Small Changes in pH Have Direct Effects on Marine Bacterial Community Composition: A Microcosm Approach

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Abstract

As the atmospheric CO2 concentration rises, more CO2 will dissolve in the oceans, leading to a reduction in pH. Effects of ocean acidification on bacterial communities have mainly been studied in biologically complex systems, in which indirect effects, mediated through food web interactions, come into play. These approaches come close to nature but suffer from low replication and neglect seasonality. To comprehensively investigate direct pH effects, we conducted highly-replicated laboratory acidification experiments with the natural bacterial community from Helgoland Roads (North Sea). Seasonal variability was accounted for by repeating the experiment four times (spring, summer, autumn, winter). Three dilution approaches were used to select for different ecological strategies, i.e. fast-growing or low-nutrient adapted bacteria. The pH levels investigated were in situ seawater pH (8.15–8.22), pH 7.82 and pH 7.67, representing the present-day situation and two acidification scenarios projected for the North Sea for the year 2100. In all seasons, both automated ribosomal intergenic spacer analysis and 16S ribosomal amplicon pyrosequencing revealed pH-dependent community shifts for two of the dilution approaches. Bacteria susceptible to changes in pH were different members of Gammaproteobacteria, Flavobacteriaceae, Rhodobacteraceae, Campylobacteraceae and further less abundant groups. Their specific response to reduced pH was often context-dependent. Bacterial abundance was not influenced by pH. Our findings suggest that already moderate changes in pH have the potential to cause compositional shifts, depending on the community assembly and environmental factors. By identifying pH-susceptible groups, this study provides insights for more directed, in-depth community analyses in large-scale and long-term experiments.

Introduction

Since the beginning of the industrial period, the oceans have taken up one-quarter to one-third of anthropogenic CO2 emissions [1,2]. This has already led to a reduction in surface ocean pH of 0.1 units, which may reach up to 0.7 units assuming the depletion of all fossil fuel reserves during the next three centuries [3]. In contrast, pH has constantly remained above 8.1 for the last 23 million years [4]. By the year 2100, atmospheric pCO2 values of 700 or 1000 ppm may lower mean surface pH in the southern North Sea to 7.82 or 7.67, respectively [5].

The effects of the anticipated rapid reduction in pH on marine organisms, and their ability to adapt, will determine future marine biodiversity and ecosystem functions. Yet the impact of ocean acidification on different groups of marine organisms remains under debate [6,7], especially regarding heterotrophic bacteria as important players in marine biogeochemical cycles. Joint et al. [8] recently argued that microbe-dependent processes will not substantially change in a more acidic ocean, as marine microbes already experience large regional, temporal and depth-dependent pH variability, and even greater pH ranges are observed in freshwater lakes. This view was challenged by a meta-analysis on microbe-related ocean acidification research, which identified nitrogen fixation, cyanobacterial photosynthesis and elemental ratios as affected by changes in seawater carbonate chemistry [9]. Concerning other microbial processes and especially heterotrophic bacteria however, results have often been inconsistent and Liu et al. [9] concluded that “more research is needed at multi-species and community scales”.

What we know about ocean acidification effects on bacterial communities predominantly stems from complex systems such as symbiotic microbial communities of corals or large-scale mesocosm experiments. At reduced pH, coral microbial communities were found to shift to bacteria associated with stressed or diseased hosts [10,11], which could however not be confirmed at natural CO2 vent sites [12]. Furthermore, a decrease in the relative abundance of Alphaproteobacteria and an increase in the relative abundance of Flavobacteriales were observed in natural biofilms from the Great Barrier Reef [13].

Knowledge on the seawater bacterial community has remained scarce though. In mesocosm experiments, only minor indications of bacterial community shifts with pH were found [14,15]. Notably, these findings relied on only one or two replicates per
pCO₂ treatment, which is a common problem in mesocosm studies. Although these experiments are biologically highly complex, involving indirect pH effects through food web interactions, they are usually carried out in low replication, due to logistical challenges and high costs. As a consequence, these experimental designs preclude a robust statistical interpretation. Furthermore, the natural variability of bacterial communities, which is characterized by seasonally recurring patterns [16–18], is not taken into account.

Therefore, a straightforward small-scale approach is needed to allow for high replication and the consideration of differently assembled communities. The problem of limited culturability of marine bacteria can be addressed by dilution experiments. A certain fraction of natural bacterioplankton is able to grow in particle-free, unenriched seawater [19], although the confinement to experimental containers causes shifts in community structure often described as the “bottle effect” [20]. To alleviate this bias and select for typical marine bacteria, the concept of dilution culture was developed [21]. Originally, this approach followed a dilution to near extinction strategy to isolate previously uncultured marine strains [22]. The idea was developed further to study diluted cultures of entire bacterioplankton assemblages, in which depending on the dilution strategy, different parts of the bacterial community are selected for [23,24]. Dilution experiments were successful in e.g. identifying selective grazing mortality of bacterial groups [25].

Here we present a highly replicated, culture-dependent investigation of direct pH effects on bacterial communities from the North Sea. The sampling station Helgoland Roads is well suited for this approach, as seasonal and long-term microbiological data are available [26–29] and dilution or enrichment experiments have been carried out before [25,30]. We used three dilution approaches to study different bacterial groups and repeated the experiment in different seasons. We show that small changes in pH have direct effects on bacterial community composition and identify Gammaproteobacteria, Flexibacteraceae, Rhodobacteraceae and Campylobacteraceae as phylogenetic groups responding most notably to differences in pH.

Materials and Methods

Ethics statement

No specific permits were required for the sampling and activities performed in the study.

Experimental set-up and sampling

We conducted the experiment in spring, summer and autumn, 2010, and in winter 2011 (Table 1) with surface water sampled at 1 m depth at sampling station Helgoland Roads (54°11.3’N, 7°54.0’E), North Sea. Water samples were subjected to three different dilution protocols and three different pH levels (see sections below), yielding a total of nine different treatments. Each treatment was replicated five times in autoclaved one liter borosilicate glass bottles. Bottles were filled with 1100 ml to keep the headspace small (volume 50–60 ml) to prevent excessive CO₂ exchange at the air-seawater interface observed in systems open to the atmosphere [31]. Bottles were incubated in the dark at approximately in situ temperature (Table 1) and were mixed daily by inversion. Samples were taken after four weeks of incubation.

We used the following three dilution protocols: ‘no dilution’, ‘serial dilution’ (diluted weekly 1:1000 starting after one week), and ‘initial dilution’ (initially diluted 1:1000). Larger organisms were removed by filtration through 10 µm Isopore™ Membrane Filters (TCTP-type, Millipore, Eschborn, Germany). For the dilutions, we used seawater collected on the respective initial sampling date, which had been repeatedly sterilized by filtration (0.2 µm Isopore™ Membrane Filters, GTTP-type, Millipore).

