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Interspecies Interactions Stimulate Diversification of the *Streptomyces coelicolor* Secreted Metabolome

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Soils host diverse microbial communities that include filamentous actinobacteria (actinomycetes). These bacteria have been a rich source of useful metabolites, including antimicrobials, antifungals, anticancer agents, siderophores, and immunosuppressants. While humans have long exploited these compounds for therapeutic purposes, the role these natural products may play in mediating interactions between actinomycetes has been difficult to ascertain. As an initial step toward understanding these chemical interactions at a systems level, we employed the emerging techniques of nanospray desorption electrospray ionization (NanoDESI) and matrix-assisted laser desorption ionization–time of flight (MALDI-TOF) imaging mass spectrometry to gain a global chemical view of the model bacterium *Streptomyces coelicolor* interacting with five other actinomycetes. In each interaction, the majority of secreted compounds associated with *S. coelicolor* colonies were unique, suggesting an idiosyncratic response from *S. coelicolor*. Spectral networking revealed a family of unknown compounds produced by *S. coelicolor* during several interactions. These compounds constitute an extended suite of at least 12 different desferrioxamines with acyl side chains of various lengths; their production was triggered by siderophores made by neighboring strains. Taken together, these results illustrate that chemical interactions between actinomycete bacteria exhibit high complexity and specificity and can drive differential secondary metabolite production.

**IMPORTANCE**

Actinomycetes, filamentous actinobacteria from the soil, are the deepest natural source of useful medicinal compounds, including antibiotics, antifungals, and anticancer agents. There is great interest in developing new strategies that increase the diversity of metabolites secreted by actinomycetes in the laboratory. Here we used several metabolomic approaches to examine the chemicals made by these bacteria when grown in pairwise coculture. We found that these interspecies interactions stimulated production of numerous chemical compounds that were not made when they grew alone. Among these compounds were at least 12 different versions of a molecule called desferrioxamine, a siderophore used by the bacteria to gather iron. Many other compounds of unknown identity were also observed, and the pattern of compound production varied greatly among the interaction sets. These findings suggest that chemical interactions between actinomycetes are surprisingly complex and that coculture may be a promising strategy for finding new molecules from actinomycetes.
teractions, i.e., that production of a siderophore that cannot be used by one strain can lead to impaired development due to a decrease in local iron availability (12, 13). Another recent work that mapped pairwise interactions among a set of 64 Streptomyces isolates concluded that outcomes of such interactions are mostly determined by metabolite production (14). Several other studies have examined interactions between streptomycetes and the common soil bacterium Bacillus subtilis (15–22). These studies have documented a number of different interaction modalities, including inhibition of streptomycete development (15, 18–20, 22), degradation of signaling compounds via secreted enzyme activity (15, 18), and alteration of streptomycete natural product biosynthesis (16, 17, 21, 22). Given that each actinomycete is probably capable of producing dozens of secreted metabolites and that interspecies interactions may influence production of these molecules, we sought an experimental framework that would allow us to systematically examine changes in the secreted metabolome during multiple pairwise interspecies interactions.

The nascent mass spectrometry (MS) techniques of nanospray desorption electrospray ionization (NanoDESI) and matrix-assisted laser desorption ionization–time of flight (MALDI-TOF) imaging have recently opened new ways of examining the exchange of secondary metabolites between interacting bacteria in situ (21–24). In the work described here, we used these tools to examine the changes in the secreted metabolome of the well-studied actinomycete S. coelicolor as it grew near other actinomycetes. We studied five such interactions, and in each case we found many metabolites that were not produced by S. coelicolor when grown as a pure culture. We also observed that the sets of metabolites associated with S. coelicolor colonies were highly variable depending on the interacting partner, suggesting a unique response in each case. Several interactions triggered the production of an extended family of acylated desferrioxamines, never before documented a number of different interaction modalities, including inhibition of streptomycete development (15, 18–20, 22), degradation of signaling compounds via secreted enzyme activity (15, 18), and alteration of streptomycete natural product biosynthesis (16, 17, 21, 22). Given that each actinomycete is probably capable of producing dozens of secreted metabolites and that interspecies interactions may influence production of these molecules, we sought an experimental framework that would allow us to systematically examine changes in the secreted metabolome during multiple pairwise interspecies interactions.

