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Notes:
Matching phylogeny and metabolism in the uncultured marine bacteria, one cell at a time

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The identification of predominant microbial taxa with specific metabolic capabilities remains one of the biggest challenges in environmental microbiology, because of the limits of current metagenomic and cell culturing methods. We report results from the direct analysis of multiple genes in individual marine bacteria cells, demonstrating the potential for high-throughput metabolic assignment of yet-uncultured taxa. The protocol uses high-speed fluorescence-activated cell sorting, whole-genome multiple displacement amplification (MDA), and subsequent PCR screening. A pilot library of 11 single amplified genomes (SAGs) was constructed from Gulf of Maine bacterioplankton as proof of concept. The library consisted of five flavobacteria, one sphaerobacterium, four alphaproteobacteria, and one gammaproteobacterium. Most of the SAGs, apart from alphaproteobacteria, were phylogenetically distant from existing isolates, with 88–97% identity in the 16S rRNA gene sequence. Thus, single-cell MDA provided access to the genomic material of numerically dominant but yet-uncultured taxonomic groups. Two of five flavobacteria in the SAG library contained proteorhodopsin genes, suggesting that flavobacteria are among the major carriers of this photometabolic system. The purM and nasA genes were detected in some 100-cell MDA products but not in SAGs, demonstrating that organisms containing bacteriochlorophyll and assimilative nitrate reductase constituted <1% of the sampled bacterioplankton. Compared with metagenomics, the power of our approach lies in the ability to detect metabolic genes in uncultured microorganisms directly, even when the metabolic and phylogenetic markers are located far apart on the chromosome.

Flavobacteria | flow cytometry | multiple displacement amplification | proteorhodopsin | single-cell genomics

The PCR- and direct cloning-based sequencing of environmental DNA extracts has revealed the enormous previously unknown phylogenetic and metabolic diversity of prokaryotes (1–9). Although yet-uncultured taxa are believed to comprise >99% of all prokaryotes, their metabolic capabilities and ecological functions remain enigmatic, largely because of methodological limitations. For example, PCR-based clone libraries are intrinsically limited to the analysis of one gene at a time, with no direct way of linking libraries of diverse genes. Large-scale environmental shotgun sequencing, although extremely productive for finding novel genes, is prohibitively expensive, and so far is limited to only partial genome assembly of the most numerically dominant taxa in complex marine microbial communities (5, 9). Genomic analyses of large environmental DNA inserts can lead to remarkable discoveries, such as proteorhodopsin genes in bacteria (2). However, large insert-based function assignment is limited to situations where the metabolic gene of interest is located near phylogenetic markers (e.g., ribosomal genes). Thus, currently available culture-independent research tools are poorly suited for identification of microorganisms with specific metabolic characteristics. This significantly limits the progress in such diverse fields as biogeochemistry, microbial ecology and evolution, and bioprospecting.

We propose sequencing multiple DNA loci in individual bacterial cells rather than environmental DNA extracts, as a more productive alternative for metabolic mapping of uncultured microorganisms. This strategy has been gaining momentum, with recent implementations of single-cell multiplex PCR in termite gut microbiota by Ottesen et al. (10) and partial genome sequencing of single cells of Prochlorococcus by Zhang et al. (11). Here we show important improvements in single-cell separation and DNA analysis protocols and demonstrate a proof-of-concept metabolic mapping of taxonomically diverse marine bacterioplankton.

One of the main challenges for single-cell studies is efficient and contamination-free separation of individual cells from other microorganisms and extracellular DNA in an environmental sample. Prior studies of DNA in individual prokaryote cells used serial sample dilution (11), dilution using microfluidics (10), or micromanipulation (12). Here we used high-speed droplet-based FACS. Compared with alternative methods, FACS offers several critical advantages, including high-throughput rates and the ability to sort targeted plankton groups, based on cell size and fluorescence signals of natural cell components and fluorochromes (13). Furthermore, cell separation by FACS creates microsamples containing the target cell and only 3–10 pl of sample around it (13). This reduces the codeposition of extracellular DNA, which in marine waters occurs at concentrations similar to cell-bound DNA (14, 15).