The pH levels were the current in situ seawater pH (8.15–8.26, Table 1), pH 7.82 and 7.67. We acidified the initial seawater samples (fraction <10 µm) and the sterile-filtered seawater used for the dilutions with 2 M HCl. To determine pH at the beginning and after the four weeks of incubation, we used a ProLab 3000 pH meter with an IoLine pH combination electrode with temperature sensor (type IL-pHT-A170MF-DIN-N), calibrated with standard buffer solutions (pH 4.01, 6.87, 9.18) (all materials: SI Analytics, Mainz, Germany). All pH measurements were carried out at the respective incubation temperature and are reported on the National Bureau of Standards (NBS) scale.

For cell counts, samples of 4 ml were fixed with formaldehyde (1% [w/v] final concentration) for 1 h at room temperature and were subsequently stored at −80°C until further analysis. Aliquots of 500 µl were stained for 10 min with 10 µl of a freshly prepared 400X SYBR Green (invitrogen™, Life Technologies, Paisley, UK) solution in sterile filtered dimethyl sulfoxide (DMSO). Directly prior to staining the cells, we added 10 µl of a diluted solution of Fluoresbrite® Polychromatic Red Microspheres 1.0 µm (Polysciences Europe, Eppelheim, Germany) as an internal counting standard [final concentration of about 10% of the expected number of cells]. We analysed samples with an Accuri C6 flow cytometer (BD Accuri Cytometers, Ann Arbor, MI, USA) with the fludrics setting “slow” for 1.5 min. To reduce noise, we set a threshold on FL1-H of 550 for winter and spring and of 600 for summer and autumn samples, respectively. The actual flow through was calibrated with BD Trucount™ Controls (BD Biosciences, San Jose, CA, USA).

Biomass was collected on 0.2 µm Isopore™ Membrane Filters (GTTP-type, 47 mm diameter, Millipore) and DNA extraction was performed as previously described [29]. DNA concentration and purity were determined by photometry using an Infinite 200 PRO NanoQuant (Tecan, Männedorf, Switzerland).

ARISA

For ARISA fingerprints, we used forward primer L-D-Bact-132-a-A-18 (5’-CCCGGTTTCCACTTGG-3’) and reverse primer S-D-Bact-1522-b-S-20 (5’-TGCGGTGAGTCCCCCTCCTT-3’) [32], the latter labeled with an infrared dye. PCR reactions were performed in volumes of 25 µl containing 10 ng template DNA, 2.5 µl Taq Buffer (10X), 5 µl TaqMaster PCR Enhancer (5X), 0.7 µl of each primer (20 µM), 0.75 µl dNTPs (2.5 mM each), and 1.4 U Taq DNA polymerase (5 Prime, Hamburg, Germany). Cycling conditions were: 95°C for 3 min, followed by 30 cycles of 95°C for 1 min, 50°C for 1 min and 68°C for 1 min, with a final step at 68°C for 5 min. Aliquots of PCR products were verified on agarose gels and depending on agarose gel band intensities, original or diluted PCR products were mixed with an equal volume of formamide containing stop mix. Samples were heated to 95°C for 2 min, subsequently kept on ice for 10 min, and 0.25 to 0.5 µl were separated in 5.5% polyacrylamide gels. Running conditions were 1500 V for 14 h on a LI-COR 4300 DNA Analyzer. A pre-run of 15 min at 45°C was carried out to precondition the gel and sequencer prior to loading the samples. As a size reference, we used a 50–1500 bp standard (all materials: LI-COR Biosciences, Lincoln, NE, USA).

ARISA gels were analyzed with the Bionumerics 5.10 software (Applied Maths, Sint-Martens-Latem, Belgium). Bands with intensities lower than 4% of the maximum value of the respective lane and bands smaller than 300 bp were neglected. Binning to
band diversity was performed according to Brown et al. [33]. Beta-
diversity was calculated with the Jaccard coefficient.

16S ribosomal amplicon pyrosequencing

Based on ARISA results, samples were selected for pyrosequ-
encing of the bacterial 16S rDNA (pH in situ) and 7.67 of all ‘no
dilution’ and ‘serial dilution’ treatments and of the ‘initial dilution’
in summer; additionally starting communities. Primers used to
construct the amplicon library were of the structure 5’-[Roche’s
adaptor for long reads (Lib-L)] - [template-specific sequence] -3’.
As template specific sequences we used forward primer GM3 (5’-
AGAGTTTGATCMTGGC-3’) and reverse primer 907 (5’-
CCGTCAATTGCMTTTGAAGGTTT -3’) [34]. To pool all 93
samples in a single run, 93 different forward primers were
constructed, each containing a different 10 bp long Multiplex
Identifier (Roche) between the adaptor sequence and the template
specific sequence. Amplifications were conducted in two duplicate
reactions of 50 µl, each containing 20 ng of template DNA, 5 µl
Taq Buffer (10 ×), 10 µl TaqMaster PCR Enhancer (5 ×), 1.25 µl
each of primer (20 µM), 5 µl dNTPs (2.5 mM each), and 2.5 U of
Taq DNA polymerase (5 Prime). Cycling conditions were: 94
°C for 10 min, followed by 20 cycles of 94°C for 1 min, 44°C for
1.5 min and 68°C for 2 min, with a final step at 68°C for 15 min.
The duplicate reactions were combined and purified with the
QIAquick PCR purification kit (Qiagen, Valencia, CA, USA).
Amplicons were pooled equimolarly, including 2 µl of a negative
control with a separate Multiplex Identifier to check for
contaminations, and sequenced on 1/2 plate on a Roche 454
GS-FLEX Titanium platform at the Institute of Clinical Molecular
Biology (IKMB), University of Kiel, Germany. Sequence data
were deposited in the NCBI Sequence Read Archive (accession
number SRP014019).

Sequences were processed with the MOTHUR software
(version 1.23.0, [35]). Flowgrams were trimmed to 360 to 800
flows and the shhh.flows command with standard settings was used
to remove sequencing noise. After the removal of primer and
barcode sequences, unique sequences were aligned to the SILVA
reference database. After removing chimeras (chimera.uchime
routine) and chloroplast sequences, 163,029 sequences remained
(20,419 of them unique). Sequences started and ended at the same
alignment position, 95% being of a length between 381 and
420 bp. A distance matrix was created and sequences were
grouped into operational taxonomic units (OTUs) on a 97% level.
The number of sequences in each sample was normalized by
randomly selecting the number of sequences present in the
smallest sample (n = 494). This resulted in a subset containing
2133 OTUs which we used for all further analyses. Sample
coverage and richness (observed number of OTUs) were obtained
as implemented in MOTHUR. OTU abundance data were
square root transformed and beta-diversity was calculated with the
Bray-Curtis coefficient.