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**RESULTS**

**Experimental system for studying actinomycete interactions.** S. coelicolor is perhaps the best-studied actinomycete. Many features of its genome, secondary metabolism, growth, development, and stress response have been characterized (25–28). Some growth media, such as R2YE, stimulate rapid multicellular development of S. coelicolor and production of a range of secondary metabolites, including the pigmented antibiotics actinorhodin and the prodiginines (29). Other growth media, including ISP2 (International Streptomycyes Project medium 2), support the growth of S. coelicolor, but robust development and secondary metabolite production occur only after prolonged incubation (i.e., >10 days). To look for interactions that stimulated such metabolite biosynthesis by S. coelicolor on ISP2 medium, we spotted 1 µl of an S. coelicolor spore suspension 5 mm from 20 similar inocula of other actinomycetes. Several actinomycetes triggered production of the red antibiotic prodiginine in S. coelicolor, in some cases as early as 3 days. For example, see the interaction with Amycolatopsis sp. AA4 in Fig. 1A. Of these 20 interactions, we focused on 5 (all shown in Fig. 1A) that provoked different temporal and phenotypic responses in S. coelicolor ranging from no stimulation of pigmentation or development (Streptomyces sp. E14 interaction) to partial or strong pigmentation (Streptomyces sp. SPB74 and Amycolatopsis sp. AA4 interactions, respectively) and both pigmentation and development (Streptomyces albus J1074 and Streptomyces viridochromogenes DSM40736 interactions). Given that developmental and natural product biosynthetic regulatory cascades are linked in S. coelicolor, we hypothesized that these different phenotypes expressed by S. coelicolor might be indicative of differences in the pattern of small molecules produced.

To test this hypothesis, we employed two metabolomic approaches, nanospray desorption electrospray ionization (NanoDESI) and microbial matrix-assisted laser desorption ionization–time of flight (MALDI-TOF) imaging mass spectrometry (IMS), to examine the chemical response of S. coelicolor in each of these interactions. Importantly, both techniques allow for sampling directly from colonies grown on an agar substrate (21, 23). NanoDESI MS is a recently developed methodology in which the solvent is delivered to the sample surface via a fused silica capillary (21). A second capillary draws the solvent (now containing dissolved analyte) off the sample surface and delivers it directly into the mass spectrometer such that a small liquid bridge (~1 µl) is constantly maintained on the sample surface between the two capillary ends. The analyte desorbed from the sample surface is then subjected to data-dependent tandem MS analysis, ultimately yielding individual tandem mass spectra for the hundreds to thousands of ions detectable within the sample. We thus used NanoDESI MS and MALDI-TOF IMS to analyze the colonies of the interactions shown in Fig. 1A. The general experimental and data analysis workflow is diagrammed in Fig. 1B. In every case, we sampled both the S. coelicolor colony and the interacting colony, which we refer to as the initiator colony, at days three, five, and seven. We also sampled control colonies of S. coelicolor that were grown in isolation on the same medium and at similar time points. All samples were prepared and analyzed in duplicate. The thousands of MS2 spectra from all time points were used to build a spectral network that allowed visualization of chemical species in both structurally familiar and temporal contexts based on statistically significant similarities between their tandem MS fragmentation patterns (21, 30).

While NanoDESI affords unparalleled sampling, microbial MALDI-TOF IMS allows high-resolution mapping of ions within a sample (23). As a complement to the NanoDESI spectral networks, we also collected IMS data sets for each interaction and S. coelicolor control colonies at day five. The combined output from these complementary techniques provided a uniquely rich data set for simultaneously assessing the response of a single organism in multiple interactions at a system-wide scale and for prospecting for novel secondary metabolites.

**Interspecies interactions cause differential secondary metabolite production in S. coelicolor.** To broadly characterize the chemical response of S. coelicolor in these interactions, we considered these data in terms of chemical inputs from the initiator colonies and chemical output from S. coelicolor. In our work flow, we chose to represent the MS2 spectra visually via spectral networking, which allows each MS2 spectrum within the entire experimental data set to be compared in a pairwise manner to all other MS2 spectra and scored based on the statistical similarity between the two fragmentation patterns. When the results of this analysis are imported into two-dimensional (2D) visualization software (such as Cytoscape), this allows similar compounds to be grouped together in 2D space separate from other compounds
exhibiting different fragmentation patterns. In this visualization, individual compounds present in the samples are represented as single nodes (circles) within the spectral network (30), with each node containing one or many MS2 spectra, as is the case when multiple MS2 spectra are judged to be identical by the spectral scoring algorithms (31). When the MS2 spectra for two nodes meet a set of criteria designed to assess structural relatedness, they are connected with an edge (line) (21, 31). Applying this algorithm to the pooled data sets yielded an aggregated parent network, which was then filtered to maximize interpretability. This
entailed removing nodes associated with several controls, including the solvent alone, agar medium with no colonies, and S. coelicolor colonies grown alone at 3, 5, and 7 days. Finally, nodes were retained only if constituent spectra were found in both runs of data collection from duplicate samples, giving a refined network composed of nodes representing ions associated with the initiator colonies and S. coelicolor colonies during the interactions (Fig. 2).

