## **UC Santa Cruz**

## **UC Santa Cruz Previously Published Works**

## **Title**

Conserved Responses in a War of Small Molecules between a Plant-Pathogenic Bacterium and Fungi

## **Permalink**

https://escholarship.org/uc/item/7nv2k7cn

## **Journal**

mBio, 9(3)

## ISSN

2161-2129

## **Authors**

Spraker, Joseph E Wiemann, Philipp Baccile, Joshua A et al.

## **Publication Date**

2018-07-05

## DOI

10.1128/mbio.00820-18

## **Copyright Information**

This work is made available under the terms of a Creative Commons Attribution License, available at <a href="https://creativecommons.org/licenses/by/4.0/">https://creativecommons.org/licenses/by/4.0/</a>

Peer reviewed





# Conserved Responses in a War of Small Molecules between a Plant-Pathogenic Bacterium and Fungi

⑤ Joseph E. Spraker,a\* ⑥ Philipp Wiemann,b\* Joshua A. Baccile,c\* Nandhitha Venkatesh,a Julia Schumacher,d Frank C. Schroeder,e ⑥ Laura M. Sanchez,f Nancy P. Kellerb,f

<sup>a</sup>Department of Plant Pathology, University of Wisconsin—Madison, Madison, Wisconsin, USA

ABSTRACT Small-molecule signaling is one major mode of communication within the polymicrobial consortium of soil and rhizosphere. While microbial secondary metabolite (SM) production and responses of individual species have been studied extensively, little is known about potentially conserved roles of SM signals in multilayered symbiotic or antagonistic relationships. Here, we characterize the SM-mediated interaction between the plant-pathogenic bacterium Ralstonia solanacearum and the two plant-pathogenic fungi Fusarium fujikuroi and Botrytis cinerea. We show that cellular differentiation and SM biosynthesis in F. fujikuroi are induced by the bacterially produced lipopeptide ralsolamycin (synonym ralstonin A). In particular, fungal bikaverin production is induced and preferentially accumulates in fungal survival spores (chlamydospores) only when exposed to supernatants of ralsolamycinproducing strains of R. solanacearum. Although inactivation of bikaverin biosynthesis moderately increases chlamydospore invasion by R. solanacearum, we show that other metabolites such as beauvericin are also induced by ralsolamycin and contribute to suppression of R. solanacearum growth in vitro. Based on our findings that bikaverin antagonizes R. solanacearum and that ralsolamycin induces bikaverin biosynthesis in F. fujikuroi, we asked whether other bikaverin-producing fungi show similar responses to ralsolamycin. Examining a strain of B. cinerea that horizontally acquired the bikaverin gene cluster from Fusarium, we found that ralsolamycin induced bikaverin biosynthesis in this fungus. Our results suggest that conservation of microbial SM responses across distantly related fungi may arise from horizontal transfer of protective gene clusters that are activated by conserved regulatory cues, e.g., a bacterial lipopeptide, providing consistent fitness advantages in dynamic polymicrobial networks.

**IMPORTANCE** Bacteria and fungi are ubiquitous neighbors in many environments, including the rhizosphere. Many of these organisms are notorious as economically devastating plant pathogens, but little is known about how they communicate chemically with each other. Here, we uncover a conserved antagonistic communication between the widespread bacterial wilt pathogen *Ralstonia solanacearum* and plant-pathogenic fungi from disparate genera, *Fusarium* and *Botrytis*. Exposure of *Fusarium fujikuroi* to the bacterial lipopeptide ralsolamycin resulted in production of the antibacterial metabolite bikaverin specifically in fungal tissues invaded by *Ralstonia*. Remarkably, ralsolamycin induction of bikaverin was conserved in a *Botrytis ci*-

Received 25 April 2018 Accepted 27 April 2018 Published 22 May 2018

Citation Spraker JE, Wiemann P, Baccile JA, Venkatesh N, Schumacher J, Schroeder FC, Sanchez LM, Keller NP. 2018. Conserved responses in a war of small molecules between a plant-pathogenic bacterium and fungi. mBio 9:e00820-18. https://doi.org/10.1128/mBio 00820-18.

**Editor** Arturo Casadevall, Johns Hopkins Bloomberg School of Public Health

**Copyright** © 2018 Spraker et al. This is an open-access article distributed under the terms of the Creative Commons Attribution 4.0 International license.

Address correspondence to Nancy P. Keller, npkeller@wisc.edu.

\* Present address: Joseph E. Spraker, School of Plant Sciences, University of Arizona, Tucson, Arizona, USA; Philipp Wiemann, Hexagon Bio, Menlo Park, California, USA; Joshua A. Baccile, Division of Chemistry and Chemical Engineering, California Institute of Technology, Pasadena, California, USA.

J.E.S. and P.W. contributed equally to the preparation of the manuscript.

<sup>&</sup>lt;sup>b</sup>Department of Medical Microbiology and Immunology, University of Wisconsin—Madison, Madison, Wisconsin, USA

<sup>&</sup>lt;sup>c</sup>Boyce Thompson Institute and Department of Chemistry and Chemical Biology, Cornell University, Ithaca, New York. USA

dInstitute for Biology and Biotechnology of Plants, Westfälische Wilhelms-Universität Münster, Münster, Germany

<sup>&</sup>lt;sup>e</sup>Department of Medicinal Chemistry and Pharmacognosy, College of Pharmacy, University of Illinois at Chicago, Chicago, Illinois, USA

fDepartment of Bacteriology, University of Wisconsin—Madison, Madison, Wisconsin, USA

*nerea* isolate carrying a horizontally transferred bikaverin gene cluster. These results indicate that horizontally transferred gene clusters may carry regulatory prompts that contribute to conserved fitness functions in polymicrobial environments.

**KEYWORDS** *Fusarium*, antimicrobial activity, bacterial wilt, bikaverin, chemical communication, microbial interactions, ralsolamycin

Although many soil bacteria and fungi are well studied individually because of their importance in agricultural settings, little is known about how they interact with one another. Many soil microbes are equipped with an arsenal of unique biosynthetic enzymes, producing bioactive molecules that are thought to help them secure a niche in their local environment. These compounds are often termed secondary metabolites (SMs) because they are seemingly dispensable in axenic culture, although the genes involved in their production can account for large proportions of some microbial genomes and are evolutionarily maintained, suggesting that they are indispensable in a natural environment. Hence, it is thought that microbial SMs play an important role in intra- and interspecific interactions, including cross-kingdom communication (1–3).

The plant-pathogenic bacterium *Ralstonia solanacearum* is a common inhabitant of soils globally. It is a devastating pathogen of well over 200 plant species, including both mono- and dicotyledonous plants (4). Although *R. solanacearum* has been studied primarily because of its severe impacts on plant hosts, it has recently been shown to impact plant-prokaryote community composition (5), and it also interacts intimately with many soil fungi through volatile and diffusible signaling (6–8). The lipopeptide ralsolamycin (synonym ralstonin A), produced by a hybrid polyketide synthase/nonribosomal peptide synthetase (PKS-NRPS) biosynthetic gene cluster (*rmy*) (7, 9, 10), was shown to induce chlamydospore formation in a phylogenetically diverse panel of plant-associated and/or soil-inhabiting fungi representing members of the Mucoromycota, Ascomycota, and Basidiomycota (7). Among the assayed fungi were members of the genus *Fusarium*.

Fusarium spp. are well known for their SM biosynthetic capacity, as they produce a number of mycotoxins, such as beauvericin, fumonisins, trichothecenes, zearalenone, and fusaric acid, which impact animal and human health (11). Aside from these well-described toxins, Fusarium spp. also produce plant growth-regulatory hormones such as gibberellic acid (12) as well as characteristic colored metabolites, including bikaverin (Fusarium fujikuroi and oxysporum species complex) (13), aurofusarin (Fusarium graminearum species complex) (14), and fusarubins/bostrycoidins (conserved in all characterized Fusarium spp. to date) (15–18); some of these SMs have potential applications in medicine (19) and biocontrol (20, 21). While fusarubins/bostrycoidins have been characterized as, perithecial pigments (15–18), the biological and ecological functions of most of these metabolites remain elusive. Surprisingly, a horizontally acquired and functional bikaverin gene cluster is present in some strains of the phylogenetically distinct, broad-range necrotrophic plant pathogen Botrytis cinerea (22).