Statistical analysis

The experiment was based on a crossed factorial design,
consisting of the three factors ‘season’ (four levels, fixed), ‘dilution’
(three levels, fixed), and ‘pH’ (three levels, fixed). For pyrose-
sequencing, the factors ‘dilution’ and ‘pH’ consisted of only two
levels and the summer ‘initial dilution’ was analyzed separately. In
case of a significant influence of ‘pH’ or of an interaction term
involving ‘pH’, post-hoc comparisons for the factor ‘pH’ within the
highest ranking interaction were performed.

Bacterial abundance and richness were analyzed through
defactorial ANOVAs [36] with Statistica 7.1 (StatSoft, Tulsa, OK,
USA), using Tukey’s HSD test for post-hoc comparisons. Beta-
diversity was analyzed by permutational multivariate ANOVA
(PERMANOVA, [37]) and similarity percentage analysis (SIM-
PER). SIMPER allowed us to calculate the total similarity within
dissimilarity between treatments, and to determine charac-
teristic and discriminatory OTUs. To visualize patterns of the
influence of ‘pH’ within ‘season’-dilution’ combinations, we
performed principal co-ordinate analysis (PCO, [38]). For all
multivariate analyses, we used Primer 6 with the add-on package
PERMANOVA+ (PRIMER-E Ltd, Plymouth, UK).

Results

Environmental data, pH manipulations, initial community
composition and bacterial abundance

In situ pH varied from 8.15 to 8.22 and seawater temperature
from 3.2 to 15.8°C (Table 1). At the end of the experiment, the
mean absolute deviations from the initial pH values were
0.10±0.06 (‘no dilution’, n=60), 0.07±0.05 (‘serial dilution’,
n=59) and 0.62±0.28 (‘initial dilution’, n=60) (see Table S1 for
pH in all ‘season’-dilution’ combinations). Initial bacterial
abundance ranged from 3.3×10^5 to 1.9×10^6 cells ml^-1
(Table 1). The starting community was more diverse in autumn
and winter (134 and 181 observed OTUs) compared to spring
and summer (69 and 54 observed OTUs). Starting communities
were dominated by sequences of Alphaproteobacteria (especially
in summer), Flavobacteria (especially in spring) and Gammaproteobacteria.
Additionally, up to 20% Bacteria that could not be assigned to
classes (autumn and winter) and 10% Betaproteobacteria (winter)
occurred (Figure 1).

After the four weeks of incubation, bacterial abundance ranged
between 1.1×10^5 and 6.1×10^6 cells ml^-1 and was significantly
influenced by ‘season’ (F_3,113 = 21.451, p<0.001), ‘dilution’
(F_2,113 = 111.279, p<0.001), and their interaction
(F_6,113 = 14.399, p<0.001), but not by ‘pH’ (F_2,113 = 1.6163,
p = 0.202) (Figure S1).
Bacterial community composition

Based on ARISA fingerprints, all three experimental factors and their interactions significantly influenced bacterial community structure (PERMANOVA, Table 2). The highest amount of variation was explained by ‘season’, ‘dilution’ and their interaction term (Sq. root, Table 2). Communities in both ‘no dilution’ and ‘serial dilution’ were significantly influenced by ‘pH’ in all seasons (Table 3), predominantly already at pH 7.82. Additionally, communities from both lowered pH treatments differed from each other, except in summer. ‘Initial dilution’-communities in contrast were influenced by ‘pH’ only in spring and summer.

For 16S ribosomal amplicon pyrosequencing, we selected all ‘season’-‘dilution’ combinations for which significant differences between communities at pH \textit{in situ} and 7.67 had been observed (Table 3). Sequencing data confirmed that community structure in both ‘no dilution’ and ‘serial dilution’ was significantly influenced by all experimental factors and interactions (PERMANOVA, Table 4). In accordance with ARISA results, ‘season’, ‘dilution’ and their interaction term were the highest sources of variation (Sq. root, Table 4). Except for the ‘serial dilution’ in autumn, all ‘pH’-dependent differences were confirmed (Table 5 and Table 6). PCO plots of ‘season’-‘dilution’ combinations are shown in Figure S2 (based on ARISA) and Figure S3 (based on 16S ribosomal amplicon pyrosequencing).

Sample coverage obtained by pyrosequencing ranged from 72.1 to 99.8% (Table S2), yielding bacterial richness estimates of 8 to 218 OTUs per sample (Figure S4). In both ‘no dilution’ and ‘serial dilution’, richness was significantly influenced by ‘season’ (F3,63 = 122.201, \(p = 0.0001\)), ‘dilution’ (F 1,63 = 1901.215, \(p = 0.0001\)) and ‘pH’ (F1,63 = 29.608, \(p = 0.0001\)), and the interactions ‘season’x‘dilution’ (F3,63 = 138.125, \(p = 0.0001\)), ‘season’x‘pH’ (F3,63 = 9.317, \(p = 0.0001\)), ‘dilution’x‘pH’ (F 1,63 = 28.395, \(p = 0.0001\)) and ‘season’x‘dilution’x‘pH’ (F3,63 = 2.880, \(p = 0.05\)). For the ‘serial dilution’, no ‘pH’-dependent differences were found, whereas for the ‘no dilution’, richness was significantly higher at pH 7.67 than at pH \textit{in situ} in spring (p = 0.001) and autumn (p<0.001, Tukey’s HSD test, 63 d.f.). In the ‘initial dilution’ in summer, richness was not significantly influenced by ‘pH’ (F1,7 = 1.093, \(p = 0.331\)).

Bacterial groups in the ‘season’-‘dilution’ combinations

Considering only pH \textit{in situ} treatments, i.e. disregarding pH effects, communities in different dilutions within a given season differed between 86 and 97% from each other. Classes predominantly discriminating between ‘no dilution’ and ‘serial dilution’ were \textit{Flavobacteria}, \textit{Alphaproteobacteria} and \textit{Gammaproteobacteria} (Figure 2; see Table S3 for similarity and Table S4 for dissimilarity analyses).

No dilution’-communities were characterized mainly by \textit{Flavobacteria}, \textit{Alphaproteobacteria} and \textit{Gammaproteobacteria} which could not be assigned to families. \textit{Flavobacteria} were the most important
group in spring and winter, while in summer, Alphaproteobacteria, especially of the SAR11-clade, dominated. In autumn, the community was more diverse. Less abundant groups characteristic for ‘no dilution’-communities included Planctomycetaceae, Betaproteobacteria, and Bacteroidetes that could not be assigned to classes.

