

# Complete Genome of *Roseobacter ponti* DSM 106830<sup>T</sup>

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**Data deposition:** The whole-genome sequence project has been deposited at DDBJ/ENA/GenBank under the accession numbers CP048788 (chromosome) and CP048789 (plasmid). The NCBI BioProject and BioSample IDs are PRJNA605959 and SAMN14082710, respectively. The raw reads have been deposited at NCBI SRA database under the accession numbers SRR11069767 (paired-end Illumina) and SRR11069766 (Oxford Nanopore).

## Abstract

Members of the *Roseobacter* group are known for their different ecologically relevant metabolic traits and high abundance in many marine environments. This includes traits like carbon monoxide oxidation, sulfur oxidation, nitrogen oxidation, DMSP demethylation, denitrification, and production of bioactive compounds. Nevertheless, their role in the marine biogeochemical cycles remains to be elucidated. *Roseobacter ponti* DSM 106830<sup>T</sup>, also designated strain MM-7<sup>T</sup> (=KCTC 52469<sup>T</sup> =NBRC 112431<sup>T</sup>), is a novel type strain of the *Roseobacter* group, which was proposed as new *Roseobacter* species. It was isolated from seawater of the Yellow Sea in South Korea. We report the complete genome sequence of *R. ponti* DSM 106830<sup>T</sup>, which belongs to the family *Rhodobacteraceae*. The genome of *R. ponti* DSM 106830<sup>T</sup> comprises a single circular chromosome (3,861,689 bp) with a GC content of 60.52% and an additional circular plasmid (p1) of 100,942 bp with a GC content of 61.51%. The genome encodes 3,812 putative genes, including 3 rRNA, 42 tRNA, 1 tmRNA, and 3 ncRNA. The genome information was used to perform a phylogenetic analysis, which confirmed that the strain represents a new species. Moreover, the genome sequence enabled the investigation of the metabolic capabilities and versatility of *R. ponti* DSM 106830<sup>T</sup>. Finally, it provided insight into the high niche adaptation potential of *Roseobacter* group members.

**Key words:** *Roseobacter ponti* DSM 106830<sup>T</sup>, genome, phylogeny, metabolism, aerobic anoxygenic phototrophy.

## Significance

The phylogeny of the *Roseobacter* group is complex and has recently been reconsidered. Members of the *Roseobacter* group are highly likely to play a key role in the ocean due to their inhabitation of various ecological niches and their general abundance. However, their role in the ocean as part of the biogeochemical cycling is not yet fully understood. Therefore, complete genomes of *Roseobacter* group members are required to tackle these questions. In particular, the here presented genome of the new type strain *Roseobacter ponti* DSM 106830<sup>T</sup> helps to improve the phylogenetic resolution by representing a potential missing link in the *Roseobacter* group. Furthermore, we provide insights into the group genomic equipment which reveals high adaptational and functional properties.

## Introduction

The marine ecosystem is highly dynamic and bacterial diversity in the oceans is stunning. Bacterioplankton is dominated by a few marine bacterial clades, including the gammaproteobacterial SAR86 clade and the alphaproteobacterial SAR11, SAR116, and *Roseobacter* clades (Rappe et al. 2000; Suzuki et al. 2001; Kirchman 2008). Recently, the latter one has been described as *Roseobacter* group (Freese et al. 2017; Simon et al. 2017; Sonnenschein et al. 2018). These microorganisms are flexible in their metabolic potential, such as heterotrophy, photoheterotrophy, or autotrophy, lifestyle, such as free-living, particle-associated, or eukaryote-associated (Luo and Moran 2014). *Roseobacter* group members are widely distributed and in some marine ecosystems they constitute 15–20% of the bacterial community (Selje et al. 2004; Suzuki, Preston, et al. 2001; Moran et al. 2007). They possess different

