Saccharomyces cerevisiae Requires CFF1 To Produce 4-Hydroxy-5-Methylfuran-3(2H)-One, a Mimic of the Bacterial Quorum-Sensing Autoinducer AI-2

Julie S. Valastyan,a,b Christina M. Kraml,c Istvan Pelczer,d Thomas Ferrante,a Bonnie L. Basslera,b

aDepartment of Molecular Biology, Princeton University, Princeton, New Jersey, USA
bHoward Hughes Medical Institute, Chevy Chase, Maryland, USA
cLotus Separations LLC, Department of Chemistry, Princeton University, Princeton, New Jersey, USA
dDepartment of Chemistry, Princeton University, Princeton, New Jersey, USA

ABSTRACT Quorum sensing is a process of cell-to-cell communication that bacteria use to orchestrate collective behaviors. Quorum sensing depends on the production, release, and detection of extracellular signal molecules called autoinducers (AIs) that accumulate with increasing cell density. While most AIs are species specific, the AI called AI-2 is produced and detected by diverse bacterial species, and it mediates interspecies communication. We recently reported that mammalian cells produce an AI-2 mimic that can be detected by bacteria through the AI-2 receptor LuxP, potentially expanding the role of the AI-2 system to interdomain communication. Here, we describe a second molecule capable of interdomain signaling through LuxP, 4-hydroxy-5-methylfuran-3(2H)-one (MHF), that is produced by the yeast Saccharomyces cerevisiae. Screening the S. cerevisiae deletion collection revealed Cff1p, a protein with no known role, to be required for MHF production. Cff1p is proposed to be an enzyme, with structural similarity to sugar isomerases and epimerases, and substitution at the putative catalytic residue eliminated MHF production in S. cerevisiae. Sequence analysis uncovered Cff1p homologs in many species, primarily bacterial and fungal, but also viral, archaeal, and higher eukaryotic. Cff1p homologs from organisms from all domains can complement a cff1Δ S. cerevisiae mutant and restore MHF production. In all cases tested, the identified catalytic residue is conserved and required for MHF to be produced. These findings increase the scope of possibilities for interdomain interactions via AI-2 and AI-2 mimics, highlighting the breadth of molecules and organisms that could participate in quorum sensing.

IMPORTANCE Quorum sensing is a cell-to-cell communication process that bacteria use to monitor local population density. Quorum sensing relies on extracellular signal molecules called autoinducers (AIs). One AI called AI-2 is broadly made by bacteria and used for interspecies communication. Here, we describe a eukaryotic AI-2 mimic, 4-hydroxy-5-methylfuran-3(2H)-one (MHF), that is produced by the yeast Saccharomyces cerevisiae, and we identify the Cff1p protein as essential for MHF production. Hundreds of viral, archaeal, bacterial, and eukaryotic organisms possess Cff1p homologs. This finding, combined with our results showing that homologs from all domains can replace S. cerevisiae Cff1p, suggests that like AI-2, MHF is widely produced. Our results expand the breadth of organisms that may participate in quorum-sensing-mediated interactions.

KEYWORDS autoinducer, collective behavior, interspecies, quorum sensing, signaling, yeast

Bacteria use chemical communication to gauge local cell population density. This process, called quorum sensing, relies on the production, release, accumulation,
**Figure 1** *S. cerevisiae* produces MHF, an AI-2 mimic. (A) Diagram showing the structure of DPD and relevant interconversions among molecules. (B) Light output by the *V. harveyi* TL-26 reporter strain in response to *S. cerevisiae* culture fluids containing yeast AI-2 mimic in PBS (squares) and in water (triangles). (C) Chromatogram depicting fractionation of yeast AI-2 mimic preparations. The area containing the active fraction is enlarged in the inset. The chromatograms show absorption at 214 (green), 254 (blue), and 280 (red) nm. The arrow depicts the peak containing the activity. mAU, milli-absorbance units. (D) Light output from the *V. harveyi* TL-26 reporter strain in response to a titration of the active 8- to 9-min fraction from C. (E) Structure of MHF. RLU denotes relative light units, which are bioluminescence/OD600 of the reporter strain, and the dotted line-labeled Max AI-2 refers to the activity from 125 nM DPD. In B and D, error bars represent standard deviations of biological replicates, \( n = 3 \).
and detection of extracellular signal molecules called autoinducers (AIs) (for recent reviews, see references 1–3). Quorum sensing enables bacteria to assess whether they are at a low or high cell density and, if the latter, engage in collective behaviors that, to be successful, require many cells acting in synchrony. For example, quorum sensing controls traits such as bioluminescence, biofilm formation, and virulence factor production.