Here, we describe interactions between *R. solanacearum* and the fungi *F. fujikuroi* and *B. cinerea*, which we show to be mediated by the bacterial lipopeptide ralsolamycin. Ralsolamycin induces visible bikaverin production in several *Fusarium* species, specifically at the bacterial-fungal interaction zone where it accumulates in the chlamydospores. Metabolic profiling of *F. fujikuroi*-ralsolamycin treatments reveals a unique fungal metabolic shift comprising not only bikaverin but also other compounds, including the bioactive metabolite beauvericin. Both bikaverin and beauvericin suppress *R. solanacearum* growth, suggesting a protective fungal function of these bacterium-induced SMs against bacterial competition. Remarkably, ralsolamycin retains its function in inducing bikaverin production in *B. cinerea* despite disparate regulation by nutritional cues of this metabolite in the two fungal species. These findings support a model where regulatory cues responsive to bacterial SM signals can be horizontally

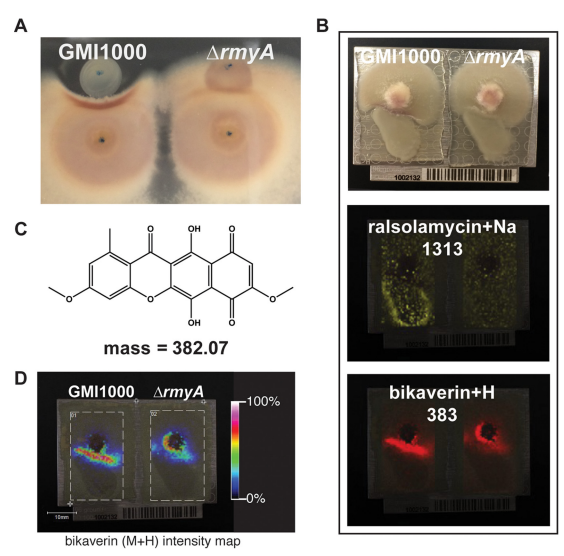


FIG 1 Ralsolamycin induces bikaverin production in F. fujikuroi. (A) When grown in coculture on an agar plate, R. solanacearum GMI1000, but not the  $\Delta rmyA$  mutant, inhibits growth of F. fujikuroi and induces the production of a red pigment in this fungus. (B) Imaging mass spectrometry experiments. (Top) Picture of cocultures mounted on MALDI plate. (Middle) Ralsolamycin (m/z = 1,313) is the only detectable difference between GMI1000 and  $\Delta rmyA$  strains. (Bottom) Imaging mass spectrometry shows that bikaverin (m/z = 383) is produced in proximity to bacterial colonies. The complete IMS data set is shown in Fig. S2. (C) Chemical structure of bikaverin and exact mass. (D) Intensity map of bikaverin shows that substantially more compound accumulates proximal to the GMI1000 colony than to the  $\Delta rmyA$  colony.

transferred in tandem with protective fungal SM gene clusters to confer increased ecological fitness.

### **RESULTS**

Bacterial ralsolamycin induces bikaverin production in Fusarium fujikuroi. The ralsolamycin-producing wild-type R. solanacearum strain GMI1000 can induce chlamydospore development in many fungi, including F. fujikuroi, F. graminearum, Fusarium verticillioides, and Fusarium oxysporum f. sp. lycopersici (7). Using an in vitro coculture assay (Fig. 1A), we found that ralsolamycin (see Fig. A1 and Table A1 in Data Set S1 in the supplemental material) also induces pigment production in three Fusarium species: F. fujikuroi, F. graminearum, and F. oxysporum f. sp. lycopersici (Fig. S1A). Conversely, the  $\Delta rmyA$  mutant did not induce chlamydospore development or localized pigment production in the coincubated fungal species (Fig. S1A).

We selected F. fujikuroi strain IMI58289 for more detailed investigations because of its well-annotated genome and SM biosynthetic potential (17). Additionally, the biosynthetic pathways and molecular cues that regulate production of the two red pigments produced by this isolate, bikaverin (13) and fusarubin (15), are fairly well understood. The interacting R. solanacearum/F. fujikuroi colonies were excised at 72 h when pigmentation was observed at the zone of interaction and analyzed via matrixassisted laser desorption ionization-time of flight imaging mass spectrometry (MALDI-TOF IMS) (23) to map the spatial distribution of metabolites to their specific location in the coculture. In total, 30 ions mapped to distinct locations within microbial cultures (Fig. S1B). As expected, R. solanacearum GMI1000 showed ions representing the sodium adduct of ralsolamycin, which were absent in  $\Delta rmyA$  cultures (7) (Fig. 1B). Nine ions mapped specifically to the area of pigment production at the GMI1000-F. fujikuroi interface (Fig. S1B). Three of these ions, m/z 383, 405, and 421, were different adducts (proton, sodium, and potassium adducts, respectively) of one compound with a molecular mass of 382 Da ( $\pm 0.5$  Da), corresponding to that of the polyketide bikaverin (13) (Fig. 1B and C and S1B). These signals were more intense at the region where F. fujikuroi was interacting with GMI1000 than where it interacted with the ΔrmyA mutant, which mainly showed an intensified signal at the center of the fungal colony (Fig. 1D). Two other ions, m/z 369 and 385, were putatively identified as sodium and potassium adducts, respectively, of gibberellic acid 3 (GA3, molecular weight [MW] of 346 [Fig. S1B]). These ions were diffused across the colony and were detected in the interaction of F. fujikuroi with both R. solanacearum GMI1000 and the  $\Delta rmyA$  mutant. Four other ions detected in the interaction zone could not be assigned to any known F. fujikuroi metabolites. We did not observe any specific signals for fusarubin production (MW of 306 [Fig. S1B]), suggesting that the red pigmentation at the coculture interface was primarily due to bikaverin production.

Bikaverin accumulates in chlamydospores and contributes to protection from invasion. Focusing on bikaverin due to its clear response to ralsolamycin, we aimed to identify the cellular location of bikaverin accumulation in Fusarium, particularly regarding chlamydospore formation. Chlamydospores are long-lived survival spores that we previously found to be induced in many fungi by ralsolamycin (7). Microscopic examination of GMI1000-F. fujikuroi cocultures revealed red-pigmented chlamydospores (Fig. 2A and S2), indicating that bikaverin accumulated specifically in these fungal survival structures. This contrasted with a near-loss of chlamydospore production and no pigmentation in the  $\Delta rmyA$  mutant-F. fujikuroi interaction (Fig. 2A).

Previous microscopic analyses of Aspergillus flavus chlamydospores showed that the spores can be invaded by R. solanacearum (7). Since bikaverin accumulates at high concentrations in the chlamydospores of F. fujikuroi, we hypothesized that this may serve a protective function for chlamydospores against bacterial invasion. Therefore, we set out to compare bacterial colonization of chlamydospores using a green fluorescent protein (GFP)-labeled strain of GMI1000 in coculture with the F. fujikuroi wild type and a bik1 mutant deficient in bikaverin production (13), using confocal laser scanning microscopy (CLSM). We found GFP-labeled bacteria in 1.95% of wild-type and 6.65% of Δbik1 chlamydospores. A chi-square analysis of colonized (containing one or more bacteria) versus uncolonized (containing no bacteria) chlamydospores of the wild-type and  $\Delta bik1$  strains showed colonization to be dependent on bikaverin production ( $\chi^2$  = 4.682, K = 1, P = 0.031 [Table S1]). Although this analysis suggests that bikaverin contributes to protecting chlamydospores from invasion, the mean total fluorescence areas inside all of the observed chlamydospores were not significantly different between the wild type and the  $\Delta bik1$  strain (p=0.315) (Fig. 2B). The wild-type F. fujikuroi strain had a few heavily colonized chlamydospores similar to those of the  $\Delta bik1$  strain but had fewer colonized with low bacterial numbers (Fig. 2B and C). In total, these data indicate that while bikaverin production can reduce bacterial invasion, it is not solely responsible for limiting bacterial growth in invaded F. fujikuroi chlamydospores and that other fungal metabolites likely contribute to protection against bacterial invasion and/or proliferation.