The resulting network contains 629 total nodes. Of these, 227 nodes were associated exclusively with S. coelicolor colonies. There were 278 nodes associated exclusively with the various initiator colonies. Nineteen nodes were found only in both S. coelicolor and a given initiator at the same time (common), and 105 nodes showed variable behavior, i.e., they were found associated with S. coelicolor and/or an initiator at different times. Only 23 compounds were associated exclusively with S. coelicolor control colonies, and 43 compounds were common to interacting and control S. coelicolor colonies (see Fig. S25 in the supplementary appendix available at http://gasp.med.harvard.edu/journals/traxler_2013_SI_nanodesi.pdf). At a broad level, the spectral network shows that these interactions triggered production of many compounds by S. coelicolor which were not observed when S. coelicolor was grown in isolation.

To see how similar the response of S. coelicolor was between interactions, we constructed Venn diagrams encompassing all S. coelicolor-associated nodes for all five interactions for each day. The diagram for day three is shown in Fig. 3A. This analysis highlighted an important trend, namely that the majority of the nodes fall into the outer, or “unique,” zones of the diagram, implying that they are interaction specific. For day three, the total number of unique nodes was 94, versus 29 that fell into the “shared” interior sections of the diagram. This trend was evident at every time point (Fig. 3B). We also note that while some interactions triggered production of relatively few compounds at a given time point (i.e., S. albus and Streptomyces sp. SPB74 at day 3), each interaction, with the exception of the S. albus interaction, underwent a time when the majority of its stimulated compounds were unique (see Venn analyses for days 5 and 7 in the supplemental appendix at the above URL). Taken together, these results suggest that the chemical response of S. coelicolor was highly idiosyncratic depending on the interacting strain and the time of sampling.

When we interrogated the network for known compounds in the S. coelicolor chemical response, we found subnetworks representing at least four major compound families: the antibiotics actinorhodin and prodiginine and the siderophores coelichelin and desferrioxamines B and E (Fig. 2). To examine the pattern of production of each of these molecular families, we considered the percentage of the total number of nodes for each family active in each interaction at each time point (heat map in Fig. 3C). For example, the prodiginine family of

FIG 2 Aggregated and refined spectral network of metabolites observed with NanoDESI during actinomycete interactions. The network is composed of nodes representing ions associated with S. coelicolor colonies grown near another actinomycete (ions found in S. coelicolor colonies grown alone were removed) and ions from initiator colonies. Nodes associated only with S. coelicolor at any time are blue. Nodes associated only with initiators are red. Nodes found only in both an initiator and S. coelicolor at the same time are yellow. Gray indicates nodes with variable behavior (i.e., found in multiple contexts). Representative structures of identified metabolites are shown.
Antibiotics is represented in the network as a subnetwork consisting of eight total nodes observed across all 5 interactions and all 3 time points. Six of these eight nodes were observed when *S. coelicolor* was interacting with *Amycolatopsis* sp. AA4 at day seven, representing robust stimulation of this compound family.

Production of each of these four compound families varied according to the interaction and sampling time. For example, *Streptomyces* sp. E14 did not stimulate prodigine or actinorhodin production (as expected, given that no *S. coelicolor* pigmentation is evident in this interaction), while *Amycolatopsis* sp. AA4, *Streptomyces* sp. SPB74, *S. viridochromogenes*, and *S. albus* all stimulated production of both pigmented antibiotics, although the timing of production varied depending on the initia- tor. The genomes of all six strains examined in these interactions have been sequenced and analyzed for their secondary metabolic potential using the antiSMASH platform (32). The results of this analysis (provided in Table S5 in the supplemental appendix at http://gasp.med.harvard.edu/journals/traxler_2013_SI_nanodesi.pdf) showed that none of the initiator strains contains genes for production of either actinorhodin or prodiginines, and accordingly, all compounds of these families associated with these interactions are presumed to originate with *S. coelicolor*. As further validation of our overall approach, we determined the spatial distribution of the prodiginines within the interacting colonies using IMS (see Fig. S1 in the supplemental appendix at the above URL). Prodigine distribution was found to match exactly the pattern expected given the visibility of red pigment and the activity of nodes within the prodiginine subnetwork.

