\(_4\), and PO\(_4\), which were either close to or below the detection limit in summer (0.00, 0.11, and 0.09 µM), were much higher in fall/winter (0.14, 0.81, and 0.16 µM).

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oligohaline stations showed an average proportion of PA bacteria between 3 and 6% independent of season. The mesohaline station, however, showed a remarkably higher proportion of PA bacteria in summer (~1% in fall/winter vs. 7% in summer) in accordance with the higher POC concentration. In contrast to the relatively low abundance of PA bacteria, bacterial protein production (BPP) (only measured in summer) was almost in the same range as for FL bacteria. Cell specific bacterial protein production (csBPP) of all PA bacteria was significantly higher than FL bacteria (~17 times) (Wilcoxon rank sum test: \( W = 36, p = 0.005 \)). For all bacteria fractions, highest csBPP occurred at the marine station, followed by the mesohaline and oligohaline stations. In parallel to the rather low total BPP, community respiration was also lowest at the oligohaline station and calculated bacterial growth efficiencies of total bacteria varied between 11 and 30% (Table 1).

**Bacterial Diversity**

The \( \alpha \)-diversity measures the OTU diversity of individual samples. The OTU distribution (Figure 2) showed a higher overall bacterial diversity in fall/winter than in summer. Out of the total 348 bacterial OTUs in summer, 184 OTUs were found at the marine, 158 OTUs at the mesohaline and 213 OTUs at the oligohaline stations. In summer, the oligohaline station with the lowest water temperature had the highest total bacterial \( \alpha \)-diversity and the mesohaline station the lowest. In fall/winter, the total number of OTUs was higher (547), with 340 OTUs at the marine, 251 OTUs at the mesohaline, and 281 OTUs at the oligohaline station. The majority of OTUs at the mesohaline station was shared with those found at the other stations. Only a small proportion exclusively occurred at the mesohaline station, with a higher proportion in fall/winter. At the mesohaline station, there was a high number of shared OTUs with either the marine or oligohaline stations. Interestingly, when comparing all bacterial fractions, high numbers of both PA and FL OTUs were related to those OTUs, which occurred at all three stations. A high number of exclusively PA OTUs occurred in almost all samples (52% of total OTUs and 59% of the OTUs in winter), with a particularly high number of exclusively PA OTUs at the marine station in both seasons (71% in winter and 53% in summer). A high number of both PA and FL OTUs were shared between oligohaline and mesohaline stations and surprisingly also between the marine and oligohaline stations. Comparison of \( \alpha \)-diversity indices (Figure 3) showed that richness was significantly higher in the PA fraction (Wilcoxon rank sum test: \( W = 268, p = 0.0008 \)), but a pairwise comparison of each station showed no significant difference for the mesohaline station in both seasons and the oligohaline station in summer (Supplemental Table S1). The “Simpson index” \( [1-D] \) indicated that PA and FL fractions were in general evenly distributed at the marine station, but not at the meso- and oligohaline stations, where single taxa dominated the system. “Shannon and Simpson indices” showed higher diversity of PA bacteria at the marine station in both seasons and at the oligohaline station in fall/winter. Evenness was higher in the FL fraction at the mesohaline station in both seasons and the oligohaline station in summer. Generally, samples with a low evenness showed also a low diversity in the Shannon and Simpson indices (raw data shown in Supplemental Table S2).

The \( \beta \)-diversity indices measure the similarity (or dissimilarity) in microbiome composition between samples. At all stations, bacterial communities in Baltic surface waters revealed a higher OTU richness in fall/winter than in summer (Figure 3). The dominant bacterial communities (\( \geq 1% \) of total bacteria) (Figure 4) were \( \alpha \)-, \( \beta \)-, \( \gamma \)-, and \( \delta \)-proteobacteria as well as Bacteroidetes. Also Flavobacteria, Actinobacteria, Planctomycetes, Cyanobacteria, and Verrucomicrobia occurred at all stations. Deferrribacteres, Firmicutes, and Spirochaetes were present only at the marine station, whereas Chloroflexi were found at the marine and oligohaline station. Chlorobi were detected at the meso- and oligohaline stations. Acidobacteria, Chlamydiae, and Candidate division OD1 were exclusively found at the oligohaline station in fall/winter. Richness was lower in summer than in fall/winter since Firmicutes, Deferrribacteres, Chloroflexi, Spirochaetes, and Chlorobi did not occur and \( \delta \)-proteobacteria and Verrucomicrobia were less abundant at the mesohaline station. Only the oligohaline station revealed no considerable difference between winter and summer.