Ralsolamycin overrides suppression of multiple SMs under high-nitrogen conditions. Although the ecological functions of most fungal SMs have not been



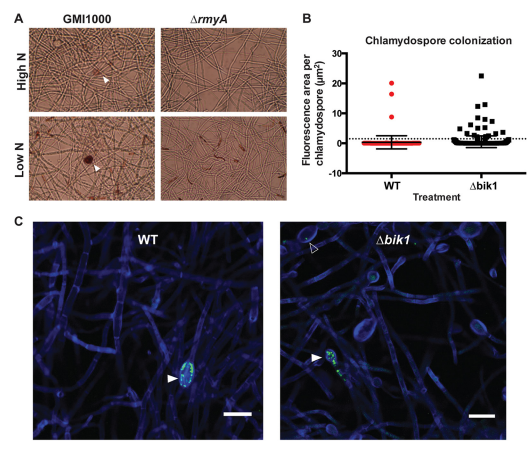


FIG 2 Bikaverin contributes to protecting chlamydospores from bacterial invasion. (A) Microscopic examination of chlamydospores shows bikaverin accumulation in chlamydospores induced by GMI1000-conditioned medium under both high- and low-nitrogen conditions. (B) Analysis of confocal microscopy images for total bacterial GFP fluorescence in all fungal chlamydospores shows no significant change in total bacterial burden (P = 0.315) between  $\Delta bik1$  (n = 301) and wild-type (WT) (n=154) F. fujikuroi. The dotted line indicates the 1.5- $\mu$ m<sup>2</sup>-area cutoff used for chi-square test of colonization differences, showing that more  $\Delta bik1$  chlamydospores are invaded, although at low bacterial counts. (C) Confocal microscopy of GFP-labeled bacteria in chlamydospores of wild-type (WT) and Δbik1 isolates of F. fujikuroi. Solid white arrowheads indicate heavily colonized chlamydospores; an empty white arrowhead indicates invasion by single bacterial cells. Fungal cell walls (blue) are stained with calcofluor white, and GMI1000 cells are constitutively expressing GFP (green). Bars, 20  $\mu$ m. All Z-sections are set to 0.5  $\mu$ m.

established, the production of many SMs, including bikaverin, in F. fujikuroi is tightly regulated by multiple abiotic signals, including pH and nitrogen availability (13, 17). To determine if ralsolamycin can override the tight nitrogen-dependent regulation of the bikaverin biosynthetic gene cluster where bikaverin is repressed by high nitrogen, we grew F. fujikuroi under either high- or low-nitrogen conditions, adding R. solanacearum-conditioned medium from either GMI1000 or the  $\Delta rmyA$  mutant. Additionally, we supplemented F. fujikuroi cultures with purified ralsolamycin to test bikaverin induction. High-performance liquid chromatography (HPLC) and expression analyses showed that, regardless of nitrogen regime, GMI1000 extracts (Fig. 3A to C) and purified ralsolamycin (Fig. 3D and E) significantly increased bikaverin production through transcriptional activation of the bik cluster genes. In fact, bikaverin levels in the high-nitrogen/GMI1000 treatment approached that of the wild type in low nitrogen. Taken together, this suggests that ralsolamycin is a primary cue in inducing bik gene expression and bikaverin biosynthesis, regardless of nutritional environment.

To further explore the fungal chemical repertoire induced by ralsolamycin, we analyzed fungal cultures using ultrahigh-performance liquid chromatography coupled to high-resolution mass spectrometry (UHPLC-HRMS) using the program XCMS (49). Principal-component analyses of all samples showed that F. fujikuroi metabolism is Spraker et al.

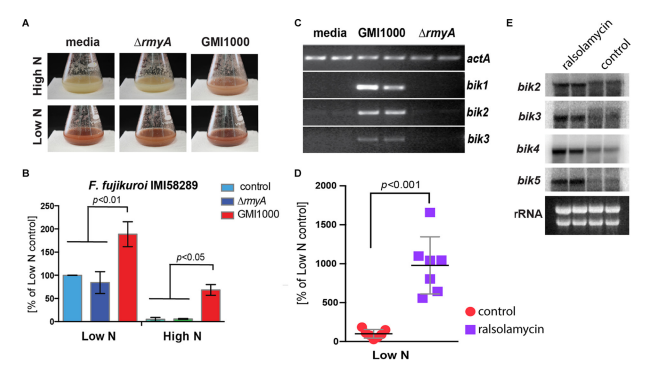


FIG 3 GMI1000-conditioned medium and purified ralsolamycin induce bikaverin production in *F. fujikuroi*. (A) GMI1000-conditioned medium, but not Δ*rmyA* strain-conditioned medium, caused dramatic shifts in pigment production in both *F. fujikuroi* strains relative to medium controls. (B) HPLC-PDA analysis of liquid cultures of *F. fujikuroi* treated with GMI1000-conditioned medium shows significantly increased bikaverin production under both high- and low-nitrogen conditions. (C) Semiquantitative reverse transcription-PCR shows that bikaverin biosynthetic genes *bik1*, *bik2*, and *bik3* in *F. fujikuroi* are induced in response to GMI1000-conditioned medium but not in response to Δ*rmyA* strain-conditioned medium or bacterial medium controls. Actin was used as a loading control. (D) Purified ralsolamycin induces bikaverin production in *F. fujikuroi* which was quantified by HPLC from seven biological replicates. Acetonitrile was used as the carrier control. (E) Northern analysis showing induction of bikaverin biosynthetic genes (technical duplicates) in response to purified ralsolamycin.

shifted most in response to GMI1000-conditioned, high-nitrogen medium (Fig. S3A). Using the metaXCMS package (24), we found that, in accordance with previous studies (17), a low-nitrogen condition induces far more spectral features (573) (Fig. S3B) than it represses (207) (Fig. S3C) compared to high-nitrogen conditions. However, treatment with GMI1000-conditioned high-nitrogen medium induced a total of 311 features compared to unconditioned high-nitrogen medium alone (Fig. S3B), restoring, as shown in Fig. 3, bikaverin. When  $\Delta rmyA$  mutant-conditioned high-nitrogen medium was assessed, only 114 features were found to be induced compared to unconditioned high-nitrogen medium (Fig. S3B), indicating that this treatment caused the least drastic shift in metabolism. Interestingly, we found the highest number of common induced features between unconditioned low-nitrogen medium and GMI1000-conditioned high-nitrogen medium compared to unconditioned high-nitrogen medium (25) (Fig. S3B).

Targeted metabolomics examination of these treatments revealed a significant increase in GMI1000 treatment of not only bikaverin (P = 0.003) but also the cytotoxic metabolite beauvericin (P = 0.008) (Table S2 and Fig. S3). Like bikaverin, the biosynthetic genes of beauvericin are known to be repressed under high-nitrogen conditions, although via a different regulatory mechanism than bikaverin (13, 17, 26).

**Antibacterial activity of bikaverin and beauvericin.** The induction of both bikaverin and beauvericin in response to the bacterial signal ralsolamycin, coupled with the accruement of bikaverin in chlamydospores, suggested a possible evolved function of these metabolites toward protection from the endosymbiotic bacteria. Indeed, both metabolites have been previously shown to be antibacterial (27–30).

To examine this hypothesis, we first used conditioned medium obtained from *F. fujikuroi* wild type and the *bik1* mutant for antibacterial activity assays and found that wild-type *F. fujikuroi*-conditioned medium reduced cell viability by 5-fold relative to the



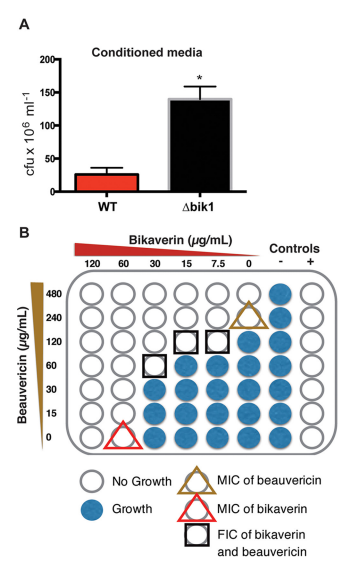
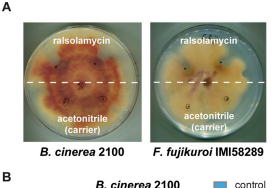


FIG 4 Fusarium SMs contribute to protecting chlamydospores from bacterial invasion. (A) F. fujikuroi (wild-type [WT])-conditioned medium significantly reduced recoverable colonies of R. solanacearum relative to conditioned medium from a  $\Delta bik1$  mutant. (B) Diagram showing the combinations of concentrations of bikaverin and beauvericin used in the checkerboard assay and the observed MICs and FICs of these compounds toward R. solanacearum.

