



The *Roseibium album* (*Labrenzia alba*) Genome Possesses Multiple Symbiosis Factors Possibly Underpinning Host-Microbe Relationships in the Marine Benthos

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ABSTRACT Here, we announce the genomes of eight *Roseibium album* (synonym *Labrenzia alba*) strains that were obtained from the octocoral *Eunicella labiata*. Genome annotation revealed multiple symbiosis factors common to all genomes, such as eukaryotic-like repeat protein- and multidrug resistance-encoding genes, which likely underpin symbiotic relationships with marine invertebrate hosts.

Roseibium *album* (1) (homotypic synonyms, *Stappia alba* [2] and *Labrenzia alba* [3]) is a Gram-negative, marine alphaproteobacterium (order *Hyphomicrobiales*, family *Stappiaceae*) that has frequently been isolated from sessile, filter-feeding invertebrates such as sponges, corals, and oysters (2, 4–6). To date, however, only a few genomes of this species are available. To illuminate the putative roles of *R. album* in association with marine animals, here we report the genomes of eight *R. album* strains that were isolated from the octocoral *Eunicella labiata* (Table 1), and we present symbiosis factors and environmental resistance traits common to all genomes (Fig. 1).

The eight *R. album* strains were isolated from *E. labiata* specimens that had been collected off the coast of Faro, Portugal, after plating of host-derived homogenates on half-strength marine agar medium followed by incubation at 18°C for 4 weeks (4). Genomic DNA was extracted from pure cultures using the Wizard genomic DNA purification kit (Promega, Madison, WI, USA) as described previously (4). The isolates were sequenced at the Joint Genome Institute (JGI) as part of the Genomic Encyclopedia of Type Strains Phase IV (KMG-V) project. Default parameters were used for all software unless otherwise specified. Genome libraries (300 bp) were prepared with the KAPA HyperPrep kit (KAPA Biosystems) and sequenced using the Illumina NovaSeq 6000 platform (S4 flow cell). Raw reads were quality filtered per JGI standard operating practice (SOP) protocol 1061 using BBTools v38.86 (<http://bbtools.jgi.doe.gov>). Filtered reads were assembled into contigs using SPAdes v3.14.1 (7) with 25, 55, and 95 k-mers, and contigs were annotated using the Integrated Microbial Genomes and Microbiomes (IMG/M) pipeline v5.0.17 (8). We report the Clusters of Orthologous Groups of proteins (COG) profiles obtained for all eight strains using the IMG/M platform (9). Average nucleotide identity (ANI) values obtained on the IMG/M platform (9) for all pairwise combinations among the strains were above 98.3%, supporting the same-species status of the strains. Genome completeness and contamination scores were assessed with the Microbial Genomes Atlas (MiGA) (10) (Table 1). AntiSMASH v6.0 (11) was used to identify secondary metabolite biosynthetic gene clusters (SM-BGCs).

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TABLE 1 General features of the *Roseibium album* (*Labrenzia alba*) genomes reported in this study