PA bacteria at the marine station were dominated by Bacteroidetes, mainly Flavobacteria, followed by \( \alpha \)-, \( \beta \)-, \( \gamma \)-, and \( \delta \)-proteobacteria, and Cyanobacteria in summer (Figure 4). In fall/winter, however, bacterial phyla were more evenly distributed among PA bacteria and PA and FL fractions were more similar. FL bacteria were greatly dominated by Alphaproteobacteria, followed by Bacteroidetes and Gammaproteobacteria. Bacterioprotein bacteria and Cyanobacteria contributed little to FL bacteria. Mesohaline PA bacteria were greatly dominated by Cyanobacteria, but otherwise they were relatively similar to FL bacteria with Bacteroidetes, Alpha- and Gammaproteobacteria being the major bacterial groups. In fall/winter, the meso- and oligohaline samples comprised a relatively high number of Actinobacteria in the FL fraction, whereas the PA bacteria showed higher numbers of Gammaproteobacteria and Planctomycetes compared to the FL fraction.

PA bacteria at the oligohaline station were dominated by Planctomycetes in summer (Figure 4), followed by Bacteroidetes and Betaproteobacteria. In contrast, the FL fraction was characterized by a dominance of Alpha- and Betaproteobacteria as well as Actinobacteria. In fall/winter, the oligohaline FL BCC was characterized by a high abundance of Actinobacteria, and a higher amount of Planctomycetes, Gammaproteobacteria, and Cyanobacteria compared to the PA fraction. Except for the high amount of Alphaproteobacteria at the oligohaline station in summer, the relative abundance of Alphaproteobacteria and Gammaproteobacteria increased with salinity, whereas Actinobacteria and Betaproteobacteria decreased. For PA bacteria there was a trend of increasing dominance of Bacteroidetes from higher to lower salinity levels, whereas Planctomycetes showed the opposite trend.

The total BCC revealed significant differences between marine, mesohaline and oligohaline stations for both studied seasons (Table 2) even though this shift was less clear in
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FIGURE 2 | Venn diagram of OTU distribution of the normalized data between marine (mar), mesohaline (meso) and oligohaline (oligo) samples in summer (A) and winter (B) (areas are proportional to OTU number). Relative abundances of all exclusive and shared OTUs in either particle-associated fraction, free-living fraction or both fractions are shown.

FIGURE 3 | Boxplots of different $\alpha$-diversity indices (Chao1, Eveness, Shannon, Simpson) of a randomized dataset (430 reads) to compare the diversity between particle-associated (PA) and free-living (FL) bacteria in summer and winter. Samples considered as mar: marine, meso: mesohaline and oligo: oligohaline.
In summer, BCC of the mesohaline station was more similar to the marine station, whereas in fall/winter it was more similar to the oligohaline station (Figures 5A,B). BCC varied with stations, fractions, and sampling time, except BCC of the oligohaline station did not significantly differ between both seasons (Table 2). Supplemental Figure S1C shows that this pattern was mainly explained by FL bacteria being highly similar in summer and fall/winter. Changes in BCC with stations revealed a stronger significance in summer \( R = 1.00, \ p = 0.002 \) than in fall/winter \( R = 0.78, \ p = 0.003 \), and differences in BCC between stations were lower when comparing the neighboring stations (mesohaline vs. marine or mesohaline vs. oligohaline) (Table 2). Differences in BCC were mainly explained by the higher overall bacterial \( \alpha \)-diversity in fall/winter compared to the summer samples and changes in the dominant phyla (Figure 3). Differences between PA and FL BCC were more significant in fall/winter than in summer (Table 2). Thereby, the mesohaline station showed, together with the oligohaline station, the lowest differences between FL and PA fractions in summer (Supplemental Figures S1A–D). At all stations there were differences in PA vs. FL BCC in summer as well as in fall/winter (Figures 5C,D). The high BCC similarity of meso- and oligohaline samples in fall/winter is based on both fractions (PA and FL). Generally, there was a significant difference between PA and FL BCC in summer as well as in fall/winter (Figures 5C,D). The high BCC similarity of meso- and oligohaline samples in fall/winter is based on both fractions (PA and FL). Generally, there was a significant difference between PA and FL BCC within stations or seasons (Table 2, Supplemental Figure S2). Comparing each sample within season and station with a Wilcoxon rank sum test revealed a significant difference between both fractions at the marine station in both seasons as well as at the oligohaline station in fall/winter. Differences between bacteria fractions at the mesohaline station, however, remained non-significant (Supplemental Table S1).

