Cocultivation of Anaerobic Fungi with Rumen Bacteria Establishes an Antagonistic Relationship

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ABSTRACT Anaerobic gut fungi (Neocallimastigomycetes) live in the digestive tract of large herbivores, where they are vastly outnumbered by bacteria. It has been suggested that anaerobic fungi challenge growth of bacteria owing to the wealth of biosynthetic genes in fungal genomes, although this relationship has not been experimentally tested. Here, we cocultivated the rumen bacteria *Fibrobacter succinogenes* strain UWB7 with the anaerobic gut fungi *Anaeromyces robustus* or *Caecomyces churrovis* on a range of carbon substrates and quantified the bacterial and fungal transcriptomic response. Synthetic cocultures were established for at least 24 h, as verified by active fungal and bacterial transcription. *A. robustus* upregulated components of its secondary metabolism in the presence of *Fibrobacter succinogenes* strain UWB7, including six nonribosomal peptide synthetases, one polyketide synthase-like enzyme, and five polyketide synthesis O-type methyltransferases. Both *A. robustus* and *C. churrovis* cocultures upregulated S-adenosyl-l-methionine (SAM)-dependent methyltransferases, histone methyltransferases, and an acetyltransferase. Fungal histone 3 lysine 27 trimethylation marks were more abundant in coculture, and heterochromatin protein-1 was downregulated. Together, these findings suggest that fungal chromatin remodeling occurs when bacteria are present. *F. succinogenes* strain UWB7 upregulated four genes in coculture encoding drug efflux pumps, which likely protect the cell against toxins. Furthermore, untargeted nonpolar metabolomics data revealed at least one novel fungal metabolite enriched in coculture, which may be a defense compound. Taken together, these data suggest that *A. robustus* and *C. churrovis* produce antimicrobials when exposed to rumen bacteria and, more broadly, that anaerobic gut fungi are a source of novel antibiotics.

IMPORTANCE Anaerobic fungi are outnumbered by bacteria by 4 orders of magnitude in the herbivore rumen. Despite their numerical disadvantage, they are resilient members of the rumen microbiome. Previous studies mining the genomes of anaerobic fungi identified genes encoding enzymes to produce natural products, which are small molecules that are often antimicrobials. In this work, we cocultured the anaerobic fungus *Anaeromyces robustus* or *Caecomyces churrovis* with rumen bacteria *Fibrobacter succinogenes* strain UWB7 and sequenced fungal and bacterial active genes via transcriptome sequencing (RNA-seq). Consistent with production of a fungal defense compound, bacteria upregulated genes encoding drug efflux pumps, which often export toxic molecules, and fungi upregulated genes encoding biosynthetic enzymes of natural products. Furthermore, tandem mass spectrometry detected an unknown fungal metabolite enriched in the coculture. Together, these
findings point to an antagonistic relationship between anaerobic fungi and rumen bacteria resulting in the production of a fungal compound with potential antimicrobial activity.

**KEYWORDS** RNA-seq, transcriptomics, cocultivation, secondary metabolism, fungi, anaerobe, anaerobic fungi

Microbial antagonism can take many forms: antibiosis (the production by an organism of a compound that inhibits or kills another organism), competition for nutrients and space, parasitism, and others (1). Although often discussed in the context of biological control agents that protect postharvest crops (1–3), microbial antagonism has also been recognized to have a profound impact on microbial communities, especially host-associated communities (4). For example, microbial antagonism can increase microbial diversity (5, 6), protect against invasion by pathogens (7), and drive genome evolution through the acquisition of genetic material from killed cells (8). Mathematical modeling suggests that communities dominated by antagonistic relationships are more stable and resilient to perturbations than those dominated by cooperative relationships (9).

Microbial relationships, especially antagonistic ones, within the rumen microbiome are complex and not well-characterized. In particular, knowledge of rumen fungi (class Neocallimastigomycetes) and their interactions with other microbial community members is lacking. Rumen fungi, also referred to as anaerobic gut fungi, thrive in the digestive tracts of large herbivores as part of a biomass-degrading consortium with bacteria, methanogenic archaea, and protozoa (10, 11). Bacteria outnumber fungi in the rumen by at least 4 orders of magnitude (10, 11). Cocultivation of fungi with bacteria suggests that the nature of the interaction between rumen fungi and bacteria depends on the specific fungal-bacterial pairing. Antagonistic relationships, in which the cellulolytic activity of the fungus was inhibited, were observed between *Ruminococcus flavefaciens* and *Neocallimastix frontalis* MCH3 or *Piromyces communis* FL (12), *Piromyces communis*, and *Selenomonas ruminantium* (13), as well as *R. flavefaciens* and *Orpinomyces joyonii* or *N. frontalis* (14). Some previous studies of rumen fungi cocultivated with the cellulolytic rumen bacteria *Fibrobacter succinogenes* have shown no effect on biomass degradation (12, 14, 15), implying neither mutualism nor antagonism between these organisms. However, Joblin and colleagues found that *F. succinogenes* inhibited the degradation of ryegrass stems by *N. frontalis* in coculture with *Methanobrevibacter smithii*, whereas the presence of *F. succinogenes* enhanced degradation by cocultures of *Caecomyces* spp. with *M. smithii* (16). In a separate study by Roger and colleagues (14), the presence of *F. succinogenes* had no impact on the degradation of wheat straw or maize stem by *N. frontalis* or *Orpinomyces (Neocallimastix) joyonii*.

Coculture transcriptomics has proven to be a valuable tool by which to investigate the nature of microbial interactions, as demonstrated in recent publications (17–21). For example, RNA sequencing (RNA-seq) of the anaerobic fungus *Anaeromyces robustus* in coculture with the methanogen *Methanobacterium bryantii* revealed that the fungus upregulated 105 genes encoding carbohydrate-active enzymes (CAZymes), representing 12% of total predicted CAZymes (18). Coculture transcriptomics and fermentation profiling of *Pecoramyces* sp. strain F1 with *Methanobrevibacter thaueri* also supported a syntrophic fungal-methanogen relationship (21). However, RNA sequencing of both fungi and bacteria in coculture remains difficult due to the technical challenge of depleting rRNA from both microbes. Here, we cocultivated pairings of rumen fungi with *Fibrobacter succinogenes* strain UWB7 (2), and performed the first dual transcriptomic characterization of a rumen bacterium and fungus in coculture. By this approach, we tested the hypothesis that the relationship between *F. succinogenes* strain UWB7 and anaerobic gut fungi is antagonistic. Furthermore, we performed untargeted non-polar metabolomics to investigate whether the introduction of *F. succinogenes* strain UWB7 triggers the production of possible defense compounds by the anaerobic fungus. Specifically, we cultured *Anaeromyces robustus* with *F. succinogenes* strain UWB7.
RESULTS AND DISCUSSION

Secondary metabolites, although not strictly necessary for the growth or survival of an organism under all growth conditions (22), are often secreted during antagonistic relationships between microorganisms (23–25). Previous work mining the high-quality genomes of anaerobic fungi revealed that anaerobic fungi are capable of synthesizing secondary metabolites (26). A. robustus and C. churrovis encode 43 and 32 biosynthetic enzymes, respectively, for various classes of secondary metabolites, including nonribosomal peptide synthetases (NRPSs) and polyketide synthases (PKSs) (26). We hypothesized that some of the secondary metabolites produced by A. robustus and C. churrovis are compounds used for regulation, defense, or competition against rumen bacteria.