Current sequencing technologies require nanogram-to-microgram DNA templates and are not capable of direct sequencing of individual DNA molecules. Thus, DNA preamplification is necessary to sequence genomes or genomes from individual cells. For the analysis of up to two loci per cell, single cell multiplex PCR has been used in medical research since the 1980s (16) and was recently used in an environmental microbiology study (10). As a more versatile alternative, allowing for analysis of an unlimited number of loci, several methods have been suggested for whole-genome amplification, including degenerated oligonucleotide-primed PCR, primer extension preamplification, ligation-mediated PCR, and multiple displacement amplification (MDA) using phi29 or Bst DNA polymerases (17, 18). Among them, Phi29-based MDA appears the most suitable for efficient whole-genome amplification with low error and bias (17, 18) and is capable of generating micrograms of genomic DNA from nanogram-sized samples (19–21). Recently, Phi29-based MDA was used on single human (18, 22, 23), Escherichia coli (24), and Prochlorococcus (11) cells. Direct

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Abbreviations: MDA, multiple displacement amplification; SAG, single amplified genome; SSU rRNA, small-subunit rRNA; HNA, high nucleic acid; T-RFLP, terminal restriction fragment length polymorphism.

Data deposition: The sequences reported in this paper have been deposited in the GenBank database (accession nos. EF202334–EF202347 and EF508145–EF508148).

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cloning and whole-genome sequencing of single-cell MDA products remains technically challenging because of template mismatch, amplification biases, self-primed DNA synthesis, and the multibranched nature of MDA products (11, 25). However, downstream PCR appears insensitive to these MDA artifacts (24), is simple and inexpensive, and can be effectively used in the analysis of multiple genes.

The aim of this study was a proof-of-concept metabolic mapping of marine bacterioplankton. We developed procedures for clean single-cell separation by FACS, cell lysis, and subsequent single-cell whole-genome MDA for downstream PCR-based analyses of multiple loci. We successfully constructed a pilot library of single amplified genomes (SAGs) from a temperate coastal marine site. We analyzed them for the presence and DNA sequences of genes representing phylogenetic markers [small-subunit rRNA (SSU rRNA)] and several significant biogeochemical functions in marine ecosystems (proteorhodopsin, bacteriochlorophyll, nitrogenase, and assimilatory nitrate reductase). Bacterial proteorhodopsins (2) and bacteriochlorophylls (26) are photometabolic systems, recently recognized for their ubiquity and likely significance in the global carbon and energy fluxes. Nitrogenase is a key enzyme in the fixation of N₂, effectively controlling primary production in vast areas of the ocean, and appears to be possessed by some heterotrophic bacterioplankton (27). Assimilatory nitrate reductase enables some heterotrophic bacteria to use nitrate and in this way compete with phytoplankton for the upwelled nitrogen (28). So far, little is known about the taxonomic composition of microorganisms carrying these genes in marine environments.

