

A comparative metabologenomic approach reveals new mechanistic insights into *Streptomyces* antibiotic crypticity

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Classification: Physical Sciences: Chemistry; Biological Sciences: Microbiology

Keywords: *Streptomyces griseus*, cryptic metabolism, regulation, metabologenomics, tetramic acid

Significance

Streptomyces genomes harbor an immense trove of biosynthetic gene clusters (BGCs) that encode for drug-like molecules. However, only a fraction of these readily yield expected products. To investigate why this is, we used polycyclic tetramate macrolactam (PTM) antibiotic production as a model system. By comparing the genomes and PTM production profiles of several closely-related *Streptomyces griseus* clade members, we uncovered two distinct mechanisms that differentiate more robust producers from weaker ones. The first involves small insertion-deletion lesions in PTM BGC promoters that significantly modulate production. The second mechanism involves biosynthetic pathway interactions where robust PTM producers unexpectedly benefit from griseorhodin co-production and weaker producers lack the pathway. We highlight comparative metabologenomics as a powerful approach to understand antibiotic crypticity.

Abstract

Streptomyces genomes harbor numerous biosynthetic gene clusters (BGCs) encoding for drug-like compounds. While some of these BGCs readily yield expected products, many do not. Biosynthetic crypticity represents a significant hurdle to drug discovery, and the biological mechanisms that underpin it remain poorly understood. Polycyclic tetramate macrolactam (PTM) antibiotic production is widespread within the *Streptomyces* genus, and examples of active and cryptic PTM BGCs are known. To reveal new insights into the causes of biosynthetic crypticity, we employed a PTM-targeted comparative metabologenomics approach to analyze a panel of *S. griseus* clade strains that included both poor and robust PTM producers. By comparing the genomes and PTM production profiles of these strains, we systematically mapped PTM promoter architecture within the group, revealed that these promoters are directly activated via the global regulator AdpA, and discovered that small promoter insertion-deletion lesions (indels) differentiate weaker PTM producers from stronger ones. We also revealed an unexpected link between robust PTM expression and griseorhodin pigment co-

production, with weaker *S. griseus* -clade PTM producers being unable to produce the latter compound. This study highlights promoter indels and biosynthetic interactions as important genetically-encoded factors that impact BGC outputs, providing mechanistic insights that will undoubtedly extend to other *Streptomyces* BGCs. We highlight comparative metabologenomics as a powerful approach to expose genomic features that differentiate strong antibiotic producers from weaker ones. This should prove useful for rational discovery efforts and is orthogonal to current engineering and molecular signaling approaches now standard in the field.

Introduction

Many therapeutics derive from natural products and their synthetic analogs (1). Historically, *Streptomyces* and related actinobacteria were heavily screened for these molecules, which resulted in numerous essential medicines. These include many clinical antibiotics (2), and there is an urgent need for new anti-infectives to counter increasing drug-resistance (3). A massive reservoir of uncharacterized biosynthetic gene clusters (BGCs) encoding drug-like molecules resides within *Streptomyces* genomes, which has triggered resurgent interest in these organisms (4). However, a large proportion of these BGCs fail to produce detectable levels of the expected compounds under laboratory conditions. This phenomenon of cryptic or silent metabolism thus poses a significant hurdle to genomics-driven drug discovery (5, 6). Silent BGCs are often thought to be transcriptionally deficient, and synthetic biology, cell signaling, and stress mechanisms are commonly used to activate silent BGCs for molecule discovery (7). Despite decades of research on the regulation of antibiotic production in *Streptomyces* (8, 9), a deeper mechanistic understanding of *Streptomyces* silent metabolism is still needed to access the full biosynthetic potential of these organisms to overcome the drug discovery gap (10).

Here, we employed a comparative metabologenomic approach to dissect why certain *Streptomyces* strains are ready antibiotic producers while others have apparently silent BGCs.

Polycyclic tetramate macrolactam (PTM) antibiotics were specifically targeted in this study because they provide an opportune model system for multi-strain comparative analyses. This is because PTM BGCs can contain as few as three genes (11), greatly simplifying regulatory studies. Another advantage of PTM BGCs is their relative commonality (12). A survey of bacterial genomes in GenBank published in June 2016 reported over 80 PTM BGCs were detected within 669 *Streptomyces* genomes available on GenBank at the time (13), and this commonality is leverageable for in-depth comparisons between PTM producers. The environmentally and biotechnologically important *Streptomyces griseus* clade (14) was identified as a particularly advantageous cohort for these comparisons because multiple strains within it had established active or cryptic PTM BGCs (12, 15), plus several additional family members with sequenced genomes and yet-unstudied PTM clusters were available from public collections. Finally, PTM biosynthesis is increasingly understood, and this foundational knowledge was necessary to enact a targeted metabolomics approach to document sensitive production differences. PTMs have experienced intense study towards understanding their unusual hybrid non-ribosomal peptide/polyketide origins, investigating PTM therapeutic potential, and leveraging their ease of manipulation via synthetic biology (13).

This study was initiated by comparing the genomes and PTM production profiles among a cohort of PTM locus-bearing *S. griseus* clade strains, which included known examples of both strong and poor PTM producers. This led to the discovery of a subclade that has consistently higher PTM production and PTM BGC promoter strengths compared to the rest of the test strains. Despite the exceptional commonality of PTM biosynthetic loci in *Streptomyces* bacteria, PTM regulation remains poorly understood. To reveal how promoter sequence variations might underpin the observed PTM phenotype differences, the promoter driving production in the robust producer *Streptomyces* sp. strain JV180 was thus mapped and compared against the rest of the clade. Overall, many promoter features appeared to be largely conserved within the group, regardless of host-strain PTM capability. Additionally, the well-characterized global regulator AdpA (16) was confirmed to play a direct positive role on PTM locus control through gene deletion, binding site mutation, and *in vitro* binding experiments. AdpA binding sites were

detected in the PTM promoters of all tested clade members, where they display a contextually unusual arrangement downstream of promoter –10 boxes. Critically, comparisons of strong and weak PTM promoters from several *S. griseus* clade members identified a 2-3 bp indel, located between the –10 box and AdpA operator site, that strongly influences PTM production strength. Yet another PTM control mechanism that differentiates weak from stronger PTM producers was discovered following the mutagenesis of strain JV180's griseorhodin BGC. Weak *S. griseus* clade PTM producers natively lack griseorhodin BGCs, and loss of the JV180 cluster severely curtailed PTM production. Further dissection revealed that strain JV180's PTM production likely benefits from PTM-griseorhodin co-expression via a yet-uncharacterized transcriptional mechanism. In sum, this work revealed two new mechanisms by which stronger and weaker PTM producers are differentiated, and it highlights the application of targeted comparative metagenomics to cohorts of related strains to successfully reveal otherwise difficult-to-detect genomic features that tune antibiotic production.

Results

The *S. griseus* clade is an ideal model group to compare and reveal genetic underpinnings of PTM crypticity

Several features of the *S. griseus* clade make the group attractive for comparison-based approaches to reveal BGC silencing mechanisms. Clade-member *Streptomyces* sp. strain JV180 readily produces PTM compounds (12, 17), while *S. griseus* subsp. *griseus* strain IFO13350 harbors a silent PTM BGC (15, 18). Nevertheless, promoter refactoring of the IFO13350 PTM BGC and its expression in a heterologous host successfully yielded PTMs with 5/5-carbocyclic ring systems, proving the functionality of its encoded enzymes. The PTM production status of most other clade members remained unknown, and to benefit from the increased analytical depth afforded by expanded cohort sizes, several additional clade-members having PTM loci in their sequenced genomes (19) were obtained from public strain repositories (JV251-JV258). To complete the cohort, the environmental clade-member *Streptomyces* sp. strain SP18CM02,

whose genome was recently reported by our group (17), (Table S1) was also included. The phylogenetic relationships of these strains were inferred through multi-locus phylogeny (14) (Fig. S1). Importantly, the PTM BGCs of these strains appeared to be orthologous. This was based on several observations, including that their PTM BGCs share identically-ordered biosynthetic genes (*ftdABCDEF*), and the chromosomal regions that immediately flank their respective PTM BGCs also have identical gene content (see Fig 1A for a PTM BGC diagram representative of all strains studied here, and the figure legend for additional detail). Finally, the PTM enzymes encoded within each studied BGC also shared high pairwise identities to those of strain JV180 (Table S4).

To assess and compare PTM production in these strains, the robust PTM-producing strain JV180 served as an archetype. Strain JV180 PTM production was extensively analyzed using approaches similar to those reported for clifednamide-type PTM analyses in other *Streptomyces* by our group (20). Putative PTM peaks were identified using ¹³C₅-labeled ornithine precursor incorporation and diagnostic daughter-ion production following collision-induced dissociation in liquid-chromatography coupled tandem mass spectrometry (LC-MS/MS). These target the conserved ornithine-derived tetramate region, which has two possible hydroxylation states (see Fig. S2B-E, S3). As expected, all PTM peaks identified through these methods were absent when the JV180 PTM BGC was deleted (Δ *ftdABCDEF*; Fig. 1A, S2A), yielding a high-confidence set of *S. griseus* clade PTM congeners for quantitative production comparisons.

