

Evolution of Secondary Metabolite Genes in Three Closely Related Marine Actinomycete Species^{∇†}

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The marine actinomycete genus *Salinispora* is composed of three closely related species. These bacteria are a rich source of secondary metabolites, which are produced in species-specific patterns. This study examines the distribution and phylogenetic relationships of genes involved in the biosynthesis of secondary metabolites in the salinosporamide and staurosporine classes, which have been reported for *S. tropica* and *S. arenicola*, respectively. The focus is on “*Salinispora pacifica*,” the most recently discovered and phylogenetically diverse member of the genus. Of 61 *S. pacifica* strains examined, 15 tested positive for a ketosynthase (KS) domain linked to the biosynthesis of salinosporamide K, a new compound in the salinosporamide series. Compound production was confirmed in two strains, and the domain phylogeny supports vertical inheritance from a common ancestor shared with *S. tropica*, which produces related compounds in the salinosporamide series. There was no evidence for interspecies recombination among *salA* KS sequences, providing further support for the geographic isolation of these two salinosporamide-producing lineages. In addition, staurosporine production is reported for the first time for *S. pacifica*, with 24 of 61 strains testing positive for *staD*, a key gene involved in the biosynthesis of this compound. High levels of recombination were observed between *staD* alleles in *S. pacifica* and the cooccurring yet more distantly related *S. arenicola*, which produces a similar series of staurosporines. The distributions and phylogenies of the biosynthetic genes examined provide insight into the complex processes driving the evolution of secondary metabolism among closely related bacterial species.

Microbial secondary metabolism is the source of many of today's most useful pharmaceutical agents. Bacteria in the order *Actinomycetales* have been particularly important in this regard, accounting for over 50% of the bioactive microbial compounds discovered as of 2002 (1). Studies of bacterial secondary metabolism have largely targeted the discovery of new compounds and the mechanisms of their biosynthesis. As a result, we know relatively little about the ecological functions of secondary metabolites or the evolutionary histories of the biosynthetic genes responsible for their production. In cases where functions have been addressed, it is clear that secondary metabolites can act as allelochemicals, signaling molecules, and siderophores (3), while it has been suggested that suites of compounds can work synergistically against competitors (2). The evolutionary histories of the associated biosynthetic genes are proving to be equally complex (17) yet are providing new opportunities to incorporate phylogenetics into the discovery process (14).

Secondary metabolites are generally produced by large gene collectives that can exceed 100 kb and include genes involved in regulation, resistance, and transport (4, 11). The horizontal exchange of genes in these pathways is well documented (18, 21, 23, 26) and provides a rapid mechanism for bacteria to test the selective advantage afforded by the small molecule product(s) of complex biosynthetic pathways (12, 23). Documenting

the distributions of specific biosynthetic pathways among bacteria has the potential to add new insight into the extent to which these pathways are exchanged and how gene clusters evolve to create new chemical diversity.

We have been studying the ecology and secondary chemistry of the marine actinomycete genus *Salinispora*. To date, *S. tropica* and *S. arenicola* have been formally described (24), while a third species, “*Salinispora pacifica*,” has been proposed (19). The three species share 99% 16S rRNA gene sequence identity (19) and thus are at the limits of resolution attainable with this taxonomic marker. They also display different geographic distributions, with reports to date indicating that *S. tropica* is restricted to the Caribbean, *S. pacifica* occurs worldwide except for the Caribbean, and *S. arenicola* is broadly distributed and cooccurs with both species (19).

The genus *Salinispora* has proven to be a robust source of secondary metabolites, which represent the largest class of functional traits differentiating the species (30). These differences include the consistent production of compounds in the salinosporamide class by *S. tropica* and in the staurosporine class by *S. arenicola* (20). Among these compounds, salinosporamide A is a potent proteasome inhibitor that is currently advancing through clinical trials as an anticancer agent (10). Salinosporamide A was originally discovered from *S. tropica* strain CNB-392 (9), which subsequently yielded additional compounds in this series (9, 36). The staurosporines are a well-known class of protein kinase inhibitors (34) that were originally discovered from a *Streptomyces* sp. (13, 34). The consistent production of specific classes of secondary metabolites by different *Salinispora* species was used to support the ecological importance of secondary metabolism and to link this functional trait to unresolved ecological differences among the

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species. Species-specific secondary metabolite production has also been documented in fungi, and it has been suggested that this trait may be used as a taxonomic marker (22).

Mining of the draft genome sequence of *S. pacifica* strain CNT-133 led to the recent discovery of salinosporamide K (7), a new compound in the salinosporamide series. Salinosporamide K lacks the chloro-ethyl side chain at the C-2 position of the salinosporamide bicyclic ring system. As expected, the associated salinosporamide biosynthetic pathway (*Sp_sal*) characterized from this strain lacked the *salL* chlorinase (8) and associated genes responsible for the creation of the unique polyketide synthase (PKS) extender unit observed in the *S. tropica* *St_sal* pathway (6). The exact replacement of these genes with transposases in *S. pacifica* strain CNT-133, coupled with the conservation of the remainder of the biosynthetic genes in the two pathways (7), suggests gene loss in *S. pacifica* as opposed to gene gain in *S. tropica*. However, it is not clear if this genetic difference is a consistent feature of *S. pacifica*. In addition, preliminary chemical studies of *S. pacifica* provided the first evidence of staurosporine production in this species. These observations, coupled with the isolation of the salinosporamide-related cinnabaramide series in a *Streptomyces* strain (32) and the detection of the staurosporine gene cluster on a *Streptomyces* giant linear plasmid (25), raised new questions about the distributions and evolutionary histories of the associated biosynthetic pathways in *Salinispora* species.

The aims of this study were to examine the distributions and phylogenies of genes involved in the biosynthesis of compounds in the salinosporamide and staurosporine classes among *Salinispora* species. The resulting data were used to assess the roles of vertical inheritance, recombination, and horizontal gene transfer in the evolutionary histories of these biosynthetic genes. The results provide evidence of the complex processes driving the evolution of secondary metabolism in three closely related bacterial species.

MATERIALS AND METHODS

Strains and nucleic acid extraction. The 61 *S. pacifica* strains used in this study were cultured from marine sediment samples and identified based on 16S rRNA gene sequencing as previously described (7, 19). In addition, six *S. arenicola* strains and four *S. tropica* strains were also included in the analyses. Genomic DNA was extracted according to the DNeasy protocol (Qiagen Inc., Valencia, CA), with previously described changes (15), and used immediately or stored at -20°C .