Genome ^a	IMG/M identifier ^b	GC content (Mb)	Genome coverage (x)	No. of contigs	No. of N ₅₀ (bp)	Read length (bp)	Completeness (%)	Contamination (%)	No. of coding sequences	No. of RNAs	No. of genes in COGs	GenBank accession no.	Assembly accession no.	SRA accession no.		
<i>Labrenzia alba</i>	2882936267	7.00	56.44	211.9	19	668,536	19,078,598	151	98.1	2.8	6,475	59	5,303	JADOU000000000	GCA_015751945.1	SRR13202144
EL_13																
<i>Labrenzia alba</i>	2880767578	7.14	56.34	210.4	76	352,437	31,261,630	151	98.1	2.8	6,732	61	5,352	JADOU000000000	GCA_015752395.1	SRR13202553
EL_126																
<i>Labrenzia alba</i>	2880560698	7.06	56.44	215.3	51	400,118	33,793,138	151	98.1	2.8	6,629	56	5,334	JADOU000000000	GCA_015752105.1	SRR13202552
EL_132																
<i>Labrenzia alba</i>	2880567409	7.29	56.31	207.9	54	363,626	31,961,096	151	98.1	2.8	6,808	66	5,460	JADOU000000000	GCA_015752355.1	SRR13202549
EL_142																
<i>Labrenzia alba</i>	2880774395	6.9	56.44	215.3	18	899,415	32,269,048	151	98.1	2.8	6,384	62	5,230	JADOU000000000	GCA_015752425.1	SRR13202554
EL_159																
<i>Labrenzia alba</i>	2880574310	6.9	56.44	214.6	18	899,415	20,727,438	151	98.1	2.8	6,387	62	5,230	JADOU000000000	GCA_015752365.1	SRR13202550
EL_162																
<i>Labrenzia alba</i>	2880580784	7.13	56.36	211.6	31	464,795	36,215,300	151	98.1	2.8	6,593	59	5,356	JADOU000000000	GCA_015752055.1	SRR13202548
EL_195																
<i>Labrenzia alba</i>	2880780866	7.06	56.44	214.8	49	402,705	22,234,602	151	98.1	2.8	6,638	56	5,334	JADOU000000000	GCA_015752025.1	SRR13202551
EL_208																

^aAll strains reported in this study were isolated from the octocoral host *Eunicella labiata*.^bUnique genome identifier at the IMG/M portal.

COG ID	Description	EL13	EL126	EL132	EL142	EL159	EL162	EL195	EL208
COG0457	Tetratricopeptide (TPR) repeat	14	18	15	15	15	15	16	15
COG0790	TPR repeat	12	9	9	11	11	11	11	9
COG0666	Ankyrin repeat	1	1	1	1	1	1	1	1
COG2319	WD40 repeat	6	4	5	4	4	4	4	5
COG1566	Multidrug resistance efflux pump	7	8	7	8	6	6	6	7
COG4300	Cadmium resistance protein CadD, predicted permease	1	1	1	1	1	1	1	1
COG3793	Tellurite resistance protein	2	3	3	2	3	3	3	3
COG4103	Tellurite resistance protein B (TerB) family	2	2	2	2	2	2	2	2
COG1275	Tellurite resistance protein TehA and related permeases	0	0	0	0	0	0	0	0
COG0726	Peptidoglycan/xylan/chitin deacetylase, PgdA/CDA1 family	3	2	2	4	3	3	3	2
COG2861	YibQ, putative polysaccharide deacetylase 2 family	1	1	1	1	1	1	1	1
COG3525	N-acetyl-beta-hexosaminidase	1	1	1	1	1	1	1	1
COG0363	Glucosamine-6-phosphate isomerase/deaminase	2	1	2	1	1	1	2	2
COG2175	Taurine dioxygenase, alpha-ketoglutarate-dependent	3	2	3	3	3	3	3	3
COG4521	ABC-type taurine transport system, periplasmic comp.	1	1	1	1	1	1	1	1
COG4525	ABC-type taurine transport system, ATPase component	1	1	1	1	1	1	1	1
COG3119	Arylsulfatase A or related enzyme	10	9	9	9	9	9	10	9
SM-BGCs	Description								
transAT-PKS	TransAT polyketide synthase cluster	1	1	1	1	1	1	1	1
T3PKS	Type III polyketide synthase cluster	1	1	1	1	1	1	1	1
NRPS	Non-ribosomal peptide synthetase cluster	1	1	1	1	1	1	1	1
RiPP-like	Ribosomally synthesized & post-translationally modified peptides	2	2	2	1	1	1	1	2
TfuA-related	Thioamidite cluster	1	1	1	2	1	1	1	1
Thiopeptide	Thiopeptide cluster	1	1	1	2	1	1	1	1
Lanthipeptide	Lanthipeptide class II cluster	1	1	1	0	0	0	0	1
Terpene	Terpene cluster	1	1	1	1	1	1	1	1
Betalactone	Betalactone cluster	1	1	1	1	1	1	1	1
Hserlactone	Homoserine lactone cluster	0	1	1	1	0	0	0	1