Cocultivation with rumen bacteria induces stress in anaerobic fungi and activates components of fungal secondary metabolism. Although not stable for many generations of batch passaging, anaerobic fungi can grow with F. succinogenes strain UW7 for a sufficient duration to capture transcriptional responses from both organisms in coculture. A. robustus or C. churrovis was grown in isolation for 24 h prior to the introduction of F. succinogenes strain UW7 (see Fig. S1 in the supplemental material). Fungi and bacteria were cocultured until the fungus reached mid-log phase, at which point the cocultures and corresponding fungal monocultures were harvested for RNA extraction and sequencing. In response to the presence of F. succinogenes strain UW7 (Fig. 1), cultures of A. robustus or C. churrovis both upregulated genes encoding stress response proteins (chaperones), indicating that the presence of the bacteria invoked a fungal stress response (Fig. 2; see also Data Sets S1 to S3, available at https://github.com/cswift3/anaerobic_fungi_Fibrobacter_co-culture). Fungal stress was observed during growth on switchgrass or Avicel PH-101. Gene set enrichment analysis (27) supported enrichment of small heat shock proteins upregulated in coculture (Data Sets S4 to S6 at the URL mentioned above). However, roughly 10 times more genes were differentially regulated comparing fungal coculture to monoculture during growth on Avicel relative to growth on switchgrass (Table S2 and Data Sets S1 to S3 at the URL mentioned above). This difference may reflect the preference of F. succinogenes strain UW7, a cellulose-degrading specialist (28), for crystalline cellulose over complex plant
matter, resulting in both more robust bacterial growth and a stronger bacterial signal to the fungi.

In addition to a general stress response, *A. robustus* upregulated six nonribosomal peptide synthetases (NRPSs) and one polyketide synthase (PKS)-like enzyme (Table 1) at least 2-fold (adjusted *P* value of <0.05) in coculture with *F. succinogenes* strain UWB7 compared to *A. robustus* monoculture (both cultures grown on Avicel). These genes and others related to fungal secondary metabolism were previously annotated (26). Notably, these genes do not have homologs in *C. churrovis*, as determined by a bidirectional protein BLAST (29) in the MycoCosm portal (30). This suggests that the aspects of the response to *F. succinogenes* strain UWB7 related to secondary metabolism are specific to the fungal strain. Neighboring genes were coregulated with a pair of NRPS genes (Fig. 3). One NRPS gene cluster was downregulated in the Avicel coculture (Fig. 3). In higher-order fungi, secondary metabolites are linked to different developmental stages of fungi (31–33), and some of the secondary metabolites from anaerobic fungi may also serve such a purpose.

In addition, five genes encoding putative polyketide O-methyltransferases were upregulated with a log2 fold change of 2.7 or greater (Table S3). Surprisingly, a predicted protein (271870) containing a condensation domain, which would normally form a modular domain on an NRPS (34), was upregulated 2-fold in coculture relative to monoculture. Taken together, these data suggest that *A. robustus* regulates polyketide and nonribosomal peptide synthesis in response to microbial challenge by *F. succinogenes* strain UWB7.
Anaerobic fungi regulate their secondary metabolism via epigenetic modifications in the presence of rumen bacteria. LaeA is reported to modulate gene expression of biosynthetic gene clusters (BGCs) via chromatin remodeling (35), and studies have found that epigenetic modifications such as histone acetylation or methylation can regulate expression of biosynthetic gene clusters in fungi (35–37). Both A. robustus and C. churrovis upregulated genes encoding S-adenosyl methionine (SAM)-dependent methyltransferases when cocultured with *F. succinogenes* strain UWB7 (Table S3). One of these proteins may perform a function similar to that of LaeA or Lae1, which act in concert with other proteins, including proteins containing velvet domains (38), as global regulators of secondary metabolism in higher-order fungi (39–41). In previous work, we identified a homolog of the velvet-containing gene vosA (42, 43) in *C. churrovis* (26). We hypothesize that anaerobic gut fungi remodel chromatin via histone modifications to modulate their secondary metabolism, similar to what has been suggested for higher-order fungi (35, 44). When cocultured with *F. succinogenes* strain UWB7, both *C. churrovis* and *A. robustus* upregulated genes with putative functions in histone methylation or acetylation (Table S3), which are both modifications known to be involved in determining heterochromatin or euchromatin locations (45). It is possible that one of the highly upregulated methyltransferases in coculture acts as a global regulator of secondary metabolism in anaerobic gut fungi, similar to LaeA and Lae1. However, the distant evolutionary relationship between Neocallimastigomycetes and higher-order fungi as well as the current

![Gene expression diagram](image-url)

**FIG 3** Genes that were coordinately regulated in coculture of *A. robustus* with *Fibrobacter* sp. strain UWB7 versus fungal monocultures. All cultures were grown on Avicel. Log₂ fold change is shown above each gene, and the MycoCosm protein ID is shown below each gene.

### TABLE 1 Differentially regulated biosynthetic genes for secondary metabolites

| MycoCosm protein ID | SM type     | Log₂ fold change | Scaffold |
|---------------------|-------------|------------------|----------|
| A. robustus (Avicel) | NRPS 193122 | 2.9              | 480      |
|                     | NRPS 294553 | 2.9              | 182      |
|                     | NRPS 271076 | 2                | 279      |
|                     | NRPS 231391 | 1.8              | 77       |
|                     | PKS-like 266215 | 1.6          | 49       |
|                     | NRPS 218823 | 1.2              | 77       |
|                     | NRPS 330657 | 1.1              | 540      |
|                     | PKS 328517* | -1.0             | 207      |
| C. churrovis (switchgrass) | PKS 17094 | -3.6             | 116      |

*Genes marked with an asterisk in the protein ID column indicate the gene is coregulated with neighboring genes (Fig. 2). Adjusted *P* value of <0.05. The log₂ fold change refers to the fungus in coculture with *Fibrobacter* compared to the fungus grown in monoculture.
lack of genetic tractability of rumen fungi makes previous approaches used to pinpoint LaeA homologs unreliable at this time.