Gene amplification and sequencing. Specific and degenerate primer sets were designed based on the ketosynthase (KS) domains identified in the salinosporamide polyketide synthase gene *salA* in *S. tropica* (CNB-440) and *S. pacifica* (CNT-133) (7, 35). The specific primer set *salAks2F* (5'-GCGGAAATCGACGATACGT-3') and *salAks2R* (5'-TCCACATAGTCTACGAGCCA-3') targeted approximately 700 bp, while the degenerate primer set *salAks3F* (5'-CATMGCRCCTGGYARCCTCG-3') and *salAks3R* (5'-TYCACRTAGCTRCGASCCA-3') targeted approximately 750 bp. Each PCR consisted of a 50- μl mixture containing 10 \times PCR buffer (Applied Biosciences, Foster City, CA), 2.5 mM MgCl_2 (Applied Biosciences), 0.7% dimethyl sulfoxide (DMSO), 10 mM deoxynucleoside triphosphates (dNTPs), 1.5 U of AmpliTaq Gold DNA polymerase (Applied Biosciences), and 18.75 μmol of each primer. The program for the PCR included a primary denaturation step at 95°C for 15 min, followed by 30 cycles of 95°C for 1 min, 60°C for 1 min (55°C for the degenerate primers), and 72°C for 1 min, followed by a final extension at 72°C for 7 min. *S. pacifica* strain CNT-133 was used as a positive control, while *S. arenicola* strain CNS-205, which does not possess the *sal* pathway (30), was used as a negative control for all PCRs. PCR products were purified using the Zymo DNA clean and concentrator purification kit (Zymo Research Incorporated, Irvine, CA) and sequenced using the *salAks3F* primer at SeqXcel, Sorrento Valley, CA. Previously published primers

targeting the *salL* chlorinase (8), which is involved in the biosynthesis of salinosporamide A, were tested on all strains that yielded a sequence-verified *salA* KS product. The PCR conditions were as described above (annealing temperature, 55°C).

The specific primer set *staD2F* (5'-TGTGGGGSCACTACAACGA-3') and *staD1R* (5'-SGGRTCGCACATCTGCCAGAT-3') was designed based on an alignment that included *staD* sequences from *S. arenicola* strain CNS-205 (NC_009953), two *Streptomyces* spp. (accession no. AB071406 and AB088119), two *S. arenicola* draft genomes, and homologs of *rebD* (accession no. AB090952) and *vioB* (accession no. GQ266676 and AF172851). The PCR reagents were the same as those listed above, with the addition of 0.5 μl of bovine serum albumin (New England BioLabs, Inc., Beverly, MA). A touchdown PCR that consisted of an initial soak at 94°C for 12 min, followed by 3 cycles of 94°C for 1 min, 67°C for 1 min, and 72°C for 1 min, followed by 27 additional cycles of 94°C for 1 min, 62°C for 1 min, and 72°C for 1 min, with a final extension at 72°C for 10 min, was used. Appropriately sized PCR products were purified and sequenced as described above using the *staD2F* forward primer.

Phylogenetic analyses. Nucleotide sequences were edited using the Sequencher software package (version 4.5; Gene Codes Co., Ann Arbor, MI), aligned using the Muscle software program (5), and visually edited using the MacClade program (version 4.07; Dave and Wayne Maddison, Sinauer Associates, Inc., Sunderland, MA). Maximum-likelihood, maximum-parsimony, and neighbor-joining phylogenetic trees were created using the PAUP (33) and Phylml (maximum-likelihood methods) (16) software programs. Bootstraps were calculated using 1,000 replicates and resampling at all sites.

Fermentation/extraction and LC-MS analyses. Initial liquid chromatography-mass spectrometry (LC-MS) analyses of five *S. pacifica* strains that tested positive for the *salA* KS domain did not reveal the presence of salinosporamide K. Of these, the strain with the best growth (CNS-863) was selected for more-detailed studies, along with strain CNT-133 (positive control), the original source of this compound. Both strains were cultured in 2.8-liter Fernbach flasks containing 1 liter of medium A1BFe+C (10 g starch, 4 g yeast extract, 2 g peptone, 1 g CaCO_3 , 40 mg $\text{Fe}_2(\text{SO}_4)_3 \cdot 4\text{H}_2\text{O}$, 100 mg KBr, and 1 liter seawater) and shaken at 230 rpm and 27°C . Autoclaved XAD-7 resin (20 g) was added to the culture after 24 h. After 6 days, the resin was collected by filtration through cheesecloth, washed with deionized water, and eluted with acetone. The acetone was removed under reduced pressure, and the resulting aqueous layer was extracted with ethyl acetate (two times, 300 ml). The ethyl acetate soluble fraction was dried under vacuum to obtain a crude extract, which was fractionated by silica gel flash chromatography, eluting with increasing amounts of acetonitrile (CH_3CN) in dichloromethane (CH_2Cl_2) (100% CH_2Cl_2 and 100:1, 50:1, 20:1, 5:1, 1:1, and 100% CH_3CN). All fractions were subjected to LC-MS analysis using a Hewlett-Packard series 1100 LC-MS system with a reversed-phase C_{18} column (Phenomenex Luna, 4.6 mm by 100 mm; pore size, 5 μm) using a solvent gradient from 5% to 100% CH_3CN over 23 min, a flow rate of 0.7 ml/min, and UV detection. Salinosporamide K production was determined by retention time and comparison of UV and mass spectral data with an authentic standard. Low-resolution mass data were obtained in the positive mode (ESI voltage, 6.0 kV; capillary temperature, 200°C ; auxiliary and sheath gas pressure, 5 units and 70 lb/in 2 , respectively).

For staurosporine production, *S. pacifica* strain CNS-863 and *S. arenicola* strain CNS-205 (positive control) were grown in 25 ml of medium A1 (10 g starch, 4 g yeast extract, 2 g peptone, and 1 liter seawater) in 125-ml Erlenmeyer flasks for 3 to 7 days prior to transfer to 1 liter of medium A1BFe+C in 2.8-liter Fernbach flasks. All cultures were grown with shaking at 230 rpm at 25 to 27°C for 6 days and then extracted once with 1 liter of ethyl acetate. The organic layers were separated, concentrated to dryness under vacuum, and fractionated by silica gel flash chromatography, eluting with increasing amounts of methanol (CH_3OH) in dichloromethane (CH_2Cl_2) (100% CH_2Cl_2 and 100:1, 50:1, 20:1, 5:1, 1:1, and 100% CH_3OH). The fractions containing staurosporines were combined and analyzed by LC-MS as described above (4.6- by 250-mm column; UV = 292 nm) using an isocratic solvent system of 35% CH_3CN in water with 0.1% trifluoroacetic acid. Compounds were identified as staurosporines by comparison with data from Antibase (Wiley-VCH Verlag GmbH & Co) and standards stored in an internal database.

