Trajectories and Drivers of Genome Evolution in Surface-Associated Marine *Phaeobacter*

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Accepted: November 27, 2017

Data deposition: This project has been deposited at GenBank under the accessions CP010588 - CP010775, CP010784 - CP010791, CP010805 - CP010810, KY357362 - KY357447.

Abstract

The extent of genome divergence and the evolutionary events leading to speciation of marine bacteria have mostly been studied for (locally) abundant, free-living groups. The genus *Phaeobacter* is found on different marine surfaces, seems to occupy geographically disjunct habitats, and is involved in different biotic interactions, and was therefore targeted in the present study. The analysis of the chromosomes of 32 closely related but geographically spread *Phaeobacter* strains revealed an exceptionally large, highly syntenic core genome. The flexible gene pool is constantly but slightly expanding across all *Phaeobacter* lineages. The horizontally transferred genes mostly originated from bacteria of the *Roseobacter* group and horizontal transfer most likely was mediated by gene transfer agents. No evidence for geographic isolation and habitat specificity of the different phylogenomic *Phaeobacter* clades was detected based on the sources of isolation. In contrast, the functional gene repertoire and physiological traits of different phylogenomic *Phaeobacter* clades were sufficiently distinct to suggest an adaptation to an associated lifestyle with algae, to additional nutrient sources, or toxic heavy metals. Our study reveals that the evolutionary trajectories of surface-associated marine bacteria can differ significantly from free-living marine bacteria or marine generalists.

Key words: population genomics, gene flow, gene transfer agent, horizontal gene transfer, ecological niche, bacterial adaptation.

Introduction

The marine environment sustains a high diversity of bacteria (Zinger et al. 2011; Sunagawa et al. 2015). So far, the genetic divergence of evolutionary and ecologically distinct but cooccurring marine microbial populations and the underlying drivers have been elucidated for only few, mostly free-living, bacterial groups (Coleman et al. 2006; Johnson et al. 2006; Hunt et al. 2008; Carlson et al. 2009; Swan et al. 2013). Yet, a considerable fraction of marine bacteria occur, grow and evolved attached to surfaces of particles and organisms, where they can constitute up to 20% of the total bacterial biomass in ocean water (Azam et al. 1983) and up to 66% of bacterial biomass during coastal algae blooms (Becquevort et al. 1998). In addition, the contribution of attached bacteria

to overall prokaryotic substrate turnover is disproportionally high due to their higher specific activity (Crump et al. 1998; Stocker 2012) and the species diversity of associated bacteria significantly exceeds that of free-living bacteria (Bižic-lonescu et al. 2015). Compared with free-living bacteria, the high cell density and proximity in surface biofilms may facilitate an increased gene transfer and spread of traits (Balcazar et al. 2015). Despite these particular features of surface-associated marine bacteria, only few studies have actually addressed their genome diversity and evolution. Previous studies focused on the genus *Vibrio* which, however, is a bacterial generalist not specifically adapted to the attached lifestyle (Hunt et al. 2008; Shapiro et al. 2012; Kirchberger et al.

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2016), as opposed to most other species detected on marine surfaces (Bižic-lonescu et al. 2015).

The Roseobacter group is adapted to the marine environment (Simon et al. 2017) and comprises chemotrophic bacteria often associate with eukaryotes (Buchan et al. 2005). Members of the group have a much higher metabolic and ecological versatility than other dominant marine bacteria (Brinkhoff et al. 2008; Newton et al. 2010). The genus Phaeobacter colonizes artificial and biotic surfaces such as different algae, bryozoan, molluscs, crustaceans, and fish, and is often associated with aquaculture systems and harbors (Rao et al. 2005; Porsby et al. 2008; Prado et al. 2009; Thole et al. 2012; Frank et al. 2015: Gram et al. 2015: Segev et al. 2015). Phaeobacter may exert a probiotic effect due to the production of the antibiotic tropodithietic acid (Brinkhoff et al. 2004; Porsby et al. 2008; Prado et al. 2009; D'Alvise et al. 2012). In association with senescent algae, Phaeobacter can switch from a symbiotic to a pathogenic lifestyle through the induction of algaecide synthesis and subsequent lysis of algal cells (Seyedsayamdost et al. 2011). Phaeobacter spp. are metabolically highly flexible and simultaneously metabolize multiple substrates under nutrientrich conditions (Zech et al. 2013). They catabolize algal osmolytes like dimethylsulfoniopropionate, and contain the sox genes for sulfur oxidation (Dickschat et al. 2010; Newton et al. 2010; Thole et al. 2012). In their natural habitat, Phaeobacter reach low abundances (Gram et al. 2015; Freese et al. 2017) but nevertheless are ecologically significant through their antilarval and antibacterial activities, preventing biofouling even at low cell densities (Rao et al. 2007). While gene sequences of *Phaeobacter* are only barely detectable by molecular methods in standard surveys of biofilms or eukaryotes and even entirely absent in the open ocean sequence databases, selective cultivation methods could recover Phaeobacter also from open ocean zooplankton (Gram et al. 2015; Freese et al. 2017). Previous comparative analysis of few isolates suggested that the genus Phaeobacter is distributed worldwide (Thole et al. 2012) but comprises disjunct, exclusively surface-associated populations (Freese et al. 2017).

A considerable number of closely related *Phaeobacter* strains have recently become available through direct isolation from different geographic regions and habitats (e.g., Hjelm et al. 2004; Porsby et al. 2008; Prado et al. 2009). In the present study, the available isolates were subjected to a detailed population genomic and phenotypic analysis to reveal the mechanisms of incipient diversification and the potential ecological niches of this surface-associated marine model bacterium.

Materials and Methods

Origin and Cultivation of Strains

The 88 *Phaeobacter* strains originated from aquacultures in Denmark, France and Spain (Hjelm et al. 2004; Porsby et al. 2008; Prado et al. 2009) and coastal marine environments in

Australia, France, and Germany (Brinkhoff et al. 2004; Rao et al. 2005) (supplementary fig. S1 and table S1, Supplementary Material online). Further details like isolation conditions are found in the references cited. All strains produce the typical brown pigment that is associated with the formation of tropodithietic acid. Strains were grown with marine broth medium (MB, Difco 2216) and preserved in liquid nitrogen after addition of 10% (v/v) glycerine.