In contrast, ‘serial dilution’-communities were dominated by known representatives of the Gammaproteobacteria. Important groups were Oceanospirillaceae (all seasons), unclassified Alteromonadales (summer, autumn, winter), Colwelliaceae (spring, winter) and Pseudolalteromonadaceae (spring). In certain ‘serial dilution’-communities, Flavobacteriaceae and Rhodobacteraceae additionally accounted for high contributions to total similarities. The ‘initial dilution’-community in summer was dominated by Gammaproteobacteria, Flavobacteriaceae and Rhodobacteraceae as well. Within the Gammaproteobacteria however, Alteromonadaceae dominated.

### Bacterial groups responding differently to pH

In ‘season’-‘dilution’ combinations with significant ‘pH’ effects (Table 5 and Table 6), communities of the two pH treatments differed between 49 and 63% from each other. The most important discriminatory classes were Gammaproteobacteria, Flavobacteria, Alphaproteobacteria and Epsilonproteobacteria (Figure 3; see Table S3 for similarity and Table S5 for dissimilarity analyses).

Gammaproteobacterial families contributing to total dissimilarities were Oceanospirillaceae, Colwelliaceae, Alteromonadaceae, Pseudolalteromonadaceae, and to a lesser extend Idiomarinaceae, Shevanellaceae and Vibrionaceae. In spring and winter, Oceanospirillaceae were more characteristic for pH in situ, while in summer, they were more characteristic for pH 7.67. Colwelliaceae contributed more to total similarities at pH in situ in the ‘serial dilution’ in spring and winter.

### Table 3. PERMANOVA pair-wise comparisons of bacterial community composition based on Jaccard dissimilarities of ARISA profiles.

<table>
<thead>
<tr>
<th>Season</th>
<th>Comparison</th>
<th>no dilution</th>
<th>serial dilution</th>
<th>initial dilution</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>t (perm)</td>
<td>p (perm)</td>
<td>t (perm)</td>
</tr>
<tr>
<td></td>
<td>pH in situ vs. 7.67</td>
<td>1.574</td>
<td>0.327</td>
<td>0.209</td>
</tr>
<tr>
<td>spring</td>
<td>pH in situ vs. 7.67 vs. 6.7</td>
<td>3.563</td>
<td>0.013</td>
<td>1.494</td>
</tr>
<tr>
<td></td>
<td>pH in situ vs. 7.82 vs. 7.67</td>
<td>0.82</td>
<td>0.32</td>
<td>0.12</td>
</tr>
<tr>
<td>winter</td>
<td>pH in situ vs. 7.82 vs. 7.67</td>
<td>2.07</td>
<td>0.045</td>
<td>1.51</td>
</tr>
</tbody>
</table>

1Significant results (p (perm)<0.05) are highlighted in bold.
Displayed are pair-wise a posteriori comparisons of the factor ‘pH’ within ‘season’-‘dilution’ combinations, with at least 125 unique permutations per comparison.

### Table 4. PERMANOVA main tests of bacterial community composition based on Bray-Curtis dissimilarities of OTUs (16S ribosomal amplicon pyrosequencing; ‘no dilution’ and ‘serial dilution’).

<table>
<thead>
<tr>
<th>Sources of variation</th>
<th>d.f.</th>
<th>SS</th>
<th>pseudo F</th>
<th>p (perm)</th>
<th>Sq. root</th>
</tr>
</thead>
<tbody>
<tr>
<td>Season</td>
<td>3</td>
<td>74205</td>
<td>0.174</td>
<td>0.001</td>
<td>20.72</td>
</tr>
<tr>
<td>Dilution</td>
<td>1</td>
<td>71118</td>
<td>0.174</td>
<td>0.001</td>
<td>20.72</td>
</tr>
<tr>
<td>pH</td>
<td>1</td>
<td>3637.6</td>
<td>0.005</td>
<td>0.001</td>
<td>8.043</td>
</tr>
<tr>
<td>Season × Dilution</td>
<td>3</td>
<td>62133</td>
<td>0.005</td>
<td>0.001</td>
<td>4.46</td>
</tr>
<tr>
<td>Season × pH</td>
<td>3</td>
<td>10129</td>
<td>0.001</td>
<td>0.001</td>
<td>15.237</td>
</tr>
<tr>
<td>Dilution × pH</td>
<td>1</td>
<td>3431.6</td>
<td>0.002</td>
<td>0.001</td>
<td>10.906</td>
</tr>
<tr>
<td>Season × Dilution × pH</td>
<td>3</td>
<td>10272</td>
<td>0.001</td>
<td>0.001</td>
<td>21.771</td>
</tr>
<tr>
<td>Residuals</td>
<td>63</td>
<td>68634</td>
<td>0.001</td>
<td>0.001</td>
<td>33.007</td>
</tr>
<tr>
<td>Total</td>
<td>78</td>
<td>303900</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

1Significant results (p (perm)<0.05) are highlighted in bold.

### Table 5. PERMANOVA pair-wise comparisons of bacterial community composition based on Bray-Curtis dissimilarities of OTUs (16S ribosomal amplicon pyrosequencing; ‘no dilution’ and ‘serial dilution’).

<table>
<thead>
<tr>
<th>Comparison</th>
<th>no dilution</th>
<th>serial dilution</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>t (perm)</td>
<td>p (perm)</td>
</tr>
<tr>
<td>pH in situ vs. 7.67</td>
<td>2.3732</td>
<td>0.009</td>
</tr>
<tr>
<td>summer</td>
<td>1.5759</td>
<td>0.042</td>
</tr>
<tr>
<td>autumn</td>
<td>1.5233</td>
<td>0.005</td>
</tr>
<tr>
<td>winter</td>
<td>1.3772</td>
<td>0.007</td>
</tr>
</tbody>
</table>

1Significant results (p (perm)<0.05) are highlighted in bold.
Displayed are pair-wise a posteriori comparisons of the factor ‘pH’ within ‘season’-‘dilution’ combinations for ‘no dilution’ and ‘serial dilution’ treatments, with at least 125 unique permutations per comparison.

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**Table 3.** PERMANOVA pair-wise comparisons of bacterial community composition based on Jaccard dissimilarities of ARISA profiles.

**Table 4.** PERMANOVA main tests of bacterial community composition based on Bray-Curtis dissimilarities of OTUs (16S ribosomal amplicon pyrosequencing; ‘no dilution’ and ‘serial dilution’).