RESULTS

***salA* KS distributions and phylogeny.** Sixty-one *S. pacifica* isolates derived from 34 independent marine sediment samples, collected from six geographically distinct sites, were screened by PCR for sequences related to the KS domains in

TABLE 1. Strains used in this study^a

Strain no.	Species	Location	Yr	16S rRNA type ^b	16S rRNA accession no. ^d	Positive for <i>salA</i> KS	KS accession no. ^d	Positive for <i>staD</i>	<i>staD</i> accession no. ^d
CNB-440	<i>S. tropica</i>	BA	1989	~ ^c	AY040617	Yes	HQ642869	No	NP
CNH-898	<i>S. tropica</i>	BA	2000	~	HQ642876	Yes	HQ642870	NT	~
CNR-699	<i>S. tropica</i>	BA	2003	~	JN161822	Yes	HQ642872	NT	~
CNS-193	<i>S. tropica</i>	BA	2004	~	JN161823	Yes	HQ642871	No	~
CNH-643	<i>S. arenicola</i>	BA	1999	~	AY371897	No	~	Yes	HQ642932
CNP-173	<i>S. arenicola</i>	USVI	2001	~	HQ642847	NT	~	Yes	HQ642933
CNR-040	<i>S. arenicola</i>	Guam	2002	~	JN161824	NT	~	Yes	HQ642934
CNR-581	<i>S. arenicola</i>	Guam	2002	~	JN161825	NT	~	Yes	HQ642935
CNS-205	<i>S. arenicola</i>	Palau	2004	~	CP000850	No	~	Yes	HQ642963
CNS-673	<i>S. arenicola</i>	Fiji	2006	~	JN161826	NT	~	Yes	HQ642936
CNT-005	<i>S. arenicola</i>	Fiji	2006	~	JN161827	NT	~	Yes	HQ642937
CNT-850	<i>S. arenicola</i>	Hawaii	2008	~	HQ642848	NT	~	Yes	HQ642938
CNR-551	<i>S. pacifica</i>	Guam	2002	A	HQ642881	Yes	HQ642854	No	~
CNR-942	<i>S. pacifica</i>	Palau	2004	E	HQ642877	Yes	HQ642855	No	~
CNS-055	<i>S. pacifica</i>	Palau	2004	A	DQ224159	No	~	Yes	HQ642939
CNS-103	<i>S. pacifica</i>	Palau	2004	~	DQ224160	Yes	HQ642856	No	~
CNS-237	<i>S. pacifica</i>	Palau	2004	B	HQ642850	No	~	Yes	HQ642940
CNS-251	<i>S. pacifica</i>	Palau	2004	~	HQ642879	Yes	HQ642857	Yes	HQ642941
CNS-735	<i>S. pacifica</i>	Fiji	2006	~	HQ642880	Yes	HQ642859	No	~
CNS-799	<i>S. pacifica</i>	Fiji	2006	C	HQ642895	Yes	HQ642860	Yes	HQ642942
CNS-844	<i>S. pacifica</i>	Fiji	2006	~	HQ642897	No	~	Yes	HQ642943
CNS-860	<i>S. pacifica</i>	Fiji	2006	C	HQ642886	No	~	Yes	HQ642944
CNS-863	<i>S. pacifica</i>	Fiji	2006	C	HQ642851	Yes	HQ642858	Yes	HQ642945
CNS-890	<i>S. pacifica</i>	Fiji	2006	C	HQ642889	No	~	Yes	HQ642946
CNS-996	<i>S. pacifica</i>	Fiji	2006	C	HQ642888	No	~	Yes	HQ642947
CNT-029	<i>S. pacifica</i>	Fiji	2006	F	HQ642852	No	~	Yes	HQ642948
CNT-044	<i>S. pacifica</i>	Fiji	2006	C	HQ642893	No	~	Yes	HQ642949
CNT-045	<i>S. pacifica</i>	Fiji	2006	C	HQ642887	Yes	HQ642861	Yes	HQ642950
CNT-084	<i>S. pacifica</i>	Fiji	2006	D	HQ642882	Yes	HQ642862	Yes	HQ642951
CNT-094	<i>S. pacifica</i>	Fiji	2006	C	HQ642894	No	~	Yes	HQ642952
CNT-124	<i>S. pacifica</i>	Fiji	2006	C	HQ642891	No	~	Yes	HQ642953
CNT-131	<i>S. pacifica</i>	Fiji	2006	~	HQ642896	No	~	Yes	HQ642954
CNT-133	<i>S. pacifica</i>	Fiji	2006	D	HQ218996	Yes	HQ642863	No	~
CNT-138	<i>S. pacifica</i>	Fiji	2006	E	HQ642853	Yes	HQ642864	Yes	HQ642955
CNT-148	<i>S. pacifica</i>	Fiji	2006	A	HQ642899	Yes	HQ642865	Yes	HQ642956
CNT-150	<i>S. pacifica</i>	Fiji	2006	B	HQ642900	No	~	Yes	HQ642957
CNT-569	<i>S. pacifica</i>	Fiji	2008	D	HQ642885	Yes	HQ642866	No	~
CNT-584	<i>S. pacifica</i>	Fiji	2008	C	HQ642890	No	~	Yes	HQ642958
CNT-603	<i>S. pacifica</i>	Fiji	2008	~	HQ642878	Yes	HQ642867	Yes	HQ642959
CNT-609	<i>S. pacifica</i>	Fiji	2008	D	HQ642883	Yes	HQ642868	Yes	HQ642960
CNT-853	<i>S. pacifica</i>	Hawaii	2008	D	HQ642884	No	~	Yes	HQ642962
CNT-854	<i>S. pacifica</i>	Hawaii	2008	C	HQ642892	No	~	Yes	HQ642961

^a Of the 61 total strains that were screened, only those that tested positive for either the *salA* KS or *staD* gene are listed. NT, not tested; BA, Bahamas; USVI, U.S. Virgin Islands.

^b The original sequence type identified for each species was not assigned a letter.

^c ~, not applicable.

^d Accession numbers are from GenBank.

the *salA* polyketide synthase genes in *S. tropica* (35) and *S. pacifica* (7). This gene is associated with the biosynthesis of salinosporamide A in *S. tropica* (6) and salinosporamide K in *S. pacifica* (7). The homolog *cinA* was recently identified in *Streptomyces cinnabarinus* as part of the pathway responsible for the biosynthesis of the salinosporamide-related cinnabaramide series (31). In total, the *salA* KS domain was detected in 15 of 61 *S. pacifica* strains, including 5 of 7 16S rRNA gene sequence types originating from three of the six locations sampled (Guam, Palau, and Fiji) (Table 1). The same 15 strains tested positive using both specific and degenerate PCR primers. *SalA* KS sequences were also obtained from three additional *S. tropica* strains derived from the Bahamas to complement that originally found in strain CNB-440 (35). The *S. pacifica* KS sequences share 84 to 85% nucleotide identity with the *S.*

tropica sequences and 99 to 100% nucleotide identity with each other. The key amino acids VDTACSSSLVAVHLACQS, involved in forming the binding pocket of the KS domain (28), are conserved in all of the sequences.