**DISCUSSION**

Particles such as phytoplankton and terrestrial POM provide important habitats for bacteria (e.g., Grossart, 2010) and drive bacterial diversity and dynamics (e.g., Rooney-Varga et al., 2005). Many studies indicate pronounced differences in community composition of PA and FL bacteria at various sampling sites (Acinas et al., 1999; Crump et al., 1999; Ortega-Retuerta et al., 2013; Bižić-Ionescu et al., 2014). Yet, it remains unclear whether PA and FL bacteria show systematic differences in diversity and dynamics when environmental conditions, in particular salinity and DOC, change (Smith et al., 2013). Therefore, we have set a particular focus on the comparison of PA vs. FL bacterial communities at 3 sampling stations with different environmental conditions during two contrasting seasons: summer vs. fall/winter. The Baltic Sea provides an
ideal model system for such studies since it is characterized by strong and relatively stable gradients in salinity and DOC from north to south resulting in decreasing amounts of organic matter and nutrients. Our study revealed that PA OTUs account for 52% of overall total OTUs and thus represent an important component of bacterial \( \alpha \)-diversity. Dynamics of PA and FL BCC differed in relation to environmental settings, in particular those related to phytoplankton bloom development. Whereas relative OTU and absolute cell abundance of PA bacteria was low (\(<10\%\)), their significantly higher csBPP rates than FL bacteria indicates that PA bacteria represent an important component of Baltic Sea bacterioplankton (Simon et al., 2002, 2014; Grossart, 2010).

**Bacterial Communities in Relation to Changes in Environmental Variables**

The first sampling cruise was performed during November/December 2011 to account for the fall/winter season. Autumn 2011, however, was unusually warm (4–5°C higher than average November temperatures during the past 20 years) and the first ice cover of the Bothnian Bay was delayed by ca. 2 weeks (8th of December). Yet, we assigned this sampling period as “fall/winter,” since it was characterized by an accumulation of nutrients, short daylight, and a relatively low phytoplankton biomass. In contrast, our “summer” sampling was performed during May/June 2012 when nutrient concentrations were already low, daylight periods long, and a bloom of cyanobacteria and diatoms occurred (Nausch et al., 2013). However, due to relatively unfavorable weather conditions (cold and windy) the phytoplankton bloom was not as intense as in the years before. Moreover, in 2012, due to a relatively cold summer and absence of the normal temperature rise in May/June water temperatures differed little between seasons and stations. The oligohaline station differed from the other stations by low temperature and bacterial activities as well as a relatively high nitrate concentration in summer, due to its ice coverage until mid-May (Nausch et al., 2013). The phytoplankton bloom development at the mesohaline station in summer can serve as an explanation for the observed high abundance and proportion of PA bacteria.

BCC and bacterial \( \alpha \)-diversity differed with salinity and between both sampling periods with generally higher species richness in fall/winter. It is known that bacterial species richness generally decreases during bloom situations in summer (Hodges et al., 2005). This is supported by recent studies in coastal surface waters which indicate the highest bacterial richness during winter (Gilbert et al., 2010; Ghiglione and Murray, 2012). In the Baltic Sea, a larger resource heterogeneity can be also expected in fall/winter when phytoplankton blooms are relatively rare, whereas resources during phytoplankton blooms, particularly in summer, seem to be more homogeneous and may lead to the observed reduction in bacterial richness in summer (Grespo et al., 2013). Differences between PA and FL BCC were also more significant in fall/winter, when precipitation and hence terrestrial runoff of POM were higher than in summer. Changes in water conditions and concentrations of particulate matter (e.g., POC,
PN) as well as dissolved variables (e.g., DOC) were shown to correlate with seasonal changes in abundant taxa of heterotrophic bacteria (Crump et al., 2003; Fortunato et al., 2013). Recently, differences in the lifestyles and metabolic capabilities of PA and FL bacterial fractions were found by metagenome studies in coastal as well as open ocean areas (Simon et al., 2014; Fontané et al., 2015).