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mechanisms for energy generation, such as utilization of organic and inorganic compounds including sulfur oxidation and catabolism of dimethylsulfoniopropionate (DMSP) to dimethyl sulfide (DMS), which is a climate-relevant gas (Wagner-Döbler and Biebl 2006). Luo and Moran (2014) compared *Roseobacter* group members based on genes mediating biogeochemical cycling including, *Roseobacter denitrificans* and *Roseobacter litoralis* which are capable of phototrophy (Shiba 1991). Both also produce a pink pigment, bacteriochlorophyll a (BchlA) and other bioactive secondary metabolites encoded by non-ribosomal peptide synthetases (NRPS) or polyketide synthase (PKS) gene clusters (Martens et al. 2007). Although various aspects of the *Roseobacter* group have been studied in recent years, complete genome sequences of isolates are lacking or limited to very few members. Thus, in depth biochemical and genomic characterization to elucidate ecological significance and evolutionary processes shaping the genomes of *Roseobacter* group members are still incomplete.

## Materials and Methods

### Isolation, Growth Conditions, and Genomic DNA Extraction

*Roseobacter ponti* DSM 106830<sup>T</sup> obtained from the “Deutsche Sammlung für Mikroorganismen und Zellkulturen” (DSMZ; Braunschweig, Germany) was originally isolated from seawater of the Yellow Sea in South Korea (Jung et al. 2017). A single colony from an active culture plate of *R. ponti* DSM 106830<sup>T</sup> was passed for 2 days in Medium 514 (DSMZ Medium 514 Bacto Marine Broth Difco 2216, Braunschweig, Germany) at 30°C and 180 rpm (Infors AG, Bottmingen, Schweiz). Cells were pelleted at 10.020× for 15 min and washed with sterile water twice. Genomic DNA was extracted by using the MasterPure complete DNA purification kit as recommended by the manufacturer (Epicentre, Madison, WI). The quality of the DNA was checked via agarose gel electrophoresis (0.8%, 50 min, 100 V) and the concentration was determined photometrically using a NanoDrop ND-1000 Spectrophotometer (Thermo Scientific, Schwerte, Germany).

### Genome Sequencing, Assembly, and Annotation

Illumina paired-end sequencing libraries were prepared using the Nextera XT DNA Sample Preparation kit (Illumina, San Diego, CA). Quality and size of the libraries were analyzed using Agilent Bioanalyzer 2100 and the Agilent High Sensitivity DNA kit as recommended by the manufacturer (Agilent Technologies, Waldbronn, Germany). Concentration of the libraries was determined using the Qubit dsDNA HS Assay kit as recommended by the manufacturer (Life Technologies GmbH, Darmstadt, Germany). Sequencing was performed using a MiSeq system and

reagent kit v3 with 600 cycles as recommended by the manufacturer (Illumina, San Diego, CA). For Oxford Nanopore sequencing, 1.5 µg high molecular weight DNA was used for library preparation employing the Ligation Sequencing kit 1D (SQK-LSK109) and the Native Barcode Expansion kit (EXP-NBD103, Barcode 2) as recommended by the manufacturer (Oxford Nanopore Technologies, Oxford, UK). Sequencing was performed for 72 h using a MinION device Mk1B and a SpotON Flow Cell R9.4.1 as recommended by the manufacturer (Oxford Nanopore Technologies) using MinKNOW software v18.12.6 for sequencing and Guppy v3.0.3 (<https://community.nanoporetech.com>, last accessed April 29, 2019) for demultiplexing. Default parameters were used for all software unless otherwise specified. Reads were quality-filtered using fastp version 0.20.0 (Chen et al. 2018) and remaining sequencing adapters were removed with cutadapt v2.5 (Martin 2011). Unicycler version 0.4.8 (Wick et al. 2017) was used for a de novo hybrid assembly in normal mode. The quality of the assembly was assessed with CheckM v1.1.2 (Parks et al. 2015) and validated with Bandage 2.1 (Wick et al. 2015). Genome annotation was performed employing the Prokaryotic Genome Annotation Pipeline (PGAP) (Tatusova et al. 2016).