The remaining *S. griseus* clade strains were then grown under several conditions, and their extracts were analyzed via LC-MS/MS and UV spectrometry for PTM production (see Methods). As expected from orthologous PTM BGC's, all producing strains gave PTM signals that largely overlapped with those established in JV180, but strain-to-strain differences in quantity and relative PTM congener ratios were observed. From these comparative PTM production data, an interesting trend emerged. Strains JV180, SP18CM02, and JV251-253 displayed robust PTM production while the remaining strains generally showed little production (Fig. 1B, S4-S6). We noted that these strains belong to a distinct subclade on the *S. griseus* clade

phylogenetic tree (Fig. S1, group VI in green), suggesting they might share a conserved genetic basis for increased PTM production versus the other studied clade-members. Interestingly, these comparative production analyses revealed a clear bias for higher PTM production on solid media over shake-flask cultures for all tested strains (Fig. 1B). The strongest PTM producers continued to have the highest production in liquid media as well. Throughout, to obtain consistent transcriptional information, liquid media was used. Further, in agreement with prior publications, PTM production could be detected by UV absorbance in JV180 (12), while strain IFO13350 produced insufficient PTMs for detection using this method (Fig. S7A). In contrast, our PTM targeted LC-MS/MS analyses revealed detectable production in all tested clade members, but the amounts varied widely by strain. This suggests prior efforts to characterize weak PTM producers may have overlooked scant actual production due to inherent UV limitations, overcome here by MS/MS methodology (Fig. S7B, C).

PTM promoter sequence heterogeneity contributes to differences in antibiotic production

Because the most robust PTM production was found in strains of the JV180 subclade, we hypothesized that comparing these genomes against other *S. griseus* clade strains could reveal specific differences that underpin the observed production disparities. PTM BGC transcriptional differences were immediately targeted as a potential mechanism. While weak transcription is often implicated in the cryptic biosynthesis literature, this assertion is often speculative or left without mechanistic investigation (21). The possibility of PTM biosynthetic enzyme defects in some strains was ruled out as a potential cause. This is because all tested strains have high PTM BGC protein-sequence identity (Table S4), and previous transcriptional refactoring experiments were able to successfully activate the cryptic PTM BGC of strain IFO13350 in heterologous hosts as a demonstration of BGC functionality (15, 18).

Possible strain-linked differences in PTM BGC transcription were initially explored by comparing the presumed PTM promoter regions of each strain tested here. *S. griseus* clade PTM BGCs have a simple and conserved gene arrangement with minimal intergenic gaps,

suggestive of a single operon (Fig. S8A). PCR amplification of cDNA intergenic junctions confirmed that the JV180 PTM BGC is transcribed as a single operon (Fig. S8B).

Phylogenetic analysis of the ~500 nt located upstream of the first gene in each studied PTM BGC, *ftdA*, formed two groups: one by JV180-like strains and one by the remaining strains (these sequences are hereafter referred to as JV180- or IFO13350-like P_{ftdA} 's, respectively; Fig. 1C, S9). To test if the sequence differences in P_{ftdA} regions are transcriptionally relevant, we leveraged the strong PTM producer JV180. JV180 represents a uniform background to eliminate complicating metabolic and genetic variables inherent to strain-to-strain comparisons. After replacing the native promoter region of JV180 with the corresponding region of the other 10 *S. griseus* clade strains in this study, we generally observed higher PTM production and *ftdB* transcription from JV180-like P_{ftdA} 's than IFO13350-like P_{ftdA} 's (Fig. 1D, E). This supports the idea that stronger PTM producers might result from inherently stronger promoters.

Surmising that the basic transcriptional machinery of all *S. griseus* clade-members should be highly similar, we also hypothesized that if JV180-like P_{ftdA} 's are stronger than IFO13350-like P_{ftdA} 's, then introducing the former promoter type into strain IFO13350's PTM BGC should increase PTM production and transcription. However, exhaustive attempts to introduce the necessary plasmids to engineer increased production in IFO13350 (and other group member JV257; Fig. S1) failed by both conjugation and protoplast transformation. Instead, the effects of P_{ftdA} sequence variation were assayed in strain IFO13350 using a *xylE* (22) reporter gene chromosomally integrated into the Φ C31 site. Controls testing the efficacy of colorimetric XylE assays in both strains JV180 and IFO13350 were successful when using the strong constitutive promoter P_{ermE^*} , but both P_{ftdA_JV180} and $P_{ftdA_IFO13350}$ failed to drive observable XylE in both strains. However, RT-qPCR assays to detect *xylE* transcripts were successful in making the desired comparison, confirming P_{ftdA_JV180} to be stronger than $P_{ftdA_IFO13350}$ in both JV180 and IFO13350 hosts (Fig. S10). Together, the above data supported a model where P_{ftdA} sequence variations might significantly influence PTM production or silence. While BGC promoter heterogeneity is a concept that remains underexplored in the biosynthetic literature, the idea of tuning BGC outputs through promoter strength is grounded in numerous

studies where silent BGCs can be activated by replacing their native promoters with stronger ones (see recent review 23).

Mapping promoter regions of the JV180 PTM BGC enables structure-function comparisons

Existing PTM regulatory knowledge is sparse (24, 25), including within the *S. griseus* clade. Our above results suggested that *S. griseus* clade PTM regulation involves *cis*-regulatory elements located directly upstream of *ftdA* (~500 nt), although we could not rule out additional influences from *trans*-regulatory elements. To define the PTM promoter architecture of these strains, the transcriptional start site (TSS) of strain JV180 was mapped via circular Rapid Amplification of cDNA Ends (5'-cRACE). The putative TSS was a cytosine residue 195 nt upstream of the predicted start codon of *ftdA* (Fig. S11). Likely -10 and -35 boxes were assigned based upon established spacing (26, 27) (Fig. 2A). The high sequence conservation between the -35 box and the TSS (82.4% - 100% pairwise sequence identity) suggested that all examined *S. griseus* clade strains share the same core promoter (-35, -10 and TSS).

To better resolve P_{ftdA} promoter architecture and probe for the presence of *cis*-regulatory signatures upstream of the predicted -35 boxes, a series of nested deletions in this region were created in JV180 (Fig. 2A). PTM transcription and production in these deletions were largely unaffected, except where the putative -35 box was disrupted (Fig. 2A, Δ -528_-31). This confirmed the position of the JV180 -35 box, and suggested this region lacks any critical regulatory residues (Fig. 2B, C). Further, we observed highly variable sequence conservation within this region upstream for all studied *S. griseus* clade strains (38.2% - 97.8% pairwise sequence identity). This contrasts with the more strict sequence conservation seen in the core promoter region (-35, -10 and TSS).

The -10 boxes and TSS residues of all examined strains are perfectly conserved, but several single nucleotide polymorphisms (SNPs) differentiate the -35 boxes of the IFO13350-like P_{ftdA} 's (Fig. 2A, red box) from the strictly conserved -35 boxes of the JV180 group (Fig. 2A, green box). It is known that changes in bacterial -35 boxes can greatly affect promoter strength

(28, 29), and could thus affect metabolite production. Therefore, a panel of JV180 mutants carrying each observed -35 box SNP in the IFO13350 group was created and tested for PTM production and transcription (Fig. S12A). Overall, the IFO13350 group -35 box SNPs failed to significantly change PTM production and only slightly decreased transcription in the chimeric JV180 hosts (Fig. S12B, C). These differences were subtle compared to the far more substantial ones seen in our ~500 nt replacements upstream of the *ftdA* gene (Fig. 1D and E), indicating P_{ftdA} strength differences must originate through mechanisms other than -35 box differences.

AdpA positively regulates *S. griseus* clade PTM BGCs

Many *Streptomyces* mRNAs contain long 5' untranslated regions (5' UTRs) (26, 27), and these can contribute to regulatory tuning. The DNA regions encoding 5' UTRs can modulate gene expression through direct regulator binding (30), and their corresponding mRNAs can further modulate expression via riboswitches (31) or other RNA secondary structures (32). A series of ~20 nt deletions were constructed (Fig. 2D) across the 195 nt 5' UTR region of JV180 to probe contributions to PTM regulation. This revealed multiple lesions with strongly decreased PTM production and transcript levels, while others had little effect (Fig. 2E, F). mFold (33) modeling to reveal possible 5'UTR mRNA secondary structures across *S. griseus* clade strains yielded several energetically favored outputs, and we surmise some of the deleterious mutational effects seen in JV180 might stem from the disruption of these types of structures (Fig 2D inset, Fig. S13-17).

While *S. griseus* clade P_{ftdA} 5' UTR folding remains to be further explored, the above deletions were particularly useful for identifying a critical region that we subsequently characterized as an AdpA binding site (Fig. 2D-F; $\Delta 29_{-48}$, Fig. S9). AdpA is a global regulator that is well-studied in strain IFO13350, where it is known to bind >500 chromosomal sites via its weak consensus sequence TGGCSNGWWY (34). *S. griseus* AdpA is involved in the hierarchical control of morphological differentiation, the production of streptomycin and other antibiotics, and several other important processes (see review (16)). Prior ChIP-Seq and RNA-seq data

indicated that AdpA might bind upstream of IFO13350's PTM BGC (35), but because prior efforts to elicit PTM production from wild-type IFO13350 were unsuccessful (15, 18), the biological significance of AdpA's interaction with the gene cluster was tenuous. Our searches for potential AdpA consensus motifs within the P_{ftdA} regions of the studied *S. griseus* clade strains revealed an imperfect inverted repeat 29-48 nt downstream of the TSS in the JV180 P_{ftdA} (within the 5' UTR region) that was conserved in all *S. griseus* clade strains examined. Because our nested deletions revealed that this sequence is essential for JV180 PTM production (Fig. 2D-F, Δ_{29_48} , Fig. S9), it renewed the notion that AdpA might positively regulate PTM expression.