Sequencing of the 16S rRNA Gene and the Internal Transcribed Spacer (ITS) Region

DNA was extracted from 1 ml culture with the DNeasy Blood&Tissue Kit (Qiagen) and the whole 16S rRNA gene plus the ITS region was amplified using the primers 27f (5'-AGA GTT TGA TCM TGG CTC AG-3'; Lane 1991) and 23S-130r (5'-GGG TTB CCC CAT TCR G-3'; Fisher and Triplett 1999). PCR products were purified with the DNA Clean & Concentrator (Zymo) and sequenced by Sanger sequencing. The 16S rRNA sequences were aligned using the aligner tool implemented in ARB 5.1 database SSURef 108 Silva NR 99 11_10_11 (Ludwig et al. 2004) and the alignment was manually refined based on secondary structure information. ITS sequences were aligned using ClustalW implemented in MEGA 5.05 (Tamura et al. 2011). After testing for the best nucleotide substitution model within MEGA (Jukes-Cantor, 16S rRNA genes; Kimura 2 with gamma distribution, ITS region), pairwise distance matrices were calculated and Maximum Likelihood phylogenetic trees were reconstructed in MEGA. Sequences were deposited in NCBI GenBank (accession numbers: KY357362-KY357447).

Genome Sequencing, Assembly and Annotation

Genomic DNA from 28 cultures in the late exponential phase was extracted with the JETFLEX Genomic DNA Purification Kit (Genomed). For preparation of SMRTbellTM template libraries, 8 μ g of genomic DNA were sheared (g-tubesTM, Covaris, Woburn, MA, USA), the size range monitored by pulse field gel electrophoresis, and DNA fragments end-repaired and ligated to hairpin adapters using P2 or P4 chemistry (Pacific Biosciences, Menlo Park, CA, USA). SMRT sequencing was carried out on the PacBio *RSII* (Pacific Biosciences). Illumina libraries were prepared with the TruSeq DNA Sample Prep Kit v2 (Illumina Inc., San Diego, CA, USA) and paired-end sequencing was performed on the HiSeq 2500 for 100 cycles (~8 million reads per genome).

PacBio reads were assembled de novo in SMRT Portal version 2.0.1 using the *RS_HGAP_Assembly.1* and *HGAP_Assembly_Advanced.1* protocols. Indel errors were corrected by mapping of Illumina reads using the Burrows– Wheeler Aligner (BWA) (Li and Durbin 2009) and the CLC Genomics Workbench 7.0.1 (CLC bio QIAGEN, Germany) for subsequent variant and consensus calling. The final assembly was trimmed, circularized and adjusted to the replication system as start point (Bunk 2016). Genome sequences were automatically annotated using Prokka 1.8 (Seemann 2014). The genomes of *P. inhibens* DSM 17395, *P. gallaeciensis* DSM 26640, and *P. inhibens* 2.10 (DSM 24558) (Thole et al. 2012; Frank et al. 2014) were retrieved from NCBI GenBank and used as primary database for annotation. *P. inhibens* T5 (DSM 16374) was available as permanent draft from IMG (Integrated Microbial Genomes & Microbiomes) (Dogs et al. 2013) and closed in this study. In total, this yielded 32 high quality, closed genomes for analysis. Genome sequences were deposited in NCBI GenBank (accession numbers: CP010588–CP010775, CP010784–CP010791, CP010805– CP010810).

Phylogenomics

Maximum likelihood and maximum parsimony phylogenomic trees were constructed from amino acid supermatrices as described by Simon et al. (2017) employing the JTT model of amino acid evolution (Jones et al. 1992) in conjunction with gamma-distributed substitution rates (Yang 1993) and empirical amino acid frequencies. To assess the stability of the phylogenomic branching, three different supermatrices were compiled from the concatenated orthologs which were 1) present in at least four sequences (4,336 genes, 1,269,031 characters), 2) present after removing uninformative genes using MARE (Meusemann et al. 2010) (3,169 genes, 1,031,284 characters), and 3) present in all 32 genomes (2,919 genes and 941,788 characters). For phylogenetic network analysis, single copy ortholog genes were identified, aligned based on their amino acid sequences and concatenated employing the ODoSE pipeline (Vos et al. 2013). The resulting matrix contained 2,821,782 characters and was fed into a NeighborNet analysis by SplitsTree 4.13.1 (Huson and Bryant 2006).

Evolution of Phaeobacter Chromosomes

Whole chromosome alignments were done with Mauve (Darling et al. 2010) and Easyfig (Sullivan et al. 2011). The gene order conservation was calculated as fraction of orthologous genes that are syntenous based on at least one shared neighbour (allowing for a gene insertion of one) for all pairwise chromosome comparisons (Yelton et al. 2011). Functions of chromosomal proteins were determined according to the COG (Clusters of Orthologous Groups of proteins) database via WebMGA (Wu et al. 2011). All phylogenetic distances and the digital DNA-DNA hybridization were inferred from pairwise comparisons of complete chromosome sequences via Genome BLAST Distance Phylogeny implemented in the GGDC 2.1 web service (Henz et al. 2005; Meier-Kolthoff et al. 2013). The resulting tree topology corresponds to that of the tree shown in figure 1 and the three main clusters are equally well supported. To test for biogeographic clustering of strains, a Pearson's product-moment correlation between pairwise genomic distances and pairwise geographic distances of sampling sites of the strains was calculated in R (R Core Team 2015). To determine the extent of the core and pan-chromosome, a gene content matrix listing the presence or absence of a gene within a certain chromosome was extracted from the OrthoMCL group files. Subsets with increasing numbers n of strains were randomly selected 100 times, and the number of orthologs present in n strains (core chromosome) as well as the numbers of orthologs present in at least one and at maximum n-1 strains (pan-chromosome) were determined using R (R Core Team 2015). Results were visualized using ggplot2 (Wickham 2009).

Gene gains and losses were quantified employing BadiRate (Librado et al. 2012). Gene families were determined using the TribeMCL algorithm (Enright et al. 2002) and a gene content matrix generated which was applied to an ultrametric tree calculated with r8s version 1.80 (Sanderson 2003). The goodness of different gene flow models was evaluated using the Akaike Information Criterion (AIC). For the *Phaeobacter* strains, the Lambda-Innovation model provided the best fit based on its lowest AIC value and yielded the size of ancestral phylogenetic nodes and the total number of gains and losses per lineage. The Lambda model assumes equal gene birth and death rates whereas innovation explicitly accounts for gene gain by HGT.