We found a dominance of the typical bacterial community in surface waters of the Baltic Sea (Herlemann et al., 2011). The number of exclusively mesohaline taxa in the PA fraction was low in summer and accounts for only half of the taxa occurring at the two other stations. In contrast, number of FL OTUs was higher at the mesohaline station than at the marine one. In fall/winter, PA fractions at the meso- and oligohaline stations were less diverse than at the marine station, whereas FL bacteria showed no distinct differences. The more pronounced differences between BCC of FL and PA bacterial fractions in fall/winter than in summer could be explained by changes in composition and hence particle quality (>5 µm) throughout the season. Differences between PA and FL bacteria are not necessarily correlated to particle concentration, but to SPM quality (Doxaran et al., 2012). Generally, PA bacteria have larger genome sizes than FL (Allen et al., 2012), and especially PA bacteria contain more transporters that can be linked to the successive decomposition of phytoplankton blooms (Teeling et al., 2012; Smith et al., 2013). This enables a flexible and rapid adaptation to changes in various environmental conditions, e.g., snow melt or phytoplankton blooms, by switching between different life-styles.

A substantial fraction (52%) of all retrieved OTUs occurred exclusively in the PA fraction of all samples. In contrast, numbers of exclusively FL OTUs were often lower than for the PA fraction indicating that particles (>5 µm) serve as important hotspots for bacterial α-diversity (Simon et al., 2002; Grossart, 2010; Rösel and Grossart, 2012; Bižić-Ionescu et al., 2014) also in the Baltic Sea. This notion is supported by the generally higher bacterial richness of PA than FL bacteria, except for the mesohaline station in summer. The increase in number of PA bacteria in summer at the marine and mesohaline stations were related to phytoplankton development. In contrast, at the relatively cold oligohaline station number of PA bacteria remained at the same high level in both seasons indicating that PA bacteria greatly benefit from riverine terrestrial input at this station, in particular in fall/winter when riverine freshwater runoff is high. This notion is supported by the higher proportion of PA bacteria at the oligohaline station in fall/winter, whereas the mesohaline station showed the opposite trend.

Since the mesohaline site contains only a small number of exclusively mesohaline taxa (ca. 20%) the mesohaline site may represent a mixture of oligohaline and marine bacteria. Herlemann et al. (2011) found that the brackish waters of the Baltic Sea are occupied by a diverse combination of freshwater and marine clades that seem to have adapted to the brackish conditions. Consequently, α-diversity changed little in both seasons indicating that PA bacteria greatly benefit from riverine terrestrial input at this station, in particular in fall/winter when riverine freshwater runoff is high. This notion is supported by the higher proportion of PA bacteria at the oligohaline station in fall/winter, whereas the mesohaline station showed the opposite trend.