Coelichelin was found sporadically across the interactions, with the exception of the *S. viridochromogenes* interaction, where it was observed at every time point. BLAST analysis of the *S. viridochromogenes* genome clearly shows that it too possesses the genes for coelichelin biosynthesis. Thus, observation of coelichelin in this interaction may reflect production by either or both of the strains. Desferrioxamine B or E was found in every interaction and at every time point with the exception of *Streptomyces* sp. SPB74 at day three. Genes for desferrioxamine production are widely distributed among actinomycetes, with *Streptomyces* sp. SPB74, *S. viridochromogenes*, and *S. albus* all containing canonical operons for biosynthesis of this siderophore. However, *Amycolatopsis* sp. AA4 and *Streptomyces* sp. E14 do not have desferrioxamine synthesis operons. Thus, in the *Amycolatopsis* sp. AA4 and *Streptomyces* sp. E14 interactions, the observed desferrioxamines likely originated from *S. coelicolor*, while either or both of the strains may be responsible for its production in the *Streptomyces* sp. SPB74, *S. viridochromogenes*, and *S. albus* interactions. The common detection of siderophores in these interactions suggests that competition for iron may be stringent in interactions between actinomycetes, a notion further underscored by our findings detailed below and previously observed under other interaction conditions (13).

**Extended family of acyl-desferrioxamines from *S. coelicolor***

The spectral network in Fig. 2 contains many nodes of unknown identity associated with *S. coelicolor* colonies. A majority of these unknown nodes fall into the outer zones of the diagram, indicating that each interaction is more unique than it is similar to other interactions. (C) Patterns of known compound production. Each compound family is observed as a subset of nodes within the larger network in Fig. 2. Heat map colors indicate the proportion of active nodes in each interaction. For example, 1.0 indicates that all the nodes associated with a given compound family are active. Three, 5, or 7 represents the sampling time in days. Numbers beneath compound names indicate number of nodes associated with those compounds in Fig. 2. Interactions are labeled as indicated in Fig. 1A. This analysis includes nodes associated with initiator colonies.

**FIG 3** Global analysis of metabolites observed with NanoDESI. (A) Venn analysis of nodes associated with interacting *S. coelicolor* colonies on day 3. Each ellipse contains nodes found in the indicated interaction. The majority of detected ions fall into the outer zones of the diagram, indicating that each interaction is more unique than it is similar to other interactions. (B) Numbers of unique and shared nodes found at each time point. For each day, the numbers of nodes found only in single interactions were summed to give the total number of unique nodes. Nodes that were found in more than one interaction were summed to find the total number of shared nodes. The number of unique nodes exceeds the number of shared nodes at every time point, suggesting that the response of *S. coelicolor* is different depending on the interaction. (C) Patterns of known compound production. Each compound family is observed as a subset of nodes within the larger network in Fig. 2. Heat map colors indicate the proportion of active nodes in each interaction. For example, 1.0 indicates that all the nodes associated with a given compound family are active. Three, 5, or 7 represents the sampling time in days. Numbers beneath compound names indicate number of nodes associated with those compounds in Fig. 2. Interactions are labeled as indicated in Fig. 1A. This analysis includes nodes associated with initiator colonies.
amine. Confirming this possibility, we found that the MS2 spectra of four of the nodes in this subnetwork, representing compounds with m/z values of 743, 729, 687, and 659, matched exactly the MS2 spectra of several recently characterized acyl-desferrioxamines (33), including promicroferrioxamine (24). These molecules contain the core desferrioxamine siderophore structure (structure 1 in Fig. 4B) with a terminal R group composed of an acyl chain with an anteiso-methyl group (structures 3 to 6 in Fig. 4B). The fragmentation patterns for these known acyl-desferrioxamines provided the basis for annotation of more molecules in these subnetworks, ultimately showing that *S. coelicolor* made at least 12 analogs of the acyl-desferrioxamines, with appendages ranging from C7 to C17 fatty acids (summarized in Table S1 at http://gasp.med.harvard.edu/journals/traxler_2013_SI_nanodesi.pdf). We also observed a node with an m/z of 727 that corresponds to a previously described version of promicrodesferrioxamine with a nonhydroxylated central hydroxamate moiety (structure 2 in Fig. 4B).

Throughout the entire network, the nodes associated with these acyl-desferrioxamines total >90 and thus account for ~40% of the total response of *S. coelicolor* observable in the spectral network. Of the four major subnetworks circled in Fig. 4A, subnetworks a1 and a2 share broad redundancy in terms of the masses of their constituent nodes, although the discrete clustering of subnetwork a2 may indicate an unknown structural feature that sets these versions apart. Subnetwork a3 encompasses larger molecules that include versions of the C12 and C13 acyl-desferrioxamines with an unknown addition of ~118 Da (see Fig. S22 and S23 at the above URL). Finally, based on high-resolution masses, subnetwork a4 likely contains sodiated and potassiated adducts of the acyl-desferrioxamines (see Fig. S3 at the above URL). These findings show that *S. coelicolor* is capable of making an extensive repertoire of siderophores in the presence of other actinomycetes.