Horizontally transferred chromosomal elements were identified for each strain based on deviations from the normalized tetranucleotide frequency. Frequencies of each tetranucleotide were calculated for the whole chromosome and across 5 kb sliding windows with 2.5 kb overlap (Riedel et al. 2013) using the Biostrings package (Pagès et al. 2016) and the similarity between local and global frequencies was calculated as Pearson correlation coefficient. The interguartile-range of the correlation coefficients yielded the threshold below which a region was regarded as horizontally transferred (Riedel et al. 2013). Genomic islands were predicted with the IslandViewer web server (Dhillon et al. 2013) using the three different incorporated methods, SIGI-HMM (measuring codon usage), IslandPath-DIMOB (identifying abnormal sequence composition or the presence of genes related to mobile elements) and IslandPick (a comparative genomics-based method). The function of the proteins was classified according to the COG database via the WebMGA (Wu et al. 2011). A COG was defined as novel for the genus Phaeobacter if it did not occur outside of transferred elements in any Phaeobacter strain.

Prophages were predicted with PHAST (Zhou et al. 2011), but were only considered if they comprised more than just a tail protein and an integrase. If two prophages were located consecutively on the genome and one contained structural genes and terminase and the other replication genes, they were combined. Furthermore, sequences of prophages were compared with identified inducible *Phaeobacter* prophages characterized by Thole et al. (2012) and if they were

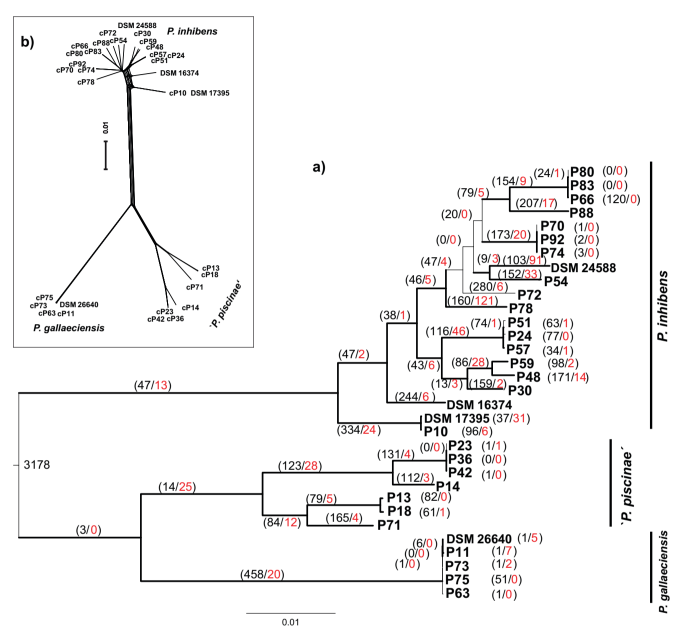


Fig. 1.—Phylogenomics of *Phaeobacter*. (a) Maximum likelihood (ML) phylogenetic tree inferred from a supermatrix of 1,269,031 aligned amino acid characters. Branches scaled according to the expected number of substitutions per site. Bold edges indicated branches with 100% bootstrapping support from all types of analysis (ML supermatrix; maximum parsimony (MP) supermatrix; ML MARE-filtered supermatrix; MP MARE-filtered supermatrix; ML coregenes matrix and MP core-genes). The numbers of inferred gene gains (black) and losses (red) are given next to the corresponding branch. Gene number of the inferred ancestral node is given at the midpoint root. (*b*) Phylogenetic network inferred by NeighborNet from the 2,821,782 aligned nucleotide characters of the concatenated single copy orthologs of 32 *Phaeobacter* strains. Scale bar, 0.01 changes per nucleotide site.

identical the start and end of the prophage sequence was corrected accordingly. Similarities of sequences (i.e., identity multiplied by coverage) were calculated using BLAST (Altschul et al. 1990) to define operational taxonomic prophage units (OTU). The OTU threshold was set to 60% since the lowest sequence similarity of the GTA present in all strains was 61.6%. Bacteriophage classification was done by VIRFAM via the head-neck-tail module genes (Lopes et al. 2014).

Phenotyping

Substrate utilization of 190 different carbon sources was determined via the Phenotype MicroArray (OmniLog PM) system using PM01 and PM02-A MicroPlates (AES Chemunex BLG 12111, BLG 12112). *Phaeobacter* strains were grown on MB agar at 25°C for 28 h, inoculated into modified medium (Buddruhs et al. 2013) and the respiration kinetics were recorded in the OmniLog Reader at 28 °C for 96 h. GENIII MicroPlates, which require a different inoculation medium did not result in reliable technical replicates for *Phaeobacter* in contrast to PM01 and PM02-A MicroPlates (data not shown). Based on prior reproducibility tests, two biological parallels were measured for each strain. After aggregation of the curve parameters for each substrate and strain using the R package *opm* (Vaas et al. 2013), the maximum height (A) as indicator for the metabolic activity was subject to a PCA using the R package *vegan* (Oksanen et al. 2015).

Statistics

All calculations were performed in R (R Core Team 2015). Significant differences in multiple comparisons of groups were calculated with Tukey procedures (function glht() in the R package *multcomp*; Hothorn et al. 2008) after an ANOVA (function aov()). Correlations were calculated using Pearson's product-moment correlation.

Results

Phylogeny, Phylogenomics, and Chromosome Structure

The overall phylogenetic diversity within the recently reclassified (Breider et al. 2014) genus Phaeobacter is very low. Maximum sequence divergence extracted from the pairwise distance matrix of 16S rRNA genes of the 88 available strains reached only 0.44% (supplementary fig. S2, Supplementary Material online). The strains clustered in four clades with an intergroup sequence divergence of <5 bp (0.37%). Two of these groups were identified as P. gallaeciensis and P. inhibens based on their phylogenetic affiliation with the respective type strains, DSM 26640^T and DSM 16374^T. The largest group of strains (47%) belonged to P. inhibens and was isolated from all five countries covered by our study (supplementary table S1, Supplementary Material online). Clustering of the ITS sequences yielded three distinct groups and revealed a 20fold higher nucleotide substitution rate compared with the 16S rRNA genes within the *P. inhibens* cluster (supplementary fig. S2, Supplementary Material online). Strain P88 and DSM 16374 contain different ITS therefore they occur more than once in the tree but the P. inhibens strains themselves fall into 11 different lineages with high bootstrap support. Although the substitution rate within another cluster, here designated "Phaeobacter piscinae" was lower for the ITS than the 16S rRNA genes, four separate ITS lineages could still be differentiated.

