

ORIGINAL ARTICLE

Contributions of ancestral inter-species recombination to the genetic diversity of extant *Streptomyces* lineages

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***Streptomyces* species produce many important antibiotics and have a crucial role in soil nutrient cycling. However, their evolutionary history remains poorly characterized. We have evaluated the impact of homologous recombination on the evolution of *Streptomyces* using multi-locus sequence analysis of 234 strains that represent at least 11 species clusters. Evidence of inter-species recombination is widespread but not uniform within the genus and levels of mosaicism vary between species clusters. Most phylogenetically incongruent loci are monophyletic at the scale of species clusters and their subclades, suggesting that these recombination events occurred in shared ancestral lineages. Further investigation of two mosaic species clusters suggests that genes acquired by inter-species recombination may have become fixed in these lineages during periods of demographic expansion; implicating a role for phylogeography in determining contemporary patterns of genetic diversity. Only by examining the phylogeny at the scale of the genus is apparent that widespread phylogenetically incongruent loci in *Streptomyces* are derived from a far smaller number of ancestral inter-species recombination events.**

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Introduction

Homologous recombination (HR), the acquisition of genetic material through genetic exchange of similar DNA sequences, is an important mechanism in the evolution of many microbial lineages. As a consequence of HR, microbial genomes contain genes having diverse evolutionary histories (Boussau *et al.*, 2008; Boyer *et al.*, 2009; Zhaxybayeva *et al.*, 2009; Caro-Quintero *et al.*, 2012), which may be different from the history of the organism itself. HR has significant impacts on the evolution and diversification of diverse taxa of *Bacteria* and *Archaea* (Hernández-López *et al.*, 2013; Paul *et al.*, 2013; Darch *et al.*, 2015; Huang *et al.*, 2015). Extensive HR also coincides with indistinct or ‘fuzzy’ boundaries between genetic clusters (Hanage *et al.*, 2005; Papke *et al.*, 2007). These observations have generated lively debates about mechanisms of microbial evolution and the concept of species in the microbial world (Baptiste and Boucher, 2009; Fraser *et al.*, 2009; Riley and Lizotte-Waniewski,

2009; Caro-Quintero and Konstantinidis, 2012; Segerman, 2012).

Discussions of microbial evolution often focus on adaptive traits as they relate to ecological niche (Koeppel *et al.*, 2008). However, neutral processes such as genetic drift comprise a cornerstone of evolutionary theory, and genetic drift can promote allopatric speciation in microorganisms (Whitaker *et al.*, 2003) independent of niche dynamics. In addition, neutral changes resulting from HR can have major impacts on the evolution of microbial genomes (Vetsigian and Goldenfeld, 2005; Fraser *et al.*, 2007; Sheppard *et al.*, 2008; Doroghazi and Buckley, 2011). Multi-locus sequence analysis (MLSA) can be used to characterize extant patterns of genetic diversity in microbial lineages and the degree to which these patterns result from HR within and between species. Knowledge of these patterns is required to infer the evolutionary processes that govern microbial diversification.

There is evidence for widespread HR both within and between *Streptomyces* species (Doroghazi and Buckley, 2010, 2014; Cheng *et al.*, 2015). *Streptomyces* use an unusual method of gene exchange in which conjugation mobilizes double-stranded DNA (Kataoka *et al.*, 1991). In the model organism, *Streptomyces coelicolor* A3(2), long considered as the genetic workhorse in *Streptomyces* biology, chromosomal markers are mobilized

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through this mechanism at a frequency of 0.1–1% (Hopwood *et al.*, 1985), with plasmid integration causing mobilization of chromosomal genes at an efficiency that approaches 100% (Chater *et al.*, 1982). However, whether this is true for other members of this genus remains unclear, as studies comparing DNA mobilization and conjugation rates in other species of *Streptomyces* are lacking. Nevertheless, this double-stranded DNA conjugation system permits inter-species recombination between isolates in the laboratory (Alacevic, 1963) and can generate hybrid strains with genomes having nearly equal genetic contributions from each parent (Wang *et al.*, 1999). Such laboratory-generated hybrids are described to display new combinations of parent phenotypes including changes in phage sensitivity (Lomovskaya *et al.*, 1977) and antibiotic production. Furthermore, acquisition of antibiotic biosynthesis and resistance genes has been documented in wild strains of *Streptomyces* (Deng *et al.*, 2011; Kinashi, 2011), suggesting that the horizontal gene exchange may have contributed to the tremendous biosynthetic diversity within the genus *Streptomyces*.

There have been few efforts to characterize the impact of recombination on the evolutionary history of *Streptomyces*. MLSA has revealed that many species of *Streptomyces* have mosaic ancestry because of widespread gene exchange (Doroghazi and Buckley, 2010). Not surprisingly, recombination rates are much higher within species than between species (Doroghazi and Buckley, 2010). High rates of gene exchange between closely related strains may act as a cohesive force that reinforces the genetic similarity of species and species clusters (Doroghazi and Buckley, 2011). For example, strains of *Streptomyces pratensis* are in linkage equilibrium over much of their genome despite having a geographic range that spans > 1000 km (Doroghazi and Buckley, 2014). This pattern of ancestry can result only if the *S. pratensis* genomes are recombining freely across a large geographic range, or if *S. pratensis* has experienced evolutionarily recent demographic expansion from an ancestral population that was in linkage equilibrium. Ecological niche differences (marine, soil, insect host and plant host) can pose barriers to gene exchange and these factors may influence lineage diversification (Cheng *et al.*, 2015), but geographic and phylogenetic factors that influence gene exchange in *Streptomyces* remain poorly described.