**Table 5.** PERMANOVA pair-wise comparisons of bacterial community composition based on Bray-Curtis dissimilarities of OTUs (16S ribosomal amplicon pyrosequencing; ‘no dilution’ and ‘serial dilution’).
and the ‘initial dilution’ in summer. In the ‘no dilution’ however, the opposite occurred. *Aberdonadaceae* dominated in the summer ‘initial dilution’, were they contributed more to total similarity at pH *in situ*. The same was found for the ‘no dilution’ in autumn, but in spring, they were more characteristic for pH 7.67. *Pseudoalteromonadaceae* contributed to total dissimilarity in spring and summer, and were always more characteristic for pH 7.67. Concerning *Gammaproteobacterial* families occurring in only one ‘season’-‘dilution’ combination, *Flavimonadaceae* contributed more to total similarity at pH *in situ*, while *Shewanellaceae* and *Vibrionaceae* contributed more to total similarity at pH 7.67.

The *Flavobacterial* family *Flavobacteriaceae* was the group with the overall highest contributions to total dissimilarities. In spring, *Flavobacteriaceae* were more characteristic for pH 7.67, but in autumn and winter, they were more characteristic for pH *in situ*. In summer, no consistent trend was found.

Within the *Alphaproteobacteria*, high contributions to total dissimilarity were attributed to *Rhodobacteraceae*, and to lesser extends to SAR11-clade, *Rhodospirillaceae* and *Sphingomonadaceae*. For *Rhodobacteraceae*, higher contributions to total similarity were generally observed at pH *in situ*. SAR11-clade contributed more to total similarity at pH *in situ* in spring, but no important differences were found in the other seasons. The other *Alphaproteobacterial* families did not show consistent trends.

Within the *Epsilonproteobacteria*, *Campylobacteraceae*, represented exclusively by the genus *Arcobacter*, contributed a third to total similarity at pH 7.67 in the ‘serial dilution’ in winter but were absent at pH *in situ*. Similar but less pronounced tendencies were found in spring and autumn.

Discriminatory groups that were less abundant or present only in one or few ‘season’-‘dilution’ combinations predominantly belonged to the classes *Sphingobacteria*, *Planctomycetaceae*, *Actinobacteria*, *Betaproteobacteria* and *Opitutae*. Notably, also various groups that could not be assigned to families contributed to pH-dependent dissimilarities.

**Discussion**

The impact of ocean acidification on bacterial communities remains under debate as studies conducted so far are highly complex but lack sufficient replication. Here we present a thorough analysis of diversely assembled bacterial communities, shaped by the experimental factors season and dilution. Season takes into account phenology, i.e. climate-driven annually recurring patterns of pelagic bacterial community composition and the ‘initial dilution’ in summer. In the ‘no dilution’ however, the opposite occurred. *Aberdonadaceae* dominated in the summer ‘initial dilution’, were they contributed more to total similarity at pH *in situ*. The same was found for the ‘no dilution’ in autumn, but in spring, they were more characteristic for pH 7.67. *Pseudoalteromonadaceae* contributed to total dissimilarity in spring and summer, and were always more characteristic for pH 7.67. Concerning *Gammaproteobacterial* families occurring in only one ‘season’-‘dilution’ combination, *Flavimonadaceae* contributed more to total similarity at pH *in situ*, while *Shewanellaceae* and *Vibrionaceae* contributed more to total similarity at pH 7.67.

The *Flavobacterial* family *Flavobacteriaceae* was the group with the overall highest contributions to total dissimilarities. In spring, *Flavobacteriaceae* were more characteristic for pH 7.67, but in autumn and winter, they were more characteristic for pH *in situ*. In summer, no consistent trend was found.

Within the *Alphaproteobacteria*, high contributions to total dissimilarity were attributed to *Rhodobacteraceae*, and to lesser extends to SAR11-clade, *Rhodospirillaceae* and *Sphingomonadaceae*. For *Rhodobacteraceae*, higher contributions to total similarity were generally observed at pH *in situ*. SAR11-clade contributed more to total similarity at pH *in situ* in spring, but no important differences were found in the other seasons. The other *Alphaproteobacterial* families did not show consistent trends.

Within the *Epsilonproteobacteria*, *Campylobacteraceae*, represented exclusively by the genus *Arcobacter*, contributed a third to total similarity at pH 7.67 in the ‘serial dilution’ in winter but were absent at pH *in situ*. Similar but less pronounced tendencies were found in spring and autumn.

Discriminatory groups that were less abundant or present only in one or few ‘season’-‘dilution’ combinations predominantly belonged to the classes *Sphingobacteria*, *Planctomycetaceae*, *Actinobacteria*, *Betaproteobacteria* and *Opitutae*. Notably, also various groups that could not be assigned to families contributed to pH-dependent dissimilarities.

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The **Table 6. PERMANOVA main tests of bacterial community composition based on Bray-Curtis dissimilarities of OTUs (16S ribosomal amplicon pyrosequencing; ‘initial dilution’ summer).**

<table>
<thead>
<tr>
<th>Sources of variation</th>
<th>d.f.</th>
<th>SS</th>
<th>pseude F</th>
<th>p (perm)</th>
<th>Sq. root</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH</td>
<td>1</td>
<td>1922.8</td>
<td>1.9165</td>
<td>0.011</td>
<td>14.384</td>
</tr>
<tr>
<td>Residuals</td>
<td>7</td>
<td>7023.1</td>
<td>31.675</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>8</td>
<td>8945.9</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Significant results (p (perm)<0.05) are highlighted in bold.*

Displayed are the tests for the factor ‘pH’ and the partitioning of multivariate variation for the ‘initial dilution’ (only summer experiment). p-values were obtained using type III sums of squares and 999 permutations under the full model. d.f.: degrees of freedom, SS: sums of squares, Sq. root: square root of the component of variation attributable to that factor in the model, in units of Bray-Curtis dissimilarities.

Direct pH Effects on Marine Bacterial Communities

We used HCl to adjust pH to values expected for the North Sea for the year 2100. Values remained relatively stable for ‘no dilution’ and ‘serial dilution’ treatments, but larger deviations were observed for the ‘initial dilution’. Interestingly, pH-dependent community shifts were least frequently found in initially diluted communities, probably at least partly due to these pH instabilities. Thus our 16S ribosomal amplicon pyrosequencing analyses and consequently our main findings and conclusions predominantly rely on ‘no dilution’ and ‘serial dilution’ treatments. Despite the relatively high pH deviations in ‘initial dilution’ treatments, we decided to report these results as well to present the complete experiment.