**Stimulation of acyl-desferrioxamine production in interspecies interactions.** Having deduced the identity of a key component of the *S. coelicolor* chemical response, we returned to the spectral network and IMS data to further examine the temporal and spatial pattern of acyl-desferrioxamine production across the five interactions. When the heat map analysis presented in Fig. 3C was extended to include 94 nodes representing the acyl-desferrioxamines, a clear pattern of production was apparent; namely, the *Amycolatopsis* sp. AA4 and *Streptomyces* sp. E14 interactions triggered robust acyl-desferrioxamine synthesis even at day 3 (Fig. 5A). In the case of the *Streptomyces* sp. SPB74 interaction, acyl-desferrioxamine production was observed starting at day 5 and for the *S. viridochromogenes* and *S. albus* interactions. These trends can be seen on a finer scale by examining the pattern of activity in different acyl-desferrioxamine subclusters within the spectral net-
FIG 5  Patterns of acyl-desferrioxamine production in interspecies interactions. (A) Heat map of acyl-desferrioxamine production. Analysis parameters are identical to those in Fig. 3C. (B) Subnetworks 1 and 2 contain the majority of acyl-desferrioxamines verified by MS2 fragmentation. Fine-scale analysis of subnetwork 1 is shown, illustrating differential acyl-desferrioxamine production in various interactions over time. Note the proximity of subnetwork 1 to desferrioxamine B (DFO B). Subnetworks 3 and 4 contain larger versions of DFOs and sodium adducts, respectively (see Fig. S3 at http://gasp.med.harvard.edu /journals/traxler_2013_SI_nanodesi.pdf). (C) Desferrioxamines are observable using IMS at day 5. m/z 561 and 601 correspond to desferrioxamines B and E, respectively. m/z 701 to 785 are representative acyl-desferrioxamines. Note the production of acyl-desferrioxamines by S. coelicolor in interactions where initiator strains do not make desferrioxamines B and/or E. Each IMS signal is scaled as a single color heat map; brighter color indicates higher signal intensity, and darker indicates lower signal intensity. (D) Acyl-desferrioxamines diffuse away from S. coelicolor colonies in three dimensions. The outermost to innermost layers (isosurfaces) correspond to 0.8, 0.88, and 0.95% ion intensity.
work, including subnetwork a1, shown in Fig. 5B. Activity in subnetworks a2 to -4 also mirrors this pattern (see the supplemental appendix at http://gasp.med.harvard.edu/journals/traxler_2013_SI_nanodesi.pdf).

We also examined the distribution of acyl-desferrioxamine production using IMS at day 5 (Fig. 5C). To serve as an internal control for the IMS experiments, an additional S. coelicolor colony was spotted 1 cm away from the S. coelicolor colony that was directly adjacent to the initiator colony. A subset of the acyl-desferrioxamines with \( m/z \) values ranging from 701 to 785 was readily detectable using this technique. Consistent with the activity within the NanoDESI spectral network, we saw production of the acyl-desferrioxamines in the Amycolatopsis sp. AA4, Streptomyces sp. E14, and Streptomyces sp. SPB74 interactions. Little to no acyl-desferrioxamine production was seen in the S. viridochromogenes and S. albus interactions; however, desferrioxamine E (\( m/z \) 601) was produced by both of the initiator strains. S. albus produced a large amount of deferrioxamine B as well (\( m/z \) 561). The near colonies in the Amycolatopsis sp. AA4 and Streptomyces sp. E14 interactions showed the most acyl-desferrioxamine production, with molecules even being detected beyond the border of the colony in several cases in the Amycolatopsis sp. AA4 interaction. To further examine the distribution of the acyl-desferrioxamines, we used a newly developed three-dimensional (3D) MALDI-TOF IMS methodology (34) (Fig. 5D). The resulting 3D renderings show that the representative acyl-desferrioxamines at \( m/z \) 743 and 757 were distributed beyond the edge of the colony not only on the agar surface but below the producing colony as well. Overall, these data imply that while the temporality and amount of acyl-desferrioxamine production varied among the interactions, all of the interactions eventually led to competition for iron. Moreover, the acyl-desferrioxamines were especially abundant in the interactions with the two strains that likely utilize siderophores other than desferrioxamine for iron acquisition (i.e., Amycolatopsis sp. AA4 and Streptomyces sp. E14).