Bacterial richness was highest at the marine station in fall/winter with 71% of all marine OTUs, and 59% of total OTUs in winter samples exclusively occurred in the PA fraction which suggests that PA bacteria are generally more diverse than FL bacteria. Several studies have shown that PA bacteria greatly contribute to the overall bacterial diversity in surface waters (LaMontagne and Holden, 2003; Crespo et al., 2013; Ortega-Retuerta et al., 2013; Bižić-Ionescu et al., 2014). For the Northwest Mediterranean, Sea Crespo et al. (2013) showed that almost half of all OTUs (49%) exclusively occur in the PA fraction, 25% were shared and 26% exclusively occur in the FL fraction. The authors have also shown that OTUs which stem exclusively from PA or FL bacteria were in general very low in abundance (6% of total abundance). Whereas most of the previous studies indicate a higher richness of FL compared to PA bacteria (Acinas et al., 1999; Hollibaugh et al., 2000; Ghiglione et al., 2009; Kellogg and Deming, 2009), our study indicates a higher richness of PA bacteria in almost all samples. This finding is supported by metagenome studies specifically differentiating between PA and FL fractions in samples from a coastal margin (Simon et al., 2014). A possible reason for this discrepancy can be related to the fact that previous studies, without next generation sequencing, may have analyzed only the most abundant taxa and hence missed a large portion of total bacterial α-diversity. The 454-pyrosequencing approach has been successfully used to estimate also tax of low abundance (Sogin et al., 2006; Pedrós-Alió, 2012) leading to a much higher bacterial α-diversity and richness, in particular of PA bacteria. Despite this advantage it should be kept in mind that sequencing errors of pyro-sequencing can bias the true number of rare taxa. Since we have quality-filtered our sequences according to Kunin et al. (2010) including removal of a) reads with one or more unresolved bases (Ns), b) errors in the barcode or primer sequence and c) atypically short or long reads as well as using d) a clustering threshold not greater than 97% identity, we think that an overestimation of OUT richness is unlikely. It should be also mentioned that the total amount of reads per sample can vary by orders of one magnitude within a single sequencing run. Randomly subsampling the same number of reads for each sample based on the smallest sample size is a common approach to allow for direct comparisons of samples (Gilbert et al., 2010; Crespo et al., 2013; Ortega-Retuerta et al., 2013). To avoid false interpretations in diversity of samples with very low reads, a rarefaction curve was performed to allow the calculation of species richness for a given number of individual samples. Though the rarefaction curves did not reach an asymptote, the most underestimated samples are the ones with already the highest diversity (Supplemental Figure S3). The function “rrarefy” generates randomly rarefied community data without replacement, so that the variance of rarefied communities is rather related to rarefaction proportion than to the sample size.

**Specific Phylogenetic Findings**

Similar to Herlemann et al. (2011), our observations revealed a higher abundance of Gammaproteobacteria with increasing salinity and the opposite trend for Actinobacteria and Betaproteobacteria (also see Bižić-Ionescu et al., 2014). Contrary to earlier observations (Herlemann et al., 2011) indicating a decreasing abundance of Alphaproteobacteria.
(mainly *Pelagibacteriales*) at lower salinities, we detected a surprisingly high abundance of FL *Alphaproteobacteria* at the oligohaline station in summer. The predominant taxa at the oligohaline station belonged to the *Pelagibacteriales* previously found in Chesapeake Bay. Chesapeake Bay is the largest estuary in the United States. Kan and colleagues have shown that the *Pelagibacteriales* (SAR11-related) clones appeared in the Chesapeake Bay and the Delaware Estuary only in the warm season (Kan et al., 2007), which correlates with our findings.

The phylogenetic distribution of bacteria at both sampling time points is in accordance with observations of Andersson et al. (2009), who analyzed the seasonal pattern of BCC (from May to October 2008) at the Landsort Deep. In this study, dynamics of *Alphah*- and *Betaproteobacteria* were closely related to a cyanobacterial bloom. *Bacteroidetes* showed a peak in summer (June) and a decrease in fall (October) when both organic matter concentrations and temperatures dropped. A similar temporal development was also found for *Actinobacteria* which were more dominant in late July until October. Contrary to our results, Herlemann et al. (2011) described a large fraction of *Verrucomicrobia* in summer (July) at the mesohaline site. This discrepancy might be related to differences in sampling years and time points. According to Andersson et al. (2009), the peak of *Verrucomicrobia* occurred from late June to August which is later than our summer and earlier than our fall/winter sampling. Therefore, it is very likely that we have missed this *Verrucomicrobia* peak with our sampling scheme. In fall/winter, the mesohaline station showed a relatively high number of *Actinomycobacteria* in the FL fraction and a dominance of *Gammaproteobacteria* and *Planctomycetes* in the PA fraction, which indicates the higher presence of oligohaline bacteria in fall/winter, probably due to higher riverine freshwater runoff and reduced phytoplankton influence.