Although acidifying seawater with HCl has been extensively applied in ocean acidification studies [31,40], it is not the most accurate approach in mimicking future changes in carbonate chemistry. In the course of ocean acidification, seawater CO2 uptake will cause a reduction in pH accompanied by an increase in dissolved inorganic carbon (DIC, composed of the species H2CO3, CO3^2-), HCO3^- and CO2(aq)) but no change in total alkalinity (TA). In contrast, additions of HCl lead to a decrease in pH at constant DIC and decreasing TA [40,41]. Still both methods lead to similar results, i.e. a decrease in pH and [CO2(aq)] and an increase in [CO3^2-] and [HCO3^-] [40]. For pCO2 values not considerably exceeding 700 ppm, all carbonate system parameters change in similar magnitude with both perturbation methods [40]. Higher acid perturbations lead to larger differences however, e.g. acidification by HCl to a pH of 7.5 leads to [HCO3^-] values being about 22% lower compared to values reached by acidification to this pH by CO2 uptake [41]. Whether the small differences in carbonate system parameters between the two methods are problematic thus depends on whether the processes investigated are sensitive to small differences in [HCO3^-] [40].

Concerning bacterial communities, the ability of certain groups to compete at different pH levels may depend both on changes in pH per se and on processes involving HCO3^- availability. Changes in pH are likely to have a physiological effect, as bacteria living in the alkaline marine milieu (with pH values generally above 8.0) have to invert energy into the homeostasis of their cytoplasmic pH (7.4–7.8) [42]. Consequently, if the difference between external and internal pH becomes smaller due to ocean acidification, bacteria may profit energetically, depending on their pH homeostasis mechanisms [43]. In a study investigating two marine bacterial strains, a *Cytophaga* strain exhibited a decreased respiration rate and thus enhanced growth efficiency at higher pCO2, which was not found for a *Roseobacter* strain though [43]. In addition, both strains exhibited higher rates of heterotrophic CO2 fixation, subsequently released as DOM, at higher pCO2 levels [43], which demonstrates that [HCO3^-] plays an important role as well. With our perturbation approach using HCl, pH-
dependent changes were likely to be detected, while the consequences of rising DIC availability may have been underestimated. Yet as all carbonate system species ([CO₂], [HCO₃⁻], [CO₃²⁻], [H⁺]) will change in the same direction with both HCl and CO₂ perturbations, biological responses are unlikely to differ substantially and results should be directly comparable [31,40]. This has already been demonstrated for other organisms in ocean acidification studies systematically comparing both methods for pCO₂ values of up to 1200 ppm, namely for corals [44] and calcifying algae [45].

**Influence of pH on bacterial community structure and abundance**

We combined ARISA and extensive 16S ribosomal amplicon pyrosequencing to permit a robust statistical analysis. The strategy

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### Table: Bacterial groups in the ‘season’-'dilution' combinations, based on SIMPER analysis of pH in situ levels

<table>
<thead>
<tr>
<th>Class</th>
<th>Family</th>
<th>spring no vs. serial</th>
<th>summer no vs. serial</th>
<th>autumn no vs. serial</th>
<th>winter no vs. serial</th>
</tr>
</thead>
<tbody>
<tr>
<td>Actinobacteria</td>
<td>Microbacteriaceae</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Sphingobacteria</td>
<td>Chitinophagaceae</td>
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<tr>
<td></td>
<td>Flammovirgaceae</td>
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<tr>
<td></td>
<td>Saprospiraceae</td>
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<tr>
<td>Flavobacteria</td>
<td>Flavobacteriaceae</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>unclass. Flavobacteriales</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>unclass. Bacteroides</td>
<td>unclass. Bacteroides</td>
<td></td>
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<tr>
<td>Lentisphaeria</td>
<td>unclass. Lentisphaeria</td>
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<tr>
<td>Planctomycetacia</td>
<td>Planctomycetaceae</td>
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<tr>
<td>Alphaproteobacteria</td>
<td>Rhodobacteraceae</td>
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<tr>
<td></td>
<td>Rhodospirillaceae</td>
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<tr>
<td></td>
<td>SAR11-clade</td>
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<td></td>
<td>Sphingomonadaceae</td>
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<tr>
<td></td>
<td>unclass. Alphaproteobacteria</td>
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<tr>
<td>Betaproteobacteria</td>
<td>Comamonadaceae</td>
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<td>unclass. Burkholderiales</td>
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<tr>
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<td>Nitrosomonadaceae</td>
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<td></td>
<td>unclass. Betaproteobacteria</td>
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<tr>
<td>Epsilonproteobacteria</td>
<td>Campylobacteraceae</td>
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<td></td>
<td>Vibrionaceae</td>
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<tr>
<td></td>
<td>Alteromonadaceae</td>
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<tr>
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<td>Colwelliaceae</td>
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<td>idiomarinaceae</td>
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<td>Gammaproteobacteria</td>
<td>Pseudoalteromonadaceae</td>
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<td></td>
<td>Legionellaceae</td>
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<td>Oceanospirillaceae</td>
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<tr>
<td>unclass. Proteobacteria</td>
<td>unclass. Proteobacteria</td>
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<tr>
<td>Opitutae</td>
<td>Puniciecoccaceae</td>
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<tr>
<td>unclass. Bacteria</td>
<td>unclass. Bacteria</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>others</td>
<td></td>
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</tbody>
</table>

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**Figure 2. Bacterial groups in the ‘season’-'dilution' combinations, based on SIMPER analysis of pH in situ levels.** Displayed are phylogenetic groups jointly contributing to 90% of the total similarity within and dissimilarity between pH in situ levels of the different dilution treatments, separately for each season (no: ‘no dilution’, serial: ‘serial dilution’, initial: ‘initial dilution’). The heat map summarizes the contributions of single OTUs (16S ribosomal amplicon pyrosequencing) on the family level. Av.S/%: average percentage contribution of the i th species to the total similarity, Av.d/%: average percentage contribution of the i th species to the total dissimilarity. Bacterial families contributing less than 1% to both totals are summarized under “others”. The amount of contribution is indicated by the color of cells, darker colors represent higher contributions. doi:10.1371/journal.pone.0047035.g002
Figure 3. Bacterial groups responding differently to pH, based on SIMPER analysis. Displayed are phylogenetic groups jointly contributing to 90% of the total similarity within and dissimilarity between pH levels, separately for each ‘season’-‘dilution’ combination (no: ‘no dilution’, serial: ‘serial dilution’, initial: ‘initial dilution’). The heat map summarizes the contributions of single OTUs (16S ribosomal amplicon pyrosequencing) on the
of simultaneously pyrosequencing a large number of samples, accepting fewer sequences instead of deep sampling, was recently applied to analyze bacterioplankton communities [46] and was highly suitable for our study regarding the comparatively low bacterial diversity in laboratory incubations.