Several experiments were carried out to investigate whether AdpA regulates P_{ftdA} , including *adpA* deletion and complementation analysis, mutating the putative AdpA binding site in P_{ftdA} , and *in vitro* binding assays. As expected from prior *adpA* studies in strain IFO13350 (36), deletion of the JV180 ortholog ($\Delta adpA$) led to the loss of morphological differentiation and pigmentation (Fig. S18). PTM production and BGC transcription were also abrogated in JV180, and these defects were rescued by ectopically expressing either native *adpA*, its IFO13350 ortholog (97% amino acid identity vs JV180 AdpA) or by replacing P_{ftdA} with the strong constitutive promoter P_{ermE^*} to drive PTM BGC expression independent of AdpA (Fig. 3A, B, S19). These results were consistent with AdpA being a transcriptional activator for P_{ftdA} . However, in bacterial regulation, regulators binding downstream of the RNA polymerase complex typically cause transcriptional down-shifts (37, 38). Thus, the location of the putative AdpA operator site downstream of the conserved -10 box is unusual for a transcriptional activator. Importantly, another AdpA-activated promoter with a downstream operator site controls *S. griseus* IFO13350's *adsA*, encoding an extracytoplasmic function sigma factor (30). This precedent thus supports the idea that AdpA could positively regulate PTM expression despite its atypical putative binding arrangement in P_{ftdA} .

AdpA-DNA co-crystallization studies indicate that the protein binds target operators as a homoduplex that recognizes a highly variable motif, containing four core invariant guanosine and cytosine nucleotides (Fig. 3C, bold residues in gray box) (34). In all *S. griseus* clade P_{ftdA} promoters studied here, these invariant residues were perfectly conserved, and introducing a

transversion mutation at any one of these nucleotides abrogated JV180 PTM production and transcription (Fig. 3C-E). This additional evidence further suggests that the identified region acts as a functional AdpA operator. Exchanging the JV180 AdpA operator site with that from IFO13350 was PTM proficient although it showed a slight decrease in PTM titer and *ftdB* transcription (Fig. 3C-E). AdpA- P_{ftdA} interactions were further examined via electrophoretic mobility shift assays (EMSA) using operator sequences from JV180 and IFO13350, plus the critical point mutants that disrupt essential DNA-AdpA interactions. As expected, recombinant histidine-tagged AdpA bound the JV180 and IFO13350 P_{ftdA} AdpA binding sites (Fig. 3F, G) but failed to shift operators with the *in vivo* tested point mutations (Fig. 3H, S20). Together, our *in vivo* and *in vitro* data strongly suggest that AdpA directly binds P_{ftdA} in both JV180 and IFO13350, but also indicate that the native SNPs (Fig. 3C, red residues) in the non-essential residues of the AdpA operator sites are not the main cause of PTM expression differences seen between P_{ftdA_JV180} and $P_{ftdA_IFO13350}$ (Fig. 3D,E).

Comparative promoter analyses reveal an indel “switch” that tunes P_{ftdA} strength and PTM production

Aside from the AdpA operator, the JV180 P_{ftdA} 5' UTR truncation experiments revealed that the 28 nucleotides between the TSS and the AdpA binding site are also critical for PTM expression (Fig. 2B-F, $\Delta 2_28$). Nucleotide alignments between promoters in this region revealed generally high conservation, except for two nucleotides (AG) that are present in all JV180-like P_{ftdA} 's but are missing from IFO13350-like P_{ftdA} 's (Fig. 4A, S8). The effects of this indel on PTM regulation were tested by deleting this AG dinucleotide from JV180, and by introducing the dinucleotide at the corresponding position in $P_{ftdA_IFO13350}$ ($P_{ftdA_IFO13350}+AG$). $P_{ftdA_IFO13350}+AG$ was tested heterologously in strain JV180. Strikingly, the dinucleotide deletion led to strongly reduced PTM production and transcription, while the amended $P_{ftdA_IFO13350}+AG$ insertion variant led to a substantial increase in JV180 PTM production and transcription compared to the wild type $P_{ftdA_IFO13350}$ sequence (Fig. 4B, C). To test if the indel's effect was sequence

specific, a JV180 transversion (Δ AG::CT) mutant was created, which exhibited reduced PTM production and transcription compared to wild-type, but was much less deleterious than Δ AG (Fig. 4B,C). This region is seemingly prone to sequence plasticity within the *S. griseus* clade; the recently isolated strain SP18CM02 contains an additional guanosine in this region compared to other JV180-like P_{ftdA} 's (Fig. 4A). Despite having otherwise high overall identity to all other JV180-like P_{ftdA} 's, $P_{ftdA_SP18CM02}$ drove slightly less transcription and resultant PTM production when heterologously introduced into JV180 (Fig. 1D, E). Together, these data suggest that of the natural sequence variants in this region, having the AG dinucleotide is important for PTM production. These data clearly reveal this indel region as a key factor in modulating natural *S. griseus* clade P_{ftdA} strength. Further work is necessary to discern how this indel region modulates promoter strength, possibly via mechanisms such as perturbed AdpA-RNA polymerase interactions or recruitment of another yet-unknown regulatory component.

Discovery of an unexpected griseorhodin biosynthetic interaction that strengthens PTM production in strain JV180

Our data thus far illustrated how leveraging within-clade comparative metabologenomics can assist regulatory region mapping, and how small easily overlooked nucleotide changes in these regions can tune antibiotic production. The use of comparative genomics to understand *Streptomyces* antibiotic production, particularly at the species level, is a relatively recent development in natural products functional genomics. Studies in this area tend to focus on BGC conservation and differentiation (e.g. comparison of the *S. albus* clade, see (39)). From these comparisons, we anticipated that our *S. griseus* clade strains would share several BGCs (beyond PTMs), and that some antibiotic BGCs in these strains would not be conserved clade-wide. We found that our *S. griseus* clade strains share a core set of 13 conserved BGCs. Some BGCs were found in only a few strains, and several BGCs were unique by strain (Fig. S21). Strikingly, strain IFO13350 is one of the oldest known producers of streptomycin (40), but all members of the JV180 group were found to lack this BGC. Likewise, we noted that JV180 group strains produce red pigments that are absent from the other studied *S. griseus* clade strains (Fig. S22). Through comparative BGC analysis and subsequent

cluster deletion in strain JV180, we attributed this pigment to the production of griseorhodin polyketide congeners (41) (Fig. 5A; $\Delta grhR2$ -V, S23A-C). Unexpectedly, this griseorhodin BGC deletion mutant exhibited several additional phenotypes, including abrogated PTM production, downregulated PTM BGC transcription, and reduced sporulation (Fig. 5B-C, S24).

The loss of PTM production in the JV180 $\Delta grhR2$ -V mutant was wholly unexpected, and the mutant's pleiotropic phenotypes suggested griseorhodin might act as a signaling molecule. To test this, four adjacent genes encoding griseorhodin polyketide synthase (PKS) assembly-line enzymes were deleted ($\Delta grhQSAB$) (Fig. 5A). This was done to specifically eliminate griseorhodin production, while leaving all other *grh* BGC genes intact. Interestingly, this mutant was PTM-proficient, arguing against the signaling idea. To further probe how the griseorhodin BGC exerts its influence, we tested a griseorhodin-enriched JV180 growth extract (see methods, Fig. S25A) and authentic γ -rubromycin (42) (a griseorhodin analog) for their ability to chemically complement the $\Delta grhR2$ -V strain. Neither sample could restore PTM production in the $\Delta grhR2$ -V mutant in flask cultures (Fig. S25B), and disk diffusion tests on agar plates failed to restore sporulation in the diffusion zone (Fig. 25C). These experiments together ruled out the griseorhodin signaling hypothesis.

To continue probing the *grh* locus for key PTM-influencing genes outside of *grhQSAB*, multiple groups of genes were deleted from the BGC (*grhR1-E*, *grhFGH*, and *grhI-P*; Fig. 5A). All three mutations caused complete loss of red griseorhodin pigmentation and showed reductions in PTM production and transcription (Fig. 5B, C, S23D, S26B). Homology-based annotations of the genes in these regions (41) led us to focus on a subset which might affect PTM biosynthesis through transcriptional or metabolic mechanisms. Genes *grhR2* and *grhR3* encode transcriptional regulators, which could influence regulatory crosstalk, *grhF* encodes a phosphopantetheinyl transferase (essential for posttranslational modification of NRPS and PKS enzymes), and *grhGH* encode β and ϵ subunits of acetyl-CoA carboxylase (ACC). ACC enzyme complexes are essential for malonyl-CoA production, a common precursor for fatty acid,

griseorhodin, and PTM biosynthesis. PTM production in JV180 Δ *grhR2-V* was not complemented by *grhR2*, *grhR3*, or *grhF*, but *grhGH* was able to restore some PTM production and transcription (Fig. 5B, C, S26A). Interestingly, expressing either *grhG* or *grhH* alone were sufficient to restore PTM production, similar to *grhGH* when expressed together (Fig S26A). A Δ *grhGH* mutant was thus created, leading to partial griseorhodin pigmentation and reduced PTM production comparable to the initial Δ *grhFGH* mutant (Fig. S23D, S26B). How the Δ *grhR1-E* and Δ *grhI-P* mutations caused decreased PTM production remains unclear. Because these lesions led to loss of putative regulatory genes (*grhR2* & *grhR3*, see (41), their phenotypes could be entangled with concomitant *grhGH* downregulation.