We use MLSA to investigate patterns of inter-species HR among diverse *Streptomyces* species from 11 species clusters. In particular, we seek to determine if patterns of inter-species HR are random or if they are associated with phylogeny and/or geography. We select five species clusters for further analysis, including three that demonstrate mosaic ancestry and two that have contributed ancestry to the mosaic clusters. We screened *Streptomyces* isolates from diverse geographic sites to identify wild-type strains representing these five species clusters.

Materials and methods

Strain isolation and sequence data

Analysis of *Streptomyces* genetic diversity and phylogeny used the MLSA scheme of Guo *et al.* (2008). Nucleotide sequences of six loci from 150 strains were obtained from the *Streptomyces* database of the pubMLST website (<http://pubmlst.org/streptomyces/> as of December 2012). These include partial sequences of six loci—16S ribosomal RNA (rRNA) gene (432 bp), *trpB* (571 bp), *gyrB* (423 bp), *rpoB* (540 bp), *atpD* (496 bp) and *recA* (504 bp). Three isolates (*S. cremeus* strain AS 4.1625 ID# 23, *S. flavidofuscus* strain AS 4.1617 ID# 21 and *S. kasugaensis* strain DSM 40819 ID# 99) were excluded from our analyses on account of missing sequence information for at least one locus. For strains with identical sequence types, only one representative was selected. We also excluded three strains (*S. atratus* strain AS 4.1632 ID# 124, *S. gelaticus* strain AS 4.1444 ID# 132 and *S. sanglieri* strain AS 4.1146 ID# 144) that have divergent *rpoB* and *trpB* genes and are more similar to other genera of *Actinobacteria* than to *Streptomyces* when compared with the non-redundant database available from the National Center for Biotechnology Information (NCBI) using the Basic Local Alignment Search Tool (BLAST; Altschul *et al.*, 1990). *S. griseoplanus* was renamed *S. pratensis* following the recent report on its taxonomical characterization (Rong *et al.*, 2013). In total, 139 strains from pubMLST representing 128 species were used in the analysis (Supplementary Table S1).

Additional strains for MLSA were derived from a geographically explicit *Streptomyces* culture collection. The culture collection consists of >1000 isolates obtained from 15 temperate grassland sites across the continental United States (Supplementary Table S2). Sites were selected to represent a narrow range of ecological characteristics including meadow, pasture or native grasslands dominated by perennial grasses and having moderately acidic soil (pH: 6.0 ± 1.0; ave. ± s.d., Supplementary Table S2). Strains were isolated using uniform conditions and this will select for strains having similar physiological characteristics. Briefly, soil was sampled from 0 to 5 cm, air dried and sieved to 4 mm. *Streptomyces* strains were isolated at room temperature on glycerol-arginine agar, pH 8.7, containing 300 mg l⁻¹ cycloheximide and 30 mg l⁻¹ Rose Bengal as previously described (Doroghazi and Buckley, 2010). The analysis of physiologically similar strains from ecologically similar sites improves our ability to detect biogeographical patterns that result from neutral processes such as genetic drift (as reviewed in Hanson *et al.*, 2012). We screened these isolates by sequencing *rpoB* to identify strains (*n* = 95; Supplementary Table S2) that belonged to species clusters 4, 5, 6, 7 and 10 (as identified in Figure 1, see Results section). These species clusters represent three mosaic species