To test whether there were differences in the amount of histone 3-lysine 4 and histone 3-lysine 27 trimethylation marks (H3K4me3 and H3K27me3, respectively) between fungal monocultures and fungal cocultures with *Fibrobacter succinogenes* strain UWB7, we performed Western blotting on monoculture and coculture cell lysates using antibodies raised to *S. cerevisiae* H3K4me3 and H3K27me3 (Fig. S2). The exposure time for the H3 loading control increased by a factor of 10 between monoculture and coculture, indicating decreased fungal biomass in coculture. However, the exposure times were nearly equivalent between H3K27me3 blots of monocultures and cocultures, indicating that despite the decreased fungal biomass in coculture, there was an increased proportion of H3K27me3 marks. H3K27me3 is known to be a repressor of transcription, whereas H3K4me3 is an activating mark (46). Consistent with the downregulation of genes due to the enhancement of H3K27me3 marks in coculture, more genes were downregulated than upregulated when comparing *A. robustus* cocultures to monocultures during growth on Avicel (Table S2). These results support that epigenetic modifications influence gene regulation when *A. robustus* is exposed to *F. succinogenes* strain UWB7. Furthermore, two genes encoding heterochromatin-associated protein HP1 were greater than 2-fold downregulated in coculture (protein identifiers [IDs] 290815 and 266437), and another gene (280338) encoding a homolog of the WSTF-ISWI chromatin remodeling complex, which has been implicated in the replication of heterochromatin (47), was 8-fold downregulated in coculture (Data Set S1 at https://github.com/cswift3/anaerobic_fungi_Fibrobacter_co-culture). In aspergilli, heterochromatin protein-1 and H3K9me3 marks have been associated with the repression of secondary metabolite gene clusters (35, 44). Taken together, these findings suggest that the secondary metabolism of anaerobic fungi is regulated via epigenetic marks and chromatin remodeling, consistent with higher-order fungi.

To further understand the fungal regulatory response to cocultivation with *F. succinogenes* strain UWB7, we analyzed the eukaryotic orthologous groups (KOGs) (48) of the differentially regulated transcripts (Fig. S3). In the case of *A. robustus* cultured with *F. succinogenes* strain UWB7 on Avicel, the percentage of significantly downregulated fungal genes in coculture was greater than that upregulated for the majority of the KOG classes (absolute log, fold change of ≥1, adjusted P value of <0.05). In other words, cocultivation repressed most fungal cellular and metabolic processes, with the exceptions of the KOG classes of posttranslational modifications and secondary metabolism. Table S4 lists the upregulated genes in coculture within the KOG class of secondary metabolism. In addition to the NRPS genes described above, 12 multidrug exporters belonging to the ABC superfamily were upregulated. The percentage of differentially regulated genes in all of the KOG classes was greater than 10%, with a median of 23% (Data Set S7 at the URL mentioned above). In contrast, cultivation on switchgrass resulted in far fewer differentially regulated genes in each KOG class between coculture and monoculture: *A. robustus* significantly regulated a median value of 1% of genes in each KOG class (Data Set S8 at the URL mentioned above), and *C. churrovis* regulated 5% (Data Set S9 at the URL mentioned above). These findings further support that the cultivation of *F. succinogenes* strain UWB7 with *A. robustus* on Avicel resulted in a stronger bacterial signal to the fungus. More broadly, it is clear that the choice of substrate, in addition to the specific organisms, has a profound impact on the gene regulation of microorganisms in cocultures.

**Rumen bacteria upregulate genes encoding components of drug efflux pumps when anaerobic fungi are present.** To further probe the relationship between *F. succinogenes* strain UWB7 and anaerobic gut fungi, we sequenced the corresponding prokaryotic mRNA in coculture and monoculture, using both eukaryotic and prokaryotic rRNA depletion (see Materials and Methods). When cocultivated with *C. churrovis* using switchgrass as the carbon source, *F. succinogenes* strain UWB7 upregulated 143 genes and downregulated 261 genes (4 and 8% of predicted genes in IMG/M (49)) (Data Set S10 at the URL mentioned above). Putative transporters comprised 12% of the upregulated genes. Table 2 summarizes upregulated genes encoding transporters (log2 fold change of ≥1.0,
TABLE 2  

| Locus tag          | Log₂ fold change | Product name                                      |
|--------------------|------------------|--------------------------------------------------|
| Ga0136279_2636     | 2.9              | ABC transporter ATP-binding protein               |
| Ga0136279_2635     | 2.2              | Putative ABC transport system permease protein    |
| Ga0136279_2405     | 2.2              | Type II and III secretion system protein          |
| Ga0136279_1390     | 1.8              | Outer membrane protein beta-barrel domain-containing protein |
| Ga0136279_1256     | 1.7              | Urea ABC transporter substrate-binding protein    |
| Ga0136279_0657     | 1.6              | Multispecies efflux RND transporter periplasmic adaptor subunit |
| Ga0136279_2085     | 1.6              | Zinc ABC transporter substrate-binding protein    |
| Ga0136279_1465     | 1.5              | Multispecies ammonium transporter                 |
| Ga0136279_2620     | 1.5              | Transporter                                       |
| Ga0136279_2553     | 1.4              | Multispecies ABC transporter substrate-binding protein |
| Ga0136279_2080     | 1.4              | Iron complex outer membrane receptor protein      |
| Ga0136279_1904     | 1.3              | General secretion pathway protein E               |
| Ga0136279_1405     | 1.2              | TonB family C-terminal domain-containing protein  |
| Ga0136279_1902     | 1.1              | Multispecies efflux RND transporter periplasmic adaptor subunit |
| Ga0136279_0818     | 1.0              | TRAP transporter large permease subunit           |
| Ga0136279_1391     | 1.0              | Calcium/sodium antiporter                        |
| Ga0136279_1901     | 1.0              | Multispecies TolC family protein                 |
| Ga0136279_2080     | 1.4              | Iron complex outer membrane receptor protein      |
| Ga0136279_1405     | 1.3              | General secretion pathway protein E               |
| Ga0136279_1902     | 1.1              | Multispecies efflux RND transporter periplasmic adaptor subunit |
| Ga0136279_0818     | 1.0              | TRAP transporter large permease subunit           |
| Ga0136279_1391     | 1.0              | Calcium/sodium antiporter                        |
| Ga0136279_1901     | 1.0              | Multispecies TolC family protein                 |

*Locus tags Ga0136279_2080 and Ga0136279_1405 are not transporters but are part of the TonB receptor complex involved in iron transport (83–85).*