In addition to the griseorhodin BGC's *grhGH*, several other *Streptomyces* PKS clusters are known to harbor additional non-housekeeping copies of ACC genes. These include *cpkKL* in the coelimycin BGC of *S. coelicolor* (43) and *jadN* in the jadomycin BGC of *S. venezuelae* (44), and it can be reasoned that these ACC copies likely assist polyketide biosynthesis through increased malonyl-CoA. Because JV180 PTM production decreased in the Δ *grhR2-V* and Δ *grhGH* mutants, but not the Δ *grhQSAB* mutant (Fig. 5C), we likewise hypothesized that PTM downshifts in these mutants might be caused, at least partially, by reduced intracellular malonyl-CoA concentration. We thus tested several additional *S. griseus* clade ACC subunit genes for their ability to rescue PTM production in the JV180 Δ *grhR2-V* strain to discern if these effects were specific to *grhGH* or are more broadly attributable across ACC subunit homologs. The tested ACC genes included housekeeping *accBE* alleles cloned from JV180 and IFO13350 (function assigned by sequence homology and gene neighborhood synteny to the *accBE* genes in *S. coelicolor* (45)) plus the previously-uncharacterized PKS-associated ACC genes SGR3280-3281 of strain IFO13350, all of which encode ACC β and ϵ subunits like *grhGH*. All of these ACC homologs partially restored PTM production in the Δ *grhR2-V* strain (Fig S26A), further supporting a role for malonyl-CoA in the PTM production defects of the Δ *grhR2-V* and Δ *grhGH* mutants.

To test if JV180 could still produce PTMs independent of the griseorhodin BGC, we constitutively expressed the strain's PTM BGC by replacing P_{ftdA} with P_{ermE^*} . This $\Delta P_{ftdA}::P_{ermE^*}$

promoter replacement in both of the $\Delta grhR2-V$ or $\Delta grhGH$ backgrounds led to increased PTM production and PTM BGC transcription, similar to a JV180 $\Delta P_{ftdA}::P_{ermE^*}$ control (Fig 5B, C). Our data show that PTM transcription and its production is affected by the lack of $grhR2-V$ and $grhGH$. It is possible that the griseorhodin BGC may affect the PTM BGC through biosynthetic malonyl-CoA availability. It is also likely that an as yet-undefined transcriptional regulatory interaction also connects the two BGCs. Malonyl-CoA responsive regulators are well-characterized in other model organisms such *B. subtilis* (46), but no such regulators are known in *Streptomyces*. Further inquiry is underway to characterize this unusual cross-cluster interaction more fully.

Intrigued by the finding that ectopic expression of just $grhG$ or $grhH$ could rescue PTM production in the JV180 $\Delta grhR2-V$ mutant, we tested if the PTM BGCs of weak-PTM producing *S. griseus* clade strains might be similarly stimulated for production. This was done by heterologously expressing $grhG$ in strains IFO13350, JV254, and JV258 (the $grhGH$ construct had low conjugation efficiency). This resulted in increased PTM production from these natively low-producing strains on agar media and also increased PTM BGC expression when tested in strain IFO13350, showing that ACC subunit overexpression can stimulate PTM production in natively poor producers (Fig. S27A, B). Combined with the above data, these observations support a model where *S. griseus* clade PTM production differences have complex origins. In the case of poor PTM producers, promoter indels and insufficient positive BGC interactions likely dampen potential production via low transcription. In contrast, better PTM producers appear to benefit from cross-BGC interactions that increase transcription from a more active P_{ftdA} variant (+AG), resulting in PTM production. Strain JV180, and possibly other members of its subclade, do not seem to be bottlenecked for PTM production at the transcriptional level based on several findings throughout this study. This includes failure of $grhG$ merodiploids to boost WT JV180 PTM production (Fig. S27A) and evidence from several recombinant strains where PTM BGC transcription was significantly upregulated by 2 to 9 fold, but PTM production increased by only up to 1.4 fold ($\Delta grhQSAB$, $adpA$ complementation, and $\Delta P_{ftdA}::P_{ermE^*}$ strains; Fig. 3A, 3B, 5B, 5C). The above findings also highlight ACC enzyme overexpression as a way to potentially upregulate

certain silent BGCs, adding a new dimension of utility for these genes in synthetic biology, where they are currently used for increasing fatty acid and polyketide titers via malonyl-CoA overproduction (47, 48).

Conclusions:

Streptomyces genomes contain many BGCs encoding for drug-like compounds, and to harness their full biosynthetic capacity for discovery, it is crucial to understand what underpins the differences between active and silent BGCs (10). Cryptic metabolism is a well-recognized problem in the field, and poor BGC transcription is often implicated as the predominant mechanism behind biosynthetic silence. However, this supposition is increasingly challenged by a growing body of research that suggests all cryptic clusters aren't necessarily transcriptionally silent (49). Further, it only addresses how biosynthetic silence could originate in a given organism, but does little towards explaining why affected BGCs show little activity to begin with. A common but difficult-to-prove hypothesis is that axenic laboratory growth deprives microorganisms of signals needed to upregulate quiescent BGCs (21). This idea is based on the complex lifestyles of *Streptomyces* bacteria, which are known to be heavily influenced by molecular and environmental cues (50). Indeed, chemical elicitor screening, co-culturing, and other regulatory manipulation strategies continue to yield new molecules from apparently silent BGCs, bolstering this idea (7, 50, 51). However, the number and types of molecular signals these organisms can sense and respond to must have a finite limit, constrained by the characteristics and capacities of their genetically encoded signal transduction pathways (10). If true, then extracellular signals can only explain one part of antibiotic crypticity, highlighting the need to understand how basic strain-to-strain genetic differences also contribute to the phenomenon.

Towards this, we leveraged the unusual commonality of PTM BGCs in *Streptomyces* to compare several highly related *S. griseus* clade strains to discover genomic features that differentiate strong and weak producers. These efforts led to the first sequence-defined PTM

promoter in the genus, confirmation that the global regulator AdpA acts directly on *S. griseus* clade PTM production by binding P_{ftdA} promoter regions in an atypical way, and that 2-3 bp lesions between -10 promoter regions and AdpA operator sites can cause substantial differences in transcription strength and biosynthetic output. To our knowledge, this type of inquiry is largely absent from the biosynthetic-regulatory literature, but similar promoter heterogeneity, with resultant tuned transcription, has been documented to control phase variation in bacterial pathogenicity (52). Further, because the AG indel discovered here resides in a poly-guanosine rich region, poly-N strand slippage (52) or a similar mechanism might plausibly explain how these promoter variants arise in *S. griseus* clade-member populations. As more *Streptomyces* are sequenced and characterized, it is likely that additional promoter region sequence variations will emerge as drivers of silent metabolism in other biosynthetic pathways. In proof, scientists working at WarpDrive Bio, Inc patented a method for activating silent BGCs encoding rapamycin-like antibiotics by “reversing” naturally-occurring short indels within LAL-family regulator binding sites (53, 54). Those regulatory indels were discovered through large-scale actinomycete genome sequence and molecule production comparisons at that company. Such knowledge is thus highly desirable to streamline targeted discovery efforts and direct new rational engineering approaches to activate select silent BGCs.

Finally, we also found that in strain JV180, a BGC encoding griseorhodin production was required for both PTM production and transcription. Part of this relationship seems to implicate malonyl-CoA, whose biosynthesis is encoded for by *grhGH*, but further research is clearly needed to reveal a definitive mechanism. The deletion of unwanted BGCs is commonly used to engineer “clean” chassis strains (55) or to uncover the metabolites of BGCs (43, 56). In light of this, our discovery of PTM dependence on griseorhodin was surprising and provides a rare cautionary example of how genome-minimization can have unintended consequences on broader strain-wide BGC function. Together, our results show that the seemingly simple monocistronic PTM BGC of strain JV180 appears to be under highly complex control, adding to a growing body of similarly complex BGC regulatory circuits and mechanisms in other *Streptomyces* (57, 58). Much remains to be learned about these types of regulatory networks in

Streptomyces, and continued inquiry is required for a more comprehensive understanding to support rational drug discovery and production.

Materials and Methods

Strains, Plasmids, Primers, Enzymes, Chemicals and General Methods

Strains, plasmids, and primers are described in Tables S1-3. Strains IFO13350 and JV251-258 were obtained from the Agricultural Research Service Culture Collection (NRRL). All primers were purchased from Integrated DNA Technologies. All restriction enzymes, Taq polymerase, and T4 ligase were purchased from New England BioLabs. PCR was generally carried out using KOD Hot Start DNA Polymerase (EMD Millipore) in FailSafe PCR 2X PreMix G (Epicentre). Taq polymerase was used for colony PCR to verify cloning and genome editing. *Streptomyces* genomic DNA was prepared for PCR by grinding a colony in DMSO as described by Van Dessel *et al* (59). Most media components and chemicals were purchased from Sigma Aldrich or Fisher Scientific unless specified. γ -rubromycin was purchased from Abcam. $^{13}\text{C}_5$ -L-ornithine was purchased from Cambridge Isotope Laboratories. Standard protocols for manipulating *E. coli* were based on those of Sambrook *et al* (60). *Streptomyces* cultures were routinely propagated on ISP2 agar and Trypticase Soy Broth (Difco) at 28°C. Glass beads were added to liquid cultures to disrupt mycelial clumps.