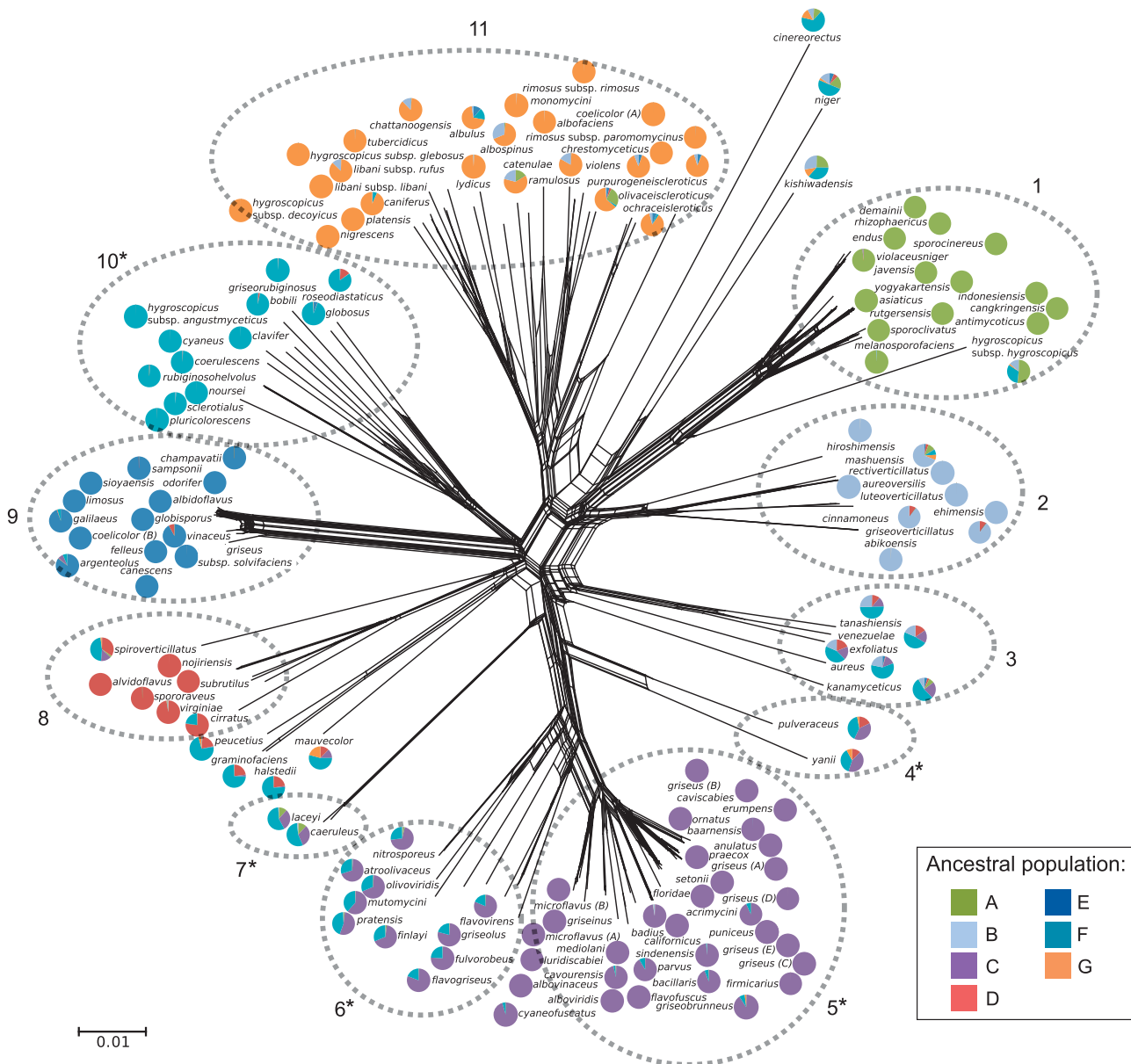


Figure 1 A phylogenetic network for *Streptomyces* generated using SplitsTree with default settings. The network was built using concatenated sequences of 16S rRNA, *trpB*, *atpD*, *gyrB*, *rpoB* and *recA* loci. The pie charts associated with each species show the ancestry of each species, with each color representing the seven ancestral lineages inferred by Structure. The dotted circles highlight distinct clusters. Asterisks indicate clusters targeted for further analysis including those with high (4 and 6) and low (5 and 10) admixture. Admixed clusters 4, 6 and 7 have genetic contributions from clusters 5 and 10. Scale bar indicates 0.01 substitutions per site.

clusters (4, 6 and 7) and two ancestral species clusters (5 and 10) identified in the analysis of pubMLST strains. We focused on these species clusters to evaluate in greater detail the phylogeny and phylogeography of strains in admixed lineages. For each isolate, we then sequenced the genes *atpD*, *gyrB*, *recA* and *trpB* using the method described in Doroghazi and Buckley (2010). We did not include the 16S rRNA gene because of its low genetic variability among taxa in our selected clusters.

Phylogenetic analyses

Multiple sequence alignments were performed using MUSCLE v3.5 (Edgar, 2004) with default parameters. A maximum likelihood phylogenetic tree using PhyML v.3.0 (Guindon *et al.*, 2010) was built for each gene under the Generalized Time Reversible (Tavaré, 1986) substitution model, estimated gamma parameter, estimated proportion of invariable sites, four substitution rate categories and 100 bootstrapped replicates. The six genes were then concatenated and used as input for the NeighborNet

program implemented in SplitsTree v4.13.1 (Huson, 1998) to further analyze species relationships. ClonalFrame was run with default settings to generate a 50% consensus tree (Didelot and Falush, 2007). We used LDhat (McVean *et al.*, 2002) to estimate the number of segregating sites and average pairwise differences for each of the five clusters in the second data set, using the concatenated alignment as input. Nucleotide diversity (π), number of synonymous substitutions per non-synonymous site (dN) and number of synonymous substitutions per synonymous site (dS) were calculated using DnaSP v5.10.01 (Librado and Rozas, 2009).

Clustering analysis

To examine the population structure and estimate the extent of genetic mixing, we utilized the model-based clustering algorithm implemented in Structure (Falush *et al.*, 2003). First, we performed clustering analysis for all 139 strains from pubMLST. We tested values of K from 2 to 10, where K is the number of inferred populations. For each specific K value, we performed 10 replicates, with each run lasting for 300 000 iterations and the initial 50 000 iterations discarded as burnin. Based on the criterion recommended by Evanno *et al.* (2005) and Prunier and Holsinger (2010), we selected the K value of 7 that best represents our data. Structure allowed us to identify species that belong to one ancestral lineage (un-admixed) and to estimate the proportion of ancestry for species that originate from more than one ancestral lineage (admixed or mosaic). The individual proportion of membership (q_i) determines the proportion of genetic ancestry that each individual derives from a given ancestral lineage. Empirical and simulation results (Barilani *et al.*, 2007) suggest $q_i \geq 0.90$ for un-admixed individuals. Using this threshold, we assigned each individual to one lineage if $q_i \geq 0.90$ or as admixed if the proportion of membership from each ancestral lineage was $q_i < 0.90$.

Tree topology tests

The program IQ-TREE v.0.9.5 (Nguyen *et al.*, 2015) was used to perform tests of phylogenetic incongruence including: the RELL method (Kishino *et al.*, 1990), the one-sided Kishino–Hasegawa test (Kishino and Hasegawa, 1989), the Shimodaira–Hasegawa test (Shimodaira and Hasegawa, 1999) and the Expected Likelihood Weight test (Strimmer and Rambaut, 2002). Significance was determined by permutation with 1000 re-samplings. The output shows the log likelihood score of all trees, the difference in log likelihood between each tree and the best tree, and the associated P -value for the probability that a given tree can explain the sequence data (Supplementary Table S3).