The 16S rRNA tree (Fig. 1) depicts the previously reported relationship of *S. tropica* and *S. pacifica* as sister taxa that share an ancestor with *S. arenicola* (20). The seven *S. pacifica* 16S sequence types included in this study (Fig. 1) represent more than half of the 13 reported to date for this species. Phylogenetic analyses of the *Salinispora salA* KS sequences reveal two well-supported lineages that are congruent with the 16S tree (Fig. 2). There is no evidence for interspecies recombination at this locus. However, the detection of clonal KS sequences among different *S. pacifica* 16S sequence types (e.g., strains CNR-551, CNS-251, CNS-799, and CNS-863) reveals consid-

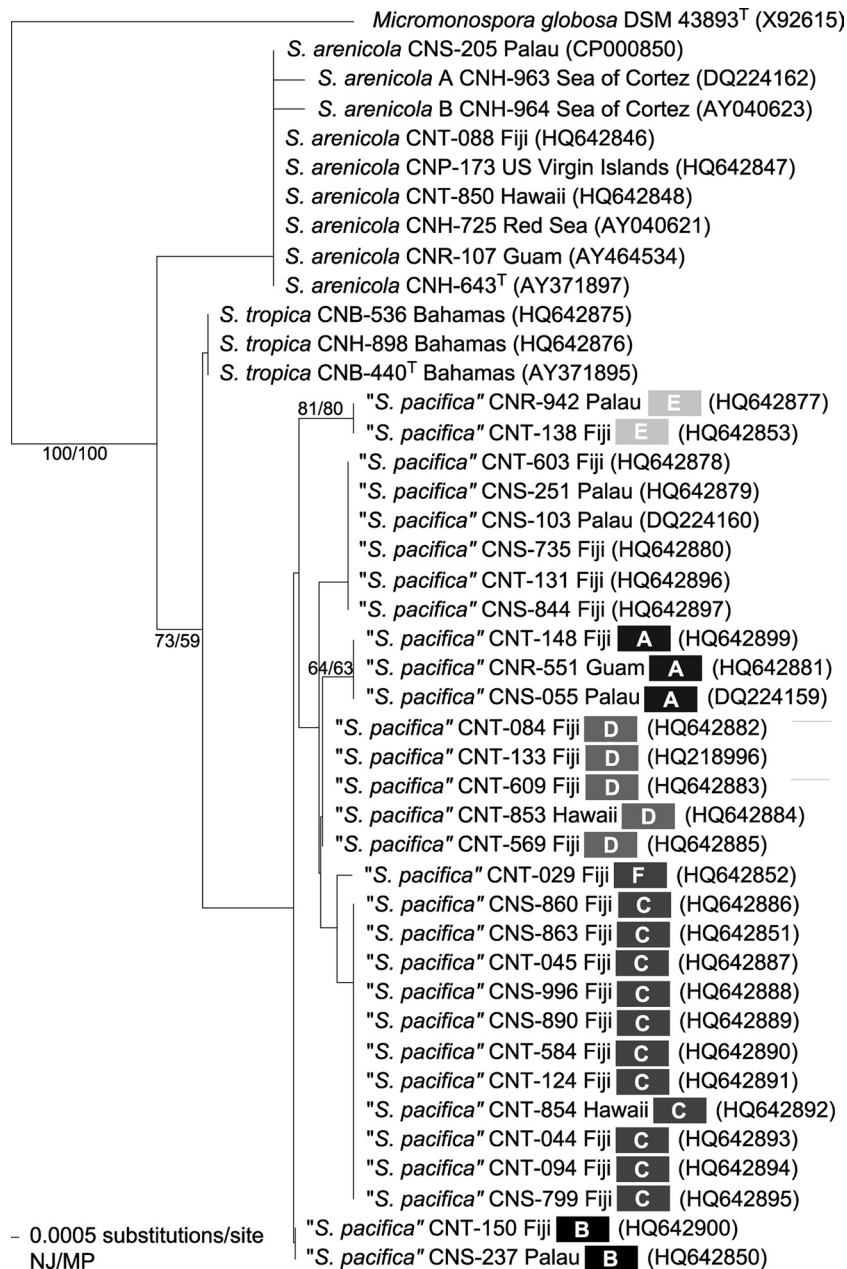


FIG. 1. 16S rRNA gene phylogeny. Neighbor-joining phylogenetic tree based on 512 nucleotide positions from strains that yielded sequence-verified *salA* KS or *staD* PCR products. Species names are followed by the strain identifier, source location, 16S sequence type (*S. pacifica* only, grayscale boxes, A to F with original sequence type not labeled), and accession number (in parentheses). Bootstrap values >60% for neighbor-joining and maximum-parsimony trees are shown for 1,000 replicates at the respective nodes. *Micromonospora globosa* was used to root the tree.

erable levels of intraspecific recombination. In addition, the KS sequence from the recently characterized cinnabaramide biosynthetic pathway (31) displays a close phylogenetic relationship with the *Salinispora* species KS sequences (75 to 78% nucleotide identity) despite originating from *Streptomyces cinnabarinus*, which belongs to a separate family in the *Actinomycetales*. KS sequences that share >68% nucleotide identity with those observed in *S. pacifica* have also been reported from taxonomically diverse actinomycetes and myxobacteria (Fig. 2) and depict a phylogeny that is highly incongruent with the

taxonomic relationships of these organisms. These results provide strong evidence that the KS sequences associated with the biosynthesis of compounds in the salinosporamide class have been subjected to horizontal gene transfer (HGT). *SalA* KS sequences were not detected in one complete (30) and two draft (unpublished data) *S. arenicola* genome sequences. In addition, compounds in the salinosporamide class were not detected among 30 *S. arenicola* strains examined in a prior study (20). Taken together, these results provide strong evidence that the *sal* pathway was acquired prior to the *S. tropica* and *S. pacifica* spe-