***Streptomyces* conjugations**

Conjugations were performed using JV36 as the general *E. coli* donor as previously described (61). *Streptomyces* sp. strain JV180 spores were collected from lawns plated on 8340 agar (1% Proflo cottonseed meal (ADM); 2% D-mannitol; 0.1% yeast extract; 0.01% KH_2PO_4 ; 0.01% $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$; 0.002% $\text{CaCl}_2 \cdot 2\text{H}_2\text{O}$; 0.2% (v/v) R2 trace elements solution (62); 2.067% MES hemi-sodium salt; 2% agar; pH 6.5) using sterile cotton swabs into TX Buffer (63). Strain JV254

and JV258 spores were collected from lawns plated on SFM (62) and ISP-S (61), respectively. Exconjugants were selected with 50 µg/mL colistin and 50 µg/mL apramycin. Successful integrations by Φ C31 integrase-based vectors were verified by colony PCR as previously described (20).

Marker-less gene deletion/promoter replacement

All gene deletions and P_{ftdA} mutants were constructed using double homologous recombination as previously described (20). To avoid undesired recombination between the wild-type and the mutant P_{ftdA} sequences, an intermediate $\Delta P_{ftdA}::tsr$ mutant was constructed, with specifics provided the Supplementary methods. This $\Delta P_{ftdA}::tsr$ mutant was used as the parent strain for the construction of most P_{ftdA} mutants. Some plasmids for genome editing were cloned using overlap-extension PCR instead of Gibson assembly as described previously (64) (see Table S2).

PTM production and analysis

Strains were streaked from -80°C glycerol stocks onto ISP2 plates and incubated at 28°C for 2-3 days. A plug was cut from the plate and used to inoculate 3 mL of Trypticase Soy Broth in 24-well deep well plates, which were shaken at 300 rpm at 28°C . One 4 mm glass bead was added per well to disrupt mycelial clumps. After 2 days of growth, 200 µL of cultures were either inoculated into 20 mL ATCC-MOPS (adapted from ATCC172: 1% dextrose; 2% soluble starch, 0.5% yeast extract; 0.5% N-Z amine; 0.63% MOPS; pH 7.2) in a 125 mL flask with 6 mm glass beads for disrupting clumps or plated on 8340 agar overlaid with cellophane and incubated at 28°C . Flask cultures were shaken at 250 rpm. After 4 days of growth, 1 mL was collected in RNALater for protein and RNA analyses (see RT-qPCR section below) and the rest of the cultures were extracted twice with equal volumes of ethyl acetate. Solid media (plate) cultures were incubated for 6 days and the mycelia and spores were collected from the cellophane for protein extraction by trichloroacetic acid as described by Bose and Newman (65) and quantification by

Bradford assay. The remaining agar was diced and extracted with ethyl acetate by soaking overnight.

The ethyl acetate extracts were dried at low pressure and re-suspended in 500 μ L of LC-MS grade methanol and syringe filtered before LC-MS analysis. PTM analysis was performed using a Phenomenex Luna C18 column (75 x 3 mm, 3 μ m pore size) installed on an Agilent 1260 Infinity HPLC connected to an Agilent 6420 Triple-Quad mass spectrometer. For each run, 10 μ L sample was injected and the chromatography conditions were as follows: $T = 0$, 5% B; $T = 4$, 45% B; $T = 12$, 53% B; $T = 16$, 100% B, $T = 20$, 100% B; A: water + 0.1% formic acid, B: acetonitrile + 0.1% formic acid; 0.8 mL/min. The diode array detector was set to measure absorbance at 320 nm. The mass spectrometer was set to precursor ion scan mode with the precursor ions m/z : 450 – 550, collision energy = 30 V, fragmentor = 70 V, and daughter ions m/z : 139.2 or 154.2. The resulting data was analyzed offline with Agilent MassHunter Qualitative Analysis software. Chromatograms were extracted for each parent-daughter ion mass transition, and the integrated areas for the major PTM congeners (see Fig. S28) were used to compare PTM production. One PTM peak had identical retention times and fragmentation spectra as an authentic standard of maltophilin (sourced from EMC Microcollections, GmbH; Fig S29), a stereoisomer of 10-*epi*-maltophilin produced by *Streptomyces* sp. strain SCSIO 40010, which has a similar PTM BGC to strain JV180 (66). The sums of PTM peak areas were normalized by total protein, and the relative PTM production was calculated relative to the appropriate control strain, typically JV180 *rpsL*. PTM production experiments were generally carried out in triplicates, unless specified. The statistical significance in the differences observed was calculated by two-tailed Student's T-test, typically relative to JV180 or its *rpsL* mutant, JV307 or otherwise indicated. * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$. Error bars represent standard deviation.

Generally, the PTM production data reported were based on liquid media cultures that were used to collect the corresponding RT-qPCR data. Relative PTM production trends were consistent between solid and liquid media. Some figures in the SI show relative PTM production on solid media, where flask culture/RT-qPCR data was not collected.

569

570 **RT-qPCR**

571 Strains were cultivated in flasks as described above. After 4 days, 1 mL of culture added to 2 mL
572 of RNAlater and vortexed to stabilize RNA. The mixture was centrifuged at 3,214 X *g* for 10
573 minutes and the supernatant was discarded. The RNA-stabilized pellet was resuspended in 250
574 μ L of 10 mg/mL lysozyme (Sigma) and incubated at 37°C for 30 minutes. To the lysate, 750 μ L of
575 Trizol reagent (Fisher) was added and protein and RNA were extracted following the
576 manufacturer's protocol from this point. Protein concentration was measured by Bradford
577 assay. The RNA was resuspended in 84 μ L of nuclease-free water. DNase treatment was carried
578 out by adding 10 μ L 10X Turbo DNase buffer, 4 μ L Turbo DNase with 2 μ L of RNAsin (Promega)
579 for approximately 6 hours at 37°C. Removal of leftover DNA was confirmed by PCR and gel
580 electrophoresis before the DNase inactivation reagent from the Turbo DNase kit (Fisher) was
581 added. RNA concentration was measured with a NanoDrop and 5 μ g of RNA was used for
582 reverse transcription with Superscript II (Thermo) following the manufacturer's protocol.

583 Primers for qPCR were designed using the IDT primerquest tool. Real-Time PCR was
584 performed on a CFX Connect Real-Time PCR Detection System (BioRad) with the following
585 program: 1 cycle at 95°C for 3 min, 40 cycles of 95°C for 10 s and 55°C for 30 s. Each reaction
586 contained 5 μ L of iTaq Universal SYBR Green Supermix (BioRad), 2 μ L of nuclease free water, 1
587 μ L of 10 μ M forward primer, 1 μ L of 10 μ M reverse primer, and 1 μ L of template cDNA. The
588 relative transcript abundance was calculated using the $\Delta\Delta C_T$ method and *hrdB* was used as the
589 housekeeping gene (67). The primer efficiency was determined as described by the qPCR
590 instrument manufacturer for several pairs of PTM BGC probes, and the *ftdB* primers YQ376-
591 180ftdB1153 and YQ377-180ftdB1278 were chosen for subsequent experiments as they had
592 the highest efficiency and produced the most consistent results. Data shown represent at three
593 technical replicates each for at two biological replicates. The statistical significance in the
594 differences observed was calculated by two-tailed Student's T-test, typically relative to JV180 or

its *rpsL* mutant, JV307 or otherwise indicated. * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$. Error bars represent standard deviation.

Expression and purification of His-tagged AdpA

The full-length *adpA* gene of *Streptomyces* sp. strain JV180 was amplified by PCR using primers pET11a-AdpA-F and pET11a-AdpA-R. The PCR product was cloned into the expression plasmid pET11a. The expression recombinant plasmid, pKN052, contained the sequence *adpA*-CTC-GAG-(CAC)₆-TGA under the control of the T7 promoter, similar to the construct reported by Yamazaki *et al* (30). *E. coli* BL21 (DE3) Rosetta harboring pKN052 was cultured in LB medium with 100 µg/mL ampicillin at 37°C overnight. 1 mL of seed culture was transferred to 150 mL of LB medium with 100 µg/mL ampicillin and incubated by shaking at 37°C at 250 rpm. When OD₆₀₀ reached 0.6-0.8, the cells were chilled on ice for 1 h. After adding IPTG to 1 mM, the culture was continued for shaking at 250 rpm for 22 h at 18°C. The cells were harvested by centrifugation at 5000 rpm for 30 min, resuspended in Tris-HCl buffer (20 mM Tris-HCl, 200 mM NaCl and 10% glycerol: pH:8.0) and stored at -80°C. To purify the protein, cells were disrupted by sonication (3 min: 10 sec on/10 sec off at 20% amplitude) and His-tagged AdpA was purified with nickel-nitrilotriacetic acid (Ni-NTA, Qiagen) resin by eluting with 250 mM imidazole. Protein expression was verified with SDS-PAGE before downstream experiments.