Not surprisingly, both methods revealed that the greatest influence on bacterial community composition could be attributed to season and dilution. Within the differently assembled communities, pH had a significant effect on all ‘no dilution’ and ‘serial dilution’ incubations (excluding pyrosequencing results for the ‘serial dilution’ in autumn). This indicates that even at moderate pH differences, community shifts can be observed. Furthermore, analysis of the complete data set by ARISA revealed that differences in community structure were predominantly observed already at pH 7.82, suggesting that a tipping point for bacterial community shifts may be reached even earlier. Communities from pH 7.82 and 7.67 also differed from each other (except in summer) indicating that slight differences in the degree of pH reduction are crucial. For ‘initial dilution’-communities however, we observed pH-dependent differences only in spring and summer. Possibly, in addition to the pH instabilities discussed above, the variability between replicates introduced by the 1000-fold dilution at the beginning masked pH effects in autumn and winter, where initial diversity was higher.

In agreement with community structure, richness was predominantly influenced by season and dilution. Differences between pH levels were only observed in ‘no dilution’-communities in spring and autumn, with higher richness at pH 7.67. Increased bacterial diversity at reduced pH has been reported for coastal microbial biofilms exposed to natural CO2 vents under low, but not under ambient light conditions [47] and for coral microbial communities [10], where it was assumed to be the outcome of a temporary disturbance of community equilibrium [48]. Although our results provide only minor evidence to support this hypothesis, a moderate reduction in pH seems at least not to reduce bacterial diversity, as observed for other environmental stressors such as chemical compounds [49]. Despite the good agreement concerning community structure, bacterial richness estimates were not consistent between 16S ribosomal amplicon pyrosequencing and ARISA (data not shown). Limitations of ARISA in estimating bacterial richness have previously been reported [50,51], but similarly, pyrosequencing can lead to an overestimation of richness due to sequencing errors and the presence of chimeric sequences [52]. Using a subsample of equal number of sequences for all analyses however, we at least “standardized” these error sources [53] and therefore consider pyrosequencing richness estimates comparable between samples of our study.

Concerning bacterial abundance, a reduction in pH down to 7.67 did not have an effect, which is in agreement with previous findings from mesocosm studies [14,15,54]. In this context, possible pH effects on nutrient availability and utilization should be considered. It has been reported that with decreasing seawater pH, bioavailability of dissolved iron decreases [55]. Additionally, significant changes in phosphate, silicate and ammonia have been predicted for the seawater pH reductions expected due to ocean acidification [56,57]. For hydrolysis reactions of dissolved organic matter such as organic acids, proteins and humic materials, a strong pH influence was predicted as well [58], but no detailed investigations exist. Furthermore, lower pH was shown to lead to an increased activity of total protease in mesocosms [54] and extracellular α- and β-glucosidase in laboratory experiments investigating direct pH effects [59]. In contrast, different responses of these enzymes were found after short-term incubation (3 h) at reduced pH [60]. In our experiment, the lack of pH-dependent differences in bacterial abundances suggests no change in overall nutrient availability or utilization at reduced pH.

### Bacterial groups in the ‘season’-’dilution’ combinations

The specific members of bacterial groups establishing in the dilutions differed with the initial seasonal starting communities, but the following general trends were observed: Both dilution strategies resulted in the selection of known representatives of culturable Gammaproteobacteria, which has previously been reported for initially diluted seawater incubations [23,25]. Dilution and unavoidable disruption of some cells during filtration result in more nutrients per individual cell than in the original sample, which Gammaproteobacteria exploit most efficiently [20,23,26,61]. Apparently, when an incubation is diluted only after one week (and then repeatedly), other groups of Gammaproteobacteria establish than in an initial dilution. The comparatively high diversity maintained in the undiluted treatment may in turn be explained by the lack of nutrient input, preventing the dominance of fast-growing bacteria. The aspect of grazing may also have played a role in structuring the communities, as neither prefiltration nor dilution can completely prevent the development of grazers [25]. Likewise, lysis of bacterial cells due to bacteriophages cannot be excluded. Whether viral pressure differed between the pH levels cannot be determined as direct pH effects on marine planktonic bacteriophages have not been investigated so far [62].

### Bacterial groups responding differently to pH

Various bacterial groups contributed to pH-dependent dissimilarities, predominantly different members of Gammaproteobacteria, Flavobacteriaceae, Rhodobacteraceae and Campylobacteraceae. Further discriminatory groups included Sphingobacteria, SAR11-clade, Planctomycetaceae, Bacteroidales, and bacteria that could not be assigned to families. Responses of bacterial groups were generally context-dependent, i.e. different in particular season and growth condition combinations.

Concerning Gammaproteobacteria, mostly the Oceanospirillaceae and families of the order Alteromonadas (Colwelliaceae, Alteromonadaceae, Pseudalteromonadaceae, Idiomarinaeeaceae, Shevanniaaceae and others) were affected by pH. Among these groups, Pseudalteromonadaceae were consistently found more characteristic for pH 7.67. In contrast, the response of Oceanospirillaceae and Alteromonadaceae was dependent on season, as Oceanospirillaceae were more characteristic for pH 7.67 in spring and winter, but more characteristic for pH 7.67 in summer; the opposite occurred for Alteromonadaceae. Colwelliaceae were more dependent on dilution, being more characteristic for pH 7.67 in undiluted treatments and more characteristic for pH 7.67 in diluted treatments. The dependence of the pH-response of Gammaproteobacteria on different environmental factors is supported by contrasting results obtained for two coral species transferred to high CO2 vent sites [12]. Although they occurred only in one ‘season’-’dilution’ combination and did not reach high abundances, it is interesting to note that Vibrionaceae were more...
characteristic for pH 7.67, as an increase in Vibrionaceae and other disease-associated bacteria at reduced pH has previously been reported in corals [10,11]. It remains to be elucidated whether this is a general trend, as the family contains human and animal pathogens [63], and is of increasing importance in temperate waters due to climate change [64–67].

Flavobacteriaceae were the family with overall highest contributions to pH-dependent dissimilarities, but no season or dilution specific trends were found. The Alphaproteobacterial groups Rhodobacteraceae and SAR11-clade were generally more characteristic for pH in situ. A change in the relative abundances of Flavobacteriaceae and Rhodobacteraceae in response to pH has previously been reported in biofilms from the Great Barrier Reef [13], with Rhodobacteraceae decreasing and Flavobacteriaceae increasing with decreasing pH. In contrast, in a laboratory experiment with coral microbial communities, an increase of the relative abundance of Rhodobacteraceae at low pH was reported [10]. Thus within both of these complex groups, pH responses differ.