Δbik1 mutant-conditioned medium (Fig. 4A). We then tested the antibacterial activity and interactions between bikaverin and beauvericin against R. solanacearum. This was performed with a checkerboard analysis (31), which showed the MICs of bikaverin and beauvericin to be 60 (157  $\mu$ M) and 240 (306  $\mu$ M)  $\mu$ g/ml, respectively. The bikaverin MIC was higher than that reported for Escherichia coli of 65  $\mu$ M (27), and the beauvericin MIC was higher than the reported 32 μM for Lactobacillus acidophilus and Staphylococcus aureus (28). The interaction between the metabolites was characterized as additive due to its fractional inhibitory concentration (FIC) index value of 0.708 (Fig. 4B; Table S3). These data, in combination with our metabolomics data, support the hypothesis that bikaverin and other SMs, such as beauvericin, act together to protect F. fujikuroi from bacterial invasion and/or proliferation.

Conservation of ralsolamycin-dependent induction of the horizontally acquired bikaverin cluster in Botrytis cinerea. We hypothesized that if ralsolamycin serves as an "alert" signal of invading endosymbiotic bacteria, then it may also induce the protective SM bikaverin in other fungi. Most bikaverin-producing fungi belong to the genus Fusarium (class Sordariomycetes), so we first confirmed that ralsolamycin



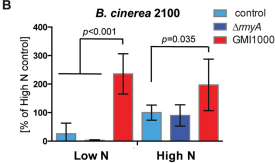


FIG 5 Ralsolamycin induces bikaverin production in both F. fujikuroi and B. cinerea. (A) Purified ralsolamycin induces bikaverin production in both F. fujikuroi and B. cinerea. The white dashed line separates treatments and controls. Solutions were applied at black dots shown on plate. (B) HPLC-PDA analysis of liquid cultures of B. cinerea treated with GMI1000-conditioned medium shows increased bikaverin production under both high- and low-nitrogen conditions. Pictures of culture flasks can be seen in Fig. S4C.

induced bikaverin production in the congeneric plant pathogen F. oxysporum f. sp. lycopersici (Fig. S4A). Interestingly, some isolates of the taxonomically distant fungus B. cinerea (class Leotiomycetes) have also acquired and maintained the bikaverin gene cluster through an ancient horizontal transfer event from Fusarium (22, 32). We therefore set out to determine if bikaverin synthesis was affected by ralsolamycin in the B. cinerea strain 2100, which contains a functional bikaverin gene cluster (22). First, we tested whether purified ralsolamycin has similar effects on bikaverin production in the two fungal species. When grown on solid medium, purified ralsolamycin caused significant growth inhibition of F. fujikuroi and B. cinerea with accumulating red pigment in the zone of interaction compared to the control (Fig. 5A). Although B. cinerea is a species that produces few to no chlamydospores, microscopic examination of the B. cinerea culture challenged with GMI1000 showed ralsolamycin-dependent accumulation of bikaverin in intra- and extrahyphal agglomerates as well as a few chlamydospores (Fig. S4B).

We next examined the effects of different nutritional conditions on bikaverin production in B. cinerea as we did for F. fujikuroi, using two nitrogen supplementation regimes, both treated with conditioned medium either from GMI1000 or ΔrmyA cultures. Analogous to the response of F. fujikuroi, GMI1000-conditioned B. cinerea cultures significantly increased bikaverin production, independent of nitrogen availability, whereas  $\Delta rmyA$  mutant-conditioned medium had no effect (Fig. 5B and S4C). In contrast, changes in the availability of nitrogen had the opposite effect on bikaverin production in B. cinerea compared to F. fujikuroi (Fig. 3B and 5B). Thus, the response to ralsolamycin and its ability to override endogenous nitrogen SM regulatory systems are conserved in two distantly related fungal genera sharing the horizontally transferred bikaverin gene cluster. These data suggest that intermicrobial signals may be an ancestral factor in the regulation of SM clusters that, furthermore, can be transferred along with the SM cluster itself in horizontal transfer events.



### **DISCUSSION**

Several recent studies of bacterial-fungal interactions have demonstrated shifts in SM production in response to coculture of bacteria and fungi (2, 3, 33–35). These shifts in metabolism indicate that SMs have profound impacts on the ability of these microbes to colonize and survive in complex biotic environments and that microbes respond to specific SM signals as part of a complex network of interactions. Although previous studies have described cross-kingdom induction of SMs, e.g., in Aspergillus nidulans/Streptomyces hygroscopicus (34) or Fusarium tricinctum/Bacillus subtilis (36) pairings, knowledge of mechanisms and molecular players underlying these interactions remains scarce. We recently described a unique Ralstonia-Aspergillus flavus interaction where ralsolamycin suppressed synthesis of an A. flavus germination and antibacterial peptide, imizoquin, suggestive of a bacterial advantage in this encounter (8). In contrast, our present study implicates ralsolamycin in the specific induction of antibacterial fungal SMs, including the known metabolites bikaverin and beauvericin, in F. fujikuroi and Botrytis cinerea. Bikaverin is specifically localized to chlamydospores in F. fujikuroi, where it may contribute to protection from bacterial invasion, and is localized to hyphal tissues in B. cinerea, a species that forms few chlamydospores. Significantly, the horizontally transferred bikaverin gene cluster in B. cinerea positively responds to ralsolamycin similarly to the bikaverin cluster in F. fujikuroi, despite having an inverse response to nitrogen sufficiency.

Since the initial realization that fungi, like bacteria, usually organize the genetic machinery for bioactive SMs in biosynthetic gene clusters (37), considerable efforts have been spent in trying to understand the origins and functions of these units. A relatively recent focus has been on horizontal transfer events where several clusters have been transferred in their (near) entirety across phylogenetically diverse taxa (22, 32, 38), including the bikaverin cluster from Fusarium (class Sordariomycetes) to the evolutionarily distant Botrytis (class Leotiomycetes) (32). The high level of conservation in terms of gene order and orientation in both lineages, as well as the lack of a bik cluster homologue in Sclerotinia sclerotiorum, a close relative of B. cinerea, allowed for a prediction of when the transfer occurred, namely, after evolutionary divergence of Sclerotinia and Botrytis (32). However, why the bikaverin cluster (or any SM cluster) would be transferred and—if transferred—whether environmental regulatory signals are maintained remained unknown. Our study provides original insights in the evolutionary rationale underlying these events by demonstrating (i) a protective role of the ralsolamycin-induced bikaverin in defending Fusarium from bacterial ingress and (ii) conservation of bikaverin induction by ralsolamycin in a distantly related fungus that anciently acquired the bik cluster via horizontal gene transfer. We propose that conserved regulation of protective, horizontally transferred SM clusters can provide fitness advantages to fungi. Although our work exemplifies a bacterial/fungal sparring match using chemicals as foils (Fig. 6), we posit that other horizontally transferred SM regulatory responses may be similarly maintained when advantageous to the recipient organism, such as melanin cluster induction by UV stress.

The conservation of the ralsolamycin response in both species extended to overriding disparate bikaverin responses to nitrogen availability in the two fungal genera. Nitrogen availability has been shown to impact the regulation of many F. fujikuroi SM clusters (17, 26), including bikaverin. For example, bikaverin (13), beauvericin (26), and gibberellins (39) are produced only under low-nitrogen conditions but repressed under high-nitrogen conditions (17, 40). The experiments conducted here demonstrate that nitrogen repression of both bikaverin and beauvericin can be overridden by ralsolamycin. Likewise, ralsolamycin overrides repressive bikaverin conditions in B. cinerea (which does not contain the beauvericin gene cluster); however, in this case the repressive conditions are nitrogen limitation, opposite to the conditions that repress bikaverin in Fusarium. Taken together, these results suggest that an entire response pathway (e.g., defensive SM production induced by a bacterial SM signal) can be transmitted along with, or in consequence of, a horizontal gene cluster transfer event.

mBio<sup>®</sup> Spraker et al.