Strains representing unique ITS lineages or isolates from different habitats were selected for genomic comparisons (marked in bold face in supplementary fig. S2, Supplementary Material online). The size of the 32 chromosomes ranged from 3.588 to 3.896 Mb and the gene counts from 3,347 to 3,713. The GC-content of all chromosomes lies between 59.9 and 60.4 mol% G + C (supplementary

table S2, Supplementary Material online). All strains contained four nearly identical rRNA operons with exception of P. inhibens DSM 16374 and P88 which possess phylogenetically differing ITS (supplementary fig. S2, Supplementary Material online). The analysis of the core chromosome yielded an asymptotic saturation curve indicating that it is robustly predicted based on this set of 32 strains. In contrast, the panchromosome, that is, the whole chromosomal gene repertoire of *Phaeobacter* did not reach saturation (supplementary fig. S3, Supplementary Material online). As much as 78-87% of the gene content of the Phaeobacter strains fell into the large core chromosome that comprised 2,920 core genes. Even considering the extrachromosomal elements the average core genome still amounted to 78.1% (3,160 core genes) and the extrachromosomal elements of *Phaeobacter* contain only 11% (449) of all functional genes.

Results of the phylogenomic analyses of the chromosomal genes were very robust and congruent with the 16S rRNA gene and ITS phylogenies. The major clades could be distinguished in all types of analysis and were supported by high bootstrap values (fig. 1). A third clade was clearly separated from *P. gallaeciensis* and *P. inhibens* which was confirmed by low values for digital DNA-DNA hybridization (39.2-45.5%, median 41.0% for chromosomes; 38.9-46.9%, median 41.0% for whole genomes). Consequently, the novel species name "P. piscinae" was assigned to the third clade which encompasses two divergent subclusters. The phylogenetic network inferred from nucleotide sequences of the core genes showed almost no conflicting phylogenies (fig. 1b), indicating a low recombination between the different strains. Strains of *P. gallaeciensis* had nearly identical genomes with only 317 polymorphic sites in the Phaeobacter core chromosome although they were isolated from different geographic regions and associated with different eukaryotes (algae, Pecten maximus, Ostrea edulis, Venerupis philippinarum; supplementary table S1, Supplementary Material online). The intraspecific diversification in the other clades was much more pronounced (P. inhibens: 191,198 sites) (fig. 1b). Overall, genomic distances of the strains did not reflect geographic distances (Pearson Correlation test r = -0.07, 95%CI(-0.16, 0.01), P > 0.5) indicating that clustering of strains is independent of their geographic distribution (supplementary fig. S4, Supplementary Material online). Furthermore, the genomic distance of Phaeobacter strains originating from the same habitat is not lower than of strains originating from different habitats (supplementary fig. S5a, Supplementary Material online). Nearly all habitats contained strains from different clades and genomic distances did not differ significantly for any habitat (supplementary fig. S5b, Supplementary Material online). Thus, genomic clades did not show specificity for the different habitats as they are presently distinguished based on standard environmental data.

All chromosomes were largely syntenic (supplementary fig. S6a, Supplementary Material online). The high synteny was

further confirmed by a strong pairwise gene order conservation (0.99–1.0, median 0.997). Distinct chromosomal rearrangements were due to inversions of long (>100 kb) chromosomal segments and were also detected among the chromosomes of very closely related strains such as *P. gallaeciensis* strain P73 versus strains DSM 26640, P11, P63, P75. Inversions were located at seven different breakpoints and mostly close to one of the rRNA operons or a tRNA (supplementary fig. S6a, Supplementary Material online). They never disrupted a gene; rather they occurred adjacent to, or replaced, a horizontally transferred element (supplementary fig. S6b, Supplementary Material online).

A highly homogenous tetranucleotide frequency distribution was also observed among all chromosomes. Their median of the local to global tetranucleotide frequency similarity ranged between 0.83 and 0.84 which was similar to the closely related *Leisingera methylohalidivorans* (NC_023135; 0.83) and *Ruegeria pomeroyi* (NC_003911; 0.84) but significantly higher than for another more distant member of the *Roseobacter* group, *Dinoroseobacter shibae* (CP000830; 0.81), and for species with high ratios of recombination to point mutation like "*Candidatus* Pelagibacter ubique" (CP00084; 0.76) and *Flavobacterium psychrophilum* (AM398681; 0.71) (Vos and Didelot 2009) (Tukey, P < 0.001). This corroborates the low rate of recombination in *Phaeobacter*.

Pan-Chromosome Structure and Evolution

Phaeobacter chromosomes have expanded continuously during the divergence of the genus and gene gains occurred over all branches. Since the last common ancestor, an average of 47 genes were gained (1.4% of the gene content) and three genes were lost (0.08% of the gene content) (fig. 1a). Both values are significantly different from zero (t-test, P < 0.001). The highest gene gain (458, 12.7% of the gene content) was detected for the branch of P. gallaeciensis, indicating a considerable evolutionary divergence of its pan-genome. Gene gains and losses on terminal branches were in the same range (median gains, 1.8%; median losses, 0.04%). The numbers of gained genes were positively correlated with the pairwise phylogenomic distance and the number of amino acid substitutions per site (Pearson correlation r = 0.71, 95%Cl(0.67, 0.75) and r = 0.62, 95%CI(0.43, 0.75), both P < 0.001) suggesting that the pan-chromosome expanded constantly and in parallel with the diversification of the nucleotide and amino acid sequences. The best fit model (lowest AIC values) explaining the gene dynamics was the Lambda-Innovation model where innovation rates (1.6×10^{-3}) innovations per branch [relative age]) significantly exceeded Lambda rates (2.9 imes 10^{-4} births/deaths per gene and branch [relative age]) by one order of magnitude (Mann-Whitney Rank Sum Test, P < 0.001). This indicates that expansion of the panchromosome was predominantly caused by horizontal gene

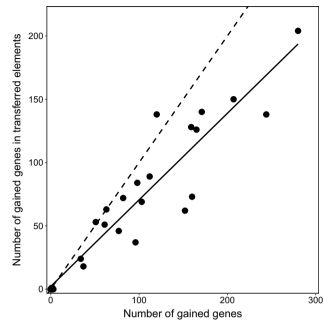


Fig. 2.—Comparison between the number of gained genes for each *Phaeobacter* strain (numbers at terminal branches in fig. 1*a*) and the number of gained genes occurring in predicted horizontally transferred elements. Solid line indicate the calculated linear regression (slope 0.68, pearson correlation r=0.95, 95%Cl(0.89, 0.97), P<0.001). Dashed line depicts theoretical correlation of r=1, that is, if all genes were gained by detectable HGT.

acquisition and/or de novo gene origin rather than by gene duplication events.