A striking growth of Campylobacteraceae was observed in the winter ‘serial dilution’ at pH 7.67, but not at pH in situ. Also in the other seasons, Campylobacteraceae were generally more characteristic for pH 7.67. This family was only represented by members of the genus Arcobacter, which has previously been reported at the study site [26]. Members of this genus are considered emerging human pathogens, often associated with fecal contamination of waters [68,69]. Arcobacter have also been described in terrestrial and marine animals, with shellfish being a potential source of human infection [70]. It may seem surprising that this group was most important in the winter experiment, but in context of food preservation, Arcobacter have previously been reported to grow at refrigeration temperatures [70]. Regarding pH tolerance, Arcobacter spp. were reported to grow at pH 5.5 to 8.0 with a growth optimum at pH 6.0 to 7.5 [71]. Moreover, members of the Campylobacteraceae generally grow within broad pH ranges, and growth especially at low pH values was often reported regarding the survival in contaminated foods and the stomach transit [71,72,73].

Concerning the less abundant groups, Opitutae (Verrucomicrobia) are noteworthy as Verrucomicrobia have only recently been reported to be nearly ubiquitously distributed and are especially abundant in waters around Helgoland [74]. They were represented by the family Planctomycetaceae and were more characteristic for pH 7.67 in spring and autumn. Similarly in corals, members of Verrucomicrobia were only present at reduced pH [10].

In summary, no consistent trend throughout all ‘season’-‘dilution’ combinations was found for the majority of bacterial groups. Notable exceptions (neglecting groups that were only present in one ‘season’-‘dilution’ combination) were Pseudoalteromonadaceae and Opitutae, which were always more characteristic for pH 7.67, suggesting that they may profit from pH reductions. The same tendency was found for Arcobacter. In contrast, groups that were generally more characteristic for pH in situ were Rhodobacteraceae and SAR11-clade, hinting at difficulties of these groups to cope with reductions in pH.

Regarding the lack of additional trends, the following aspects have to be considered: First, bacterial families such as the Rhodobacteraceae are highly diverse and different members occur at certain time points, while being of very low importance in other seasons [27,75]. Thus, supposing that different responses to pH occur at the genus, species, or even ecotype level, an analysis on the family level can only depict very rough patterns. Second, different environmental factors are likely to influence the response of bacterial groups to pH. In our experiment, a dependence on season could - besides the identity aspect mentioned above - hint at temperature effects, while a dependence on dilution could be associated with nutrient effects. Additionally, the co-occurring species must be taken into account. The high contribution to total similarity of Arcobacter at pH 7.67 in the winter ‘serial dilution’-community for instance, may in turn have influenced the comparatively lower contribution of Oceanospirillaceae at this pH and vice versa.

In conclusion already moderate changes in pH have the potential to cause shifts in bacterial communities, depending on the pool of bacterial groups constituting this community and on the prevailing environmental conditions. We identified members of the Gammaproteobacteria, Flavobacteriaceae, Rhodobacteraceae and Campylobacteraceae as susceptible to changes in pH and confirmed some results obtained in previous studies. To ultimately draw conclusions about functional implications, the question remains whether a change in species composition observed in in vitro incubations starting with an abrupt change in pH is extendable to natural bacterioplankton communities and how adaptation will affect long-term consequences. Therefore, to test whether the patterns observed also exist in more complex environments, we suggest that the groups identified in this study deserve special attention in future long-term and mesocosm experiments.

**Supporting Information**

**Figure S1** Bacterial abundance in the different treatments. No significant pH effect on bacterial abundance was found (ANOVA).

**Figure S2** Influence of the factor ‘pH’ on bacterial community composition (ARISA). Displayed are principal co-ordinate analysis plots (PCOs) for each ‘season’-‘dilution’ combination based on Jaccard dissimilarities of ARISA profiles. Symbol shape represents the pH level (circles: pH in situ, squares: pH 7.82, triangles: pH 7.67).

**Figure S3** Influence of the factor ‘pH’ on bacterial community composition (16S ribosomal amplicon pyrosequencing). Displayed are principal co-ordinate analysis plots (PCOs) for each ‘season’-‘dilution’ combination based on Bray-Curtis dissimilarities of OTUs (16S ribosomal amplicon pyrosequencing). Symbol shape represents the pH level (circles: pH in situ, triangles: pH 7.67).

**Figure S4** Bacterial richness in the different treatments. Richness was determined based on the number of OTUs (16S ribosomal amplicon pyrosequencing) in the standardized subsample (n = 494 sequences) and was compared by ANOVA, with Tukey’s HSD test for post-hoc comparisons. Asterisks (*) represent significant differences (p<0.05) between pH levels within a ‘season’-‘dilution’ combination.

**Table S1** pH after the four weeks of incubation.

**Table S2** Sample coverage obtained by 16S ribosomal amplicon pyrosequencing, based on the standardized subsample (n = 494 sequences).

**Table S3** Results of the SIMPER analysis giving the similarities within ‘season’-‘dilution’-‘pH’ combinations. Displayed are the OTUs (16S ribosomal amplicon pyrosequencing) that predominantly contributed to 90% of to
the total similarity. \(\text{Av.A}_i\): average abundance of the \(i\) th species over all samples of the treatment, \(\text{Av.S}_i\): average contribution of the \(i\) th species to the total similarity, \(\Sigma\text{Av.S}_i/\text{SD}\): the average value of the \(i\) th species as a typifying species, \(\text{Av.S}_i\%\): average cumulative contribution to the total similarity.

\(\text{PDF}\) Table 54 Results of the SIMPER analysis giving the dissimilarities between pH in situ levels of the different dilution treatments, separately for each season. Displayed are the OTUs (16S ribosomal amplicon pyrosequencing) that predominantly contributed to 90% of to the total dissimilarity. 

\(\text{Av.A}_i\): average abundance of the \(i\) th species over all samples of the treatment (no: ‘no dilution’, serial: ‘serial dilution’, initial: ‘initial dilution’), \(\text{Av.S}_i/\text{SD}\): the average contribution of the \(i\) th species to the total dissimilarity, \(\Sigma\text{Av.S}_i/\text{SD}\): the average contribution of the \(i\) th species as a typifying species, \(\text{Av.S}_i\%\): average percentage contribution of the \(i\) th species to the total similarity, \(\Sigma\text{Av.S}_i\%\): average cumulative contribution to the total dissimilarity.

\(\text{PDF}\) Table 55 Results of the SIMPER analysis giving the dissimilarities between the pH levels in situ and 7.67 of ‘season’-dilution’ combinations significantly influenced by ‘pH’ according to PERMANOVA. Displayed are the OTUs (16S ribosomal amplicon pyrosequencing) that predominantly contributed to 90% of to the total dissimilarity. 

\(\Sigma\text{Av.S}_i/\text{SD}\): the average value of the \(i\) th species as a discriminating species, \(\text{Av.S}_i\%\): average percentage contribution of the \(i\) th species to the total dissimilarity, \(\Sigma\text{Av.S}_i\%\): average cumulative contribution to the total dissimilarity.

\(\text{PDF}\) References