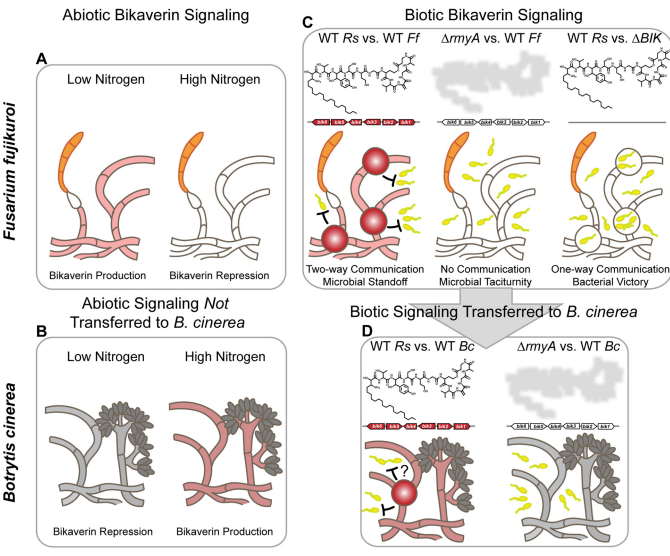


FIG 6 Model of horizontally transferred microbial weaponry. (A) In F. fujikuroi IMI58289, bikaverin biosynthesis is induced under nitrogen-limiting conditions. (B) This abiotic signaling effect on bikaverin production is not conserved in B. cinerea 2100, where the bikaverin production is repressed under nitrogen-limiting conditions. (C) F. fujikuroi IMI58289 induces chlamydospore formation and bikaverin production in response to ralsolamycin produced by R. solanacearum GMI1000 in contrast to the nonproduction mutant (ΔrmyA). R. solanacearum GMI1000 is better able to survive in the presence of a bikaverin deletion mutant (\(\Delta BIK\)) than the wild type. (D) In contrast to the nonconserved response to nitrogen availability, the response to ralsolamycin is conserved in \(B. \) cinerea 2100. Rs, R. solanacearum (yellow cells); Ff, Fusarium fujikuroi (orange spores); Bc, B. cinerea (gray spores).

In addition to bikaverin and beauvericin, we found that ralsolamycin induced a multitude of unknown F. fujikuroi metabolites under repressive conditions. These data suggest that the ralsolamycin defense signaling pathway can activate other SM clusters which may work additively or synergistically with the antibacterial activity of bikaverin and beauvericin.

This study illustrates microbial small-molecule communications, subsequent microbial responses, and conservation of response in a horizontal transfer event, thereby presenting an archetype of a coupled bacterial-fungal chemically driven feedback network. This work extends our understanding of the intricacies of microbial chemical exchange and how it may shape microbial community dynamics. Although our focus was solely on bacterial-fungal interactions in this work, we consider it likely that such chemical warfare also impacts interactions with plant hosts, potentially driving important plant phenotypes such as nutrient acquisition, growth promotion, and disease outcomes. Ongoing work is aimed at understanding these complex, multipartite interactions and how they may be leveraged to improve plant health.



#### **MATERIALS AND METHODS**

**Strains.** For bacterial strains, *R. solanacearum* strain GMI1000, the  $\Delta rmyA$  mutant, and the green fluorescent protein-labeled strain were described previously (7). For fungal strains, *F. fujikuroi* wild-type strain IMI58289 (17) and the  $\Delta bik1$  (previously  $\Delta pks4$ ) mutant were described previously (41). *F. oxysporum* f. sp. *lycopersici* strain 4287 has been described previously (42). *F. graminearum* PH1 was obtained from the Fungal Genetic Stock Center. *Botrytis cinerea* strain 2100 (CECT2100; Spanish Type Culture Collection, Universidad de Valencia) was described previously (43).

Medium and growth conditions and coculture experiments. Ralstonia solanacearum strains were routinely grown at 30°C on Casamino Acids-peptone-glucose (CPG) agar (44) supplemented with 1 g/liter yeast extract and on tetrazolium chloride (TZC) (Chem-Impex). Red, mucoid colonies were selected for use in experiments. Liquid bacterial cultures were grown overnight in CPG at 30°C and 250 rpm. Overnight liquid cultures of *R. solanacearum* were pelleted by centrifugation; washed in equal volumes of sterilized, double-distilled water; quantified using values of optical density at 600 nm (OD<sub>600</sub>); and adjusted to ~2 × 108 cells ml $^{-1}$ . For solid-plate experiments, 5  $\mu$ l (1 × 106 cells) of bacterial suspension was spotted at each point. For *R. solanacearum*-conditioned medium (both GMI100 and Δ*rmyA* strain) experiments, 1 ml of overnight culture was inoculated into a 500-ml flask containing 250 ml CPG broth and was grown for 24 h at 30°C at 250 rpm. Cells were pelleted by centrifugation in a Sorvall RC 6 Plus centrifuge at 10,000 rpm. Supernatants were sterilized by vacuum filtration through a Nalgene Rapid-Flow 0.2- $\mu$ m filter unit (Thermo Fisher, Rochester, NY) before being added to *F. fujikuroi* cultures.

Fusarium spp. were routinely grown at 30°C on potato dextrose agar (PDA), and three ~0.5-cm agar plugs were used as inoculum for liquid cultures. For all cultures, F. fujikuroi was grown for 72 h at 28°C in 300-ml flasks containing 100 ml Darken medium (45) and shaken at 180 rpm. Darken starter cultures (DSC) were used as initial inoculum onto plates as well as into liquid medium for bikaverin analysis. For plate assays and MALDI-TOF IMS experiments, 20  $\mu$ l of DSC was spotted onto the International Streptomyces Project (ISP2) agar 2 cm away from the R. solanacearum spot on CPG plates using wide-end pipette tips. The droplets were dried in a Nuaire biological safety cabinet (NU425-400) to prevent running, and then plates were wrapped with Parafilm once and incubated at room temperature for 72 h prior to imaging and MALDI-TOF IMS experimentation. When assaying purified ralsolamycin, fungi were handled as previously described and either a solution of ralsolamycin in acetonitrile (ACN) or acetonitrile carrier control was applied next to the growing culture of fungi. Droplets were allowed to dry in a laminar flow hood before being incubated, to prevent running. For liquid cultures, 500  $\mu$ l of DSC was used to inoculate 300-ml flasks containing 100 ml of ICI medium (Imperial Chemical Industries Ltd.) (25) containing 6 mM NH<sub>4</sub>NO<sub>3</sub> (low-nitrogen medium) or 60 mM NH<sub>4</sub>NO<sub>3</sub> (high-nitrogen medium). For conditioned-medium experiments, 30 ml of cell-free Ralstonia medium (either GMI1000 or ΔrmyA strain) was added to each flask of 100 ml ICI medium, and 30 ml of sterile CPG medium was used for control experiments. For liquid cultures of F. fujikuroi with purified ralsolamycin, purified ralsolamycin was added at a final concentration of 10  $\mu$ g/ml (half the average final concentration found in the supernatant) in a 12-ml total volume. Details of DNA and RNA extraction protocols are provided in Text S1 in the supplemental material.

Imaging mass spectrometry experiments. Bacteria and fungi were cultured as described above for solid-plate assays. The region of interest was excised from the agar and transferred to the same MALDI MSP 96 anchor plate (Bruker Daltonics, Billerica, MA). A photograph was taken, and the aerial hyphae of F. fujikuroi were removed gently with a cotton swab dampened with acetonitrile (ACN) (46). Following the removal of the aerial hyphae, another photograph was taken and a 50:50 mixture of recrystallized  $\alpha$ -cyano-4-hydroxycinnamic acid (CHCA) and 2,5-dihydroxybenzoic acid (DHB) (Sigma-Aldrich) was applied using a 53- $\mu$ m stainless steel sieve (Hogentogler & Co., Inc., Columbia, MD). The plate was then transferred to an oven and desiccated at 37°C overnight. Following adherence to the MALDI plate, another photographic image was taken and the sample was subjected to MALDI-TOF mass spectrometry (Autoflex; Bruker Daltonics, Billerica, MA) for IMS acquisition. Data were acquired in positive reflectron mode, with a 500- $\mu$ m laser interval in the XY and a mass range of 200 to 2,500 Da. The resulting data were analyzed using Flex imaging software, v. 4.0.