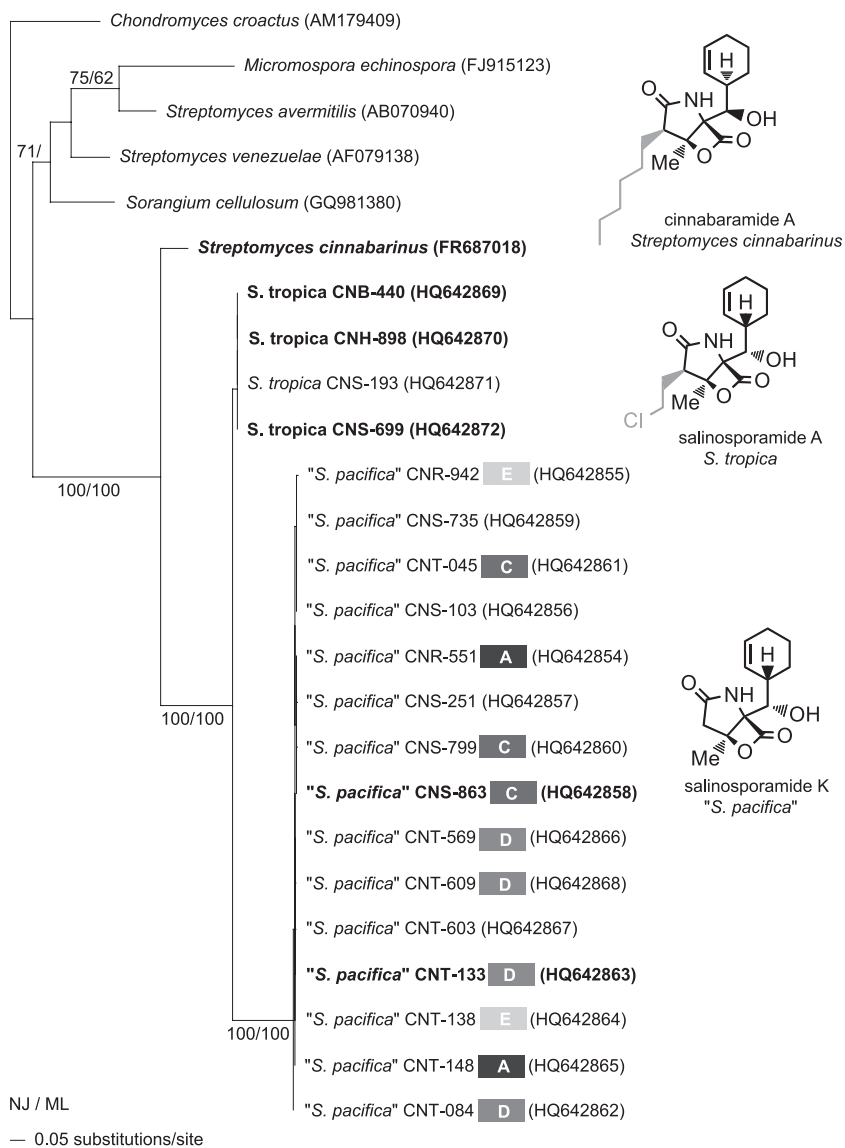


FIG. 2. *salA* KS phylogeny. Neighbor-joining phylogenetic tree based on 625 nucleotide positions. Species names are followed by the strain identifier, 16S sequence type (*S. pacifica* only, grayscale boxes, A and C to E with original sequence type not labeled), and accession number (in parentheses). Structures of compounds produced by *S. tropica*, *S. pacifica*, and *Streptomyces cinnabarinus* are shown next to their respective lineages, with differences in the C-2 substitution patterns indicated in gray. Confirmed producing strains are in bold. Bootstrap values >60% for neighbor-joining and maximum-parsimony trees are shown for 1,000 replicates at the respective nodes. A KS sequence from *C. croactus* was used to root the tree.

ciation event and subsequently evolved independently in these two lineages.

Staurosporine *staD* distributions and phylogeny. Twenty-four of the 61 *S. pacifica* strains examined, including five of seven 16S sequence types, generated sequence-verified PCR products using primers specific for the dichlorochromopyrrolic acid synthetase gene *staD* (Table 1), which is involved in the biosynthesis of the staurosporine aglycone (29). The six *S. arenicola* strains tested also yielded sequence-verified *staD* products, as expected given that this species is known to consistently produce compounds in this class (20). The *S. arenicola* sequences complement those originally observed in the CNS-205 genome (accession number NC_009953) and subsequently

in a draft genome sequence of *S. arenicola* strain CNH-643. None of the four *S. tropica* strains tested yielded a product with the *staD* primer set employed, nor was this gene observed in the genome sequence of *S. tropica* strain CNB-440 (accession number NC_009380).

The phylogeny of the *S. arenicola* and *S. pacifica* *staD* sequences reveals two well-supported clades, as would be expected if they had been inherited from a common ancestor and subsequently evolved independently in the two lineages (Fig. 3). Unlike the *salA* KS sequences, however, the *staD* sequences observed in *S. pacifica* display considerable diversity. More importantly, 7 examples of interspecies recombination are observed out of 32 sequences examined (22%). In each case (e.g., strains CNT-603