*Adjusted P value of less than 0.05. Product names were taken from the protein details for RefSeq NZ_FRC00000000.1 or the gene product name in IMG/M (49).*

adjusted P value of <0.05). Notably, a predicted TolC family protein (locus tag Ga0136279_1901) was 2-fold upregulated in coculture relative to *F. succinogenes* strain UWB7 monoculture (adjusted P value of 1.5 × 10⁻¹⁵). TolC proteins are components of efflux pumps in Gram-negative bacteria, and these pumps can transport a wide array of molecules, including antibiotics (50). In addition, two genes encoding the adaptor subunits of RND efflux pumps (Ga0136279_1902 and Ga0136279_0657) were 2- and 3-fold upregulated in coculture (adjusted P value of ≤10⁻¹⁵), suggesting that they are part of a regulon. Upon inspection of the gene neighborhoods (Fig. S4), the TolC family protein encoded by Ga0136279_1901 and adapter subunit Ga0136279_1902 were neighboring genes. The correlation of genes encoding components of multidrug efflux pumps has been previously reported. In *Enterobacteriaceae*, the genes encoding the multidrug efflux pump AcrAB-ToIC (acrA, acrB, and toIC) form a regulon (51), although toIC is not colocalized with acrA and acrB. The 2-fold upregulated gene Ga0136279_2553, annotated as an ABC transporter substrate binding protein, also bordered a gene encoding a TolC family protein (Ga0136279_2554), although the gene encoding the putative TolC protein was not differentially regulated in coculture. For effective efflux directly into the external environment, both the outer membrane channel, such as TolC, and the periplasmic adaptor are necessary (52). Therefore, it is significant that both the TolC and adaptor protein homologs are upregulated when *F. succinogenes* strain UWB7 is cocultivated with *C. churrovis*.

Besides drug efflux pumps, *F. succinogenes* strain UWB7 also upregulated at least 2-fold six genes encoding chaperones (Data Set S10 at the URL mentioned above), supporting the induction of a bacterial stress response by cocultivation with anaerobic fungi. Notably, *F. succinogenes* strain UWB7 also upregulated 32-fold a putative HicB antitoxin (Ga0136279_0693), which could be part of a toxin-antitoxin system (53). In addition, *F. succinogenes* strain UWB7 2-fold upregulated a gene encoding a putative abortive phage resistance protein (Ga0136279_0760), typically part of an RNA toxin-antitoxin system (54).

Coculture of anaerobic fungi with bacteria points to the secretion of unique metabolites. To further test the hypothesis that cocultivation of anaerobic gut fungi with *F. succinogenes* strain UWB7 triggers the production of fungal defense compounds, we performed untargeted nonpolar metabolomics analysis on fungal-bacterial cocultures and the respective fungal or bacterial monocultures. We constructed a
principal-component analysis (PCA) plot using MetaboAnalyst (55) (Fig. 4 and Fig. S5). In the three-dimensional scores plot (Fig. 4), bacterial monocultures and fungal-bacterial cocultures grown on switchgrass show a high degree of overlap. However, the cultures grown on Avicel (A. robustus monoculture, F. succinogenes strain UWB7 monoculture, and coculture) were distinct from each other. This separation is apparent in a three-dimensional PCA scores plot (Fig. 4) but not in the two-dimensional scores plot (Fig. S5). Taken together, these data suggest that, in contrast to cultures grown on switchgrass, the metabolic profiles observed in the cocultures of A. robustus with F. succinogenes strain UWB7 on Avicel are distinct from those observed in the respective monocultures.

We further investigated the distribution of nonpolar metabolites between monocultures and cocultures by constructing molecular networks using Global Natural Products Social Molecular Networking (GNPS) (56) and visualizing the networks in Cytoscape (57) with three-way coloring (58) (Fig. 5). This approach highlighted a group of unknown metabolites unique to F. succinogenes strain UWB7 that were not observed in fungal monocultures. These metabolites were not enriched by cocultivation with anaerobic fungi and therefore likely represent constitutively produced bacterial metabolites. The closest known node in the cluster matched 1-palmitoyl-sn-glycero-3-phosphoethanolamine (m/z 454.2791), a lysophospholipid (LPL), which comprises a low concentration of Gram-negative bacterial cell membranes (59). Structure and molecular class predictions from SIRIUS (60) and CANOPUS (61) suggested that the unknown nodes represent glycerophosphoethanolamines. Overall this cluster of nodes likely represents components of the bacterial membrane that are released into the supernatant after cell death and lysis.

By comparing the peak height ratios of features between culture conditions, we searched for metabolites that were enriched by cocultivation with F. succinogenes strain UWB7. We screened these metabolites for those that were consistently observed across all four biological coculture replicates. We applied an enrichment threshold of 4-fold, reasoning that a 4-fold enhancement would most likely not be caused by the simple addition of bacterial and fungal biomass in coculture but rather increased production by one or both organisms. Four features matched these stringent criteria (Table 3). All four features are single nodes truncated from the molecular network.
Cultivation of anaerobic fungi with *Fibrobacter succinogenes* strain UWB7 reveals diverse shared metabolites as well as a group of bacterial metabolites. Combined feature-based molecular network was created by GNPS (56) from positive- and (Continued on next page)
depicted in Fig. 5 and, thus, are not likely to be part of a family of structurally related compounds. Notably, the feature m/z 244.227 was not observed in *F. succinogenes* strain UWB7 monoculture but was 12-fold enriched in coculture relative to *A. robustus* monoculture on Avicel, which suggests that this is a unique fungal metabolite with enhanced production in response to the presence of *F. succinogenes* strain UWB7.

**Conclusions.** We have demonstrated using dual transcriptomics that, despite previous reports that *Fibrobacter succinogenes* had no interaction with rumen fungi (12, 14), as assessed by extent of biomass degradation in cocultures compared to monocultures, cocultivation of the close relative *F. succinogenes* strain UWB7 with *A. robustus* or *C. churrovis* resulted in drastic changes to both bacterial and fungal transcriptomes, including upregulation of bacterial drug efflux pumps and fungal chaperones, polyketide O-methyltransferases, PKSs, and NRPSs. Furthermore, fungal genes encoding putative histone-modifying enzymes were upregulated in coculture. Histone 3-lysine 27 trimethylation marks increased and heterochromatin-associated protein-1 was downregulated in coculture. Together, these results suggest that, similar to higher-order fungi, anaerobic fungi regulate their secondary metabolism via chromatin remodeling. These data support that anaerobic gut fungi activate their secondary metabolism via epigenetic and transcriptional regulation when challenged by rumen bacteria. The metabolic outcome of these transcriptional changes may be the production of a fungal defense compound produced by a PKS or NRPS. Consistent with this hypothesis, untargeted nonpolar metabolomics supports that at least one unique fungal metabolite is enriched by cocultivation with *F. succinogenes* strain UWB7. As a consequence, anaerobic fungi and the antagonistic relationships of the rumen microbiome may prove to be a valuable source of antibiotics.

**MATERIALS AND METHODS**

**Isolation and cultivation of anaerobic gut fungi.** *Anaeromyces robustus* was isolated via reed canary grass enrichment from the fecal pellet of a Churro sheep at the Santa Barbara Zoo, as described previously (62, 63). *Caecomyces churrovis* was isolated similarly (64). Both fungi were cultivated anaerobically in Hungate tubes at 39°C with reed canary grass as the carbon source in a modified formulation (MC-) of complex medium C (65), containing 0.25 g/liter yeast extract (before boiling), 0.5 g/liter Bacto Casitone (before boiling), and 7.5 vol% clarified rumen fluid. The medium was supplemented with vitamins after autoclaving as described by Teunissen and colleagues (66). Cultures were passaged every 3 to 4 days into fresh media via a 1.0-ml sterile syringe.