The bioluminescent marine bacterium *Vibrio harveyi* is a model organism used to study quorum sensing. *V. harveyi* employs three AIs, namely, AI-1, CAI-1, and AI-2, that enable intraspecies, intragenera, and interspecies communication, respectively (4–7). Germane to this report is that AI-2 is bound by the receptor LuxP, and LuxP-AI-2 binding initiates a signal transduction cascade, the output of which is bioluminescence (5, 8, 9). AI-2 is produced and detected by diverse bacterial species (6, 10–12).

Furthermore, human epithelial cells secrete an AI-2 mimic (here designated “mammalian AI-2 mimic”) of unknown structure that can be detected by LuxP, suggesting that the AI-2 signaling pathway could underpin interdomain communication (13).

Regarding AI-2 biosynthesis, the AI-2 precursor 4,5-dihydroxy-2,3-pentanedione (DPD) (Fig. 1A) is produced by the LuxS synthase from S-ribosylhomocysteine (SRH), a metabolic intermediate in S-adenosylmethionine (SAM)-dependent methylation pathways (6). DPD, the precursor to all AI-2 moieties, rapidly interconverts between different forms, and these rearranged structures can show preferences for binding to a particular bacterial quorum-sensing receptor. To activate the vibrio LuxP receptor, DPD must cyclize and coordinate borate to form the active AI-2 moiety (25,4S)-2-methyl-2,3,3,4-tetrahydroxytetrahydrofuran-borate (S-THMF-borate) (Fig. 1A) (8). The marine environment is borate rich, favoring formation of this final, borated signal molecule employed by *V. harveyi* and other vibrios. In boron-limited terrestrial environments, DPD rearranges to form (2R,4S)-2-methyl-2,3,3,4-tetrahydroxytetrahydrofuran (R-THMF), the active AI-2 moiety detected by enteric bacteria via a LuxP homolog called LsrB (Fig. 1A) (14). Another family of receptors, of which each contains a pCACHE domain, was recently discovered, expanding AI-2 detection mechanisms and the breadth of bacterial species that apparently respond to AI-2 (15).

Fungi also rely on quorum sensing to control behavior. In *Saccharomyces cerevisiae*, production of phenylethanol and tryptophol drives filamentous growth via activation of expression of *FLO11* encoding a glycoprotein required for flocculation and biofilm formation (16). *Candida albicans* uses two quorum-sensing molecules to regulate the transition from yeast to filamentous growth; tyrosol promotes the development of germ tubes required for hyphal growth, while farnesol inhibits this transition (17–20). Farnesol also mediates interdomain interactions by preventing toxin production by the bacterial pathogen *Pseudomonas aeruginosa* when in a mixed population with *C. albicans* (21, 22). Finally, farnesol is reported to modulate the human immune response in a *C. albicans* infection model (23–25). These findings hint at chemically mediated interdomain communication between fungi and other prokaryotic and eukaryotic organisms.

Here, we report a new interdomain quorum-sensing interaction. *S. cerevisiae* produces 4-hydroxy-5-methylfuran-3(2H)-one (MHF), a compound that mimics bacterial AI-2. Using *V. harveyi* as a reporter of AI-2 activity, we show that detection of and response to MHF require the LuxP receptor and signal transduction through the canonical *V. harveyi* quorum-sensing pathway. Screening of the *S. cerevisiae* deletion library revealed *CFF1* as a gene essential for MHF production. Cff1p, the protein encoded by the *CFF1* gene, is uncharacterized, but its crystal structure shows homology to sugar epimerases and isomerases (26). Mutation of a predicted catalytic residue, glutamic acid at position 44, eliminates MHF production by *S. cerevisiae*, suggesting that Cff1p may function as the MHF synthase. Many putative *CFF1* homologs exist in viral, archaean, bacterial, fungal, and higher organismal genomes, and in the majority of cases we tested, the *CFF1* genes could complement a *cff1Δ* *S. cerevisiae* mutant and restore MHF production. Alignment of Cff1p homologs showed that the key glutamic acid residue is
conserved, and in our test cases, it is required for activity. In summary, MHF has the ability to mimic Al-2, and MHF production may be prevalent in both prokaryotic and eukaryotic organisms. These findings highlight the expanding possibilities for interdomain signaling through Al-2 quorum-sensing pathways.