Electrophoretic mobility shift assay

Double-stranded DNA probes containing the 20 bp putative P_{ftdA} AdpA binding site plus 15 bp additional flanking sequences were synthesized by IDT. The DNA was resuspended in TEN buffer (10 mM Tris, 1 mM EDTA, 0.1 M NaCl, pH 8.0). For each probe, 50 nM probes were prepared using the Dig Gel Shift Kit, 2nd generation (version 10, Roche) with the following conditions: 20 µl DNA (from 100 nM stock), 8 µl 5X labeling buffer (1 M potassium cacodylate; 125 mM Tris-HCl; 1.25 mg/ml bovine serum albumin; pH 6.6), 8 µl CoCl₂ (25 mM), 1.5 µl Digoxigenin-11-

DDUTP (1 mM), 1.5 µl terminal transferase, and water up to 40 µl. After brief mixing and centrifugation, reactions were incubated at 37°C for 30 min and then chilled on ice. The reaction was quenched by adding 2 µl of 200 mM EDTA (pH 8.0).

The protein-DNA interaction assay protocol was based on that described by Ming *et al*(34). Purified AdpA-His was serially diluted in protein buffer (10 mM Tris-HCl, pH7.5; 100 mM NaCl; 2.5% (w/v) glycerol; and 0.25 mM DTT). The reaction mixtures contained 2 µl of binding buffer (200 mM Tris-HCl, pH 7.5; 1 M KCl; 2.5 mg/mL bovine serum albumin (BSA); and 1% nonidet P-40), 1.2 µl of 1 ng/µl poly[d(I-C)], 1.8 µl DNA probe (4.5 nM), the desired volume of purified protein, and water up to 20 µl. Samples were incubated for 1 h on ice. Gel electrophoresis was performed with a 5% native acrylamide gel (Bio-RAD, mini protein TBE precast gel) in 0.5X Tris-borate-EDTA buffer (TBE, 10X concentration: 890 mM Tris; 80 mM boric acid; and 20 mM EDTA, pH 8.3) at 85 V for 145 min. Electro-blotting was performed using a BioRad Turbo Transfer System (Trans-Blot Turbo™ System) on positively charged nylon membrane (Sigma-Aldrich). Crosslinking was performed by baking the nylon membrane at 120°C for 30 min. Subsequently, the chemiluminescent detection was done as mentioned in the DIG-KIT protocol with very slight modification (overnight blocking, 2 hrs anti-digoxigenin-AP treatment, and washing for 30 min each time) and the imaging was done with a LICOR (ODYSSEY Fc) imaging system (model # 2800).

Acknowledgements

We thank Jahdiel Berrios, Mayra Banuelos, and Adam Robinson for their bioinformatics assistance, Brandon Chia and John D'Alessandro for cloning assistance, and Prof. Doug Chalker for technical advice. We also thank Prof. Arpita Bose & members of the Bose Lab for useful discussions and proofreading assistance. This work was supported by WUSTL new faculty start-up funds and the National Science Foundation under NSF-CAREER 1846005 to J. A. V. Blodgett.

Contributions

647 J. A. V. B. conceived the project. Y. Q. and K. K. N. performed experiments. J. A. V. B., Y. Q., and
648 K. K. N. analyzed the data. J. A. V. B., Y. Q., and K. K. N. wrote the manuscript.

649 **Competing Interest Statement**

650 J. A. V. B. was a former employee, consultant and advisory member of WarpDrive Bio, Corp.
651 WarpDrive Bio is now a wholly- owned subsidiary of Revolution Medicines where J.A.V. B. is a
652 minority shareholder.

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Figure Legends

Figure 1. Differences in *S. griseus* clade PTM production correlate with putative PTM promoter region sequence variation. (A) In all tested *S. griseus* clade strains, the PTM locus is comprised of a conserved set of genes *ftdA-F*. The non-PTM upstream and downstream genes in all tested strains respectively encode a cysteine desulfurase (ORF –1) and mandelate racemase (ORF +1). (B) PTM production is most robust on solid media (blue bars, n = 3) versus when strains are grown liquid (brown bars, n = 4) media. n.d. = not detected. (C) Neighboring joining tree of the 500 bp upstream of each strain's *ftdA* gene. This tree indicates that P_{ftdA} sequences of JV180-like strains form a separate group from the other (IFO13350-like) P_{ftdA} 's. The analogous P_{ftdA} region of *Streptomyces albus* strain J1074 was used as an outgroup.

Sequences were aligned with MUSCLE and the tree was built using the Kimura substitution model in CLC Main Workbench. Branches supported by less than 50% of 500 bootstrap simulations were collapsed. **(D)** PTM production resulting from non-JV180 P_{ftdA} regions after introduction into JV180. The X-axis labels indicate which strains were used to source each heterologously-tested promoter. The strain numbers in parentheses specify which strains in Table S1 were used in each experiment ($n = 3$). **(E)** Relative *ftdB* transcript abundance from JV180 promoter-swapped strains in panel D ($n = 6$, except JV1178 where $n = 3$). From these data, a clear trend emerges where strains and their P_{ftdA} -region promoters sourced from the subclade that includes JV180 (green highlights, panels A-D) yield higher PTM production and transcription compared to the rest of the tested *S. griseus* clade strains (red highlights, panels A-D). The statistical significance in the differences in PTM production or transcription were calculated by two-tailed Student's T-test, relative to JV180 or its *rpsL* mutant, JV307. * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$. Error bars represent standard deviation.

Figure 2. Mapping of P_{ftdA} and identification of cis-regulatory elements by deletion analysis.

(A) Diagram of the JV180 P_{ftdA} region, with the zones deleted upstream of P_{ftdA} diagrammed below. Inset: alignment of the core promoter sequence (note: the sequences are identical in JV180-like P_{ftdA} 's). **(B)** Relative PTM production from JV180 mutants with deletions in the upstream region of P_{ftdA} ($n = 3$). **(C)** Relative *ftdB* transcript abundance from JV180 mutants with deletions in the upstream region of P_{ftdA} ($n = 6$). **(D)** Diagram of the P_{ftdA} untranslated region (UTR) downstream of the transcript start site. Regions deleted downstream of P_{ftdA} are diagrammed below. Inset: cartoonized representation of the JV180 P_{ftdA} UTR secondary structure predicted by mFold. **(E)** Relative PTM production from P_{ftdA} UTR truncation mutants ($n = 3$). **(F)** Relative *ftdB* transcript abundance from P_{ftdA} UTR truncation mutants ($n = 6$). The statistical significance in the differences in PTM production or transcription were calculated by two-tailed Student's T-test, relative to the JV180 *rpsL* mutant, JV307. * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$. Error bars represent standard deviation.

Figure 3. AdpA is required for PTM expression and directly binds P_{ftdA} *in vivo* and *in vitro*. (A)

Relative PTM production from JV180, its $\Delta adpA$ mutant, and complementation of PTM