*Aerococcus succinogenes* strain UWB7. The strain *F. succinogenes* strain UWB7 was a generous gift from Garret Suen at the University of Wisconsin-Madison. Details of the isolation of this strain are described in Neumann and Suen (28). *F. succinogenes* strain UWB7 was cultivated at 39°C anaerobically in Hungate tubes containing 9.0 ml of MC- medium supplemented with vitamin solution, prepared as

**FIG 5 Legend (Continued)**

negative-ion mode LC-MS/MS data from the nonpolar metabolites of *A. robustus*, *C. churrovis*, and *F. succinogenes* strain UWB7 cocultures and monocultures grown on Avicel or switchgrass substrates. Self-looping nodes were truncated. Three-way coloring (58) was used to visualize features in the *A. robustus* monocultures, *A. robustus*-F. succinogenes strain UWB7 cocultures, and *F. succinogenes* strain UWB7 monocultures (all grown on Avicel). Transparency of the nodes was set to emphasize nodes with high intensity in the coculture of *A. robustus*-F. succinogenes strain UWB7.

| TABLE 3 Metabolites enriched in fungal-bacterial cocultures relative to monocultures a |
|---------------------------------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| m/z | RT | Avicel (A. robustus) | C. churrovis | Switchgrass | A. robustus | C. churrovis | A. robustus | C. churrovis |
|-----|----|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
|-----|----|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| 196.040 | 2.39 | 12.89 | 19.97 | 12.35 | 8.95 | 4.52 | 13.20 | 4.98 | 4.96 |
| 244.227 | 5.80 | 16.88 | 10.51 | 12.13 | 4.98 | 4.96 | 4.98 | 4.96 |
| 380.277 | 4.98 | — b | — b | — b | — b | — b | — b | — b |
| 408.308 | 7.43 | — b | — b | — b | — b | — b | — b | — b |

a Untargeted nonpolar metabolomics features enriched at least 4-fold in coculture of *A. robustus* with *Fibrobacter succinogenes* strain UWB7 on Avicel compared to both bacterial and fungal monocultures (one-tailed Student’s t statistic, <0.05). All features were detected in positive-ion mode.

b —, feature was not detectable in UWB7 monoculture.

Swift et al.® July/August 2021 Volume 12 Issue 4 e01442-21 mbio.asm.org 10
described above, and 0.1 g of Avicel PH-101 (Sigma-Aldrich, St. Louis, MO). Each Hungate tube was inoculated with 1.0 ml of cryostock or live F. succinogenes strain UWB7 culture.

**Overview of the cocultivation conditions of anaerobic gut fungi with *Fibrobacter succinogenes* strain UWB7.** An overview of the cocultivation pairings, carbon substrates, and cocultivation incubation times is depicted in Fig. S1 in the supplemental material. Briefly, *A. robustus* was cocultivated with *Fibrobacter succinogenes* strain UWB7 on both Avicel PH-101 (Sigma-Aldrich, St. Louis, MO) and milled switchgrass (gift from U.S. Department of Agriculture), whereas *C. churrovis* was cocultivated with *F. succinogenes* strain UWB7 on switchgrass only due to the slow growth of *C. churrovis* on Avicel (64). Since *A. robustus* and *C. churrovis* are expected to grow more slowly than *F. succinogenes* strain UWB7, as evidenced by the order-of-magnitude larger specific growth rate of *F. succinogenes* compared to *C. churrovis* on soluble sugars (64, 67), both strains of anaerobic gut fungi were allowed to grow for 24 h prior to inoculation with *F. succinogenes* strain UWB7. Cocultures were subsequently allowed to grow for an additional 24 to 72 h prior to harvesting for RNA extraction. The length of incubation for the cocultivation pairings was set by time necessary for the fungus to reach mid-log growth phase, as assessed by cumulative pressure (68) of fungal monocolonies as well as visual assessment.

**Cocultivation of anaerobic fungi with *Fibrobacter succinogenes* strain UWB7.** Anaerobic liquid growth medium MC- was prepared by following the recipe for complex medium C (65), with yeast extract, Bacto Casitone, and clarified switchgrass culture filtrate added to the medium (65). A 100-ml capacity serum bottle was filled with 80 ml of MC- liquid medium and 0.8 g of switchgrass or Avicel PH-101 (Sigma-Aldrich, St. Louis, MO). The serum bottle and its contents were flushed with CO2, autoclaved, and supplemented with 0.8 ml of 100× vitamin solution (66). The serum bottle was preheated to 39°C, and then a seed culture of *A. robustus* was added into inoculating 1.0 ml of the routinely passaged *A. robustus* described above into the liquid medium using a 1-ml sterile syringe. The seed culture was immediately vented following inoculation and then incubated at 39°C for 4 days. This seed culture was used to inoculate cultures to be harvested for subsequent RNA-seq. Cultures were prepared in replicates of four by inoculating 1.0 ml of *A. robustus* into 8.0 ml (coculture) or 9.0 ml (monoculture) of MC- containing 0.1 g switchgrass or Avicel PH-101 (Sigma-Aldrich, St. Louis, MO). Prior to inoculation, the medium and substrate were autoclaved, supplemented with 0.1 ml of 100× vitamin solution (66) postautoclaving, and preheated to 39°C. The fungal culture was grown at 39°C for 24 h, and then cocultures were started by inoculating 1.0 ml of *F. succinogenes* strain UWB7 into each of four replicates. The seed culture of *F. succinogenes* strain UWB7 was grown for 24 h at 39°C from 1.0 ml of inoculum in a serum bottle containing 80 ml of MC-, 0.8 g of Avicel PH-101 (Sigma-Aldrich, St. Louis, MO), 0.6 ml of 100× vitamin solution (66), and a CO2 headspace. A. robustus-F. succinogenes strain UWB7 cocultures were incubated for an additional 24 h (switchgrass substrate) or 48 h (Avicel). The contents of each Hungate tube were transferred to 15-ml Falcon tubes (Fisher Scientific, Waltham, MA) and centrifuged at 3,200 g at 4°C using a swinging-bucket rotor (Eppendorf A-4-81) for 10 min. The supernatant was saved at −80°C for subsequent liquid chromatography-tandem mass spectrometry (LC-MS/MS), and 1.0 ml of RNAlater (Sigma-Aldrich, St. Louis, MO) was added to the pellet to preserve the RNA. The pellet from each culture was frozen at −80°C until lysis.

The cocultivation of *C. churrovis* with *F. succinogenes* strain UWB7 was performed identically to the cocultivation of *A. robustus* with *F. succinogenes* strain UWB7, except that the length of the cocultivation incubation was 40 h.