### Results and Discussion

**Taxonomic Composition of the SAG Library.** The SSU rRNA gene was successfully PCR-amplified and sequenced from 12 of 48 single-cell MDA reactions (Table 1). The SAG MS024–2C was identified as a contaminant and excluded from further analyses [see supporting information (SI) Text]. The remaining SAG library consisted of five flavobacteria, one spingobacterium, four alphaproteobacteria of the *Roseobacter* lineage, and one gammaproteobacterium, all most closely related to marine isolates and clones. Diverse representatives of the *Roseobacter* lineage are readily isolated and are relatively well studied (29, 30). Accordingly, SSU rRNA genes of the four alphaproteobacterial SAGs were 99% identical to existing isolates. In contrast, all flavobacterial, spingobacterial, and gammaproteobacterial SAGs were phylogenetically distant from established cultures, with 88–97% identities in the SSU rRNA gene. Flavobacteria as a group are proficient degraders of complex biopolymers, including cellulose, chitin, and pectin (31). Thus, certain Flavobacteria taxa may play important and specialized roles in microbial food webs and may be attractive for bioprospecting. Single-cell MDA provided access to the unique genomic material of these yet-uncultured taxa at the individual organism level. Different from single-cell multiplex PCR (10, 16), which enables analysis of up to two loci per cell, our approach generated a large quantity of high molecular-weight whole-genome amplification products. This material can be used in a virtually unlimited number of downstream PCRs (see below) and hybridization analyses and may be suitable for genomic sequencing (11).
Only high nucleic acid (HNA) bacterioplankton was analyzed in this study, which comprised 57% of all heterotrophic heterotrophic bacterioplankton, and comprised 9% of all heterotrophic bacterioplankton in this study. However, the predominance of Bacteroidetes in the SAG library was unexpected. Alphaproteobacteria typically dominate SSU rRNA sequence libraries of marine surface bacterioplankton, whereas Bacteroidetes constitute ~3% of all marine clones (33). In contrast, studies employing fluorescent in situ hybridization (34, 35), quantitative PCR (36), and metagenomics (37, 8, 9) suggest a higher proportion of Bacteroidetes (particularly Flavobacteria), in some cases ~70% of the total bacterioplankton (31). This contradiction may be caused by PCR and/or cloning biases against Flavobacteria (31). Interestingly, the ratio of Alphaproteobacteria vs. Bacteroidetes in our SAG library was 0.7 (Table 1), whereas the corresponding ratio of community terminal restriction fragment length polymorphism (T-RFLP) analyses peak areas at 55 bp (assumed Roseobacter lineage) and at 90–96 bp (assumed Bacteroidetes) was 1.0 (100 cell MDA-PCR) and 3.5 (1,000-cell seminested PCR) (SI Fig. 5). This discrepancy is supportive of a PCR bias against Bacteroidetes, which would affect T-RFLP profiles, especially those based on two rounds of seminested PCR. On the other hand, the construction of the SAG library was insensitive to PCR biases and did not involve cloning. Thus, scaled-up SAG libraries may become an ultimate tool for quantitative bacterioplankton analyses at high phylogenetic resolution. The advantage of SAG screening over fluorescent in situ hybridization, another taxon-specific quantification method, is demonstrated by the extraction of high-resolution phylogenetic information through sequencing of the entire SSU rRNA gene, as well as protein-encoding loci (see below).

Proteorhodopsin genes were detected by PCR and confirmed by sequence analysis in 2 of 11 SAGs (Fig. 1). In addition, PCR-screening detected proteorhodopsin genes in all 12 100-cell MDA reactions. Accordingly, Sabehi et al. (37) estimate that 13% bacterioplankton in the photic zone of the Mediterranean Sea and the Red Sea carried proteorhodopsin genes. Our study provides further evidence that proteorhodopsin-containing microorganisms comprise a significant fraction of marine bacterioplankton. Interestingly, both proteorhodopsin-positive SAGs were Flavobacteria, providing evidence that proteorhodopsins are common in the numerically abundant representatives of this taxonomic group (Fig. 1). The presence and photometabolic functionality of proteorhodopsins in Flavobacteria were recently confirmed by genome sequencing of four isolates (38). The first indication of proteorhodopsins in Flavobacteria was obtained from shotgun sequencing of Sargasso Sea microbes, where a proteorhodopsin gene was found on a scaffold also containing a DNA-directed RNA polymerase sigma subunit (rpoD) typical of Bacteroidetes (5). Bacteria with proteorhodopsins were first discovered by screening environmental BAC libraries (2). Using this technique, several Gammaproteobacteria, Alphaproteobacteria, and Euryarchaea were identified as proteorhodopsin hosts (39, 40). However, this approach has so far not indicated the presence of proteorhodopsins in Flavobacteria, possibly because the proteorhodopsin and SSU rRNA genes are too far apart. Studies based on community proteomics (41), community PCR (42, 43), community shotgun sequencing (5, 9), and PCR screening of metagenomic BAC libraries (39) demonstrated high diversity of proteorhodopsins in the ocean, although the vast majority of their hosts remain unknown. So far, only five marine isolates have been reported to contain proteorhodopsin genes, including alphaproteobacterium Pelagibacter ubique (44) and four Flavobacteria (38). Here we demonstrate how single-cell MDA-PCR can provide a powerful and relatively inexpensive tool for the phylogenetic mapping of this biogeochemically important gene, independent of the gene’s position on the chromosome or host cultivability.