RESULTS

Purification and identification of MHF as an AI-2 mimic produced by S. cerevisiae. We previously reported that human tissue culture cells of epithelial origin, when starved or subjected to tight junction disruption, produce a mimic of the bacterial quorum-sensing AI called Al-2 (13). These earlier findings inspired us to examine whether other eukaryotes produce Al-2 mimics. Here, we focused on the yeast S. cerevisiae for two reasons. First, evolutionarily, S. cerevisiae and humans diverged ~1 billion years ago (27), possibly yielding insight into whether Al-2 mimic production does or does not occur widely across eukaryotes. Second, S. cerevisiae can be easily cultured, grows in virtually unlimited quantities, and survives in water, which are features predicted to accelerate purification and identification of interesting compounds (28).

We first tested whether S. cerevisiae makes a molecule that can mimic Al-2. S. cerevisiae MY8092 (hereafter called S. cerevisiae) was grown in synthetic defined (SD) medium with 2% glucose as the carbon source. Following 48 h of incubation, cell-free culture fluids were prepared and assessed for an activity capable of inducing light production in the V. harveyi AI-2 reporter strain called TL-26 (29). V. harveyi TL-26 produces maximum light in response to supplementation with 125 nM pure Al-2 (S-THMF-borate) (see Fig. S1A, dotted line, in the supplemental material). High-level activity was present in the S. cerevisiae cell-free culture fluids, suggesting that S. cerevisiae produces an Al-2 mimic (Fig. S1A).

Based on our finding that human epithelial cells produce the mammalian Al-2 mimic when starved in PBS, and again with the goal of facilitating purification, we next assessed Al-2 mimic production under starvation conditions. S. cerevisiae was grown to saturation in rich medium, washed twice, resuspended in either water or PBS, and incubated overnight at 30°C. Al-2 mimic activity was present in the collected fluids following resuspension of S. cerevisiae in both PBS (Fig. 1B, squares) and water (Fig. 1B, triangles). Addition of ~25% (vol/vol) of the preparation made in water was toxic to V. harveyi TL-26 (Fig. S1B). Toxicity is due to V. harveyi sensitivity to low salt, as mimic fractions supplemented with NaCl were not toxic (Fig. S1C).

To discover whether Al-2 mimic production occurred broadly among wild yeasts or was restricted to laboratory S. cerevisiae, a panel of wild S. cerevisiae isolates obtained from different environments ranging from clinical settings to vineyards was tested for production of activity using the V. harveyi TL-26 reporter (Fig. S1D) (30, 31). All the production profiles mirrored that of laboratory S. cerevisiae, suggesting that the Al-2 mimic is broadly made by S. cerevisiae strains.

To garner sufficient yeast Al-2 mimic for structural analysis, we tested the limit to which we could concentrate the activity. At the final step of the above preparation procedure, the washed S. cerevisiae cells were resuspended in water at different cell densities, from an optical density at 600 nm (OD600) of 1 to 128. Following an overnight incubation, cell-free culture fluids were analyzed for activation of light production in V. harveyi TL-26. Yeast Al-2 mimic activity increased with increasing S. cerevisiae cell density (Fig. S1E). Moreover, the activity was specific to the Al-2 quorum-sensing pathway, as light production was not induced by the preparations when supplied to a V. harveyi reporter strain (TL-25) that is incapable of detecting Al-2 but, rather, responds exclusively to the V. harveyi quorum-sensing Al called 3-hydroxy-C4-homoserine lactone (Al-1) (29, 32) (Fig. S1F).
To identify the chemical structure of the yeast AI-2 mimic, washed *S. cerevisiae* cells were resuspended in water at an OD<sub>600</sub> of 100. Following an overnight incubation, the cell-free fluids were collected and concentrated by lyophilization. Activity-guided HPLC fractionation on a Luna C<sub>18</sub> reverse-phase column revealed one peak at 8.3 min that exhibited absorption at 254 nm (blue trace, Fig. 1C, arrow and inset) and 280 nm (red trace, Fig. 1C, arrow and inset). The material did not absorb significantly at 214 nm (green trace, Fig. 1C, arrow and inset). The peak contained high levels of yeast AI-2 mimic activity, as judged by the *V. harveyi* TL-26 reporter strain (Fig. 1D). We pooled this peak from multiple such column runs and prepared the sample for nuclear magnetic resonance (NMR) and mass spectral analyses as described in the Materials and Methods.