Recombination analyses

We used two programs in examining HR in *Streptomyces*. The Recombination Detection Program (RDP v3.44), which implements nine different detection methods, was used to analyze both data sets (Martin *et al.*, 2010). Detection was performed using default settings. We included only those recombination events that are supported by at least three out of the nine methods in RDP to avoid dependence on a single methodology (RDP, BOOTSCAN, GENECONV, MAXCHI, CHIMAERA, SISCAN, PhylPro, LARD and 3Seq). Previous recombination studies using RDP have used the 3/9 criterion, with the expectation that true recombination events will likely be detected using different approaches (Bilhère *et al.*, 2009; Cadillo-Quiroz *et al.*, 2012; Alvarez-Pérez *et al.*, 2013). The strength of RDP is its implementation of a variety of detection approaches to both identify and characterize recombination events, such as phylogenetic conflicts clustering of nucleotide substitutions and fitting of substitutions to an expected statistical distribution. Hence, using a 3/9 criterion will give us a stronger support for a recombination event, and a higher threshold will only slightly improve the results. Reported P -values were calculated using the Bonferroni correction within the program and statistical significance was set at $P < 0.05$ level. Using the alignment of the concatenated genes of each of the five clusters in the second data set as input, we also ran the Pairwise Homoplasy Index (PHI) test for recombination implemented in SplitsTree v4.13.1 (Huson, 1998).

Results and Discussion

Contemporary and ancestral inter-species recombination

We evaluated patterns of HR among *Streptomyces* species by analyzing sequences of 16S rRNA, *trpB*, *atpD*, *gyrB*, *rpoB* and *recA* loci for 139 *Streptomyces* species available from the pubMLST database (www.pubmlst.org). Seven ancestral populations were chosen to represent the collective ancestry of these species. The criteria for identifying the most likely number of ancestral populations has been described previously (Evanno *et al.*, 2005; Prunier and Holsinger, 2010); briefly we selected the smallest number of populations that explained the greatest probability of the data whereby successive increases in population number had only a marginal impact on the probability of the data. The species were then grouped into 11 species clusters on the basis of phylogenetic affinity (Figure 1). These 11 clusters roughly correspond to 16S rRNA gene clusters identified in a recent study of 615 *Streptomyces* species (Labeda *et al.*, 2012). Phenotypic conservation by species in these gene clusters is generally unremarkable (as assessed from Labeda *et al.*, 2012), although species clusters 1 and 11 contain strains that primarily have rugose spores (corresponding to

clusters 75–80 in Labeda *et al.*, 2012) and species cluster 9 contains species that produce smooth-surfaced, olive-buff colored spores carried in flexuous chains (corresponding to cluster 112 in Labeda *et al.*, 2012). Biosynthetic gene clusters generally differ between species of *Actinobacteria* (Doroghazi and Metcalf, 2013; Ziemert *et al.*, 2014), and such gene clusters are often exchanged horizontally within and between species (Gontang *et al.*, 2010; Ziemert *et al.*, 2014), although conservation of these gene clusters within species clusters remains uncharacterized.

Individual gene trees for the six loci all exhibit conflicting topologies (Supplementary Figure S1, Supplementary Table S3) supporting the finding of widespread inter-species HR. Similarly, analysis of genetic ancestry using Structure (Falush *et al.*, 2003) suggests widespread admixture between species clusters (Figure 1). Using the recombination detection software RDP, we detected 32 significantly supported HR events in the concatenated sequence alignment (Figure 2). A total of 98 of the strains (71%), and 122 of the 834 total genes (15%), were found by RDP to have evidence of inter-species HR,