The two SAG proteorhodopsin were most closely related (up to 71% identity) to proteorhodopsins from four Flavobacteria isolates and to a group of environmental clones from the North Atlantic (Fig. 1). Consistent with the flavobacterial isolates and near-surface environmental sequences, both SAGs had methionine at amino acid position 105 (eBAC31A08 numbering), indicative of absorption maxima near 530 nm (green light) (38). In general, phylogenetic relationships among proteorhodopsins and SSU rRNA genes mirrored each other, providing no evidence for recent cross-taxa horizontal transfer events like those observed in Archaea (40). On the other hand, the presence of proteorhodopsin genes was inconsistent among some closely related Flavobacteria, e.g., Polaribacter filamentus 215 and P. turgessii 23-P (Fig. 1), suggesting recent proteorhodopsin gene losses. Interestingly, proteorhodopsin genes closely related to Flavobacterial SAGs and isolates were present among environmental clones from the North Atlantic, Mediterranean Sea, and Red Sea, indicating that Flavobacteria may be major carriers of proteorhodopsin genes in diverse marine environments.

**Other Genes.** The pufM and nusA were not detected in any of the single-cell MDA products. However, they were present in six
(puF) and three (nasA) 100-cell MDA reactions (of a total of 12), indicating that <1% of bacterioplankton in the sample carried either of these genes. Accordingly, bacteriochlorophyll was previously found to be expressed (infrared fluorescence) in \( \sim 1\% \) of bacteria in coastal Maine waters at this time of year (45). All six puF were 100% identical to each other and were most closely related to bacteriochlorophylls from the Roseobacter lineage (Fig. 2A). Thus, it appears that a single Roseobacter taxon dominated bacteriochlorophyll-containing bacterioplankton in the studied sample. Two nasA were most closely related to assimilatory nitrate reductases in Roseobacter lineage, whereas one nasA was most closely related to marine Gammaproteobacteria (Fig. 2B). The pilot SAG library failed to unambiguously identify these relatively rare but biogeochemically important microorganisms. Screening of a larger SAG library would be an ideal tool for this task. Alternative community genomics-based analyses have proven less effective to match SSU rRNA, and functional genes in such rare taxa.

Genes encoding archaeal SSU rDNA and nitrogenases were not detected in any of the sorted wells, suggesting that Archaea and nitrogen-fixing organisms were extremely rare or absent in the analyzed heterotrophic HNA bacterioplankton. Eukaryote SSU rRNA genes were also not detected, confirming effective separation of prokaryotes from protists by FACS.

**Conclusions**

We demonstrate how a combination of single-cell FACS, MDA, and PCR can be used in metabolic mapping of taxonomically diverse uncultured marine bacterioplankton. Large quantities of high molecular-weight whole-genome amplification products were obtained from individual cells, allowing for a virtually unlimited number of downstream analyses. In this proof-of-concept study, we detected protophytes genes in two of five flavobacteria, providing evidence that Flavobacteria are major carriers of this photometabolic gene. We also determined that Flavobacteria were a major component of HNA bacterioplankton in the analyzed coastal sample. Fewer than 1% of the analyzed cells carried nasA, puF, and nhF.

We used standard configuration flow cytometry instrumentation that is available on most major research campuses and increasingly used aboard oceanographic research vessels. Working at the single-cell level requires especially stringent instrument cleaning, sample handling, and quality-control methods to prevent DNA contamination. We show that our methods were able to achieve sufficiently low DNA blank controls (see SI Text). The cost of MDA and subsequent PCR sequencing is on the order of tens of U.S. dollars per cell and thus is significantly less expensive than metagenomic sequencing. In addition to high-throughput screening by PCR or hybridization, SAG libraries may provide material for genomic sequencing of selected uncultured microorganisms. Two of our SAGs are currently in the process of whole-genome sequencing.