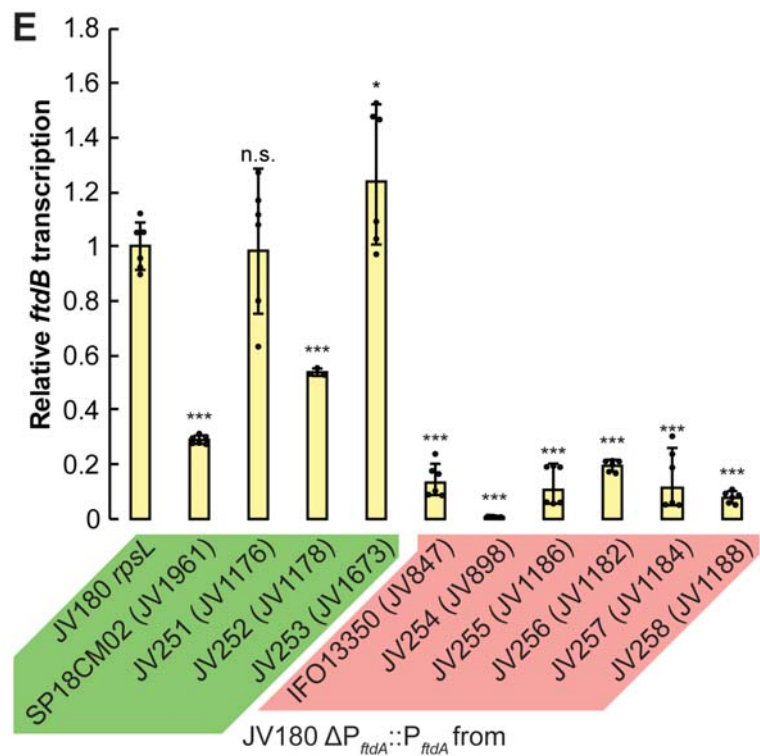
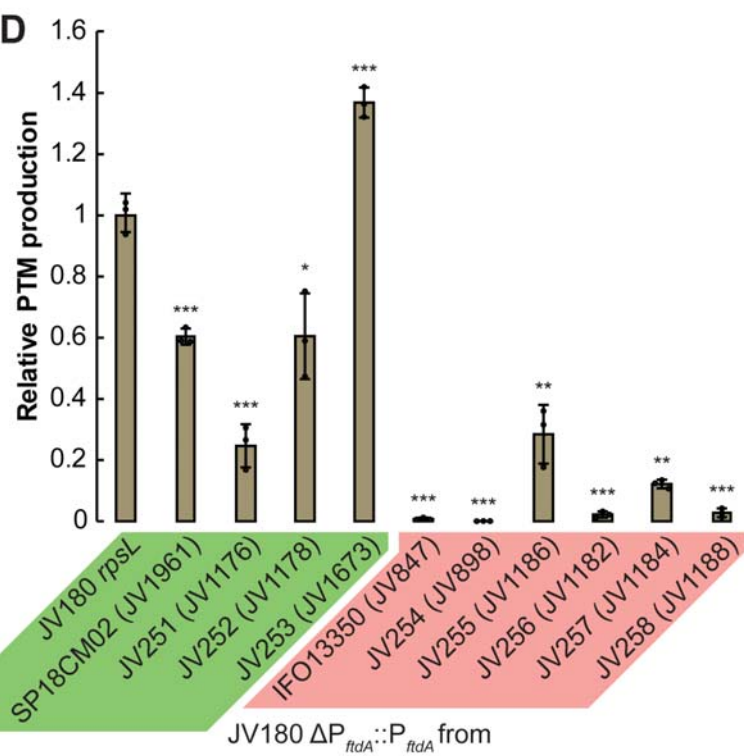
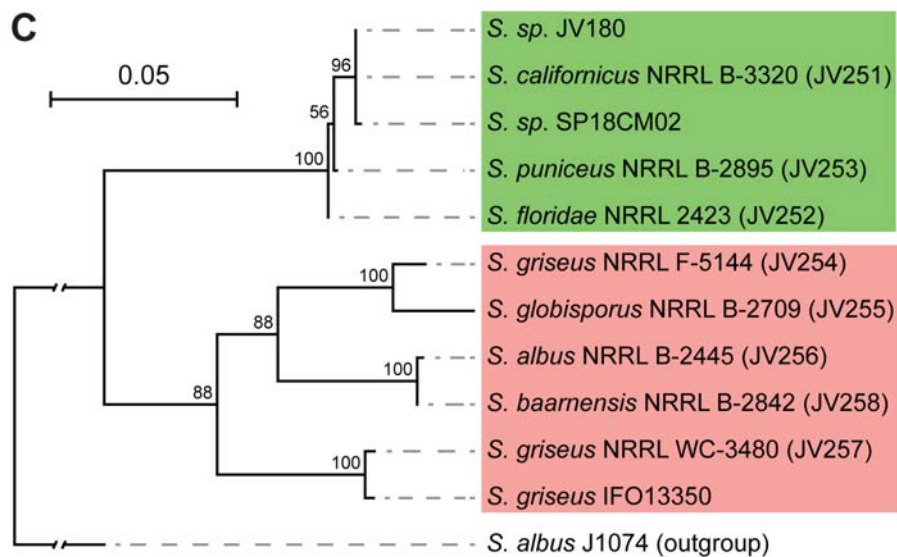
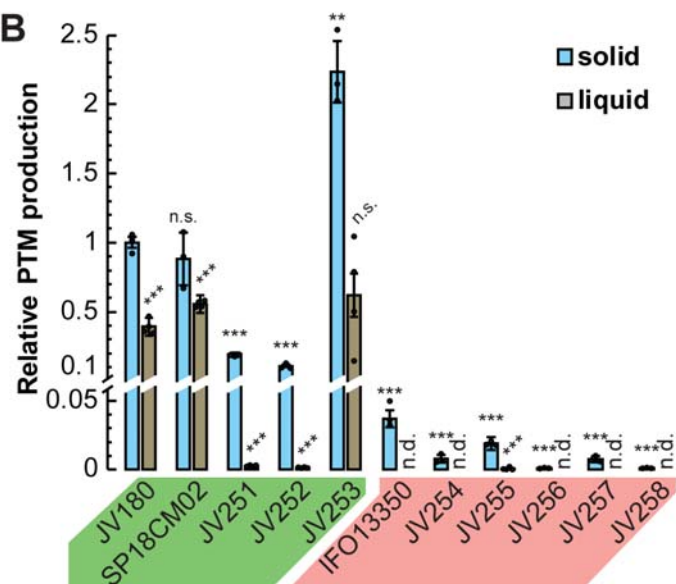
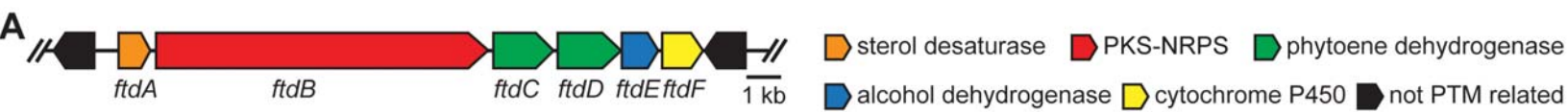
production by *adpA* or replacement of P_{ftdA} with the constitutive P_{ermE^*} (n = 3). **(B)** Relative *ftdB* transcript abundance from JV180, its $\Delta adpA$ mutant, and complementation of PTM transcription by *adpA* or replacement of P_{ftdA} with the constitutive P_{ermE^*} (n = 6). n.d. = not detected. **(C)** Alignment of the P_{ftdA} AdpA binding sites, with non-conserved nucleotides highlighted in red (note: the sequences are identical in the JV180-like P_{ftdA} 's). **(D)** Relative PTM production from JV180 P_{ftdA} AdpA binding site mutants (n = 3). **(E)** Relative *ftdB* transcript abundance from JV180 P_{ftdA} AdpA binding site mutants (n = 6). EMSA assays for His-tagged JV180 AdpA interaction with the **(F)** JV180 P_{ftdA} binding site, **(G)** IFO13350 P_{ftdA} binding site, or **(H)** a JV180 P_{ftdA} binding site containing a transversion mutation in one of the nucleotides (red, underlined) directly interacting with AdpA, which was necessary for protein-DNA interaction. The lane marked (+) is a positive control using a wild-type JV180 probe sequence. The statistical significance in the differences in PTM production or transcription were calculated by two-tailed Student's T-test, relative to the JV180 *rpsL* mutant, JV307. *p < 0.05, **p < 0.01, ***p < 0.001. Error bars represent standard deviation.

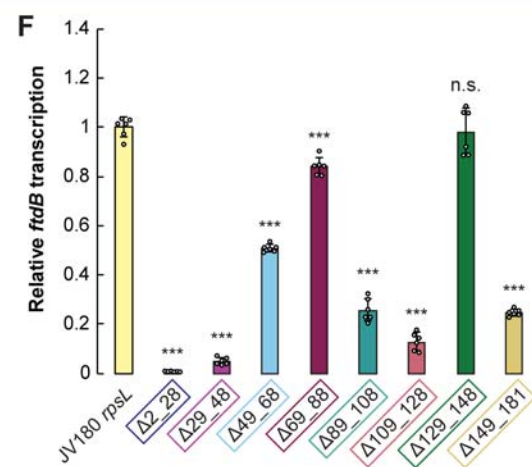
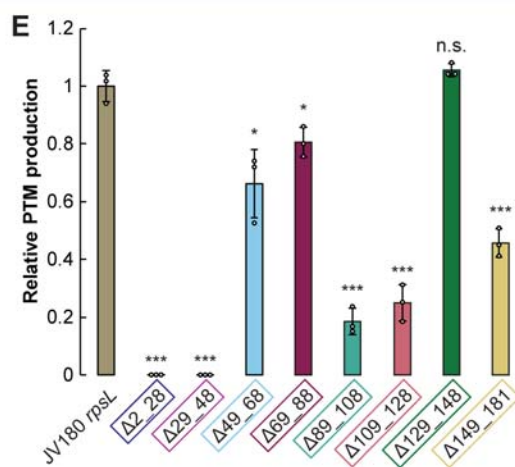
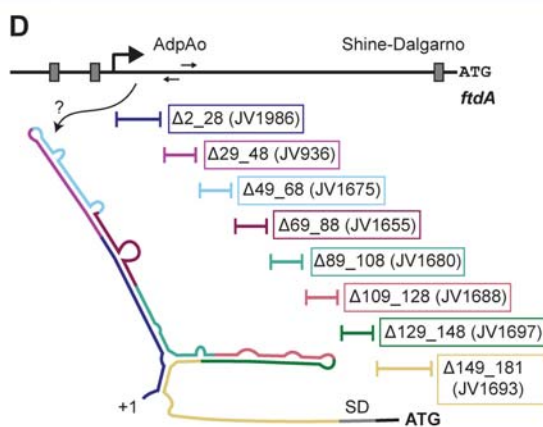
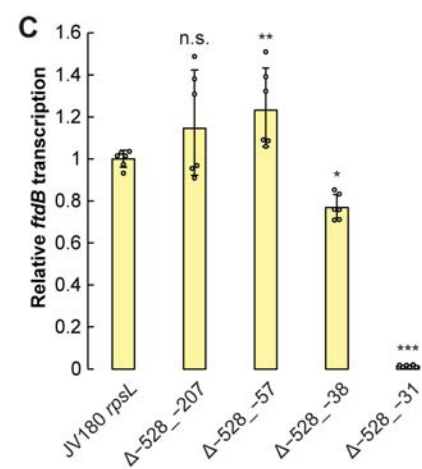
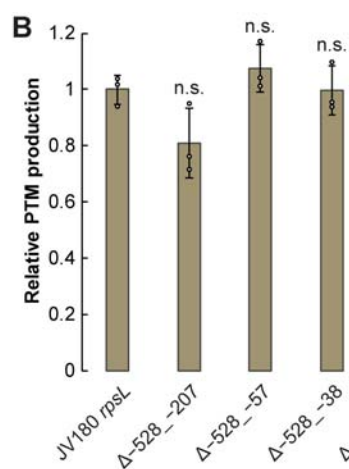
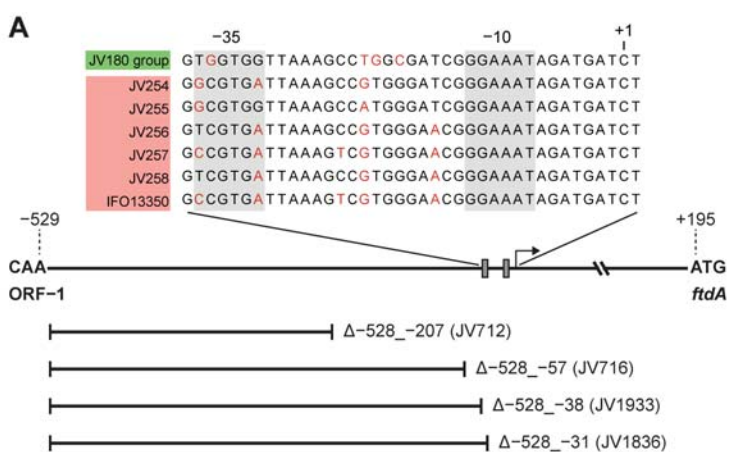
Figure 4. A conserved AG(G) indel tunes strong and weak P_{ftdA} 's. **(A)** Alignment of the P_{ftdA} UTR section containing the conserved AG(G) indel between the TSS and the AdpA binding site. Non-conserved nucleotides are indicated in red. **(B)** Relative PTM production from JV180 $P_{ftdA}\Delta AG$ mutants and JV180 $\Delta P_{ftdA}::P_{ftdA_IFO13350}\pm AG$ mutants (n = 3). **(C)** Relative *ftdB* transcript abundance from JV180 $P_{ftdA}\Delta AG$ mutants and JV180 $\Delta P_{ftdA}::P_{ftdA_IFO13350}\pm AG$ mutants (n = 6, except for JV1032 where n = 3). The statistical significance in the differences in PTM production or transcription were calculated by two-tailed Student's T-test, relative to the JV180 *rpsL* mutant, JV307, or otherwise indicated. *p < 0.05, **p < 0.01, ***p < 0.001. Error bars represent standard deviation.