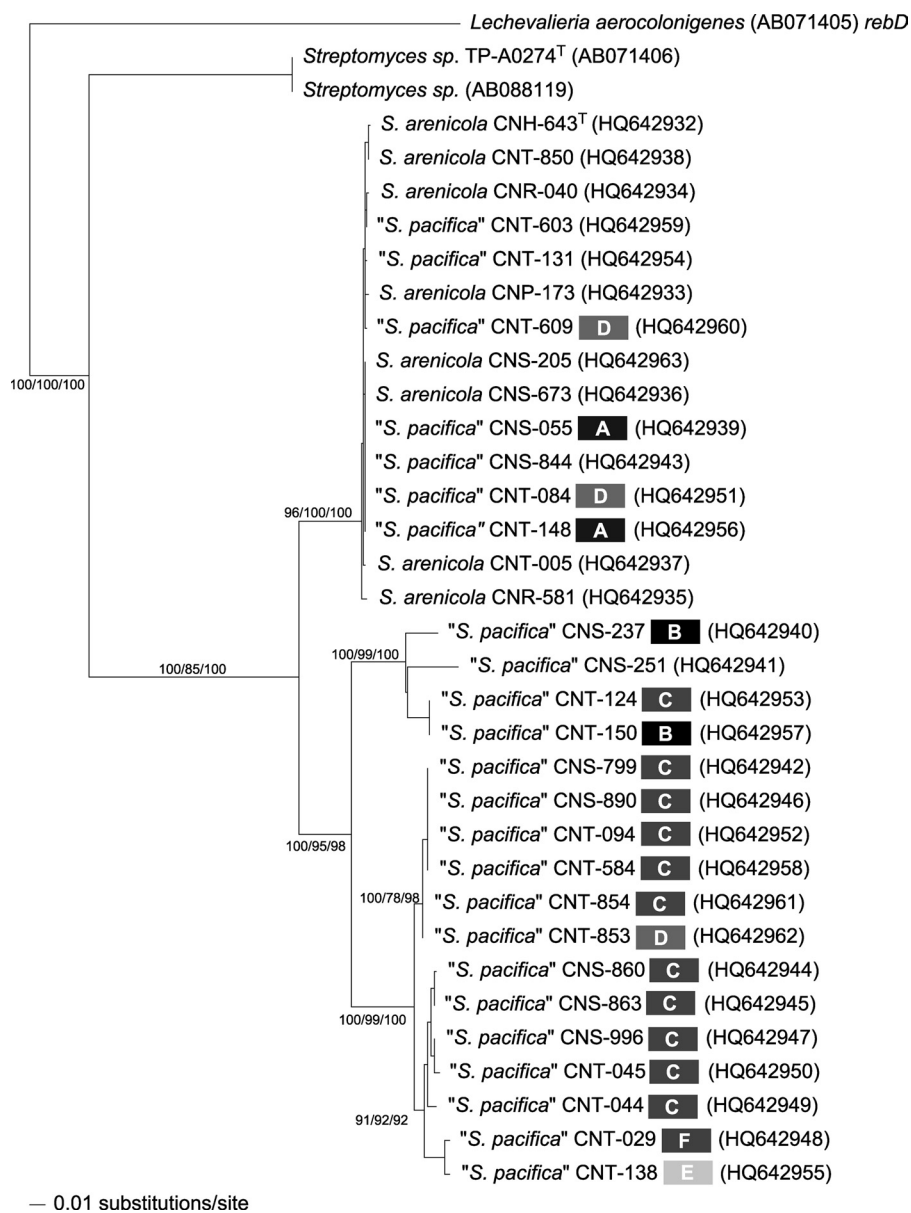


FIG. 3. *staD* phylogeny. Neighbor-joining phylogenetic tree based on 741 nucleotide positions. Species names are followed by the strain identifier, 16S sequence type (*S. pacifica* only, grayscale boxes, A to F with original sequence type not labeled), and accession number (in parentheses). Bootstrap values >60% for neighbor-joining, maximum-likelihood, and maximum-parsimony trees are shown for 1,000 replicates at the respective nodes. A homolog associated with rebeccamycin biosynthesis (*rebD*) was used to root the tree.

and CNT-131), an *S. pacifica* 16S sequence type groups within the *S. arenicola* *staD* lineage and, in six of seven cases, maintains an allele that is also observed in *S. arenicola*. There were no examples where a strain identified as *S. arenicola* based on 16S sequence maintained an *S. pacifica* *staD* allele or grouped within the *S. pacifica* lineage. The previously documented cooccurrence of *S. arenicola* and *S. pacifica* (19) provides spatial opportunities for these interspecies recombination events to occur. As in the *salA* KS tree, there are multiple examples where different *S. pacifica* 16S sequence types maintain the same *staD* allele (e.g., strains CNT-124 and CNT-150), providing evidence that intraspecific recombination has also occurred at this locus. The remarkably high level of *staD* and 16S rRNA gene sequence conservation

within *S. arenicola* (>99% at both loci) relative to *S. pacifica* provides evidence of extensive recombination or a recent period selection event in this species.

LC-MS screening. Among the 15 *S. pacifica* strains that tested positive for the *salA* KS sequence, salinosporamide K production was confirmed in strain CNS-863 (Fig. 4), in addition to strain CNT-133, the original source of this compound (7). Due to low yields, fractionation of the crude extract was required before compound production could be confirmed based on retention time, UV spectrum, and mass, all of which matched an authentic standard. The production of salinosporamide A was not observed in either strain. Probing all 15 KS-positive *S. pacifica* strains failed to detect *sall*, encoding