**Siderophores from other strains trigger acyl-desferrioxamine production in S. coelicolor.** The apparent competition for iron induced in these interactions prompts the hypothesis that siderophores from initiator strains might locally decrease the iron available to \( S. \) coelicolor and thus stimulate acyl-desferrioxamine synthesis. Our previous work with Amycolatopsis sp. AA4 showed that it makes a unique siderophore, amychelin, which cannot be used by \( S. \) coelicolor (12, 13). Several amychelin adducts were observed in the spectral network as a subnetwork associated with the Amycolatopsis sp. AA4 interaction (Fig. 6A, listed in Table S3 at http://gasp.med.harvard.edu/journals/traxler_2013_SI_nanodesi.pdf). Amycolatopsis sp. AA4 strains with mutations in the amcG locus are unable to make amychelin (13). To determine if amychelin might play a role in activating acyl-desferrioxamine synthesis in \( S. \) coelicolor, we grew a colony of the Amycolatopsis sp. AA4 \( ΔamcG \) strain adjacent to \( S. \) coelicolor and examined molecule production using IMS. Production of four representative ions from the acyl-desferrioxamine family are shown in Fig. 6B in interactions with both wild-type Amycolatopsis sp. AA4 and the \( ΔamcG \) strain. Much less acyl-desferrioxamine, both in area and in abundance, was produced by \( S. \) coelicolor in the interaction with the \( ΔamcG \) strain than in the wild-type Amycolatopsis sp. AA4 interaction. To further examine if iron starvation alone could trigger production of the acyl-desferrioxamines, we tested the ability of the iron chelator 2,2′-dipyridyl to stimulate production of these molecules (see Fig. S26 in the supplementary appendix at the above URL). Indeed, we found that when 2,2′-dipyridyl was included at 200 \( \mu \)M, acyl-desferrioxamines were produced. These results confirm that iron competition caused by xenosiderophores can lead to induction of acyl-desferrioxamine biosynthesis in \( S. \) coelicolor and that iron starvation alone is sufficient to trigger production of these siderophores.

**DISCUSSION**

In natural environments, such as the soil, bacteria live surrounded by a multitude of other organisms, including other bacteria. Actinomycetes, whose genomes contain numerous gene clusters for making complex secreted metabolites, have clearly evolved many and diverse chemical means to affect other nearby residents of the soil (3, 5). Humans have long been the beneficiaries of this evolutionary process, since the majority of natural products used clinically originate with actinomycete bacteria (1, 2). However, how actinomycetes deploy their arsenals of secondary metabolites in the presence of other species has remained largely mysterious.

In this work, we used two mass spectrometry techniques, NanoDESI and MALDI-TOF imaging, to profile the chemical output from the actinomycete \( S. \) coelicolor in interactions with five other species of actinomycetes. Importantly, these complementary techniques allowed interrogation of the secreted metabolites directly from the bacterial colonies. The resulting data sets comprise the most comprehensive chemical view of a related set of microbial interactions to date. The spectral network yielded by our analyses includes some 629 compounds. Importantly, many of these molecules are of unknown identity. Surprisingly, we found that the set of compounds associated with \( S. \) coelicolor colonies varied dramatically from interaction to interaction, suggesting a largely specific response in each case (Fig. 3). Several interactions triggered production of an extended family of acyl-desferrioxamine siderophores, never before observed from \( S. \) coelicolor (Fig. 4). This result illustrates that interspecies interactions can lead to unexpected biosynthetic shifts in secondary metabolic pathways. Together, these findings indicate that during interspecies interactions, the chemical landscape exploited by actinomycetes is both vast and dynamic. Moreover, the great number of unidentified metabolites present in these data sets hints that interspecies interactions may represent a new path to accessing the rich chemical diversity encoded in actinomycete genomes.

\( S. \) coelicolor produces several compounds representative of therapeutically important chemical families. These include desferrioxamines (used to treat iron toxicity [35]), prodiginines (currently being investigated as immunosuppressant anticaner agents [36, 37]), and the calcium-dependent antibiotic (similar to the drug daptoycin [38]). Here, we observed a large number (227) of compounds differentially associated with \( S. \) coelicolor colonies across the five interactions. Of these, 50% (114) are readily classified as related to known \( S. \) coelicolor metabolites, including the prodiginines, actinorhodins, coelichelins, and the newly observed acyl-desferrioxamines. \( S. \) coelicolor has many other gene clusters for uncharacterized metabolites (25), some of which likely account for the many unknown compounds we detected. It is also possible that some of the compounds found exclusively associated with \( S. \) coelicolor colonies may have had precursors made by the initiator colonies that were later modified directly or indirectly by \( S. \) coelicolor. NanoDESI and IMS analyses of \( S. \) coelicolor strains...
with mutations in various gene clusters will be required to positively link the many unidentified compounds observed here with specific genes. As such, our current work sets the stage and provides a solid foundation for future studies to characterize the currently cryptic secondary metabolome of this streptomycete and many others like it.