Metabolite extraction and quantification. Liquid cultures were grown as described above, collected at 72 h, and extracted with 200 ml of ethyl acetate acidified with 1 ml of 25% HCl. From each sample, 15 ml of ethyl acetate layer was transferred to a scintillation vial and dried down in vacuo. Liquid cultures with purified ralsolamycin were collected at 72 h and extracted with 8 ml ethyl acetate acidified with 0.005% HCl, 5 ml of which was dried down in vacuo. Dried sample residues were reconstituted in ACN-water-formic acid (99:99:2, vol/vol/vol) and separated on a Zorbax Eclipse XDB-C<sub>18</sub> column (Agilent; 4.6 mm by 150 mm with a 5-μm particle size) using a binary gradient of 1% formic acid as solvent A and ACN-1% formic acid as solvent B delivered by a Flexar binary liquid chromatography (LC) pump (PerkinElmer) coupled to a Flexar LC autosampler (PerkinElmer) and a Flexar Photodiode Array (PDA) Plus detector (PerkinElmer). The high-performance LC (HPLC) program started with an isocratic step at 80% A for 2 min followed by a linear gradient to 0% A over 15 min. After each run, the column was washed for 3 min using 100% B and was equilibrated for 3 min using 80% A. The flow rate was set to 1.5 ml min<sup>-1</sup>. Bikaverin was detected at 510 nm. Identification and relative quantification of metabolites were performed using Chromera Manager (PerkinElmer). Extraction and enrichment of ralsolamycin are described and nuclear magnetic resonance (NMR) experimental details are provided in Text S1 in the supplemental material.

For our UHPLC-MS-guided metabolomics experiments, conditioned medium and extraction protocols were used as described above. High-resolution UHPLC-MS was performed on a Thermo Scientific-Vanquish UHPLC system connected to a Thermo Scientific Q Exactive Orbitrap operated in electrospray

negative ionization mode (ESI). A Zorbax Eclipse XDB-C $_{18}$  (2.1 by 150 mm, 1.8- $\mu$ m particle diameter) column was used for all samples with a flow rate of 0.2 ml/min. The solvent system was water with 0.5% formic acid (solvent A) and acetonitrile with 0.5% formic acid (solvent B) with the following gradient: 20% to 100% B from 0 to 15 min, 100% B from 15 to 20 min, 100% to 20% B from 20 to 21 min, and 20% B from 21 to 25 min. Nitrogen was used as the sheath gas. Data acquisition and processing for the UHPLC-MS were controlled by Thermo Scientific Xcalibur software. Further details of data processing and analyses are provided in Text S1 in the supplemental material.

Microscopy. Cocultures were examined using an LMS 225R dissecting scope (Olympus) and a CX31 compound scope (Olympus). Images were collected on a mounted Canon EOS Rebel T3 and processed in Adobe Photoshop CC 2015 (Adobe Inc.). For assessment of endofungal colonization by R. solanacearum, green fluorescent protein (GFP)-labeled strains of GMI1000 were cocultured with F. fujikuroi in 3 ml of CPG medium for 1 week at 28°C at 180 rpm. Resultant cultures were placed on Miracloth and washed with 15 ml double-distilled H<sub>2</sub>O to remove excess bacterial cells. Hyphae were collected and stained with calcofluor white at a final concentration of 1 mg/ml for 1 min prior to microscopy. Stained samples were wet mounted and imaged on a Zeiss Elyra PS.1 LSM 780 confocal laser scanning microscope (Carl Zeiss, Oberkochen Germany) equipped with a 40× 1.10C-Apochromat lens. To verify endohyphal localization of bacteria, 0.5-µm Z-sections were taken across multiple planes of view. An argon ion laser was used to excite GFP-labeled cells as well as the calcofluor white-labeled fungal cell walls. All confocal images were processed in the open-source Fiji software package, and all overlays of multiple fluorescence channels are of single confocal Z-planes.

For analyses of bacterial invasion, regions of interest (ROIs) were drawn around all calcofluor white-labeled chlamydospores. A custom macro was used to adjust thresholds to 3,500 in the GFP channel to remove background fluorescence in all images and then quantify total bacterial (GFP) fluorescence area in each ROI (https://github.com/spraker/Chlamydospore\_colonization-counts). These data were copied to GraphPad Prism and analyzed for significance using a t test. Chlamydospores containing a contiguous fluorescence area of  $\geq$ 1.5 or <1.5  $\mu$ m<sup>2</sup> were binned as colonized or uncolonized, respectively, for chi-square analyses.

**Antibacterial assays.** For assessing antibacterial activity of conditioned medium, 200  $\mu$ l of cell-free supernatants from low-nitrogen F. fujikuroi (both wild-type and  $\Delta bik1$ ) cultures was inoculated with GMI1000 cells at a final optical density (OD) of 0.01, transferred into a 96-well plate, and grown for 48 h at 28°C. To assess bacterial survival, cells were dilution plated on CPG plus TZC and quantified microscopically. These experiments were done in triplicate, and the data were analyzed using Student's t test to determine significance. To assess antibacterial activity of purified bikaverin and beauvericin (Adipogen), similar protocols were used except that bacteria were grown in Boucher's minimal medium (47). Stock solutions of bikaverin (0.5 mg/ml) and beauvericin (5 mg/ml) were made in dimethyl sulfoxide (DMSO), and appropriate volumes were added to each well to achieve the final concentrations. Equal volumes of DMSO were added to control wells to ensure that activity was attributable to bikaverin and/or beauvericin and not the carrier solvent. For the checkerboard assay using a microdilution approach in a 96-well plate, minimum concentrations of the compounds that showed zero growth based on visible scoring and  $OD_{600}$  values after 48 h were chosen as the MICs. The sum of the fractional inhibitory concentrations (FICs) was calculated as follows for assessing the interaction:  $\Sigma$ FIC = FIC<sub>A</sub> + FIC<sub>B</sub>, where  $FIC_A = (MIC \text{ of compound A in combination with B/MIC of A alone)}, where "MIC of A alone" is the MIC$ of compound A when used as the sole agent. The "MIC of compound A in combination" is the MIC of compound A when used in combination with agent B. The following values were used as cutoffs based on EUCAST recommendations: 0.5 for synergism, 0.5 to 1 for additivity, 1 to 4 for indifference, and 4 for antagonism.

Statistical analysis. All experiments for bikaverin quantification were carried out in biological triplicates and technical duplicates unless otherwise stated in the figure legends. Statistical analysis was performed by using GraphPad Prism software using Student's t test or analysis of variance (ANOVA) followed by Tukey-Kramer analysis to show significant differences. Untargeted metabolomics analyses were performed from four replicates within the XCMS and metaXCMS software packages implemented in the R language (https://github.com/spraker/Spraker\_XCMS\_analysis\_Fusarium). For analyses of known F. fujikuroi metabolites, the MAVEN software package (48) was used to align peaks, and the area top measurement (average intensity of the top three points of a peak) was used for statistical analyses between treatments. ANOVAs followed by Tukey post hoc analyses were performed using GraphPad Prism software. Confocal microscopy images were processed in Fiji, and t tests of bacterial fluorescence in chlamydospores were performed using GraphPad Prism.

## SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/mBio .00820-18.

TEXT S1, DOCX file, 0.02 MB.

FIG S1, TIF file, 7.4 MB.

FIG S2, TIF file, 3.3 MB.

FIG S3, TIF file, 1.6 MB.

FIG S4, TIF file, 4.6 MB.

TABLE S1, PDF file, 0.04 MB.

TABLE S2, PDF file, 0.1 MB.



TABLE S3, PDF file, 0.3 MB.
TABLE S4, PDF file, 0.04 MB.
DATA SET S1, PDF file, 0.2 MB.