**Materials and Methods**

**Sample Collection and Single-Cell Sorting.** The coastal water sample was collected from Boothbay Harbor, Maine, from 1-m depth at the Bigelow Laboratory dock (43°50’40”, 69°38’27”W) on March 28 at 9:45 a.m. during high tide (water temperature 7.0°C). The unmanipulated sample was 10-fold diluted with filtered (0.2 μm pore size) sample water and stained with 5 μM (final concentration) SYTO-13 nucleic acid stain (Invitrogen, Carlsbad, CA) for prokaryote detection as in delGiorgio et al. (46). Individual bacterioplankton were sorted into 96-well plates containing 5 μl per well of PBS. Only HNA cells were sorted to reduce the probability of depositing dead cells with partially degraded genomes. Single cells were sorted into four of the eight rows on each plate. Of the remaining rows, two were dedicated to background controls, consisting of single drops generated from a sort gate drawn in the “noise” area in the lower left corner of the side scatter/green fluorescence plot. One row of 12 wells was dedicated to blanks with no drop deposition, and one row received 100 HNA bacterioplankton cells per well. Sorting was done with a MoFlo (Dako Cytomation, Carpentry, CA) flow cytometer equipped with the Cylcone robotic arm for sorting into plates, using a 488-nm argon laser and a 70-μm nozzle orifice. The cytometer was triggered on side scatter, the sort gate was based on side scatter and SYTO-13 fluorescence, and the “purify 0.5” sort mode was used for maximal sort purity. Extreme care was taken to prevent sample contamination by any non-target DNA. New sheath fluid lines were installed before each sort day. Sheath fluid and sample lines were cleaned by a succession of warm water, 5% bleach solution, and an overnight flush with DNA-free deionized water. Sheath fluid was prepared by dissolving combusted (2 h at 450°C) NaCl in DNA-free deionized water for a final concentration of 1%. Sorted plates were stored at \(-80\)°C until MDA.

**Lysis and MDA.** We compared three protocols for cell lysis, DNA denaturing, and MDA in this study.
Protocol A. Three cycles of heating to 97°C and cooling to 8°C were used for cell lysis and DNA denaturing, after which 18-h MDA was performed by using REPLI-g Mini (Qiagen, Chatsworth, CA) Phi29 polymerase and reaction buffer. For each well containing 5 μL of PBS, we used 0.5 μL of polymerase, 14.5 μL of buffer, and 5 μL of DNA-free deionized water.

Protocol B. Alkaline lysis on ice and 18-h MDA were performed by using REPLI-g Mini (Qiagen) kit reagents and following the manufacturer’s protocol for blood samples.

Protocol C. As protocol B, except that REPLI-g Midi kit (Qiagen) was used, and PicoGreen DNA stain (Invitrogen) was added to the reaction at 0.5× (final concentration). DNA synthesis was monitored with IQ5 real-time PCR system (Bio-Rad, Hercules, CA). Duplicate standards containing 0.05, 5, 500, and 50,000 fg of human genomic DNA (Promega, Madison, WI) were amplified simultaneously with the sort samples.

Initially, each of the three protocols were applied on 24 wells: 12 with single cells, 3 no-drop controls, 6 background controls, and 3 with 100 cells. Protocols A, B, and C were used after 7, 8, and 94 days of sorted cell storage at −80°C, respectively. An additional 24 wells were analyzed by using protocol B after 350 days of cell storage.

The DNA concentration in MDA reactions was determined by using a ND-1000 spectrophotometer (Nanodrop) after a cleanup with MinElute PCR Purification Kit (Qiagen).

PCR-Based Analyses of MDA Products. The MDA products were diluted 10-fold (protocols A and B; REPLI-g Mini kit products) or 200-fold (protocol C; REPLI-g Midi kit products). Two microliters of the dilute products served as templates in 25 μL of PCR. Previously described primers and PCR conditions were used to amplify genes encoding bacterial, archaeal, and eukaryal SSU rRNA, proteorhodopsin, bacteriochlorophyll, nitrogenase, and assimilative nitrate reductase (SI Table 2). The PCR products were cleaned with QiAquick (Qiagen). For the T-RFLP of bacterial SSU rRNA genes, PCR amplicons obtained with 27F–FAM and 907R primers were digested with either HhaI or BsuRI (HaeIII) restriction endonucleases (Fermentas, Hanover, MD). Sequencing and fragment analyses were performed with a 3730xl analyzer (Applied Biosystems, Foster City, CA) at the W. M. Keck Center for Comparative and Functional Genomics, University of Illinois at Urbana—Champaign, IL. For T-TRLF P, GeneScan 1000 ROX Size Standard (Applied Biosystems) was used.

Taxonomic identification of SAGs was achieved by SSU rRNA gene analysis with GenBank BLASTN (47) and Ribosomal Database Project (RDP) Classifier and Seqmatch search tools (48). The SSU rRNA sequences were checked for chimeras by using the RDP Chimera Check tool. Protein-encoding sequences were translated with National Center for Biotechnology Information ORF Finder and their identities verified by GenBank BLASTP searches for closest relatives. Evolutionary trees were constructed by using PHYLIB (49) after an automatic sequence alignment with ClustalX (50).