**RNA extraction and QC.** Samples were thawed on ice and then centrifuged at 3,220 × g at 4°C using a swinging-bucket rotor (Eppendorf A-4-81) for 10 min. RNAlater was decanted. The pellets were transferred into 2-ml screw-cap tubes containing 1.0 ml of autoclaved 0.5-mm zirconia/silica beads (Biospec) and 600 µl buffer RLT (Qiagen, Hilden, Germany) with 1 vol% 2-mercaptoethanol (Sigma-Aldrich, St. Louis, MO). The tube was briefly vortexed, and then the cells were lysed using the Biospec Mini-Beadbeater-16 for 45 s. The tubes were placed on ice for 30 s. Following lysis, the tubes were centrifuged for 3 min at 13,000 × g and 20°C using a microcentrifuge (Eppendorf 5424). The supernatant was removed using gel-loading tips (Fisher Scientific, Waltham, MA) to maximize yields and centrifuged again to remove residual cell debris for 2 min at 20,000 × g. The supernatant from each tube was then transferred into 2-ml round-bottom sample tubes (Qiagen catalog number 980381). RNA was extracted using a QIAcube by following the RNeasy Mini protocol for animal cells with QiAshredder homogenization and optional on-column DNase digest.

Quantity and quality of RNA was assessed by a Qubit fluorometer and TapeStation (Agilent), respectively. All RNA integrity number equivalents (RIN*) were above 6.0, assessed by either eukaryotic or prokaryotic ribosomal markers for cocultures.

**RNA library preparation and sequencing.** In fungal monocolonies, fungal mRNA was selectively enriched by capturing polyadenylated RNA using poly-T beads. For bacterial monocolonies, rRNA was depleted using the Illumina Ribo-Zero rRNA removal kit (Yeast) spiked into the Illumina Ribo-Zero gold rRNA removal kit (Epidemiology). To obtain both bacterial and fungal libraries from the cocultures, each sample was divided, and 200 ng was used as the input into each alternative pipeline: (i) poly(A) selection for the fungal library or (ii) ribosomal depletion by an Illumina Ribo-Zero rRNA removal kit (yeast) spiked into the Illumina Ribo-Zero gold rRNA removal kit (Epidemiology) for the library enriched in bacterial mRNA. Stranded RNA-seq libraries were created by the Joint Genome Institute and quantified by quantitative PCR (qPCR). Libraries were sequenced by paired-end 150-bp reads using a NovaSeq (Iluminia, San Diego, CA).

**RNA-seq data analysis.** Raw reads were evaluated for artifact sequences using BBDuk (69). Detected artifacts identified using kmer matching (kmer = 25) were trimmed from the 3’ end of reads. Reads were further filtered by removing RNA spike-in reads, PhiX reads, and reads containing any N’s. Reads were
trimmed for quality using the phred trimming method (set at Q6). Following trimming, short reads of less than 50 bases were removed. Filtered reads were assigned to the reference genome for the respective organism (fungal genomes available on the MycoCosm portal [30] and F. succinogenes strain UW87 genome GenBank accession no. GCA_900142945.1) using HISAT2 version 2.1.0 (70). Raw gene counts were generated by featureCounts (71) using the gene annotation files available in MycoCosm (30) for A. robustus or C. churovis and IMG (49) for Fibrobacter sp. strain UW87.

The effectiveness of poly(A) selection and ribosomal depletion methods was quantified using SortMeRNA (72) and is discussed in Text S1 in the supplemental material. Differential expression analysis was performed using DESeq2 (version 1.18.1) (73), with a minimum absolute log, fold change of 1.0 and statistical significance threshold of adjusted P value of less than 0.05 (Benjamini and Hochberg method). Gene set enrichment analysis (27) was performed with regard to gene sets comprised of eukaryotic organism (fungal genomes available on the MycoCosm portal [30] and Fibrobacter sp. strain UW87 the input in RNK format. The following parameters were used: the number of permutations was set to 1,000, and maximum and minimum sizes of gene sets to exclude were set to 500 and 5, respectively.

**Extraction and LC-MS/MS.** Two milliliters of ethyl acetate was added to 1.5 ml of fungal supernatant, vortexed, sonicated for 10 min in a water bath (room temperature), and centrifuged (5 min at 5,000 rpm), and then the top ethyl acetate layer was removed to another tube. To serve as extraction controls, tubes without sample were extracted by following the same procedure. Extracts were dried in a SpeedVac (SPD111V; Thermo Scientific) and stored at −80°C.

In preparation for LC-MS analysis, 150 μl LC-MS-grade methanol containing 1 μg/ml internal standard (2-aminobromo-5-methylbenzoic acid; Sigma) was added to dried extracts, followed by a brief vortex and sonication in a water bath for 10 min; 150 μl of resuspended extract was then centrifuged (2.5 min at 2,500 rpm) using a 0.22-μm filter (UFC409V05; Millipore) and transferred to a glass autosampler vial. Reverse-phase chromatography was performed by injecting 2 μl of sample into a C18 chromatography column (2.1 by 50 mm, 1.8 μm; Agilent ZORBAX Eclipse Plus C18) warmed to 60°C with a flow rate of 0.4 ml/min equilibrated with 100% buffer A (100% LC-MS water with 0.1% formic acid) for 1 min, followed by a linear gradient to 100% buffer B (100% acetonitrile with 0.1% formic acid) for 7 min and then held at 100% B for 1.5 min. MS and MS/MS data were collected in both positive- and negative-ion mode using a Thermo Q Exactive HF mass spectrometer (ThermoFisher Scientific, San Jose, CA), with full MS spectra acquired ranging from 80 to 1,200 m/z at 60,000 resolution, and fragmentation data were acquired using an average of stepped collision energies of 10, 20, and 40 eV at 17,500 resolution. Orbitrap instrument parameters included a sheath gas flow rate of 50 (au, arbitrary units), auxiliary gas flow rate of 20 (au), sweep gas flow rate of 2 (au), 3-kV spray voltage, and 400°C capillary temperature. Sample injection order was randomized and an injection blank of methanol only run between each sample. Raw data are available for download at https://genome.jgi.doe.gov/portal/ under the JGI Project ID 1294405.

**Metabolomics data analysis:** creation of molecular network via Global Natural Products Social Molecular Networking. The Feature-Based Molecular Networking (FBMN) workflow (74) on GNPS (56) (https://gnps.ucsd.edu) was used to construct a molecular network. First, peak finding was performed with MZmine (version 2.39) (75). An MZmine workflow was used to generate a list of features (m/z, residence time values obtained from extracted ion chromatograms containing chromatographic peaks within a narrow m/z range) and filtered to remove isotopes, adducts, and features without MS/MS. AOMA (chromatogram builder and deconvolution modules were used (76). The exact parameters used are available in an XML document upon request from the corresponding author. This document describes the batch operations that can be read by MZMine directly. The molecular networking GNPS job can be publicly accessed at https://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=abeeb3b1a8fac4b67b54b6f171a305f3.