#### **ACKNOWLEDGMENTS**

This research was funded by an NSF graduate research fellowship under grant no. DGE-1256259 as well as an NSF postdoctoral research fellowship under award no. 1612169 to J.E.S. Other agencies contributing to this work included NIH CBI training grant T32GM008500 to J.A.B., NIH R01GM112739-01 to N.P.K. and F.C.S., in part USDA Hatch Formula Fund WIS01710 to N.P.K., and K12HD055892 from the National Institute of Child Health and Human Development (NICHD) and the National Institutes of Health Office of Research on Women's Health (ORWH) and UIC startup funds to L.M.S.

We thank Bettina Tudzynski for providing F. fujikuroi IMI58289 and  $\Delta bik1$  strains, Hans-Ulrich Humpf for providing an authentic bikaverin standard, Antonio Di Pietro for providing F. oxysporum f. sp. lycopersici 4287, and Atul Jain and Terry Moore for assistance with recrystallization of DHB and CHCA.

#### **REFERENCES**

- 1. Davies J. 2013. Specialized microbial metabolites: functions and origins. J Antibiot (Tokyo) 66:361–364. https://doi.org/10.1038/ja.2013.61.
- Han J, Wang F, Gao P, Ma Z, Zhao S, Lu Z, Lv F, Bie X. 2017. Mechanism of action of AMP-jsa9, a LI-F-type antimicrobial peptide produced by Paenibacillus polymyxa JSa-9, against Fusarium moniliforme. Fungal Genet Biol 104:45–55. https://doi.org/10.1016/j.fgb.2017.05.002.
- Schulz-Bohm K, Tyc O, de Boer W, Peereboom N, Debets F, Zaagman N, Janssens TKS, Garbeva P. 2017. Fungus-associated bacteriome in charge of their host behavior. Fungal Genet Biol 102:38–48. https://doi.org/10 .1016/j.fqb.2016.07.011.
- Hayward AC. 1991. Biology and epidemiology of bacterial wilt caused by Pseudomonas solanacearum. Annu Rev Phytopathol 29:65–87. https://doi.org/10.1146/annurev.py.29.090191.000433.
- Wei Z, Hu J, Gu Y, Yin S, Xu Y, Jousset A, Shen Q, Friman V-P. 2018. Ralstonia solanacearum pathogen disrupts bacterial rhizosphere microbiome during an invasion. Soil Biol Biochem 118:8–17. https://doi.org/ 10.1016/j.soilbio.2017.11.012.
- Spraker JE, Jewell K, Roze LV, Scherf J, Ndagano D, Beaudry R, Linz JE, Allen C, Keller NP. 2014. A volatile relationship: profiling an inter-kingdom dialogue between two plant pathogens, *Ralstonia solanacearum* and *Aspergillus flavus*. J Chem Ecol 40:502–513. https://doi.org/10.1007/s10886-014-0432-2.
- Spraker JE, Sanchez LM, Lowe TM, Dorrestein PC, Keller NP. 2016. Ralstonia solanacearum lipopeptide induces chlamydospore development in fungi and facilitates bacterial entry into fungal tissues. ISME J 10:2317–2330. https://doi.org/10.1038/ismej.2016.32.
- Khalid S, Baccile JA, Spraker JE, Tannous J, Imran M, Schroeder FC, Keller NP. 2018. NRPS-derived isoquinolines and lipopeptides mediate antagonism between plant pathogenic fungi and bacteria. ACS Chem Biol 13:171–179. https://doi.org/10.1021/acschembio.7b00731.
- Baldeweg F, Kage H, Schieferdecker S, Allen C, Hoffmeister D, Nett M. 2017. Structure of ralsolamycin, the interkingdom morphogen from the crop plant pathogen *Ralstonia solanacearum*. Org Lett 19:4868–4871. https://doi.org/10.1021/acs.orglett.7b02329.
- Murai Y, Mori S, Konno H, Hikichi Y, Kai K. 2017. Ralstonins A and B, lipopeptides with chlamydospore-inducing and phytotoxic activities from the plant pathogen Ralstonia solanacearum. Org Lett 19: 4175–4178. https://doi.org/10.1021/acs.orglett.7b01685.
- D'Mello JPF, Placinta CM, Macdonald AMC. 1999. Fusarium mycotoxins: a review of global implications for animal health, welfare and productivity. Anim Feed Sci Technol 80:183–205. https://doi.org/10.1016/S0377-8401(99)00059-0.
- 12. Mander LN. 1992. The chemistry of gibberellins: an overview. Chem Rev 92:573–612. https://doi.org/10.1021/cr00012a005.
- Wiemann P, Willmann A, Straeten M, Kleigrewe K, Beyer M, Humpf HU, Tudzynski B. 2009. Biosynthesis of the red pigment bikaverin in *Fusarium fujikuroi*: genes, their function and regulation. Mol Microbiol 72:931–946. https://doi.org/10.1111/j.1365-2958.2009.06695.x.
- 14. Frandsen RJN, Nielsen NJ, Maolanon N, Sørensen JC, Olsson S, Nielsen J,

- Giese H. 2006. The biosynthetic pathway for aurofusarin in *Fusarium graminearum* reveals a close link between the naphthoquinones and naphthopyrones. Mol Microbiol 61:1069–1080. https://doi.org/10.1111/j.1365-2958.2006.05295.x.
- Studt L, Wiemann P, Kleigrewe K, Humpf HU, Tudzynski B. 2012. Biosynthesis of fusarubins accounts for pigmentation of *Fusarium fujikuroi* perithecia. Appl Environ Microbiol 78:4468–4480. https://doi.org/10.1128/AEM.00823-12.
- Hansen FT, Gardiner DM, Lysøe E, Fuertes PR, Tudzynski B, Wiemann P, Sondergaard TE, Giese H, Brodersen DE, Sørensen JL. 2015. An update to polyketide synthase and non-ribosomal synthetase genes and nomenclature in Fusarium. Fungal Genet Biol 75:20–29. https://doi.org/10 .1016/j.fgb.2014.12.004.
- 17. Wiemann P, Sieber CMK, von Bargen KW, Studt L, Niehaus EM, Espino JJ, Huß K, Michielse CB, Albermann S, Wagner D, Bergner SV, Connolly LR, Fischer A, Reuter G, Kleigrewe K, Bald T, Wingfield BD, Ophir R, Freeman S, Hippler M, Smith KM, Brown DW, Proctor RH, Münsterkötter M, Freitag M, Humpf HU, Güldener U, Tudzynski B. 2013. Deciphering the cryptic genome: genome-wide analyses of the rice pathogen Fusarium fujikuroi reveal complex regulation of secondary metabolism and novel metabolites. PLoS Pathog 9:e1003475. https://doi.org/10.1371/journal.ppat .1003475.
- Frandsen RJN, Rasmussen SA, Knudsen PB, Uhlig S, Petersen D, Lysøe E, Gotfredsen CH, Giese H, Larsen TO. 2016. Black perithecial pigmentation in *Fusarium* species is due to the accumulation of 5-deoxybostrycoidinbased melanin. Sci Rep 6:26206. https://doi.org/10.1038/srep26206.
- Limón MC, Rodríguez-Ortiz R, Avalos J. 2010. Bikaverin production and applications. Appl Microbiol Biotechnol 87:21–29. https://doi.org/10 .1007/s00253-010-2551-1.
- Son SW, Kim HY, Choi GJ, Lim HK, Jang KS, Lee SO, Lee S, Sung ND, Kim JC. 2008. Bikaverin and fusaric acid from *Fusarium oxysporum* show antioomycete activity against *Phytophthora infestans*. J Appl Microbiol 104:692–698. https://doi.org/10.1111/j.1365-2672.2007.03581.x.
- Kwon HR, Son SW, Han HR, Choi G, Jang K, Choi Y, Lee S, Sung N, Kim J. 2007. Nematicidal activity of bikaverin and fusaric acid isolated *Fusarium oxysporum* against pine wood nematode, *Bursaphelenchus xylophilus*. Plant Pathol J 23:318–321. https://doi.org/10.5423/PPJ.2007.23.4.318.
- Schumacher J, Gautier A, Morgant G, Studt L, Ducrot PH, Le Pêcheur P, Azeddine S, Fillinger S, Leroux P, Tudzynski B, Viaud M. 2013. A functional bikaverin biosynthesis gene cluster in rare strains of *Botrytis cinerea* is positively controlled by VELVET. PLoS One 8:e53729. https://doi.org/10.1371/journal.pone.0053729.
- Yang JY, Phelan VV, Simkovsky R, Watrous JD, Trial RM, Fleming TC, Wenter R, Moore BS, Golden SS, Pogliano K, Dorrestein PC. 2012. Primer on agar-based microbial imaging mass spectrometry. J Bacteriol 194: 6023–6028. https://doi.org/10.1128/JB.00823-12.
- 24. Patti GJ, Tautenhahn R, Siuzdak G. 2012. Meta-analysis of untargeted