Between 15 and 25 elements per genome were predicted to be of foreign origin based on tetranucleotide frequency (9-20 elements), the analysis of prophages (1-5 elements) and genomic islands (7-17 elements) (supplementary fig. S6b, Supplementary Material online). On average these elements together constituted 7.5% of the chromosomal gene content and contained an average of ten genes. The size of the nonprophage elements were on average 7.5 kb. The elements were distributed over the whole chromosome without clade specific patterns (supplementary fig. S6b, Supplementary Material online). The majority (71%) of the gained genes estimated via the Lambda-Innovation model were detected in the transferred elements. In addition, the numbers of gained genes were tightly correlated to their numbers in transferred elements (fig. 2), emphasizing HGT as the major source for the genome expansion in all lineages of *Phaeobacter* alike. Most transferred proteins were closely related to those of the related genera Ruegeria, Leisingera, and Roseobacter, which represent typical generalists within the Rhodobacteraceae (supplementary table S3, Supplementary Material online).

A total of 84 putative prophages were detected. One type of prophage was present in all 32 chromosomes and identified as the gene transfer agent (GTA) (cf. Thole et al. 2012). The phylogeny of GTA largely corresponded to the branching pattern of the core genome (supplementary fig. S7, Supplementary Material online), indicating an early acquisition of GTA before the divergence of the three Phaeobacter species and a subsequent coevolution with their chromosomes. In addition, all strains carried the essential competence genes required for GTA-mediated gene uptake (Brimacombe et al. 2015) (supplementary table S4, Supplementary Material online). The 52 remaining prophages were classified as Myoviridae, Siphoviridae, and Podoviridae. Based on the established OTU threshold, they clustered into 20 OTUs of which seven (Pro1–Pro7) occurred in multiple strains and 13 were unique (supplementary table S5, Supplementary Material online). In contrast to GTA, the prophage pattern of the 32 Phaeobacter strains did not follow their genome phylogeny (supplementary fig. S8, Supplementary Material online). Pro1, the second most frequent prophage type in Phaeobacter, occurred in 12 genomes and could therefore be analyzed in detail. The nucleotide phylogeny of Pro1 was entirely congruent with the chromosome tree for members of P. inhibens, suggesting an integration of this phage before the radiation of this species and subsequent repeated losses (supplementary fig. S9, Supplementary Material online). However, the occurrence of two different Pro1 prophages in the genomes of "P. piscinae" suggests at least two independent transfers of Pro1 into the chromosome of "P. piscinae" (colored branches in supplementary fig. S9, Supplementary Material online). In the host chromosome, Pro1 always occurred at the same position with the exception of strains DSM16374 and P48. No Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR) were detected in any of the Phaeobacter strains indicating low selection pressure from lytic phages.

Diversification of Functional Genes and Phenotypes

In all Phaeobacter chromosomes, functional genes of amino acid transport and metabolism (COG category E) were dominant, followed by genes encoding transcription proteins (category K; 81% of which represent regulatory proteins) and by genes encoding energy production and conversion (category C) (fig. 3a). All chemotaxis genes belonging to COG category T and numerous genes involved in biofilm formation (e.g., motility, polysaccharide metabolism, and export) were found in the strains as expected based on their phenotypic properties. The quantitative distribution of COGs across different categories was similar (Mann-Whitney Rank Sum Test: no significant differences, P > 0.33; Pearson's test: significantly positive correlation, P > 0.88, P < 0.001) to that of other Alphaproteobacteria available in IMG like the facultatively anaerobic soil bacterium Paracoccus denitrificans or the oligotrophic free-living marine Planktomarina temperata (fig. 3a). The distribution of COG categories within the foreign, transferred elements in Phaeobacter was significantly dependent of the chromosomal COG category distribution in Alphaproteobacteria (Chi-square tests, P 0.02-<0.001). Aside from poorly characterized COGs, most COGs fell also into characterized categories C, E, and K (fig. 3a). In contrast to the chromosomal COG distribution in Phaeobacter, a significantly higher proportion of category L (replication, recombination and repair) was present in transferred elements (paired t-test, P < 0.001) but 29% of the COGs were identified as transposases which can be expected in transferred elements. In order to guantify HGT events leading to evolutionary innovations, the proportion of COGs in transferred elements, which were novel for all Phaeobacter chromosomes was determined (fig. 3b, triangles). Only few of the COGs represent innovations (supplementary table S6, Supplementary Material online). The percentages correspond to an average of 5.1 and 3.4 novel COGs of categories L and K, respectively, that were transferred per Phaeobacter genome, and one novel COG of categories D (cell cycle control, cell division, chromosome partitioning), E and M (cell wall/ membrane/envelope biogenesis) transferred per genome.

Principal component analysis (PCA) of functional gene content identified sets of specific functional genes commensurate with the clustering of Phaeobacter species (fig. 4). In particular, P. gallaeciensis contains an ABC type cobalt transport system, additional catalases, a complete Type IV secretion system previously described for plasmid pDSHI10 of D. shibae (Petersen et al. 2013), and a 1-aminocyclopropane-1-carboxylate (ACC) deaminase (gr8, gr10; fig. 4, supplementary table 57, Supplementary Material online). In most cases these genes were found on transferred genomic islands. Outside of P. gallaeciensis, the functional genes for the Type IV secretion system and ACC deaminase were only detected in the most deeply branching strains of P. inhibens (DSM 17395, P10) where it occurred in an element of foreign origin. A functional gene unique for P. inhibens is a divalent cation transporter (gr4). Heme utilization proteins (gr2), which are characteristic for other members of the Roseobacter group (Roe et al. 2013) are missing in this species. "P. piscinae" lacks a betalactamase class A and a Na⁺/glutamate symporter (gr3). Notably, members of this group that fell into two separate phylogenetic subclades also differed with respect to the presence of copper resistance proteins (gr6, gr12) in one subgroup (strains P14, P23, P36, P42; compare fig. 1) and the absence of polyketides and nonribosomal peptides synthases (COG2091, gr16) in the second subgroup (strains P71, P13, P18). Again, the copper resistance genes were located on genomic islands.