### Phylogenetic Analysis

The Genome Taxonomy Database Toolkit (GTDB-Tk) v1.0.1 (Chaumeil et al. 2019), was used to provide an initial taxonomic placement. Afterwards, the genome sequence was uploaded to the Type (Strain) Genome Server (TYGS) (<https://tygs.dsmz.de>), for a whole-genome-based taxonomic analysis (Meier-Kolthoff and Göker 2019). Analysis was performed for the 16 closest relatives (supplementary table S1, Supplementary Material online) at 16S rRNA gene level and with whole-genome sequences as described by TYGS using default parameters (as of January 28, 2020). In addition, an extended 16S rRNA gene analysis including sequences of not fully genome-sequenced type strains was performed by employing TYGS (Meier-Kolthoff and Göker 2019). The fast homology search tool AAI-profiler (Medlar et al. 2018) was also used to assess the phylogeny by using the deduced proteome of *R. ponti* DSM 106830<sup>T</sup>.

### Metabolism and Secondary Metabolites

To investigate the metabolic potential of *R. ponti* DSM 106830<sup>T</sup>, BlastKOALA version 2.2 (Kanehisa et al. 2016) was used to get an overview of pathways. Putative secondary metabolites and putative phage regions were identified with AntiSMASH 5.1.0 (Blin et al. 2019) and PHASTER (Arndt et al. 2016), respectively.

## Results and Discussion

### Genomic Features

The genome of *R. ponti* DSM 106830<sup>T</sup> was sequenced using both, long-read (Oxford Nanopore) and short-read technology (Illumina). After quality-filtering, 201,474 long-reads with a mean length of 7,642 bp (Oxford Nanopore) and 2,980,230 paired-end Illumina reads (2 × 300 bp) were obtained. Unicycler version 0.4.8 (Wick et al. 2017) was used for a de novo hybrid assembly. This resulted in two circular contigs representing a chromosome and a plasmid with a total average coverage of 559-fold (chromosome) and 764-fold (plasmid). The assembly was manually validated with Bandage version 2.1 (Wick et al. 2015). CheckM detected a completeness of 99.25% and a contamination rate of 0.48%. The genome comprises one circular chromosome (3,861,689 bp) and one circular plasmid (100,942 bp) with a GC-content of 62.92% and 61.51%, respectively. Genome features are summarized in table 1.

In total, 52.1% of genes were annotated by BlastKOALA and classified into 23 functional categories according to the KEGG Orthology. Among all categories, the environmental information processing (11.8%) and carbohydrate metabolism (11.6%) were the most abundant.

### Phylogeny of *R. ponti* DSM 106830<sup>T</sup>

GTDB-Tk (Chaumeil et al. 2019) revealed that this strain is novel and placed *R. ponti* DSM 106830<sup>T</sup> taxonomically into the family *Rhodobacteraceae*, based on average nucleotide identity (N/A) and relative evolutionary divergence values (~0.97). Phylogenetic assignment at genus level was not possible. To refine the phylogenetic position of *R. ponti* DSM 106830<sup>T</sup> a phylogenetic tree based on 16S rRNA gene sequences and whole-genome sequence was constructed. The 16S rRNA gene-based tree grouped *R. ponti* DSM 106830<sup>T</sup> together with *R. litoralis* OCh 149, *R. denitrificans* OCh 114, and *R. denitrificans* DSM 7001 into the genus *Roseobacter* (supplementary fig. S1A and S1C, Supplementary Material online). The phylogenetic tree based on Genome Blast Distance Phylogeny takes the whole-genome sequence into account and resulted in a different clustering (supplementary fig. S1B, Supplementary Material online). Briefly, *R. ponti* DSM 106830<sup>T</sup> was most closely assigned to *Sulfitobacter marinus* DSM 23422 based on GBDP calculations for the 16 closest relatives (supplementary table S1A and S1B, Supplementary Material online). The tree emphasizes the challenge to make valid phylogenetic classifications of new isolates from the *Roseobacter* group. The whole-genome tree supports the classification of *R. ponti* DSM 106830<sup>T</sup> into the *Roseobacter* group, but due to deep branching and low bootstrap values genus assignments are not supported (supplementary fig. S1B, Supplementary Material online). Additionally, proteome AAI-profile analysis further validates this finding with AAI values <80%. AAI-profiler assigned