For each feature, the most intense fragmentation spectrum was uploaded to GNPS. All MS/MS fragments were removed within ±17 Da of the precursor m/z. Window filtering was achieved by selecting only the top 6 fragment ions in the ±50-Da window throughout the spectrum. Parameters were set as a precursor ion mass tolerance of 0.05 Da and MS/MS fragment ion tolerance of 0.05 Da. The edges of the molecular network were specified to a cosine score greater than 0.70 and more than 6 matching peaks. The edges were further filtered such that an edge was permitted if and only if the joined nodes were present in the other respective node’s top 10 most similar nodes. Lowest-scoring edges were removed from molecular families such that no family contained more than 100 nodes. All spectra within the molecular network were queried against GNPS spectral libraries (56). Each library spectrum was filtered by following the same procedure as that applied to the input data. The minimum criteria for a match between a network spectrum and a library spectrum were that the score be greater than 0.7 and that at least 6 peaks match. MS/MS spectra were annotated by DEREPLICATOR (77).

It should be noted that a spectrum match to a database spectrum is not a definitive identification of the feature. It could be an isomer with a similar fragmentation, an ion with a close but not exact m/z but similar fragmentation pattern, or an in-source degradation product of another larger molecule (the degradation product may look similar to the database match).

GNPS positive- and negative-mode networks were merged using a custom Python script to group nodes having a retention time difference of less than 0.15 min and an m/z difference of less than 20 parts per million, assuming the negative mode species ionized as [M-proton]− and the positive mode species ionized as [M+proton]++. The resulting network is available as a pdf (Data Set S11 at https://github.com/
Coculture of Anaerobic Fungi with Rumen Bacteria

cswift3/anaerobic_fungi_Fibrobacter_co-culture) or in GRAPHML format (for direct visualization in Cytoscape [57]) upon request from the corresponding author.

Finally, the molecular network was visualized using Cytoscape [57] and three-way coloring. Given three numerical values to compare, the corresponding hue for each value can be calculated according to Baran and colleagues [58] using a custom Python script. The transparency of each node is determined by the value of each normalized to the minimum and maximum of the set of values. In this case, the three values to compare were the GNPS-normalized peak areas of each feature (averaged across four biological replicates) for three different treatments: (i) A. robustus monoculture (Avicel substrate), (ii) F. succinogenes strain UW87 monoculture (Avicel substrate), and (iii) A. robustus-F. succinogenes strain UW87 coculture (Avicel substrate). The method of per-sample normalization selected in the GNPS job was “row sum normalization (per file sum to 1,000,000),” and the mean was chosen as the aggregation method per group (treatment). For the molecular network depicted in Fig. 5, the transparency of each node was normalized with respect to the minimum and maximum GNPS-normalized peak areas of the coculture condition. For example, features that are many orders of magnitude more intense in one group than another will not be transparent (high alpha). A square root normalization was applied to the intensity difference to calculate transparency values that emphasize the most important features. In comparison, features that have approximately the same intensity in all treatment groups will have high transparency (low alpha).

Structure and class prediction of the unknown bacterial metabolites (glycophosphoethanolamines; Fig. 5) was performed by SIRIUS 4.0 (60) and CANOPUS (61) by the MS-GF+ (1.3.0) workflow on the ProteoSAFe web server from the Center for Computation Mass Spectrometry. The job may be viewed and cloned from https://proteomics.ucsd.edu/ProteoSAFe/status.jsp?task=e4914be8624c4e4eb03b3f3e609918ab6c.

PCA of metabolomics data using MetaboAnalyst 4.0. MetaboAnalyst 4.0 (78) was used to construct principal-component analysis (PCA) plots from the peak-heights feature table of all samples, generated using MZmine2 (75, 79). Sample normalization was set to “normalization by sum.” The “promp” normalization in R (80), which requires the package “chrometrics,” was used internally within MetaboAnalyst to perform the PCA.

Growth of fungal cultures for Western blotting of fungal epigenetic modifications. To prepare the A. robustus seed culture, 1.0 ml of A. robustus from routine cultivation was transferred into a 60-ml glass serum bottle preheated to 39°C containing 40 ml of MC- medium with 0.4 g of Avicel PH-101 (Sigma-Aldrich, St. Louis, MO) and supplemented with 0.4 ml of 100× vitamin solution (66) after autoclaving. The seed culture was grown for 4 days. From the seed culture, 1.0 ml was inoculated into each of six 80-ml cultures of MC- with 0.8 g Avicel PH-101 (Sigma-Aldrich, St. Louis, MO) supplemented with 0.8 ml of 100× vitamin solution (66). Three of these cultures were incubated at 39°C for 24 h, and then each was fixed with 1.0 ml of F. succinogenes strain UW87 seed culture. The remaining three bottles were also incubated at 39°C, but they were not inoculated with F. succinogenes strain UW87. The F. succinogenes strain UW87 seed culture was prepared by inoculating 1.0 ml of active culture into a 60-ml serum bottle containing 40 ml of MC- supplemented with vitamin solution (66) and 0.4 g of Avicel PH-101 (Sigma-Aldrich, St. Louis, MO). The active culture was F. succinogenes strain UW87 revived from cryostock 1 week prior and passaged one time. The cocultures and monocultures were grown for a total of 72 h following the fungal inoculation. The cultures were then transferred into 50-ml Falcon tubes and centrifuged using a fixed-angle rotor (Eppendorf F-34-6-38) at 4°C and 3,000 × g for 10 min. The cell pellets were stored at −80°C until lysis.

Extraction of fungal cultures for Western blotting of fungal epigenetic modifications. The frozen cell pellets prepared above were resuspended in 3 ml of 2 M NaOH (Fisher Scientific, Waltham, MA, USA) with 10% (vol/vol) beta-mercaptoethanol (Sigma-Aldrich, St. Louis, MO, USA). The solution was gently mixed and incubated on ice for 5 min to promote hydrolysis of the fungal cell wall. The solution was then centrifuged at 14,000 × g for 30 s at 4°C. The resulting pellet was resuspended in 3 ml of high-salt extraction buffer containing 40 mM HEPES, pH 7.5 (Fisher Scientific, Waltham, MA, USA), 350 mM NaCl (Fisher Scientific, Waltham, MA, USA), 0.1% (wt/vol) Tween 20 (Bio-Rad, Hercules, CA, USA), and 10% (vol/vol) glycerol (Fisher Scientific, Waltham, MA, USA). The solution was immediately centrifuged at 14,000 × g for 30 s at 4°C. The resulting pellet was resuspended in 3 ml of 2 × SDS sample buffer, 0.1 M Tris-HCl, pH 6.8 (Fisher Scientific, Waltham MA, USA), 4% (wt/vol) SDS (Fisher Scientific, Waltham, MA, USA), 0.2% (wt/vol) bromophenol blue (Sigma-Aldrich, St. Louis, MO, USA), 20% (vol/vol) glycerol (Fisher Scientific, Waltham, MA, USA), and 10% (vol/vol) beta-mercaptoethanol (Sigma-Aldrich, St. Louis, MO, USA). Cells pellets incubated at 100°C for 10 min, prior to being centrifuged at 14,000 × g for 30 s at 4°C. The supernatants were stored at −20°C until further use.