Identification of the bioactive molecule relied on a comparison of results from liquid chromatography-mass spectrometry (LC-MS), NMR, and gas chromatography (GC)-MS. LC-MS analysis showed that two components were present in the active fraction. From our initial HPLC fractionation, these components correspond to the peak with absorption at 254 nm and 280 nm (the active peak) and the peak with absorption at 214 nm (an inactive contaminant), which could not be completely separated for LC-MS. The bioactive component had an exact mass of 115.039 (M<sup>+</sup>H) and a putative molecular formula of C<sub>5</sub>H<sub>6</sub>O<sub>3</sub>. These data, combined with an analysis of key peaks in the <sup>13</sup>C NMR spectra (signals at δ 194, 172, and 134 ppm), led us to consider structures A to D (see Fig. S2A in the supplemental material). Definitive evidence for structure A was obtained by GC-MS analysis, including matching of the fragmentation pattern of the active component against a database of known structures (Fig. S2B). The yeast AI-2 mimic structure A was identified as MHF (Fig. 1E, Fig. S2A). Indeed, a comparison of mass spectral, NMR, and HPLC analytical data confirmed that MHF purified from *S. cerevisiae* was identical to an authentic commercial sample of MHF (Fig. S2C and D).

**The mammalian AI-2 mimic is not MHF.** With the MHF structure in hand, we could investigate whether the *S. cerevisiae* and the previously reported mammalian AI-2 mimic are identical or not. As noted earlier, the mammalian AI-2 mimic has not been identified, so we did not have purified compound (13). Rather, we made a preparation from Caco-2 cells containing high-level mammalian AI-2 mimic activity in PBS (13). To determine the elution pattern for MHF in such Caco-2 cell preparations, we spiked commercial MHF into the mammalian AI-2 mimic preparation prior to HPLC fractionation. MHF eluted at 14 min (see Fig. S3A, arrow, in the supplemental material) in the context of Caco-2 culture fluids. Samples from Caco-2 cells that had not been spiked did not have a peak at the expected elution time for MHF (Fig. S3B, arrow). To eliminate the possibility that MHF was present in the Caco-2 cell preparations but at a level below the UV detection limit on the HPLC instrument, we tested all of the fractions for activity in the *V. harveyi* TL-26 reporter assay. While the reporter assay showed that the mammalian AI-2 mimic was indeed present in the non-MHF-spiked Caco-2 preparations (Fig. S3C, black), there was no activity in the 12- to 14-min HPLC fraction (Fig. S3C, red). Collectively, these data demonstrate that the mammalian AI-2 mimic is not MHF. In future studies, we will focus on identification of the mammalian AI-2 mimic.

MHF agonizes LuxP with a Nanomolar EC<sub>50</sub>. We next assessed the quantity of MHF present in *S. cerevisiae* cell-free fluids. To do this, we grew *S. cerevisiae*, pelleted and washed the cells, and then resuspended the cells at three different cell densities in water. Following overnight incubation, we removed the cells and compared the activities in each of the cell-free fluids to known quantities of the commercial MHF standard. We used two different MHF quantitation methods (Fig. 2A). First, different concentrations of commercial MHF were assessed by HPLC, and the areas under the MHF peaks were used to generate a standard curve. The *S. cerevisiae* preparations were likewise subjected to HPLC analysis, and the amount of MHF present in each sample was calculated by interpolating the area under the HPLC peak to that from the standard curve (Fig. 2A, black bars). Second, commercial MHF was assessed in the *V. harveyi* TL-26 reporter assay at different concentrations to generate an activity-based standard
curve. The *S. cerevisiae* cell-free fluids were identically assayed, and the MHF concentration in each preparation was estimated from the activity standard curve (Fig. 2A, white bars). The concentrations of MHF in the preparations calculated by the two methods were in close agreement. Assuming MHF production has a linear relationship with OD$_{600}$ values, we can use our data to estimate that *S. cerevisiae* produced 1.2 $\pm$ 0.4 mM MHF per OD$_{600}$ of cells. In the context of detection by the *V. harveyi* quorum-sensing apparatus, the 50% effective concentration (EC$_{50}$) for AI-2 is 3 nM and that for MHF is 300 nM (Fig. 2B). Thus, while both compounds exert activity in this system within the concentration range reported for bacterial AIs (33–36), the LuxP receptor prefers AI-2 over MHF.