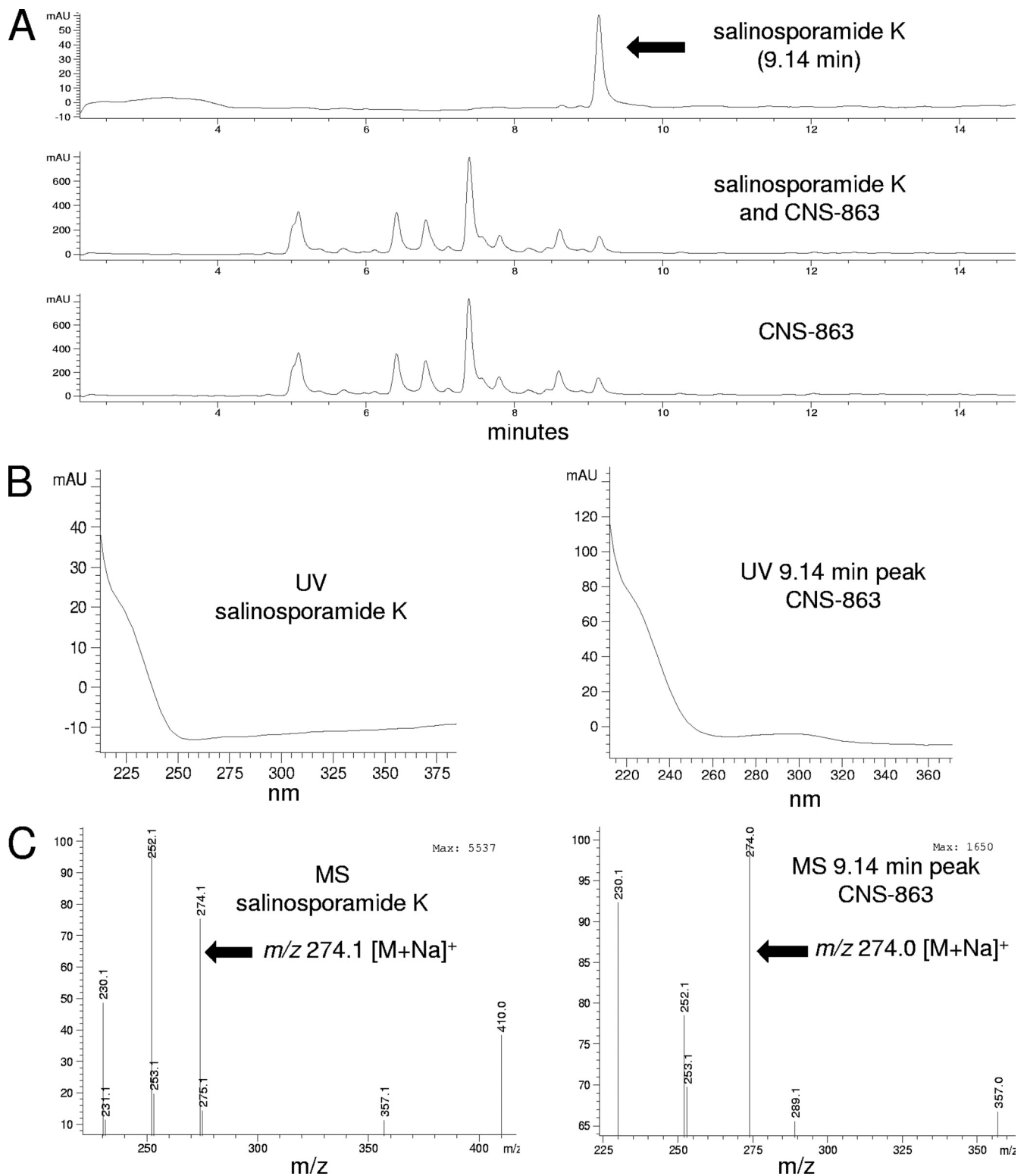


FIG. 4. LC-MS data supporting salinosporamide K production. (A) LC trace for authentic salinosporamide K, a chromatography fraction obtained from a culture extract of strain CNS-863 coinjected with salinosporamide K, and the CNS-863 fraction alone. (B) UV trace of salinosporamide K and the peak observed at 9.14 min in strain CNS-863. (C) Low-resolution mass data of authentic salinosporamide K and the 9.14-min peak.

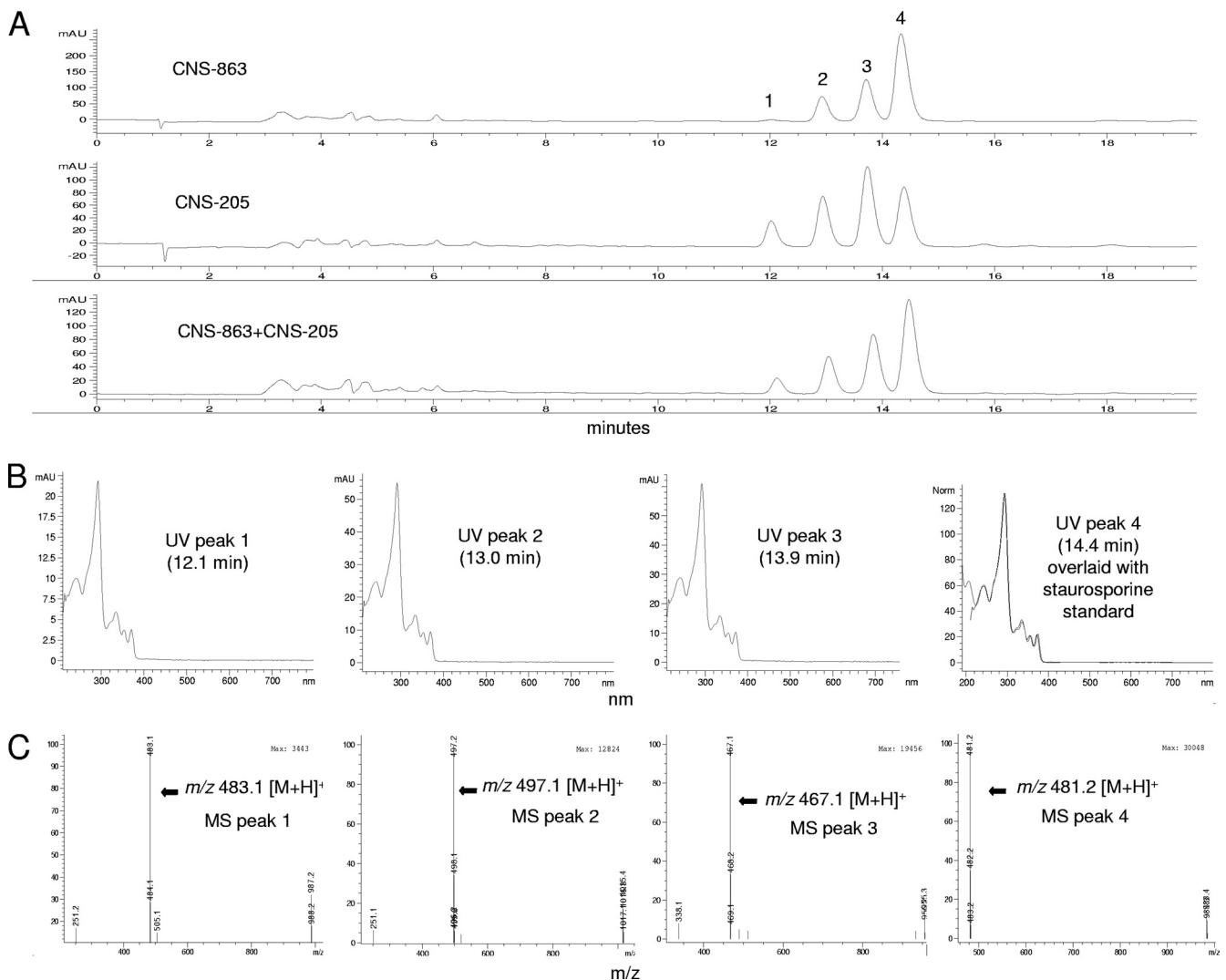


FIG. 5. LC-MS data supporting staurosporine production. (A) LC trace for *S. pacifica* strain CNS-863 and *S. arenicola* CNS-205 (positive control) with detection at the UV maximum for staurosporines (292 nm). The similar retention times of the four peaks in both strains supports the production of the same compounds. (B) UV data for peaks 1 to 4 in strain CNS-863. The UV spectrum for peak 4 is overlaid with that of a staurosporine standard with a mass of 480. (C) MS data for peaks 1 to 4 in strain CNS-863. All of these masses correspond to previously reported staurosporines. Similar UV and mass data were recorded for CNS-205.

the chlorinase associated with the biosynthesis of the chlorinated extender unit in salinosporamide A (6, 8). This result provides additional support for the occurrence of the salinosporamide K, as opposed to the salinosporamide A, pathway in the 15 *S. pacifica* strains.

Of the 24 *S. pacifica* isolates that tested positive for *staD*, an organic extract from strain CNS-863 was analyzed by LC-MS to confirm the presence of compounds in the staurosporine class. This extract contained three major peaks (Fig. 5), with masses that correspond to those of various staurosporine analogs (see Fig. S1 in the supplemental material). Although peak 1 is not clearly visible in the CNS-863 trace, the UV and mass data confirm the presence of the same compounds that were observed in strain CNS-205. Thus, both strains appeared to produce the same staurosporines but in different relative amounts. The UV characteristics of all of these compounds were identical and matched with 99% identity to a staurospo-

rine standard (Fig. 5). Because each peak could be associated with as many as five staurosporine analogues that possess identical mass and UV properties, it was not possible to assign precise structures to the compounds produced by strain CNS-863 using the methods employed.