Number of COG entries or SM-BGCs:

0 1 2-3 4-6 7-10 11-15 16-20

FIG 1 Selected COG functions characteristic of the *Roseibium album* (*Labrenzia alba*) genomes described here, as well as SM-BGCs present in all genomes. Values for each entry represent the numbers of coding sequences assigned to COG functions per genome (top) and the numbers of SM-BGCs coding for major compound classes (such as polyketides and terpenes) per genome (bottom).

Analysis of the *R. album* genomes revealed the presence of various COGs important for the establishment of symbiotic relationships, including eukaryotic-like WD40, ankyrin, and tetratricopeptide repeats (Fig. 1). COG functions related to drug (e.g., COG1566) and heavy metal resistance were common to all eight genomes, including traits specific for tellurite resistance that are usually encoded on plasmids (12) (Fig. 1). The eight strains harbor genes for the utilization of chitin, a trait that was previously reported for sponge and coral symbionts (13), as revealed by the presence of polysaccharide deacetylases and exochitinases. Other features included a coding potential for a possible role in sulfur cycling, e.g., through the catabolism of the sulfolipid cerebroside 3-sulfate and of the amino-sulfonic acid taurine, two compounds that are widely synthesized in animal tissue and utilized by bacterial symbionts (14). Moreover, all *R. album* genomes possess the potential to produce a diverse range of secondary metabolites. Indeed, we found a variety of SM-BGCs encoding terpenes, *trans*-AT-polyketide synthases and type III polyketide synthases, nonribosomal peptide synthetases, and several ribosomally synthesized peptides in this genome pool (Fig. 1).

Data availability. The genome sequences of the eight *R. album* (*L. alba*) strains have been deposited in GenBank by the JGI. Accession numbers are listed in Table 1.