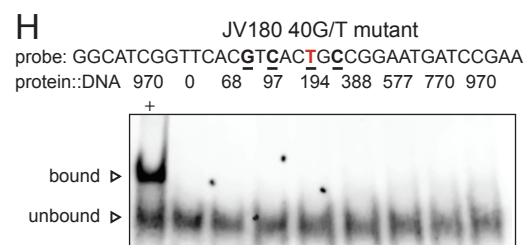
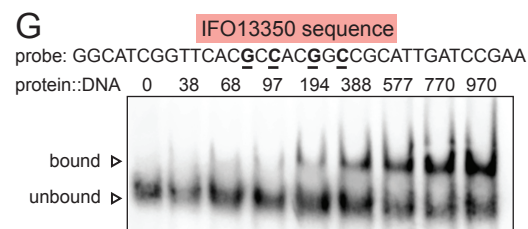
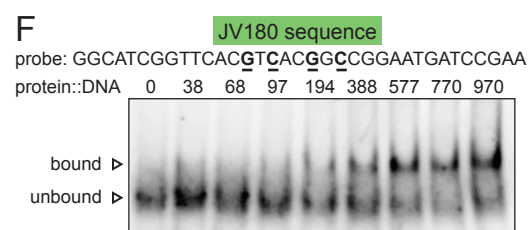
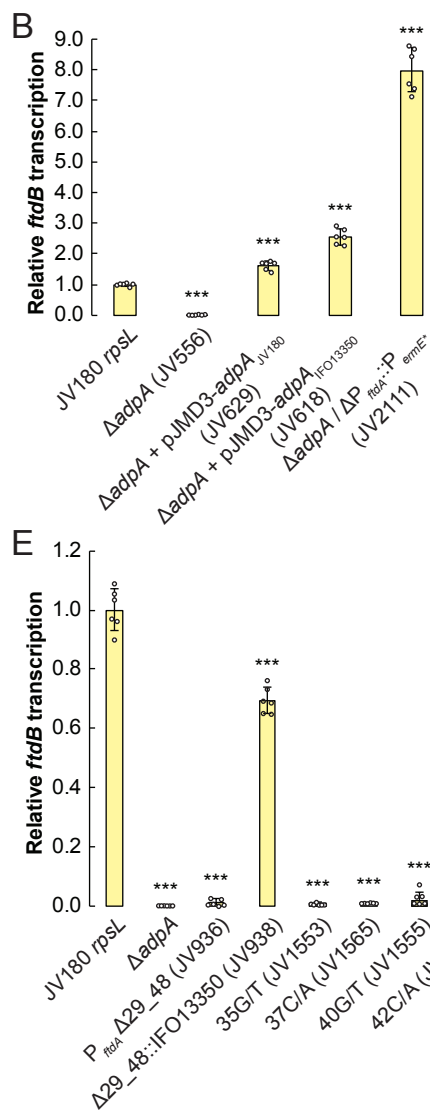
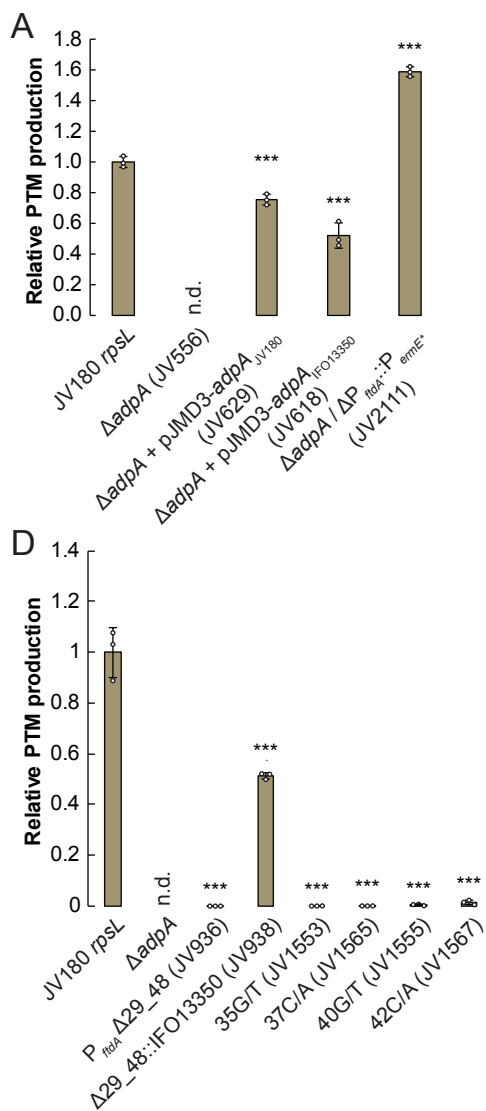
Figure 5. Deletions in the JV180 griseorhodin (*grh*) BGC often negatively impact PTM production and transcription by P_{ftdA} . **(A)** Diagram of the JV180 griseorhodin BGC with gene ranges deleted in various mutants mapped below as corresponding colored bars. **(B)** Relative PTM production from various *grh* mutants in JV180 *rpsL* (white background) and JV180 $\Delta P_{ftdA}::P_{ermE^*}$ background strains (grey background, n = 3). n.d. = not detected. **(C)** Relative *ftdB*

873 transcript abundance from various *grh* mutants in JV180 *rpsL* and JV180 $\Delta P_{ftdA}::P_{ermE}$
874 background strains as noted in panel B, (n = 6). The statistical significance in the differences in
875 PTM production or transcription were calculated by two-tailed Student's T-test, relative to the
876 JV180 *rpsL* mutant, JV307, or otherwise indicated. *p < 0.05, **p < 0.01, ***p < 0.001. Error
877 bars represent standard deviation.

878







A	+1	AdpA binding site	
JV180	TCTGGGGC	CGATGGGGGTCAG	GGGGGGCATCGGTTTCACGTCAC
SP18CM02	TCTGGGGC	CGATGGGGGTCAG	GGGGGGGCATCGGTTTCACGTCAC
JV251	TCTGGGGC	CGATGGGGGTCAG	GGGGGGCATCGGTTTCACGTCAC
JV252	TCTGGGGC	CGATGGGGGTCAG	GGGGGGCATCGGTTTCACGTCAC
JV253	TCTGGGGC	CGATGGGGGTCAG	GGGGGGCATCGGTTTCACGTCAC
JV254	TCTGGGGCTGATGGGGGTC	- - -	GGGGGGCATCGGTTTCACGCCAC
JV255	TCTGGGGCTGATGGGGGTC	- - -	GGGGGGCATCAGTTTCACGCCAC
JV256	TCTGGGGCTGATGGGGGTC	- - -	GGGGGGCATCAGTTTCACGCCAC
JV257	TCTGGGGCTGATGAGGTC	- - -	GGGGGGCATCGGTTTCACGCCAC
JV258	TCTGGGGCTGATGGGGGTC	- - -	GGGGGGCATCAGTTTCACGCCAC
IFO13350	TCTGGGGCTGATGAGGTC	- - -	GGGGGGCATCGGTTTCACGCCAC