**Table 1**

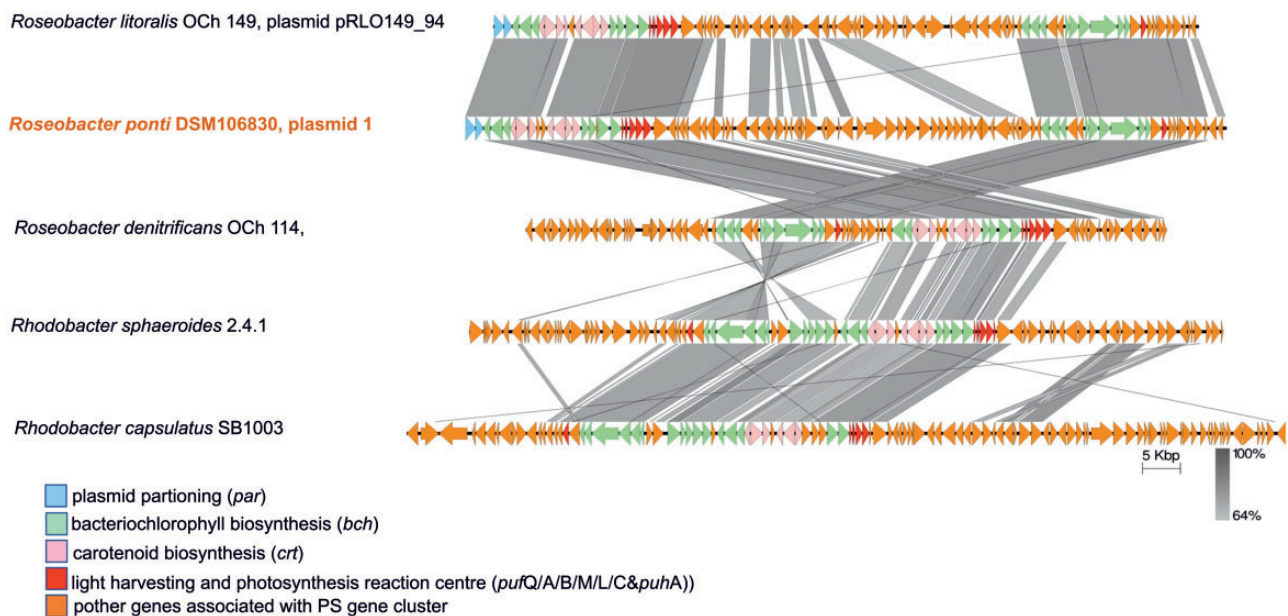
Genomic Features of *Roseobacter ponti* DSM 106830<sup>T</sup>

Features	Chromosome	Plasmid (p1)
Genome size (bp)	3,861,689	100,942
GC content (%)	60.52	61.51
Gene number	3,715	97
rRNA genes	3	0
tRNA genes	42	0
ncRNA	2	0
tm-rRNA genes	1	0
Regulatory RNA	8	0
CRISPR	0	0
Phage	1	0

strain *R. ponti* DSM 106830<sup>T</sup> inconclusively to a *Rhodobacteraceae* bacterium, though the closest sequence neighbors belong to both genera *Roseobacter* and *Sulfitobacter*. Consequently, new isolates including the here presented *R. ponti* DSM 106830<sup>T</sup> of the *Roseobacter* group should not be solely classified on the basis of the 16S rRNA gene as they share >89% 16S rRNA gene identity (Buchan et al. 2005), which can lead to incorrect phylogenetic assignments. Recently, several genus reassignments of the *Roseobacter* group are ongoing (Wirth and Whitman 2018). Based on the here presented data, we suggest *R. ponti* DSM 106830<sup>T</sup> as a missing link between the *Sulfitobacter* and *Roseobacter* genus rather than a new species within the *Roseobacter* genus.