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REFERENCES

- Hördt A, López MG, Meier-Kolthoff JP, Schleuning M, Weinhold LM, Tindall BJ, Gronow S, Kyrpides NC, Woyke T, Göker M. 2020. Analysis of 1,000+ type-strain genomes substantially improves taxonomic classification of *Alphaproteobacteria*. *Front Microbiol* 11:468. <https://doi.org/10.3389/fmicb.2020.00468>.
- Pujalte MJ, Macián MC, Arahal DR, Garay E. 2005. *Stappia alba* sp. nov., isolated from Mediterranean oysters. *Syst Appl Microbiol* 28:672–678. <https://doi.org/10.1016/j.syapm.2005.05.010>.
- Biebl H, Pukall R, Lünsdorf H, Schulz S, Allgaier M, Tindall BJ, Wagner-Döbler I. 2007. Description of *Labrenzia alexandrii* gen. nov., sp. nov., a novel alphaproteobacterium containing bacteriochlorophyll a, and a proposal reclassification of *Stappia aggregata* as *Labrenzia aggregata* comb. nov., and of *Stappia alba* as *Labrenzia alba* comb. nov., and emended descriptions of the genera *Pannonibacter*, *Stappia* and *Roseibium*, and of the species *Roseibium denhamense* and *Roseibium hamelinense*. *Int J Syst Evol Microbiol* 57:1095–1107. <https://doi.org/10.1099/ijss.0.64821-0>.
- Keller-Costa T, Eriksson D, Gonçalves JMS, Gomes NCM, Lago-Lestón A, Costa R. 2017. The gorgonian coral *Eunicella labiata* hosts a distinct prokaryotic consortium amenable to cultivation. *FEMS Microbiol Ecol* 93:fix143. <https://doi.org/10.1093/femsec/fix143>.
- Rodrigues GN, Lago-Lestón A, Costa R, Keller-Costa T. 2018. Draft genome sequence of *Labrenzia* sp. strain EL143, a coral-associated alphaproteobacterium with versatile symbiotic living capability and strong halogen degradation potential. *Genome Announc* 6:e00132-18. <https://doi.org/10.1128/genomeA.00132-18>.
- Karimi E, Keller-Costa T, Slaby BM, Cox CJ, da Rocha UN, Hentschel U, Costa R. 2019. Genomic blueprints of sponge-prokaryote symbiosis are shared by low abundant and cultivatable *Alphaproteobacteria*. *Sci Rep* 9:1999. <https://doi.org/10.1038/s41598-019-38737-x>.
- Bankevich A, Nurk S, Antipov D, Gurevich AA, Dvorkin M, Kulikov AS, Lesin VM, Nikolenko SI, Pham S, Prjibelski AD, Pyshkin AV, Sirotkin AV, Vyahhi N, Tesler G, Alekseyev MA, Pevzner PA. 2012. SPAdes: a new genome assembly algorithm and its applications to single-cell sequencing. *J Comput Biol* 19:455–477. <https://doi.org/10.1089/cmb.2012.0021>.
- Huntemann M, Ivanova NN, Mavromatis K, Tripp HJ, Paez-Espino D, Palaniappan K, Szeto E, Pillay M, Chen IM, Pati A, Nielsen T, Markowitz VM, Kyrpides NC. 2015. The standard operating procedure of the DOE-JGI Microbial Genome Annotation Pipeline (MGAP v.4). *Stand Genomic Sci* 10:86. <https://doi.org/10.1186/s40793-015-0077-y>.
- Chen IA, Chu K, Palaniappan K, Ratner A, Huang J, Huntemann M, Hajek P, Ritter S, Varghese N, Seshadri R, Roux S, Woyke T, Eloe-Fadrosh EA, Ivanova NN, Kyrpides NC. 2021. The IMG/M data management and analysis system v.6.0: new tools and advanced capabilities. *Nucleic Acids Res* 49:D751–D763. <https://doi.org/10.1093/nar/gkaa939>.
- Rodríguez-R LM, Gunturu S, Harvey WT, Rosselló-Mora R, Tie die JM, Cole JR, Konstantinidis KT. 2018. The Microbial Genomes Atlas (MiGA) web-server: taxonomic and gene diversity analysis of Archaea and Bacteria at the whole genome level. *Nucleic Acids Res* 46:W282–W288. <https://doi.org/10.1093/nar/gky467>.
- Blin K, Shaw S, Steinke K, Villebro R, Ziemert N, Lee SY, Medema MH, Weber T. 2019. AntiSMASH 5.0: updates to the secondary metabolite genome mining pipeline. *Nucleic Acids Res* 47:W81–W87. <https://doi.org/10.1093/nar/gkz310>.
- Chasteen TG, Fuentes DE, Tantaleán JC, Vásquez CC. 2009. Tellurite: history, oxidative stress, and molecular mechanisms of resistance. *FEMS Microbiol Rev* 33:820–832. <https://doi.org/10.1111/j.1574-6976.2009.00177.x>.
- Raimundo I, Silva R, Meunier L, Valente SM, Lago-Lestón A, Keller-Costa T, Costa R. 2021. Functional metagenomics reveals differential chitin degradation and utilization features across free-living and host-associated marine microbiomes. *Microbiome* 9:43. <https://doi.org/10.1186/s40168-020-0970-2>.
- Karimi E, Slaby BM, Soares AR, Blom J, Henstchel U, Costa R. 2018. Metagenomic binning reveals versatile nutrient cycling and distinct adaptive features in alphaproteobacterial symbionts of marine sponges. *FEMS Microbiol Ecol* 94:fiy074. <https://doi.org/10.1093/femsec/fiy074>.