Phaeobacter strains were also phenotypically compared based on their metabolic activity utilizing 190 different carbon substrates. The analysis of these data by PCA revealed distinct substrate utilization patterns of *Phaeobacter* clades (fig. 5). *P. gallaeciensis* was separated from the other strains based on its significantly higher metabolic activity utilizing the different Tween compounds, α -hydroxybutyric acid, melibionic acid, and α -methyl-D-galactoside. In addition, group wise

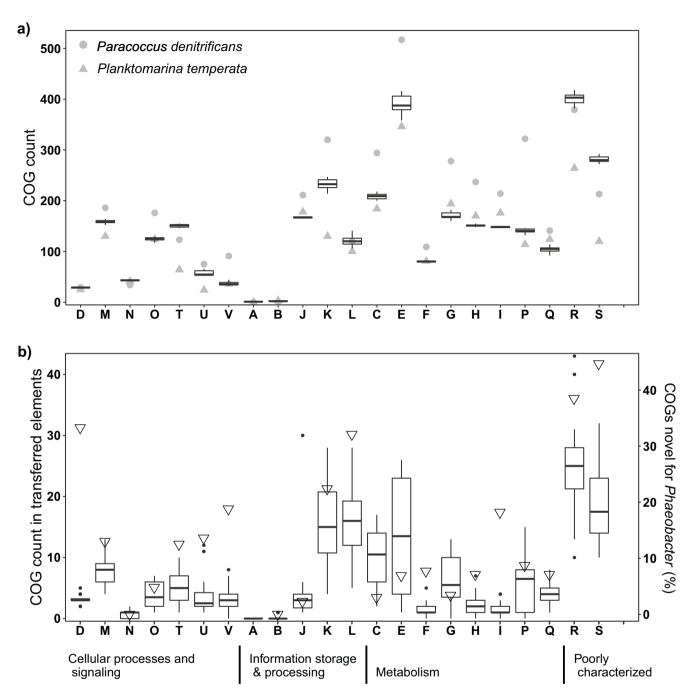


Fig. 3.—Abundance of different COGs present in the 32 *Phaeobacter* chromosomes in comparison to *Paracoccus denitrificans* PD1222 and *Planktomarina temperata* RCA23, DSM 22400, both from IMG, sorted by categories (a) and number of COGs gained through HGT (b). Open triangles indicate percentage of COGs gained by HGT that are novel for the genus *Phaeobacter*. Box plots show median, 25% and 75% percentiles, whiskers the 1.5*interquartile range and all values outside the range are shown as outliers. Letters indicate COG categories: A (RNA processing and modification), B (chromatin structure and dynamics), C (energy production and conversion), D (cell cycle control, cell division, chromosome partitioning), E (amino acid transport and metabolism), F (nucleotide transport and metabolism), G (carbohydrate transport and metabolism), H (coenzyme transport and metabolism), I (lipid transport and metabolism), J (translation, ribosomal structure and biogenesis), K (transcription), L (replication, recombination and repair), M (cell wall/ membrane/envelope biogenesis), N (cell motility), O (posttranslational modification, protein turnover, chaperones), P (inorganic ion transport and metabolism), R (general function prediction only), S (function unknown), T (signal transduction mechanisms), U (intracellular trafficking, secretion, and vesicular transport), and V (defense mechanisms).

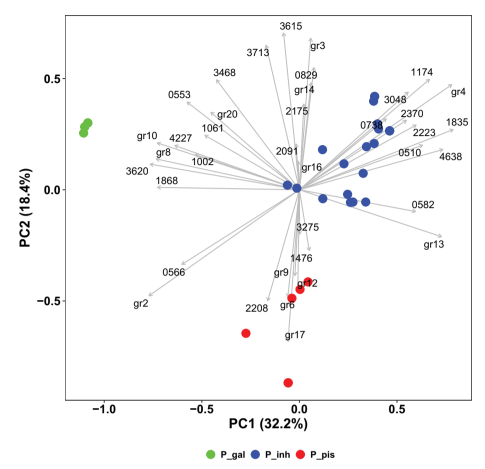


Fig. 4.—Principal component analysis of the presence of COGs in the 32 *Phaeobacter* chromosomes. Strains were colored according to their phylogenetic lineage (P_gal: *P. gallaeciensis*, P_inh: *P. inhibens*, P_pis: "*P. piscinae*"). Only COGs differing significantly between the clades (Tukey test with $P \le 0.02$) are indicated, COGs at the same position were grouped (gr10: COG0310, 0338, 0376, 0412, 0543, 0619, 1122, 1231, 1269, 1508, 1757, 2072, 3284, 3437, 3531, 3886, 4206, 5266; gr12: COG1276, 2132; gr13: COG2957, 2994, 5183; gr14: COG1144, 1271, 3203; gr16: COG3319, 3321; gr17: COG0423, 3448; gr2: COG3720, 4558, 4559, 4771; gr20: COG2932, 3409; gr3: COG0786, 2367, 3290; gr4: COG0598, 3457, 4067; gr6: COG2610, 3667; gr8: COG0417, 0501, 2515, 2948, 3451, 3504, 3702, 3704, 3736, 3838; gr9: COG1816, 2198, 2746).

(Tukey) tests revealed a preferential utilization of α -ketobutyric acid, sorbitol, methyl lactate, and unfavored utilization of tyramine and glucosamine of this species (P < 0.05). "*P. piscinae*" utilized N-acetylglutamate significantly better than the other strains, but α -ketobutyric acid and L-homoserine enabled a lower metabolic activity. *P. inhibens* strain P88 was a strong outlier because it exclusively was capable of utilizing xylitol and L-arabitol (fig. 5, insert) probably due to the exclusive presence of a D-xylulose reductase (PhaeoP88_01862, K05351) and xylokinase (PhaeoP88_01866, K00854).

Discussion

Evolution of the Phaeobacter Genome

The high fraction of the core genome of all three *Phaeobacter* species and its high synteny was rather unexpected based on the genome size and high metabolic versatility of *Phaeobacter* (Brinkhoff et al. 2008; Seyedsayamdost et al. 2011;

Gram et al. 2015). So far, high values for synteny (>0.9) have been reported for the SAR11 subclade Ia which is characterized by a small streamlined genome (Grote et al. 2012). A high relative fraction of the core genome (~80%) was only determined for bacterial species with specialised lifestyles or ecological niches, such as the subclade Ia of the free-living SAR11 (Grote et al. 2012), the obligate intracellular *Chlamydia psittaci* group (Voigt et al. 2012) and symbiotic *Vibrio fischerii* (Bongrand et al. 2016). In bacterial generalists with a comparable high 16S rRNA gene similarity (>99%) and genomic similarity the core genome constitutes a much smaller fraction, e.g., 52% in *Pseudomonas syringae* or 38–44% in *Vibrio cholerae* (Thompson et al. 2009; Nowell et al. 2014).