T-RFLP Profiling of Bacterioplankton Communities. Triplicate 1.000 cell aliquots of HNA bacterioplankton were sorted as above into microcentrifuge tubes preloaded with 5 μL of lysine-N-Go (Pierce, Rockford, IL) and then stored at −80°C. Cell lysis was performed according to lysine-N-Go instructions. Entire lysate volumes were used as templates in 50–μL 30-cycle PCRs by using primers 27F and 1492R (SI Table 2). Two-microliter aliquots of these PCR products served as templates in a second semi nested 25–μL and 30-cycle PCR with primers 27F–FAM and 907R. PCR products were cleaned and digested, and fragment analyses were performed as above.

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Supporting Information

**Fig. 3.** Gel electrophoresis of multiple displacement amplification (MDA) products obtained using Protocol C: molecular weight ladder (lanes A and H), no-drop controls (lanes B-C), single bacterioplankton cells (lanes D-E), and 100 bacterioplankton cells (lanes F-G). The 0.75% TAE agarose gel was loaded with 3 ml of high molecular weight ladder (Invitrogen) and 10x diluted products of MDA reactions and then electroporated at 0.5 V/cm for 5 h.
Fig. 4. Real-time monitoring of multiple displacement amplification reactions of (A) standards prepared from human genomic DNA and (B) flow-cytometrically sorted material. Inset in A shows the log-linear standard curve for DNA concentration.
Fig. 5. T-RFLP profiles of bacterial SSU rRNA genes obtained by (A) single-cell MDA-PCR, (B) 100-cell MDA-PCR, and (C) 1,000-cell seminested PCR. Panel A is a composite of six single-cell profiles. Insets show discrimination of the peaks near 90 bp with an expanded scale. All profiles were obtained using 27F-FAM and 907R primers and Hhal restriction endonuclease.

Table 2. PCR primers

<table>
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<th>Primers</th>
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<td>Assimilatory nitrate reductase, <em>asA</em></td>
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**SI Text**

**Quality control of single-cell sorting and whole-genome amplification**

**Multiple displacement amplification.** Approximately 0.5 and 5 mg of genomic DNA was synthesized in single cell REPLI-g Mini (protocols A and B) and REPLI-g Midi (protocol C) reactions, respectively, with the apparent dominant product size >10 kbp (SI Fig. 3). Most MDA reactions, including all but two blank treatments, resulted in DNA synthesis. The synthesis of DNA in MDA negative controls has been observed before and is likely caused by Phi29 self-priming, DNA contamination, or both (1-3). Real-time monitoring of MDA demonstrated that reaction speed depended upon template amount when template was above 5 fg DNA (SI Fig. 4A). The template-dependent dynamics of MDA observed in this study suggested that 6-8 h were necessary to complete MDA of fg-level templates using REPLI-g Midi kit (QIAGEN). This agrees with a similar analysis by Zhang et al. (2) but contradicts Spits et al. (4), who suggested 2-h reactions for MDA of single-cell genomes.

A log-linear standard curve was produced from the 5-fg, 500-fg, and 50-pg DNA treatments (SI Fig. 4A Inset), and used to calculate tentative DNA concentrations in the sorted samples. The average amount of DNA in the no-drop controls, background controls, single-cell, and 100-cell treatments were estimated to be 0.07 (range 0.0-0.2), 0.78 (0.0-1.8), 2.0 (0.0-10.3) fg (SI Fig. 4B). The average for single cells was 4.8 fg, if wells with virtually no amplification (<0.3-fg DNA content estimate) were excluded from the calculation. The estimated amount of DNA in the single- and 100-cell treatments was in agreement with published marine bacterioplankton genome size estimates based on fluorometry (2.5 fg) (5), flow cytometry (1.5 fg) (6), and whole-genome sequencing data (7, 8). The low-DNA content in no-drop controls (mean, 0.07 fg) suggests that sample contamination from handling and reagents was below 4% of an average single-cell genome. The background "noise" controls were sorted drops from the low side scatter and fluorescence region. They showed significantly higher DNA (mean, 0.78 fg)
than the no-drop controls. This DNA could have originated from large viruses, DNA debris, and small (low nucleic acid) bacteria. No DNA was detected in sheath fluid collected at the end of instrument lines. Thus, our sample handling stringency was adequate to prevent significant sample contamination with extraneous genomic DNA.