Western blotting of fungal epigenetic modifications. Thirty microliters of previously frozen cell lysates prepared in SDS sample buffer was gently mixed prior to loading on a 15% polyacrylamide gel. Candida glabrata whole-cell lysate, extracted as mentioned for the extraction of fungal cultures for Western blotting, was used as the histone H3 nuclear loading control as well as a positive control for H3K4me3 Western blots. Pirrromyes sp. strain UH3-1 whole-cell lysate was used as a loading control for H3K27me3 Western blots. Gel electrophoresis occurred for 65 min at 150 V under constant voltage at room temperature. The gel and Immobilon polyvinylidene difluoride (PVDF) membrane (Fisher Scientific, Waltham, MA, USA) were briefly washed with 100% methanol (Fisher Scientific, Waltham, MA, USA) prior to being washed with 1 × Towbin buffer containing 25 mM Tris, pH 8.3 (Thomas Scientific, Swedesboro, NJ, USA), 192 mM glycine (Sigma-Aldrich, St. Louis, MO, USA), and 10% (vol/vol) methanol. The gel and membrane were overlaid on top of 9 pieces of 3MM chromatography paper (Fisher Scientific, Waltham, MA, USA) that were soaked in 1 × Towbin buffer. After overlaying the gel and membrane on top of the 9 sheets of chromatography paper, an additional six sheets of chromatography paper already saturated with 1 × Towbin buffer were overlaid on top of the gel. Both membrane and chromatography paper were previously cut to dimensions of 5.5 cm
by 8.5 cm in order to match the dimensions of the resolving gel. After rolling out the transfer sandwich to remove air bubbles, the proteins were transferred under semidy conditions using a Hoefer Hsi Semi-plot TE70 semidy transfer unit (Holliston, MA, USA) for 90 min at 42 mA under constant amperage at room temperature. Membranes were blocked overnight with 3% (wt/vol) milk (Great Value, Bentonville, AR, USA) with 0.15% (wt/vol) sodium azide (Sigma-Aldrich, St. Louis, MO, USA) dissolved in 1× Tris-buffered saline (TBS), pH 7.5 (Fisher Scientific, Waltham, MA, USA), at 4°C. The following day, membranes were washed at room temperature for 30 min with 1× TBS buffer, pH 7.5, exchanging the buffer every 10 min. Primary antibodies were diluted in 10 ml of 1× TBS buffer and incubated with the membranes for approximately 3 h at room temperature on a rotter at slow speed. Rabbit anti-histone H3 antibody (ab1791; Abcam, Cambridge, MA, USA) was diluted 1:10,000 in 1× TBS buffer and used as a nuclear loading control. Rabbit anti-H3K4me3 antibody (39016, ActiveMotif, Carlsbad, CA, USA) was diluted 1:50,000 in 1× TBS buffer and used as a nuclear loading control. Rabbit anti-H3K27me3 antibody (07-449; Upstate-Millipore, Lake Placid, NY, USA) was diluted 1:5,000 in 1× TBS, pH 7.5. After 3 h, the blots were washed with 1× TBS, pH 7.5, for 30 min, exchanging the buffer every 10 min. Horseradish peroxidase-conjugated goat anti-rabbit secondary antibody (111-035-003; Jackson ImmunoResearch, West Grove, PA, USA) was diluted 1:10,000 in 1× TBS, pH 7.5, and 10 ml of this solution was added to each blot, which was incubated on a rotter for 3 h at room temperature. The blots were washed with 1× TBST buffer (10 mM Tris, pH 7.5, 100 mM NaCl, 0.1% [wt/vol] Tween 20) for 30 min, exchanging the buffer every 10 min. A volume of 300 μl of Crescento horseradish peroxidase reagent (Fisher Scientific, Waltham, MA, USA) was added to each blot, and the blots were imaged in a Bio-Rad Chemidoc imager under default chemiluminescence settings and autoadjusted exposure time. For the H3 blots, the positive loading control was masked during autoadjusted exposure to avoid overwhelming the sample signals.

**Helium ion microscopy.** The A. robustus-F. succinogenes strain UWB7 coculture was prepared as described above. The cell pellet, including the Avicel growth substrate, was harvested and suspended in phosphate-buffered saline (PBS, pH 7.5) in a 15-ml Falcon tube, to which glutaraldehyde (Sigma-Aldrich, St. Louis, MO) was added to a final concentration of 2 vol%. The tubes were incubated at room temperature for 1 h on a rotator. The tubes were then centrifuged at 700 × g for 10 min at 4°C, and the buffer was removed. The pellet was resuspended in 10 ml of 25 vol% ethanol and incubated for another hour. This process of suspension, incubation, and centrifugation was repeated for a stepwise ethanol dehydration series for 1 h on a rotator. The tubes were finally resuspended in 700 μl of 100% ethanol. The pellets were then dried via critical point drying with an Autosamdri-815 (Tousimis, Rockville, MD) and carbon dioxide gas. The samples were then coated with gold in a Hummer-2 (HUM-2000; Tokyo Electron) sputter coater at 200 mA and imaged on an FEI Helios Nanolab 600i DualBeam microscope (Carl Zeiss Microscopy, Peabody, MA).

**Data availability.** Supplementary data sets are available at the following Github repository: https://github.com/cswift3/anaerobic_fungi_Fibrobacter_co-culture. All sequencing reads have been deposited in the Sequencing Read Archive (SRA) and are associated with NCBI BioProject PRJNA666900. The raw mass spectrometry data were deposited on the MassIVE public repository (MSV000086033). Raw data are also available for download at https://genome.jgi.doe.gov/portal/ under the JGI Project ID 1294405. The molecular networking GNPS job can be publicly accessed at https://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=aeeb3b1a8fac4b67b54b6f1171a3053f. The ProteoSAFE job may be viewed and cloned from https://proteomics2.ucsd.edu/ProteoSAFe/status.jsp?task=e49148e8624c4e4cb0c3f0e09918ab6c.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**TEXT S1**, DOCX file, 0.02 MB.

**FIG S1**, DOCX file, 0.2 MB.

**FIG S2**, DOCX file, 0.6 MB.

**FIG S3**, DOCX file, 0.2 MB.

**FIG S4**, DOCX file, 0.2 MB.

**FIG S5**, DOCX file, 0.1 MB.

**TABLE S1**, DOCX file, 0.02 MB.

**TABLE S2**, DOCX file, 0.01 MB.

**TABLE S3**, DOCX file, 0.02 MB.

**TABLE S4**, DOCX file, 0.01 MB.

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