The accessory genome typically contains the genes relevant for the adaptation to particular environments (Kettler et al. 2007). Commensurate with the large core genome, the net gain of *Phaeobacter* genomes in accessory genes was small (average gains and losses, 1.8% and 0.04% of the

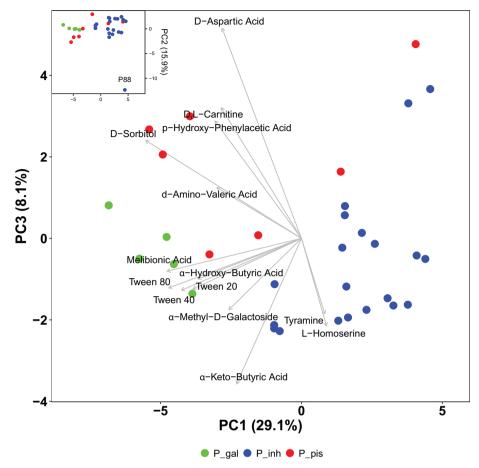


Fig. 5.—Principal component analysis of the utilization patterns of 190 substrates determined for the 32 *Phaeobacter* strains on OmniLog PM01 and PM02A plates. Strains were colored according to their phylogenetic lineage (P_gal: *P. gallaeciensis*, P_inh: *P. inhibens*, and P_pis: "*P. piscinae*"). PCA based on two biological replicates for each strain which mean position is shown. Substrates depicted contribute more than average to the ordination of data (Borcard et al. 2011). Results for principal component (PC) 1 and PC 3 are shown. Insert shows analysis for PC1 and PC2; its full scale plot, and an analysis along PC1 and PC4 are shown in supplementary fig. S10, Supplementary Material online).

chromosome, respectively) and lower than in other bacteria (up to 20% and 9.2% net gain in Prochlorococcus and Pseudomonas syringae, respectively) (Kettler et al. 2007; Nowell et al. 2014). The majority of the accessory genes of Phaeobacter were gained through transferred elements and likely were drawn from other members of the Roseobacter group. Like other members of the Roseobacter group, all Phaeobacter strains investigated contained GTAs, which can mediate high rates of interspecific gene transfer (Lang and Beatty 2007; McDaniel et al. 2010; Luo and Moran 2014). The GTA was acquired early before divergence of Phaeobacter and in contrast to prophages was always maintained in and continuously evolved with the chromosome. GTAs transfer small DNA fragments randomly generated from the host genome to recipient bacteria containing the necessary competence genes (Hynes et al. 2012; Brimacombe et al. 2015). The small size of the horizontally acquired elements in Phaeobacter, their origin in other Roseobacter genera containing GTA, as well as the presence of competence genes in *Phaeobacter* all are commensurate with HGT events mediated by GTAs. These findings indicate that GTAs constitute an important driver of HGT, diversification and niche adaptation in the genus *Phaeobacter*. One third of the accessory genome was identified as prophages including a *Myoviridae* phage. Obviously, the diversity of phages within the *Roseobacter* group extends well beyond the few documented *Podoviridae* and *Siphoviridae* (Huang et al. 2011; Ji et al. 2015; Liang et al. 2016). In contrast to GTA, vertical transmission of prophages seems to be rare in *Phaeobacter* and loss of prophages may have occurred repeatedly, in contrast to other bacterial groups such as the *Enterobacteriacae* (Bobay et al. 2014). The rapid turnover of prophages may also be caused by deletion during the transformation events mediated by GTA (Brimacombe et al. 2015; Rocha 2016).

Despite their comparatively limited acquisition of functional genes and high genome conservation, *Phaeobacter* encompasses three clearly separated phylogenomic clusters that are only marginally affected by homologous recombination. This finding is not due to sampling artifacts or cultivation bias as demonstrated by our phylogenetic analysis of all the 12 environmental 16S rRNA gene sequences that are available in public databases. All these sequences were found to group with one of the three *Phaeobacter* clusters but did not branch off between them (not shown). Even P. gallaeciensis strains isolated from the open ocean zooplankton fell into the existing sequence cluster (Freese et al. 2017). Since we did not find evidence for geographic isolation as the driver of diversification, the three Phaeobacter clades likely constitute distinct ecotypes (Cohan and Perry 2007). Different ecotypes with different physiologies and cluster-specific functional genes have been identified in the marine Prochlorococcus and the SAR11 clade (Venter et al. 2004; Kettler et al. 2007; Rusch et al. 2007), but in the latter cases differed by as much as > 4% (Carlson et al. 2009) and only rarely by < 1% (Hunt et al. 2008) in their 16S rRNA gene sequences. Since the differentiation in Phaeobacter occurred on a lower level of phylogenetic divergence (<0.5%) we sought to identify potential mechanisms of this micro-diversification.

Potential Drivers of Speciation

While no habitat preference was obvious for the Phaeobacter clades, specific gene functions and physiological traits distinguished the clades from each other, suggesting that they are linked to the evolutionary diversification within the genus. Due to the low population census sizes, and the failure to detect Phaeobacter at its low abundances by cultivationindependent approaches in most marine samples (see Introduction) prompted us to apply a reverse ecology approach in combination with a laboratory analysis of phenotypic properties to predict differences in the ecology of the clades. The large phylogenetic distance of P. gallaeciensis to the other clusters enabled the identification of a particularly large number of gained genes for this species. Algae, unlike copepods or fish, rapidly take up cobalt (Nolan et al. 1992). Therefore, the presence of the ABC-type cobalt transporter to acquire sufficient amounts of the essential trace element (Rodionov et al. 2006) may give P. gallaeciensis a competitive edge when growing in association with algae. In line with this hypothesis, the high number of catalases detected in P. gallaeciensis would protect the cells against damage and inhibition by reactive oxygen species produced by algae (Palenik et al. 1987; Oda et al. 1997). The complete, chromosomal type IV secretion system in P. gallaeciensis strains may allow the transfer of effector macromolecules to the host (Christie et al. 2005). Plant-associated terrestrial bacteria degrade the plant hormone precursor 1-aminocyclopropane-1-carboxylate (ACC) which led to promoted plant growth (Nascimento et al. 2014). In marine algae, ethylene is also produced via ACC (Maillard et al. 1993; Plettner et al. 2005) and the acquisition of an ACC deaminase by P. gallaeciensis as well as its superior utilization of the deamination product α -ketobutyric acid suggest that this species may exert a growth promoting effect on algae. P. gallaeciensis is also more competitive to utilize the algal osmoprotectant sorbitol as well as methyl lactate which may act as an antimicrobial component produced by microalga (Santoyo et al. 2009). Although these traits also appear in bacteria from other environments, their combined occurrence suggest that P. gallaeciensis has an advantage in algal associations over the other clades. However, our data indicate that the traits determined do not confer exclusive habitat specificity as for instance P. gallaeciensis can also occur with clam larvae (Ruiz-Ponte et al. 1998; supplementary table S1, Supplementary Material online). However, a metagenomics study of Roseobacter group members associated with an Emiliania huxleyi bloom revealed highest relative abundances of *P. gallaeciensis* which were 3.5 times higher than of P. inhibens (Segev et al. 2016), supporting our conclusions on the selective advantages of *P. gallaeciensis* in algal associations.