- metabolomic data from multiple profiling experiments. Nat Protoc 7:508–516. https://doi.org/10.1038/nprot.2011.454.
- Geissman TA, Verbiscar AJ, Phinney BO, Cragg G. 1996. Studies on the biosynthesis of gibberellins from (-)-kaurenoic acid in cultures of Gibberella fujikuroi. Phytochemistry 5:933–947.
- Niehaus EM, Studt L, von Bargen KW, Kummer W, Humpf HU, Reuter G, Tudzynski B. 2016. Sound of silence: the beauvericin cluster in Fusarium fujikuroi is controlled by cluster-specific and global regulators mediated by H3K27 modification. Environ Microbiol 18:4282–4302. https://doi .org/10.1111/1462-2920.13576.
- Deshmukh R, Mathew A, Purohit HJ. 2014. Characterization of antibacterial activity of bikaverin from *Fusarium* sp. HKF15. J Biosci Bioeng 117:443–448. https://doi.org/10.1016/j.jbiosc.2013.09.017.
- Sondergaard TE, Fredborg M, Christensen AMO, Damsgaard SK, Kramer NF, Giese H, Sørensen JL. 2016. Fast screening of antibacterial compounds from fusaria. Toxins 8(12):E355. https://doi.org/10.3390/toxins 8120355.
- Castlebury LA, Sutherland JB, Tanner LA, Henderson AL, Cerniglia CE. 1999. Use of a bioassay to evaluate the toxicity of beauvericin to bacteria. World J Microbiol Biotechnol 15:119–121. https://doi.org/10 .1023/A:1008895421989.
- Ali-Vehmas T, Rizzo A, Westermarck T, Atroshi F. 1998. Measurement of antibacterial activities of T-2 toxin, deoxynivalenol, ochratoxin A, aflatoxin b1 and fumonisin B1 using microtitration tray-based turbidimetric techniques. Zentralbl Veterinarmed A 45:453–458. https://doi.org/10 .1111/j.1439-0442.1998.tb00848.x.
- European Committee for Antimicrobial Susceptibility Testing (EUCAST) of the European Society of Clinical Microbiology and Infectious Diseases (ESCMID). 2000. EUCAST definitive document E.Def 1.2, May 2000: terminology relating to methods for the determination of susceptibility of bacteria to antimicrobial agents. Clin Microbiol Infect 6:503–508. https://doi.org/10.1046/j.1469-0691.2000.00149.x.
- 32. Campbell MA, Rokas A, Slot JC. 2012. Horizontal transfer and death of a fungal secondary metabolic gene cluster. Genome Biol Evol 4:289–293. https://doi.org/10.1093/gbe/evs011.
- 33. König CC, Scherlach K, Schroeckh V, Horn F, Nietzsche S, Brakhage AA, Hertweck C. 2013. Bacterium induces cryptic meroterpenoid pathway in the pathogenic fungus Aspergillus fumigatus. Chembiochem 14: 938–942. https://doi.org/10.1002/cbic.201300070.
- Schroeckh V, Scherlach K, Nützmann HW, Shelest E, Schmidt-Heck W, Schuemann J, Martin K, Hertweck C, Brakhage AA. 2009. Intimate bacterialfungal interaction triggers biosynthesis of archetypal polyketides in *Aspergillus nidulans*. Proc Natl Acad Sci U S A 106:14558–14563. https://doi.org/ 10.1073/pnas.0901870106.
- 35. Nützmann HW, Reyes-Dominguez Y, Scherlach K, Schroeckh V, Horn F, Gacek A, Schümann J, Hertweck C, Strauss J, Brakhage AA. 2011. Bacteria-induced natural product formation in the fungus Aspergillus nidulans requires Saga/Ada-mediated histone acetylation. Proc Natl Acad Sci U S A 108:14282–14287. https://doi.org/10.1073/pnas.11035 23108.

- Ola ARB, Thomy D, Lai D, Brötz-Oesterhelt H, Proksch P. 2013. Inducing secondary metabolite production by the endophytic fungus *Fusarium* tricinctum through coculture with *Bacillus subtilis*. J Nat Prod 76: 2094–2099. https://doi.org/10.1021/np400589h.
- Keller NP, Hohn TM. 1997. Metabolic pathway gene clusters in filamentous fungi. Fungal Genet Biol 21:17–29. https://doi.org/10.1006/fgbi.1997.0970.
- Slot JC, Rokas A. 2011. Horizontal transfer of a large and highly toxic secondary metabolic gene cluster between fungi. Curr Biol 21:134–139. https://doi.org/10.1016/j.cub.2010.12.020.
- 39. Michielse CB, Pfannmüller A, Macios M, Rengers P, Dzikowska A, Tudzynski B. 2014. The interplay between the GATA transcription factors AreA, the global nitrogen regulator and AreB in *Fusarium fujikuroi*. Mol Microbiol 91:472–493. https://doi.org/10.1111/mmi.12472.
- Rodríguez-Ortiz R, Limón MC, Avalos J. 2009. Regulation of carotenogenesis and secondary metabolism by nitrogen in wild-type *Fusarium fujikuroi* and carotenoid-overproducing mutants. Appl Environ Microbiol 75:405–413. https://doi.org/10.1128/AEM.01089-08.
- Linnemannstöns P, Schulte J, Del Mar Prado M, Proctor RH, Avalos J, Tudzynski B. 2002. The polyketide synthase gene pks4 from *Gibberella fujikuroi* encodes a key enzyme in the biosynthesis of the red pigment bikaverin. Fungal Genet Biol 37:134–148. https://doi.org/10.1016/S1087-1845(02)00501-7.
- 42. Ortoneda M, Guarro J, Madrid MP, Caracuel Z, Roncero MI, Mayayo E, Di Pietro A. 2004. *Fusarium oxysporum* as a multihost model for the genetic dissection of fungal virulence in plants and mammals. Infect Immun 72:1760–1766. https://doi.org/10.1128/IAI.72.3.1760-1766.2004.
- Birkenbihl RP, Diezel C, Somssich IE. 2012. Arabidopsis WRKY33 is a key transcriptional regulator of hormonal and metabolic responses toward *Botrytis cinerea* infection. Plant Physiol 159:266–285. https://doi.org/10 .1104/pp.111.192641.
- 44. Hendrick CA, Sequeira L. 1984. Lipopolysaccharide-defective mutants of the wilt pathogen *Pseudomonas solanacearum*. Appl Environ Microbiol 48:94–101.
- 45. Darken MA, Jensen AL, Shu P. 1959. Production of gibberellic acid by fermentation. Appl Microbiol 7:301–303.
- Moree WJ, Phelan VV, Wu CH, Bandeira N, Cornett DS, Duggan BM, Dorrestein PC. 2012. Interkingdom metabolic transformations captured by microbial imaging mass spectrometry. Proc Natl Acad Sci U S A 109:13811–13816. https://doi.org/10.1073/pnas.1206855109.
- 47. Genin S, Gough CL, Zischek C, Boucher CA. 1992. Evidence that the hrpB gene encodes a positive regulator of pathogenicity genes from *Pseudomonas solanacearum*. Mol Microbiol 6:3065–3076. https://doi.org/10.1111/j.1365-2958.1992.tb01764.x.
- Melamud E, Vastag L, Rabinowitz JD. 2010. Metabolomic analysis and visualization engine for LC-MS data. Anal Chem 82:9818–9826. https:// doi.org/10.1021/ac1021166.
- Smith CA, Want EJ, Maille GO, Abagyan R, Siuzdak G. 2006. XCMS: Processing mass spectrometry data for metabolite profiling using nonlinear peak alignment, matching, and identification. Anal Chem 78: 779–787.