In our study, there was a high proportion of PA *Planctomycetes* and *Flavobacteria* which greatly differ in their life-style. Interestingly, we found a dominance of PA *Planctomycetes* exclusively in the oligohaline samples (35% of PA bacteria in summer and 19% in fall/winter), which mainly consisted of CL500-3 (95% of PA *Planctomycetes*). CL500-3 was also the most significant driver of the separation between fractions (SIMPER: 5%) and between the separation of the oligohaline and the mesohaline as well as the marine station (SIMPER: 7 and 6%). This group is typically found in freshwater and was first described in an oligotrophic crater lake (Urbach et al., 2001). They tend to a PA lifestyle and thus drive the separation of BCC’s in different habitats (Jackson et al., 2014). *Planctomycetes* seem to be involved in the mineralization of algal biomass, which is supported by the positive correlation between their abundance and chl concentrations (Pizzetti et al., 2011). *Bacteroidetes-Flavobacteria* were less abundant in these samples, but dominated at the marine station. *Flavobacteria* occur in marine and freshwater habitats as typical PA bacteria since they represent metabolic generalists with a broad range of polysaccharide degrading enzymes (Jeske et al., 2013; Walsh et al., 2013). The marine station in summer was dominated by *Polaribacter*, *Flavicola* and *Formosa* genera; both *Polaribacter* and *Flavicola* were previously found on heterogenous particles (Bižić-Ionescu et al., 2014). *Formosa* was identified as an early colonizer of a decaying diatom bloom (Teeling et al., 2012). *Flavobacterium* was mainly found in the oligohaline and mesohaline stations, but seemed to be quite stable in both seasons (around 4%) at the oligohaline station. However, this group was an important member in structuring the fractions (SIMPER 4%).

In conclusion, strong differences in BCC as well as in $\alpha$— and $\beta$-diversity occurred among stations with different salinities and seasons revealing significantly higher species richness in fall/winter than in summer. PA bacteria comprise a large number of less abundant OTUs, especially at the oligotrophic marine site. At the marine station, $\alpha$-diversity of PA bacteria was in both seasons higher than of FL bacteria and substantially contributed to overall bacterial diversity. Interestingly, key PA bacteria consisted of *Planctomycetes* at the oligohaline station and of *Flavobacteria* at the marine station, which differ greatly in their life-style. PA bacteria—although usually less abundant (<10% of total bacteria)—were characterized by higher cell specific activities and consequently greatly contribute to overall BPP. Thus, PA bacteria represent an important component of overall bacterial $\alpha$- and $\beta$-diversity and presumably activity in the Baltic Sea.

**ACKNOWLEDGMENTS**

Sequencing was performed at the Berlin Center for Genomics in Biodiversity Research. We thank all partners from the ATKiM project, the crew of MS Meteor (M86-1a, b; M87-3a) and Thomas Hornick for their technical support during sample collection. We also thank Helge Arz, Matthias Labrenz and Detlef Schulz-Bull for cruise logistic and nutrient analyses. Ulrich Struck of the Museum für Naturkunde Berlin is thanked for contributing POC data, and Susan Mbedi and the Berlin Center for Genomics in Biodiversity Research for 454 sequencing. The Aquatic Microbial Ecology group at IGB Stechlin and Kirsten Pohlmann are acknowledged for providing statistical advice and many helpful comments on earlier drafts of the paper. The study was funded by the Leibniz Association as part of the ATKiM project (SAW-2011-IOW-3). We also thank the German Science foundation for financially support for the cruise M86-1a, 1b, and M87-3a of MS Meteor.

**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb.2015.01297

Supplemental Figure S1 | Unweighted-pair group method with arithmetic mean (UPGMA) dendograms based on Bray-Curtis dissimilarities of PA and FL bacteria distribution at different stations [marine (A), mesohaline (B), oligohaline (C), and for all samples (D)].

Supplemental Figure S2 | Unweighted-pair group method with arithmetic mean (UPGMA) dendograms based on Bray-Curtis dissimilarities of the distribution of PA and FL bacteria at different stations in summer and winter marine (A,D), mesohaline (B), oligohaline (C,F).

Supplemental Figure S3 | Rarefaction curves for free-living and particle-associated samples of the total dataset.
Supplemental Table S1 | Wilcoxon Rank sum test between all samples within station and season and a factor (F) from 0 to 1, which compares the groups [triplicates (a, b, c) vs. fractions (PA, FL)] within one station (mar, meso, oligo) and season (summer, fall/winter) to estimate the difference between the fractions, with 1 as highest possible difference.

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Supplemental Table S2 | Number of OTUs, reads, Chao1, Shannon [H], and Simpson [1–D] and Buzzas and Gibson's Evenness [e−H/S] diversity indexes of averages of samples after normalization to 430 reads per sample, PA, particle-attached; FL, free-living; PA, particle-associated fraction; FL, free-living fraction; Tot, total bacteria.
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