The family of desferrioxamines described here has fatty acid appendages ranging from 7 to 17 carbon units. While several acyl-desferrioxamine versions have been found from at least two bacterial species (24, 33), the range of the acyl-desferrioxamines produced by *S. coelicolor* is unprecedented. Interestingly, both the promicroferrioxamines and the other known examples of acyl-desferrioxamines were isolated from actinobacteria sampled from intertidal habitats, initially suggesting that desferrioxamine acylation might be an adaptation to life in marine environments (24, 33). Further reinforcing this notion, marine bacteria from diverse genera are known to produce suites of acylated siderophores (39), including the marinobactins (40), aquachelins (40), amphibactins (41), ochrobactins (42), and synechobactins (43). While a few other acylated siderophores are known from terrestrial bacteria (44–46), the finding that *S. coelicolor* makes a very large suite of acylated desferrioxamines strongly suggests that the advantages

![Figure 6](image-url)

**FIG 6** Acyl-desferrioxamine production by *S. coelicolor* is stimulated by a siderophore from a nearby actinomycete. (A) The actinomycete *Amycolatopsis* sp. AA4 produces the siderophore amychelin. Sodiated and potassiated adducts of amychelin are visible as an *Amycolatopsis* sp. AA4-associated subnetwork. (B) Five days IMS of *S. coelicolor* grown near wild-type *Amycolatopsis* sp. AA4 and a mutant lacking the gene *amcG*, which does not produce amychelin. Ion abundance is visualized as a heat map. When grown near the ΔamcG strain, which does not make amychelin, *S. coelicolor* produces much less of the acyl-desferrioxamines.
associated with modulating siderophore solubility are not limited to marine environments.

Siderophores with relatively long fatty acid tails, such as mycobactin T and the amphibactins (14 to 21 carbons in length), have been found associated with cell membranes (41, 47), while siderophores such as marinobactins and aquachelins, with 12 to 16 carbons in their acyl chains, are known to partition as micelles and vesicles into the aqueous environment (40). Other siderophores, with acyl chain lengths of 10 carbons or less, such as rhizobactins and carboxymycobactins, diffuse relatively freely (39, 47). Moreover, synthetic acylation of desferrioxamine B, with appendages of up to 7 carbons in length, was shown to alter their solubility and membrane permeability (48). The range of acyl moieties found in the S. coelicolor desferrioxamines, from a single carbon in desferrioxamine B, 3 carbons in desferrioxamines E and G, and 7 to 17 carbons for the new versions found here, encompasses the entire range of solubility. We note that versions of desferrioxamines with appendages ranging from C11 to C14 were clearly present beyond the edges and below S. coelicolor colonies (Fig. 5C and D, Amycolatopsis sp. AA4 interaction), possibly hinting that micelle-mediated iron acquisition may be occurring here. We also observed that production of the acyl-desferrioxamines was largely stimulated by production of a competing siderophore by one of the initiator strains (Fig. 6). Together, these findings point to a remarkably diversified foraging strategy in the face of fierce competition for iron.

The biosynthetic pathway for desferrioxamine is well documented (49, 50). The third step of the proposed pathway entails the acylation of N-hydroxycadaverine by the enzyme DesC to yield N-hydroxy-N-succinyl-1,5-diaminopentane (hsDAP), three units of which then serve as building blocks for the final trimeric desferrioxamine structure. In organisms that make desferrioxamines B and G (including S. coelicolor), DesC is proposed to have relaxed substrate specificity, allowing it to incorporate acetylcoenzyme A (CoA) or succinyl-CoA into desferrioxamine precursors (50). The acyl moieties found in place of the normal acetyl-CoA or succinyl-CoA additions suggest that the relaxed substrate specificity of S. coelicolor DesC may extend much further than originally thought, allowing it to incorporate these long acyl chains into the normal desferrioxamine biosynthetic pathway.