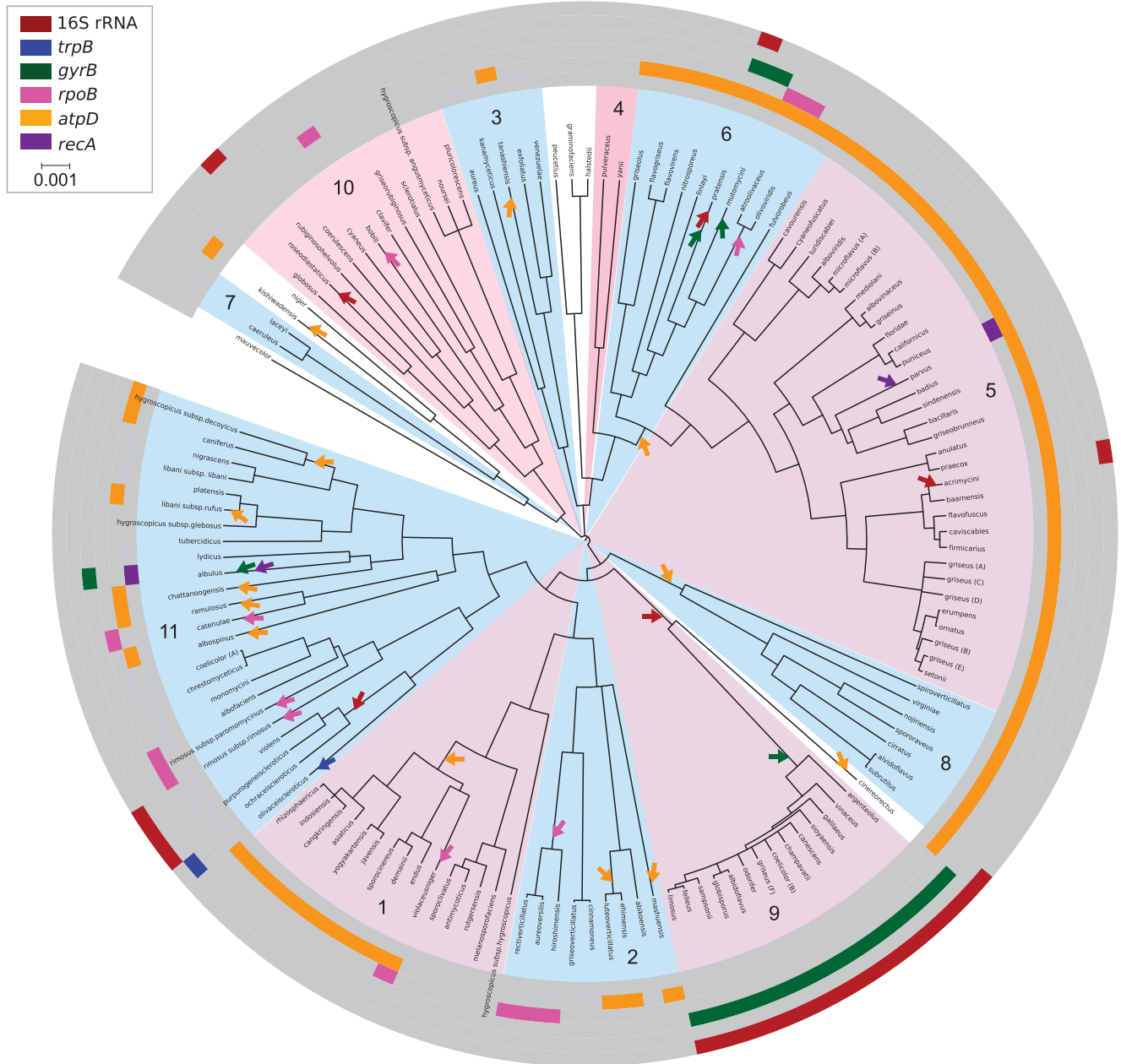


Figure 2 Clonal phylogeny for *Streptomyces* generated using ClonalFrame (50% consensus tree). Numbers and alternating blue and pink colors in the middle of the tree correspond to species clusters from Figure 1. Arrows indicate HR events at nodes of the tree as inferred by RDP and their position relative to the length of each branch is arbitrary. Colored bars in the outer rings indicate genes that are affected by HR as determined by RDP analysis. The loci inferred to be acquired by HR are identified by color as indicated in the scale. The scale bar in the ClonalFrame genealogy represents time in coalescent units.

and these results are broadly consistent with prior observations (Doroghazi and Buckley, 2010). However, the statistical criteria used by RDP to detect these HR events are insensitive to exchange of short similar sequences, and hence should be viewed as a conservative lower bound on the actual number of events that have occurred.

Genes observed to have evidence of HR can be derived both from contemporary and ancestral HR events. The fixation of ancestral HR events in ancestral lineages will result in an indefinitely large number of contemporary lineages that have inherited 'horizontally acquired' genes through vertical descent. Thus, widespread evidence of HR within a lineage can result from both prolific contemporaneous gene exchange and/or accumulation of ancestral HR events over evolutionary time. We mapped the evolutionary history of HR events by examining patterns of shared ancestry (that is, recombination donors and recipients inferred by RDP) among horizontally acquired genes (Figure 2). Of the 32 HR events detected, 9 are inferred to have taken place in ancestral lineages. These ancestral HR events included four that occurred in the last common ancestors of species clusters (subsequently inherited by 63 strains) and five that occurred in the last common ancestors of various subclades (subsequently inherited by 23 species). Hence, ancestral HR accounts for 86 of the 98 strains (88% of strains) in which HR was detected by RDP analysis. The remaining 23 HR events were unique to a single strain (note that sums are not additive because several strains have multiple HR events). Furthermore, it is likely that many of the HR events detected in a single lineage may have also occurred ancestrally, but are observed at the tips of the tree because related strains that share these HR events remain unsampled. Hence, to more rigorously evaluate the role of ancestral HR on the genetic diversity of *Streptomyces*, it is necessary to more rigorously sample from species clusters affected by ancestral HR.

Phylogenetic relationships of targeted species clusters

To further evaluate the role of ancestral HR in determining the extant genetic diversity of *Streptomyces*, we focused upon three admixed species clusters (4, 6 and 7) and two additional species clusters (5 and 10) that contain extant representatives of the ancestral populations that contributed genetic ancestry to these admixed clusters (Figure 1). Specifically, we sought to determine whether the HR events that contribute to phylogenetic incongruence were due to contemporary or ancestral HR events. The pubMLST database did not contain sufficient numbers of strains to permit robust analysis of these five species clusters; hence, additional strains that belonged to these clusters were identified from a collection of wild-type strains that we isolated from a

geographically diverse set of soil samples from temperate grassland sites across the United States (as described in Materials and methods section). We identified and sequenced loci from 95 new strains bringing the total number of strains representing the five clusters to 152. The concatenated sequence alignment for the five genes is 2534 bp. Pairwise nucleotide identity of MLSA loci within species clusters ranged from 91.9% to 100%, whereas pairwise nucleotide identity between species clusters ranged from 87.7% to 95.5% (Supplementary Figure S2).