### *Roseobacter ponti* DSM 106830<sup>T</sup> Metabolic Versatility

*Roseobacter* group members are equipped with a diverse toolkit of metabolic capabilities, which partly explains their success in colonizing a broad range of different marine habitats (Buchan et al. 2005). Some metabolic properties of *R. ponti* DSM 106830<sup>T</sup> were studied by Jung et al. (2017). The analysis of the genome sequence of *R. ponti* DSM 106830<sup>T</sup> revealed specific traits and metabolic adaptations to ecological niches. Jung and coworkers screened the strain for presence of genes encoding photosynthetic reaction center proteins (*pufL*, *pufM*, and *pufA*), enabling aerobic anoxygenic phototrophy (AAP), by PCR and detected *pufA* but not *pufL* and *pufM*. However, the whole-genome sequence confirmed that all three genes responsible for AAP are present (G3256\_19015, G3256\_18695, and G3256\_18700). In addition, the genome harbors putative genes important for a functional photosynthetic gene cluster (PGC) and the production of BchlA (Zheng et al. 2011). These included the *bch* genes important for BchlA biosynthetic pathways (G3256\_18600, G3256\_18655, G3256\_18670, G3256\_18675, G3256\_18980), the *puf* operon involved in formation of the reaction center (G3256\_18680–G3256\_18705), *pufA* participating in reaction center assembly (G3256\_19015), and *crt* genes (G3256\_1860,



**FIG. 1.**—Synteny comparison of *Roseobacter ponti* DSM 106830<sup>T</sup> plasmid 1 to closest relatives. References include *Roseobacter litoralis* OCh 149, plasmid pRLO149\_94 (CP002624.1), and cutted regions of *Roseobacter denitrificans* OCh 114 chromosome (CP000362.1), *Rhodobacter sphaeroides* 2.4.1, chromosome 1 (CP000143.2), and *Rhodobacter capsulatus* SB1003, chromosome (CP001312.1). The comparison was performed with Easyfig (Sullivan et al. 2011) using BlastN percent identities. Synteny between related regions is indicated by vertical gray-shaded areas and black lines. The legend indicates the biological categories of genes involved in the photosynthesis cluster.

G3256\_18620, G3256\_18640, G3256\_18645, G3256\_18650) responsible for carotenoid biosynthesis and regulation (Pradella et al. 2004; Zheng et al. 2011; Petersen et al. 2012; Chi et al. 2015). The *puf* operon (*pufQBALMC(X)*) analysis of Petersen et al. (2012) to was extended to determine the closest related synteny to the *R. ponti* DSM 106830<sup>T</sup> *puf* operon (fig. 1). The *puf* operon comprises six genes encoding a cytochrome subunit (*pufQ*), light harvesting proteins (*pufA* and *pufB*), the photosynthetic reaction center subunits L and M (*pufL* and *pufM*, respectively) the photosynthetic reaction center cytochrome C (*pufC* or *pufX* in *Rhodobacter*) (Kortlüke et al. 1997; Zheng et al. 2011). *Roseobacter ponti* DSM 106830<sup>T</sup> contains *pufC*, which is also present in *R. denitrificans* OCh 114 and *R. litoralis* OCh 149 (fig. 1) (Kortlüke et al. 1997). The analysis showed that the *puf* operon of *R. ponti* DSM 106830<sup>T</sup> is most similar to a *puf* operon of *R. litoralis* OCh 149 (fig. 1). In both organisms, the operon is encoded by plasmids (Pradella et al. 2004) and share PGC genes. Plasmid genes that are unrelated to the *puf* operon or PGC are genetically not conserved. Remarkably, in other members of the *Roseobacter* group, such as *Rhodobacter encapsulatus* and *R. denitrificans*, the *puf* operon is encoded by the chromosome (Kortlüke et al. 1997; Petersen et al. 2012), indicating an evolutionary adaptation to a specific ecological niche of some *Roseobacter* group members via plasmid acquisition.