DISCUSSION

Secondary metabolites are nonessential metabolic products believed to impart significant positive effects on the fitness and ecology of the bacterial populations that produce them. These compounds are the products of large gene collectives that are subject to horizontal gene transfer (12, 18) and whose distributions among closely related bacterial populations remain largely unknown. Here we report the distributions and phylogenetic relationships of genes involved in the biosynthesis of two classes of biologically active secondary metabolites in the

marine actinomycete genus *Salinispora*. The results provide evidence for both HGT and vertical inheritance and a glimpse into the complexity of secondary metabolite evolution in closely related bacterial species.

The genus *Salinispora* has proven to be an interesting model with which to address questions about secondary metabolite production and its relationship to species-level taxonomic assignments. The recent discovery of salinosporamide K from *S. pacifica* strain CNT-133 (7) was unexpected considering that compounds in this class had previously been found exclusively from *S. tropica* (20). Given that *S. tropica* and *S. pacifica* are sister taxa, this observation provided the opportunity to test the hypothesis that the associated biosynthetic pathway was inherited from a common ancestor and subsequently evolved independently in the two species. The occurrence of the *salA* KS domain in a majority of the *S. pacifica* 16S rRNA sequence types, coupled with the congruence of the 16S (Fig. 1) and KS (Fig. 2) trees, provides support for this hypothesis. The detection of salinosporamide K in *S. pacifica* strain CNS-863 provides a link between the *salA* KS sequence and the presence of the complete *Sp_sal* biosynthetic pathway. It also provides additional support for the lineage specificities of salinosporamides A and K in *S. tropica* and *S. pacifica*, respectively. Further support for lineage specificity comes from a prior study in which salinosporamide A production was observed in 6 *S. tropica* strains but not in 41 *S. pacifica* and *S. arenicola* strains (20). Finally, the *salL* chlorinase, which is essential for the production of salinosporamide A in *S. tropica* (6), was not detected in any of the 15 KS-positive *S. pacifica* strains. These results support the hypothesis that the *S. pacifica* lineage maintains a salinosporamide pathway that is devoid of the genes associated with the biosynthesis of the ethyl chloride moiety observed in salinosporamide A. Although it cannot be determined if these genes were lost in *S. pacifica* or gained in *S. tropica*, the presence of transposases in the *S. pacifica* *Sp_sal* pathway at the precise locations where the genes responsible for the biosynthesis of the PKS extender unit occur in the *S. tropica* *St_sal* pathway (7) provides support for gene loss in *S. pacifica*.

Despite evidence for the divergence of the salinosporamide A and K pathways in *S. tropica* and *S. pacifica*, respectively, salinosporamide K and the associated KS sequence proved difficult to detect in many of the *S. pacifica* strains examined. Conversely, salinosporamide A and the associated KS sequence have been consistently observed in *S. tropica*. A number of possible explanations could account for these differences, including failure to access KS templates (14) or differences in the regulatory mechanisms of the two pathways. Another interesting possibility is that the sporadic distribution of the salinosporamide K pathway is linked to the decreased cytotoxic potency of this compound (7) and any associated loss in fitness advantage it confers relative to salinosporamide A. At present, however, the ecological functions of these compounds remain undefined, and therefore any links between cytotoxic potency and selective advantage remain highly speculative (7).

The lack of recombination between the *S. tropica* and *S. pacifica* *salA* KS sequences provides an unexpected line of support for both culture-dependent (19) and culture-independent (27) reports indicating that these two species do not

cooccur. The basal position of the recently characterized and closely related KS sequence from the cinnabaramide biosynthetic pathway (Fig. 2) provides intriguing evidence that the *sal* pathway was acquired prior to the divergence of *S. tropica* and *S. pacifica* and that these pathways have subsequently maintained independent evolutionary trajectories in the two *Salinispora* lineages. Regardless of the evolutionary history of this pathway, the results provide support for a link between geographic isolation and the diversification of secondary metabolite biosynthesis in two closely related species. Based on these results, it can be proposed that the successful cultivation of new *Salinispora* lineages may lead to the discovery of new chemical diversity in the salinosporamide series.

The detection of compounds in the staurosporine class among strains of *S. pacifica* was also surprising given that these compounds had previously been observed exclusively in *S. arenicola* (20). An analysis of the *staD* genes associated with staurosporine biosynthesis in the two *Salinispora* species reveals two distinct lineages, as would be expected of a pathway inherited from a common ancestor. However, the incongruence with the 16S tree includes seven recombination events, which suggests considerable allelic exchange among these two cooccurring species (19). These recombination events were recorded between strains that originated from distant locations, e.g., *S. arenicola* CNS-673 from Fiji and *S. pacifica* CNS-055 from Palau, providing insight into the geographic scales on which these events can be observed. However, given that only 1 of 14 genes in the staurosporine biosynthetic pathway was examined (29), it is not possible to determine the extent to which these recombination events affected the entire operon.

The close relationship of the *staD* gene sequence in a *Streptomyces* sp. and those observed in both *Salinispora* spp. provides strong evidence that this pathway has been exchanged horizontally; however, it is not clear at what point in the *Salinispora* phylogeny it may have been acquired. The *staD* gene was not detected in the *S. tropica* CNB-440 genome (35), in three draft genomes of the closely related genus *Micromonospora*, or in the recently released and closely related *Verrucosipora maris* genome (GenBank accession number NC_015434). Staurosporine also was not observed in previous studies of *S. tropica* secondary metabolism (20). These observations suggest that if *staD* was acquired prior to the divergence of *S. arenicola*, it was subsequently lost in *S. tropica*. Alternatively, it may have been acquired by *S. arenicola* and then horizontally transferred to *S. pacifica*. Regardless of the evolutionary history of this pathway, both lineages produce the same four staurosporine analogs (by LC-MS), indicating that the differences observed in the *staD* sequences (Fig. 3) do not appear to be linked to the production of different compounds.

The persistence of pathways acquired by HGT is due to the selective advantage their small-molecule products confer to the host (12, 23). The incongruence of HGT and species-specific secondary metabolite production suggests a complex interplay between gene acquisition and natural selection that creates mechanisms for the generation of new structural diversity and the fixation of adaptive products via periodic selection. Inferring the evolutionary histories of the biosynthetic pathways associated with secondary metabolism remains complex but provides opportunities to understand how nature creates new

structural diversity and the extent to which this diversity is linked to specific taxonomic groups. The three *Salinispora* species provide a well-defined model system within which to assess secondary metabolite gene evolution. The results from the present study reveal considerable levels of genetic exchange and clues to the mechanisms of secondary metabolism evolution in closely related taxa.

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