The individual gene phylogenies show significant incongruence across the five species clusters (Figure 3 and Supplementary Figure S4, Supplementary Table S4). As expected, and as described below, many of these inter-species HR events took place in a common ancestor of a species cluster or subclade and were inherited vertically by extant strains. For example, strains from cluster 7 are most similar to cluster 10 strains at the loci *trpB*, *gyrB*, *rpoB* and *recA* (Supplementary Figure S4), and across these four loci they differ considerably from cluster 6 strains (>10% substitutions per site). However, an incongruent result is obtained with *atpD*, as cluster 7 strains are instead more similar to cluster 6 strains (<5% substitutions per site; Supplementary Figure S4) than to cluster 10 strains. This result suggests that a recent common ancestor of cluster 7 exchanged either part or all of its *atpD* gene with an ancestor of cluster 6. Similarly, strains in cluster 4 share affinity with those from clusters 5 and 6 for *trpB*, *gyrB*, *rpoB* and *recA* but affiliate with cluster 10 at *atpD* (Supplementary Figure S4). In addition, *S. olivoviridis* and *S. atroolivaceus*, each have four genes that associate with cluster 6, but have *gyrB* genes that affiliate with cluster 5 (Supplementary Figure S4).

We also observed 11 species and strains with ambiguous cluster affiliation intermediate between clusters 4, 5 and 6, including: *S. finlayi*, *S. nitrosporeus*, *S. fulvorobeus*, *S. flavogriseus*, *S. flavovirens*, *S. griseolus*, or75, or89, wa23, wa1022 and wa1068 (taxa labeled in black; Figures 3 and 4, Supplementary Figures S3 and S4). The latter eight of these form a cohesive group (99% bootstrap value) in the concatenated phylogeny and in all gene trees (Figure 3, Supplementary Figures S3 and S4); however, the placement of this group varies considerably in the different gene trees (Supplementary Figure S4). For example, this group of eight is most closely related to cluster 6 in the *gyrB* and *recA* gene trees, to cluster 5 in the *atpD* and *rpoB* gene trees, and to cluster 4 in the *trpB* tree (Supplementary Figure S4). Again, these results indicate that the most recent common ancestor of this group acquired multiple genes by HR and extant species subsequently inherited these genes from this common ancestor. The placement of the remaining three species, *S. finlayi*, *nitrosporeus*, and *S. fulvorobeus* varies considerably between clusters

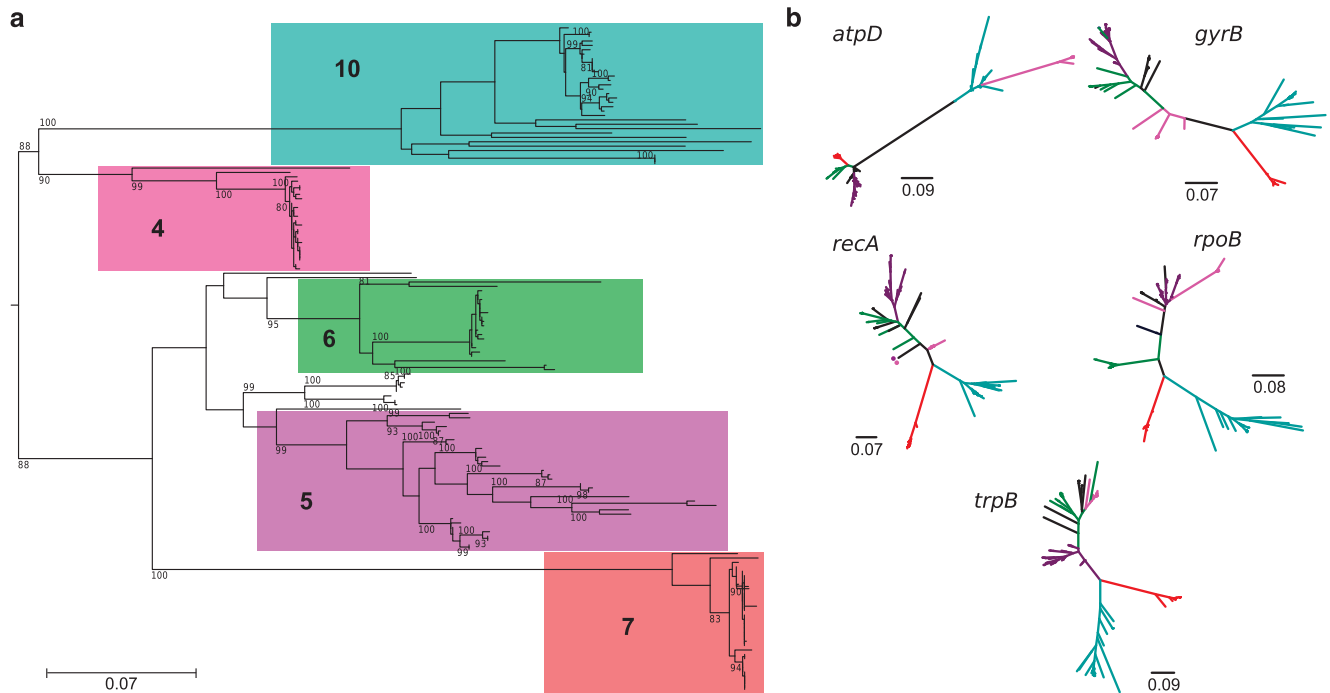


Figure 3 Maximum likelihood phylogeny of species clusters 4, 5, 6, 7 and 10, with the addition of isolates collected from diverse geographic sites ($n = 152$). **(a)** Phylogeny calculated from the concatenated alignment of the five housekeeping genes *trpB*, *gyrB*, *rpoB*, *atpD* and *recA* (see Supplementary Figure S3 for names of taxa). Only bootstrap values $\geq 50\%$ are shown. **(b)** Individual gene trees for each of the five genes (see Supplementary Figure S3 for names of taxa). Taxa labeled in black indicate ambiguous cluster membership. Scale bars indicate substitutions per site.