The genome analysis by BlastKOALA resulted in a variety of pathway, such as genes involved in biogeochemical cycling

(Luo and Moran 2014) including the dissimilatory nitrite reductase (*nirK*), dimethylsulfoniopropionate demethylase (*dmdA*), sulfur oxidation protein complex (*soxB*), and large subunit of carbon monoxide dehydrogenase (*coxL*). In comparison to *R. denitrificans* OCh 114 *nasA*, *nirB*, *napA*, *narG*, and *nirS* involved in the nitrogen metabolism are absent in the genome of *R. ponti* DSM 106830<sup>T</sup>. It is indicated that *R. ponti* DSM 106830<sup>T</sup> is capable of degrading DMSP and performing carbon monoxide oxidation.

Additionally, the search for gene clusters involved in secondary metabolite synthesis identified six putative clusters of which five were encoded by the chromosome and one by the plasmid. Interestingly, the plasmid harbors one putative terpene cluster which encodes the synthesis of the carotenoid spheroidenone, which is the main light-harvesting carotenoid of *Roseobacter* group members (fig. 1) (Wagner-Döbler and Biebl 2006). Pigment gene clusters are a typical feature of AAP bacteria and cell color can range from yellow/orange over brown or pink/red to purple (Zheng et al. 2011). In addition to the photosynthetic apparatus including the *puf* operon, the carotenoids are encoded on the plasmid. Carotenoids are, amongst others, protective against harmful radicals, such as oxygen and radiation (Chi et al. 2015) and could be advantageous from an evolutionary point of view.

#### *Roseobacter ponti* DSM 106830<sup>T</sup> Horizontal Gene Transfer

The *Roseobacter* group members occur in a wide variety of different ecological niches in the marine oceans, indicating a

high adaptation potential (Wagner-Döbler and Biebl 2006; Brinkhoff et al. 2008). Evolutionary driving forces for genetic diversification by horizontal gene transfer (HGT) and mechanisms of DNA exchange include phages (transduction), plasmids (conjugation), and virus-like particles or gene transfer agents (GTA) (Wall et al. 1975; Pal et al. 2005; Zhan et al. 2016). To investigate the evolutionary potential of *R. ponti* DSM 106830<sup>T</sup> the genome was screened for indicators of HGT. This revealed one putative chromosomal phage region (17.4 kb, 2,122,003–2,139,468). PHASTER classified the completeness as questionable and the typical insertion sites attL/attR were not identified. It is likely that this element is rather a GTA, a virus-like particle which is proposed to originate from ancient prophage remnants (Lang and Beatty 2000). Since gene equipment and content of GTAs are similar to phages, detection by PHASTER is expected. Previous genomic studies showed that nearly all genomes of the Roseobacter group possess GTAs (Lang and Beatty 2000; Newton et al. 2010; Huang et al. 2011). In detail, the detected region comprises 19 putative phage-associated genes of which three were annotated as GTA.

In conclusion, the analysis of the *R. ponti* DSM 106830<sup>T</sup> genome sequence shows that phylogenetic classifications of new Roseobacter group members should not be performed by 16S rRNA gene but on the whole-genome comparisons. We suggest *R. ponti* DSM 106830<sup>T</sup> as a missing link between the genera *Sulfitobacter* and *Roseobacter* rather than as a new species in the genus *Roseobacter*. Notably, the plasmid p1 of *R. ponti* DSM 106830<sup>T</sup> encodes the AAB operon, which was described for only six *Rhodobacteraceae* members including *R. litoralis* OCh 149 (Pradella et al. 2004; Brinkmann et al. 2018). Additionally, a putative GTA was detected in the chromosome. Both indicate the adaptive capabilities to a specific ecological niche and the oligotrophic marine environment by HGT via plasmids and GTAs. Finally, the genome analysis confirms previous studies (Jung et al. 2017) and is the foundation for future physiological analyses of *R. ponti*. Additionally, the data provided here will be valuable for studies targeting PGC within the Roseobacter group.

## Supplementary Material

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

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