The experimental framework utilized here yields data sets that afford unique opportunities for the discovery of natural products. Indeed, the differential chemical output observed from S. coelicolor implies that this bacterium has great flexibility in the expression of its secondary metabolol. While competition for iron provides a straightforward rationale for the induction of siderophore synthesis in S. coelicolor interactions with other actinomycetes, the physiological cues underlying the induction of antibiotics and other metabolites remain to be elucidated. Understanding the mechanisms of such interactions is of great interest since they may offer a new door to accessing "cryptic" metabolites from other actinomycetes.

An appendix containing supplemental figures and chemical annotations for the molecules detected in this study can be accessed at http://gasp.med.harvard.edu/journals/traxler_2013_SI_nanodesi.pdf.

MATERIALS AND METHODS

Strains and growth conditions. Strains used in this study are listed in Table S4 in the supplemental appendix at http://gasp.med.harvard.edu/journals/traxler_2013_SI_nanodesi.pdf. All experiments were done on ISP2 agar (10 g malt extract, 4 g glucose, 4 g yeast extract, 15 g agar, 1 liter Milli-Q H2O). For all experiments, 10 ml of agar was added to standard 100-mm petri plates to yield an agar surface ~2 mm thick, which is suitable for preparation for IMS. Strains were inoculated onto agar in 1-µl aliquots from frozen spore suspensions. S. coelicolor and the initiator strains were spotted 3 mm apart at the same time. For IMS experiments, a second colony of S. coelicolor was spotted 1 cm away from the first S. coelicolor colony to serve as an internal control. Petri plates were incubated at 30°C until the appropriate time points in large sealable bags to prevent desiccation.

MALDI-TOF imaging mass spectrometry. IMS was carried out as described previously (22, 24). At the appropriate time point, colonies and the surrounding agar were cut and removed from petri plates and transferred to Bruker MSP 96 anchor plates. The samples were then sprinkled with Universal MALDI Matrix (Fluka 50149) using a 53-µm sieve. Once a thick, uniform layer of matrix was deposited on the sample, it was placed at 37°C for 4 h or until it was completely desiccated. Excess matrix was blown off with compressed air, and any residue remaining on exposed surfaces of the MALDI plate was wiped away with methanol. Spectra were acquired using a Bruker Autoflex MALDI-TOF MS, and ions were visualized using FlexImaging software. 3D IMS was performed as described previously (34). Briefly, an 8-µm-thick section of agar was excised and cut widthwise at room temperature using microtome blades into 1.0-mm sections with each section placed on its side on the MALDI plate. 2D MALDI-TOF IMS analysis was conducted. Then, data were imported in the Matlab software program, where sections were aligned and processing was performed. The resulting 3D volume data set corresponding to an m/z value was visualized using 50%, 75%, and 90% semitransparent iso-surfaces, indicating 80%, 88%, and 95% relative abundances of the molecular compound within the imaging area, respectively.

NanoDESI mass spectrometry. NanoDESI mass spectrometry was carried out essentially as described elsewhere (21). The NanoDESI source was coupled to a Thermo LTQ-FT-ICR (linear trap quadrupole Fourier transform ion cyclotron resonance) MS. Briefly, at the desired time points, samples were placed on the NanoDESI sample stage, and the liquid bridge was placed in contact with the sample for 15 min. During this time, the liquid bridge was placed at several locations on the colonies themselves and on the agar adjacent to the colonies located on the opposite side from the interacting strain. All data were collected in the positive ion mode in the range of 100 to 2,000 m/z using a solvent consisting of 65% acetonitrile and 35% water containing 0.05% formic acid. The data acquisition mode included ions for 10 min once they had been trapped and fragmented three times.

Computation of mass spectral networks. Spectral networks were assembled largely as described elsewhere (21). MS2 spectra were clustered using the MS-Clustering software program to build consensus spectra for repeatedly observed ions (31). Pairs of consensus spectra were aligned if both spectra fell within the top 10 alignments for each of the respective spectra and the cosine of their peak match scores was ≥0.6. The algorithm assumed a peak mass tolerance of 1.0 Da and an MS2 peak tolerance of 0.5 Da. The networks were visualized in the software program Cytoscape (51), where consensus spectra are represented as nodes connected by edges to aligning nodes. In order to maximize chemical/biological interpretability, nodes found in several control data sets were removed. These included nodes found associated with the solvent, the agar substrate, and S. coelicolor colonies grown in isolation at 3, 5, and 7 days. Finally, only nodes that were reproducible in both runs of data collection were retained.

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**ADDITION IN PROOF**

While this manuscript was in review, another paper appeared that also observed the production of acylated desferrioxamines from *S. coelicolor* served the production of acylated desferrioxamines from

**REFERENCES**