4, 5 and 6 for each of the different genes (Supplementary Figure S4) suggesting that these lineages may have been recipients of a number of HR events from species in these different clusters.

Contributions of geography to recombination

Geographical and ecological factors have been previously shown to influence patterns of gene flow in diverse microorganisms (Reno *et al.*, 2009; Cadillo-Quiroz *et al.*, 2012; Shapiro *et al.*, 2012). It is possible to evaluate whether patterns of admixture exhibit geographical partitioning in *Streptomyces*, as the geographic origin of our strains is known. Analysis of strains with similar characteristics (all strains were isolated under the same conditions on the same medium, see Materials and methods section) from similar habitats maximizes our ability to observe neutral genetic variation associated with biogeography (as discussed in Hanson *et al.*, 2012). HR in housekeeping genes would generally be expected to result in neutral variation, which can readily be detected through MLSA (Falush *et al.*, 2003).

We observe that species clusters 4 and 7 are found almost exclusively (98% of strains, Figure 4) in higher latitude sites (WA, ME and OR), whereas species clusters 6 and 10 occurred almost exclusively (95% of strains, Figure 4) in lower latitude sites (TX, FL, MS, NC and WI). This pattern corresponds to historical patterns of glaciation as all higher latitude sites were subject to late

Pleistocene glaciation (Peltier, 1994). We note that much of Wisconsin (WI) was within the driftless zone and did not experience glacial cover during the last glacial maximum (Holliday *et al.*, 2002). Species clusters 4 and 7 have mosaic ancestry with asymmetric contributions from ancestral populations represented by strains in clusters 5, 6 and 10 (Figures 1 and 3b, Supplementary Figure S4). Species clusters 4 and 7 also have lower nucleotide diversity and higher dN/dS than those of clusters 6 and 10 (Table 1). In addition, significant intra-cluster recombination is observed within clusters 6 and 10 but not within clusters 4 and 7 (PHI test, Table 1). These results suggest that the fixation of ancestral HR events in species clusters 4 and 7 may have occurred as the result of rapid demographic expansion into virgin habitat exposed by glacial retreat, as discussed below.

Glaciation has had strong impacts on the biogeography and genetic diversity of a range of terrestrial species (Conroy and Cook, 2000; Milá *et al.*, 2006; Wilson and Eigenmann Veraguth, 2010; Pointing *et al.*, 2014; Newman and Austin, 2015) and has been shown to impact microbial diversity in soil (Vercken *et al.*, 2010; Boyd *et al.*, 2011; Park *et al.*, 2011; Eisenlord *et al.*, 2012; Stibal *et al.*, 2012; Hamilton *et al.*, 2013). Rapid demographic expansion resulting from glacial retreat has had profound impacts on the genetics of species (Hewitt, 1996; Conroy and Cook, 2000; Hewitt, 2000, 2004; Wilson and Eigenmann Veraguth, 2010). In particular,