The other Phaeobacter ecotypes were characterized by a faster utilization of glucosamine and of the biogenic amine tyramine. Tyramine occurs in many organisms but elevated concentrations occur by oxygen-limited decomposition of protein-rich organic matter like fish or other seafood (Prester 2011), whereas glucosamine is most abundant in chitinous organisms like crustaceae (Benner and Kaiser 2003). Marine animals or environments rich with animal resource patches may thus constitute preferred environments for P. inhibens and "P. piscinae." Selective advantages of an association of "P. piscinae" with animals is further suggested by the lack of a specific transporter for glutamate, since the latter is mainly produced by plant and algae (Matsunaga et al. 1988; Tapiero et al. 2002), and also by the significantly more rapid utilization of N-acetylglutamate by "P. piscinae." N-acetylglutamate is a metabolic intermediate in the arginine synthesis and hence occurs in many organisms, but particularly high levels are present in fish where it acts as an important cofactor in the urea cycle (Caldovic and Tuchman 2003). Notably, one of the two "P. piscinae" subclusters has exclusively acquired genes homologous to CopABCD. These genes facilitate copper homeostasis and resistance in copper rich environments (Bondarczuk and Piotrowska-Seget 2013) and may represent a specific adaptation of members of the subcluster to copper compounds introduced in aguaculture environments through antifouling coatings or feeding supplements (Lorentzen et al. 1998; Stickney and McVey 2002). The distinct patterns of functional genes in the two "P. piscinae" subclusters indicate an incipient diversification of two different ecotypes in this particular cluster.

The loss of the heme acquisition/degradation system that distinguishes *P. inhibens* from the majority of the *Roseobacter* group (Roe et al. 2013) and also from the other *Phaeobacter* clusters suggests an adaptation to the utilization of iron ions through siderophores (Thole et al. 2012; investigated strains were reclassified from *P. gallaeciensis* to *P. inhibens*). Within

P. inhibens the 20 strains did not reveal a prevalence of any particular functional gene in different strains, suggesting that the ongoing radiation within this particular cluster was selectively neutral.

It has to be emphasized that most of the phenotypic differences detected in carbon substrate utilization concerned the efficiency of substrate utilization. These quantitative rather than qualitative differences seem to be of particular relevance for the evolution of *Phaeobacter*, but would have remained undetected by conventional testing. Our approach allowed the prediction of potential niches of different *Phaeobacter* clades that can now be tested in targeted phenotypic or environmental approaches.

Hallmarks and Implications of the *Phaeobacter* Population Genomics

Whereas the evolution of the entire Roseobacter group has been characterized by a steady net genome reduction (Luo et al. 2013), our data revealed that the more recent evolution of the three *Phaeobacter* species occurred through a slow, but continuous expansion of their chromosomes. They evolve in a manner different from free-living marine bacteria like the SAR11 clade and Prochlorococcus which were shaped by genome streamlining (Luo et al. 2011; Grote et al. 2012). The large core genome, high level of genome synteny, and the low proportion of chromosomal gene flow indicates that the three clusters of Phaeobacter have to be considered as "young" bacterial species (Nowell et al. 2014) which nevertheless already may constitute potential ecotypes. The traits for their versatile life-style are still conserved among all strains investigated. Within the surface-associated Phaeobacter, the evolutionary processes thus differ from that of marine generalists which are also capable of growing in the attached mode, such as Vibrio which maintains a large flexible genome leading to strong variations in gene content between closely related strains of the same population (Polz et al. 2006). Additional research is needed to better understand if these are general principles of evolution and niche adaptation of surface-associated bacteria which can constitute up to 66% of bacterial biomass in coastal marine environments (Becquevort et al. 1998).

Similar to the general evolutionary trend of the entire *Roseobacter* group (Luo et al. 2013), innovation in the genus *Phaeobacter* occurred mostly through acquisition of gene families involved in amino acid transport and metabolism (COG category E), gene regulation (K), and replication/recombination/repair (L). Based on our genome analysis, the particular adaptation of members of the *Roseobacter* group to the utilization of ephemeral nutrient patches (Luo et al. 2013) was further strengthened during the recent and ongoing differentiation of *Phaeobacter* species. We actually do not know which environmental factors triggered the first steps of incipient diversification. However, our results suggest that, at the

present stage of evolutionary diversification, the limited number of clade-specific laterally transferred genes provides adaptive advantage to different niches existing on surfaces of marine organisms and particles and drive ongoing diversification.

Given the distinct, but phylogenetically closely related clusters (>99.5% 16S rRNA gene sequence similarity) that were maintained during the evolution of *Phaeobacter*, our results not only confirmed that distinct geno- and ecotypic diversity is often hidden in bacterial groups with similar 16S rRNA gene sequences (Jaspers and Overmann 2004) but actually indicate mechanisms how populations with highly similar rRNA gene sequences diversify. Furthermore, the presently employed general cutoff for species delineation (97% or 98.6%) appears too coarse and should only be applied after careful consideration of the overall genomic divergence.

Taken together, the cluster-specific adaptations and genomic diversification revealed by the present work indicates that acquisition of functional genes reinforce speciation that may even be ongoing in "*P. piscinae*" whereas the recent genome divergence within *P. inhibens* so far has remained largely neutral. Our research further indicates that GTA likely mediates a large part of HGT and is an important driver of genome expansion in *Phaeobacter*. However, it remains to be investigated if this is a general characteristic of the surface-dwelling roseobacters.

Supplementary Material

Supplementary data are available at *Genome Biology and Evolution* online.

Acknowledgments

The work was supported by the Transregional Collaborative Research Centre "*Roseobacter*" (TRR 51/2 TA07), funded by the Deutsche Forschungsgemeinschaft. Cisse H. Porsby, Susanna Prado Plana and Jean-Louis Nicolas provided strains and information regarding their habitat. Rüdiger Pukall organized the strains, checked their quality and preserved them. We thank Markus Göker for fruitful comments and discussions regarding the design of phylogenetic analysis. We thank Isabel Vogt for correcting the genome annotations according to the NCBI requirements and Anika Methner, Alicia Geppert, Nicole Heyer, and Simone Severitt for excellent technical assistance.

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Associate editor: Bill Martin