**PCR of the bacterial SSU rRNA gene.** Initially, 12 wells with single cells were subjected to each of the lysis-MDA protocols A, B, and C. Using MDA products as templates in PCR, SSU rRNA genes were successfully amplified from one, six, and two of the wells, respectively (Table 1). Thus, protocol B (cold alkaline lysis and MDA using REPLI-g Mini kit) resulted in the highest success rate (50%) of MDA-PCR. According to manufacturer's statements, we expected similar cell lysis and MDA success rate with REPLI-g Mini and Midi kits, with the Midi kit producing higher DNA yields per sample. The lower success rate of protocol C may be in part explained by possible DNA degradation during prolonged sample storage (94 vs. 8 days). After a 350-day storage, the application of protocol B on 12 additional single-cell wells resulted in a 25% success rate. Diverse factors may have contributed to the less than 100% success rate in single-cell MDA-PCR, including failed deposition of some cells during FACS, DNA degradation during post-FACS storage, incomplete cell lysis, incomplete MDA, and mismatches to some bacterioplankton groups in the "universal" PCR primers used in this study. It is noteworthy that the 25-50% success rate in MDA-PCR achieved with protocol B was similar to the single cell PCR success rate achieved by Ottesen et al. (9) and to FISH success rate using "universal" SSU rRNA probes (10).

The SSU rRNA PCR products were obtained from all 12 100-cell wells and from one background control (of 24 total). No PCR products were obtained from the 12 no-drop controls. High-quality nonchimerical sequences of near-complete SSU rRNA genes were obtained from all single-cell MDA-PCR products. The SSU rRNA genes of 11 of the 12 SAGs were most closely related to marine isolates and clones (Table 1). One exception, MS024-2C, was identified as *Delftia acidovorans*, characteristic for soils, freshwaters, and anthropogenic environments. An identical sequence was also retrieved from one of the background controls. This suggests that the sequence is likely a contaminant originating from handling or the reagents. Thus, the overall rate of apparent contamination with bacterial SSU rRNA gene was two of 84 wells (2%) of single cells, background controls, and no-drop controls combined.

For additional quality control, T-RFLP profiles were generated for the SSU rRNA gene of each PCR-positive single cell and control MDA reaction, using two alternative restriction endonucleases. All profiles contained single peaks (Table 1, SI Fig. 5A), further confirming that only one cell type was deposited and amplified in each well with no apparent contamination with DNA from other bacterial taxa. It is possible that in some cases aggregates of genetically identical cells, which were optically similar to single cells (e.g., partially divided cells), were sorted into the same well. However, this would not have any adverse effect on the downstream molecular analyses or interpretation.

The HhaI-based T-RFLP profiles generated from 100-cell MDA reactions were dominated by a peak at 55 bp and a group of peaks at around 90 bp (SI Fig. 5B). Similar T-RFLP profiles were also obtained from 1,000 cell aliquots lysed with Lyse-N-Go protocol and amplified using seminested PCR (SI Fig. 5C). This suggests that the diverse cell lysis and DNA amplification methods used in this study were targeting the same bacterioplankton taxa. The T-RFs found in 100- and 1,000-cell profiles corresponded to those found in SAG profiles of Alphaproteobacteria (55 bp) and Bacteroidetes (around 90 bp; SI Fig. 5A). The 203-bp *Delftia* fragment was absent in community profiles, further suggesting its origin as a sample handling contaminant. Conspicuously, the 575-bp T-RF, characteristic to the marine gammaproteobacterium MS024-3A, was also absent in community profiles, either due to PCR biases or rarity of the taxon. On the other hand, 92- and 864-bp peaks, present in all 100- and 1,000-cell community profiles, were not represented in the SAG library. Considering the relatively small size of the pilot SAG library and the fact that all but two (MS024-1C and MS190-2A) SAG SSU rRNA gene sequences were unique, it is clear that this library covered only a fraction of biodiversity in the bacterioplankton community.