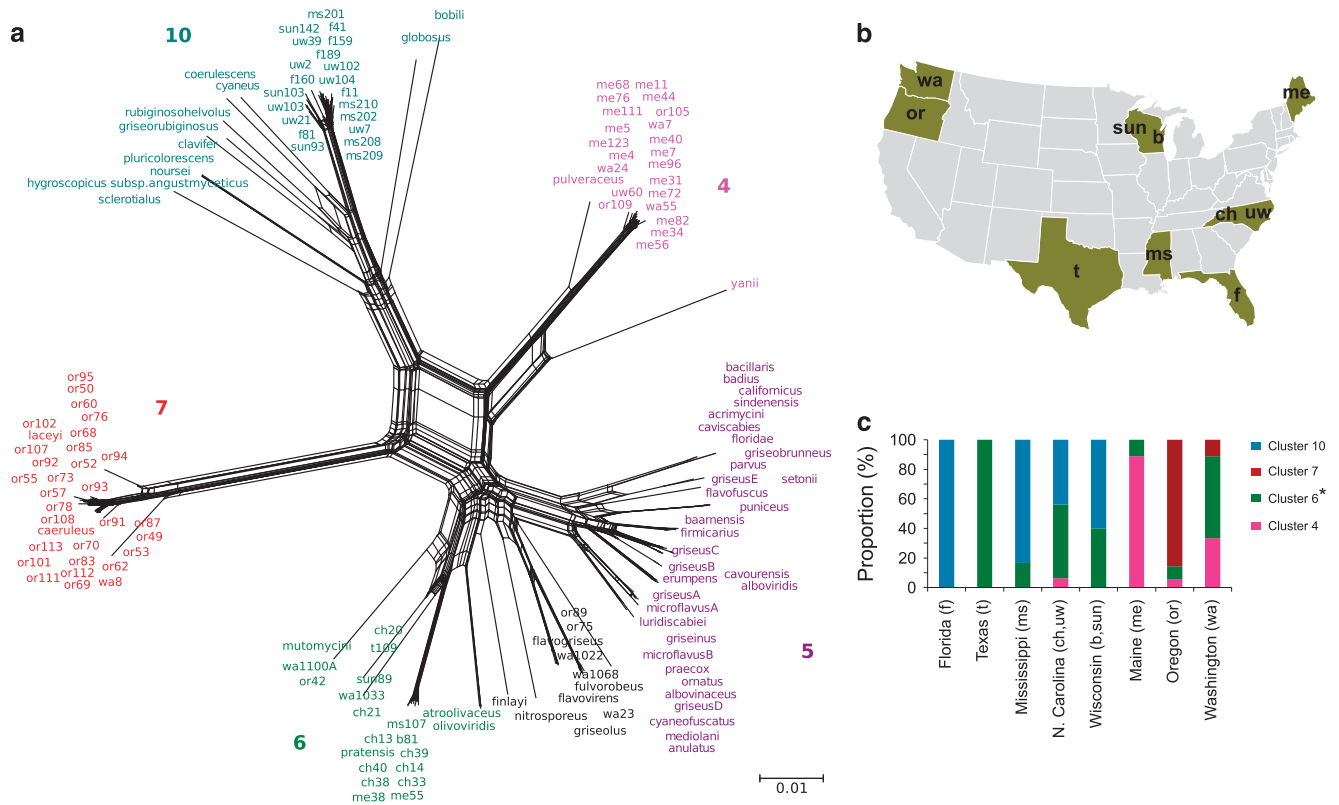


Figure 4 Evolutionary relationships and geographic sources of isolates from the second data set, consisting of species clusters 4, 5, 6, 7 and 10. (a) SplitsTree network of the five selected clusters, genetic relationships were calculated from the concatenated alignment of the five housekeeping genes *trpB*, *gyrB*, *rpoB*, *atpD* and *recA*. Colors correspond with Figure 3. Taxa labeled in black indicate ambiguous cluster membership from Figure 3. Scale bar indicates 0.01 substitutions per site. (b) Location of sites from which strains were isolated (see also Supplementary Table S2). The labels for each site correspond to the sampling names used for the isolates. (c) Distribution of the five clusters in each state in the United States. Colors of the bars correspond to those found in panel a. Cluster 6 consists of isolates labeled in green and black in the SplitsTree.

Table 1 Estimated population and evolutionary parameters

Cluster	Number of sequences	Length	No. segregating sites	Average pairwise differences	Nucleotide diversity π	dN/dS	PHI test P-value
4	22	2528	34	5.874	0.00232	0.6637	0.1293
6*	22	2528	292	78.515	0.03108	0.6250	0.0
7	30	2534	98	12.577	0.00496	0.6643	0.6412
10	21	2519	91	21.238	0.00844	0.583	4.65E-03

Abbreviation: PHI, Pairwise Homoplasmy Index. Values for number of segregating sites and average pairwise differences were calculated by LDhat. The P-value for the PHI test was calculated under the null hypothesis of no recombination. Nucleotide diversity (π), defined as the average number of nucleotide differences per site, as well as dN and dS values were calculated using DnaSP.

demographic expansion should manifest in lower nucleotide diversity (Hewitt, 1996), the potential for relaxed selection because of founder effects (Templeton, 2008) and less recombination because of epidemic population growth (Smith *et al.*, 2003). These predictions are broadly consistent with our observations for species clusters 4 and 7 (as described in the paragraph above).

We hypothesize that inter-species HR has occurred at relatively low frequency between species clusters 5 and 10 (relative to intra-cluster HR), but

demographic expansion caused genes acquired by inter-species HR to reach fixation in the populations that ultimately founded species clusters 4 and 7. Furthermore, even if mosaic (or hybrid) strains resulting from inter-species HR are maladapted because of genetic incompatibility (Berg and Kurland, 2002), demographic expansion would create an opportunity for maladapted stains to increase in frequency in the absence of competition from ancestral strains (as described in Templeton, 2008). Over time, compensatory mutations would be

expected to accumulate, eliminating fitness drag from maladapted alleles (Bouma and Lenski, 1988).

The Pleistocene was characterized by repeated glacial–interglacial cycles (Cox and Moore, 2000), generating many opportunities for lineage divergence because of demographic expansion and contraction (Hewitt, 1996). Indeed, glacial–interglacial cycles of population expansion and contraction have been widely recognized for their impacts on hybridization and lineage divergence of terrestrial species (Swenson and Howard, 2005). It may be possible that species cluster 6, which has mosaic ancestry (Figure 1) but is found in southern sites and has higher nucleotide diversity than species clusters 4 and 7, was generated in a prior glacial–interglacial cycle. Species cluster 6 also exhibits multiple patterns of incongruence across strains, whereas clusters 4 and 7 have each inherited a single incongruent locus (Supplementary Figure S4).

There is widespread evidence of mosaicism across *Streptomyces* species (Doroghazi and Buckley, 2010), but most of the incongruence in gene phylogenies is due to relatively few ancestral recombination events that have been inherited by many extant strains (Figure 2). Looking only at six loci, it appears that even if inter-species and inter-cluster recombination was relatively rare on a per generation basis, such events could accumulate over evolutionary time leading to complex patterns of genetic mosaicism. Furthermore, we hypothesize that the fixation of incongruent loci is a consequence of evolutionary dynamics associated with historical biogeography and that such dynamics may explain the ancestral formation of mosaic lineages. If the fixation of inter-species recombination events is associated with historical demographic events, then it means that the frequency of genetic mosaicism in *Streptomyces* may be less a function of inter-species recombination frequency and more a function of historical biogeography. Future tests of this hypothesis will require analysis of additional strains and sites to better establish whether species and species clusters have distinct geographical range, to establish range boundaries, and to perform comparative population genomic analysis to evaluate the frequency of ancestral HR and the extent of genetic mosaicism in relation to phylogeography.

Conflict of Interest

The authors declare no conflict of interest.

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