**Identification of CFF1 as an *S. cerevisiae* gene essential for MHF production.** To identify the component(s) responsible for MHF production in *S. cerevisiae*, we screened the yeast deletion library for an *S. cerevisiae* mutant that was defective in MHF production (37–39). As described in the Materials and Methods, cell-free fluid preparations were made from >5,000 *S. cerevisiae* mutants and incubated with the *V. harveyi* TL-26 reporter strain. Bioluminescence was measured to assess the ability of each *S. cerevisiae* mutant to make MHF (Fig. 3A). Mutants were identified that elicited at least two standard deviations less light from the reporter strain than the mean amount of light production elicited from all strains (Fig. 3A). Eight putative mutants were retested for the ability to make activity (Fig. 3B). Two mutants, namely, *cff1*Δ and *rps1b*Δ, failed to activate the reporter strain. Cff1p (systematic name, YML079wp) is a cupin superfamily protein (26). Rps1bp (systematic name, YML063wp) is a component of the 40S ribosomal subunit (40). The six other potential mutants proved to have been false positives upon reassessment (Fig. 3B).

To verify the phenotypes of the mutants, clean deletions of *CFF1* and *RPS1B* were constructed in *S. cerevisiae*. The *cff1*Δ mutant displayed no defect in growth rate (see Fig. S4A, circles, in the supplemental material; compare to wild-type [WT] growth shown by the squares). As previously reported (41), the *rps1b*Δ mutant had a growth defect (Fig. S4A, triangles). Neither mutant exhibited sensitivity to overnight incubation in water (Fig. S4B). Importantly, culture fluids prepared from the clean *rps1b*Δ mutant produced nearly the wild-type level of AI-2 mimic activity, as determined by the ability to induce light production in the *V. harveyi* TL-26 reporter (Fig. 3C, triangles). In contrast, preparations made from the *cff1*Δ mutant had no activity (Fig. 3C, circles). PCR analysis revealed that the mutant annotated as *rps1b*Δ in the yeast deletion library, in fact, possesses a deletion in *CFF1*, explaining its inability to stimulate the reporter strain as well as the ability of our newly constructed *rps1b*Δ mutant to produce activity. Thus, *CFF1* is the only gene revealed by our screen to be required for production of the activity we are monitoring.
To confirm that the yeast AI-2 mimic activity produced by the protein encoded by CFF1 is MHF, we prepared and fractionated cell-free culture fluids from the cff1Δ strain using the identical procedure we used for isolation of MHF from wild-type S. cerevisiae. No MHF peak could be detected in the cff1Δ mutant preparation (Fig. 3D; compare to Fig. 1C, inset). Consistent with this finding, the relevant HPLC column fraction had no activity in the V. harveyi TL-26 reporter assay (Fig. S4C). These data suggest that Cff1p has a required role in MHF biosynthesis in yeast.
Our finding that Cff1p is required for MHF production is surprising. The presence of MHF in fermented food products made from *S. cerevisiae* has been reported; however, the suggested route to MHF is either spontaneous starting from D-ribulose-5-phosphate (42–45) or under extreme conditions, via the Maillard reaction (46, 47). Quite to the contrary, our data suggest that MHF production in *S. cerevisiae* is enzyme catalyzed and under physiological conditions. Cff1p has not been characterized. However, there does exist a crystal structure (26). It shows a putative ligand binding pocket containing amino acid residues identical to those required for catalysis by epimerases and isomerases that share the cupin fold (48). Specifically, the conserved E44 residue is proposed to have a catalytic role. We made an E44A substitution in Cff1p and assayed the mutant protein for MHF production. The substitution did not alter Cff1p stability as judged by visualization of a fused HALO tag (Fig. S4D); however, culture fluids prepared from the *S. cerevisiae* Cff1p-E44A mutant elicited 100-fold less light from the *V. harveyi* TL-26 reporter strain than preparations from WT *S. cerevisiae* (i.e., less than 1% activity remained) (Fig. 3E), showing that the glutamate residue at position 44 is key for the presumptive enzymatic activity that generates MHF.

**CFF1 homologs exist in organisms from all domains.** Cff1p has structural similarity to sugar isomerases and epimerases, and the Cff1p amino acid sequence is similar to proteins of unknown function (26). Recently, Tourneroche et al. reported 12 wild fungal species that exist as endomicrobiota of kelp and that possess AI-2 activity, as judged by a *V. harveyi* reporter system analogous to the one we use here (49). The molecule(s) responsible for the AI-2 activity have not been identified. We wondered whether these fungal species might make MHF. Examination of their proteomes revealed Cff1p homologs in 2 of the 12 species, namely, *Trametes versicolor* and *Botrytis cinerea*, with approximately 50% similarity and 35% identity, respectively, to *S. cerevisiae* Cff1p at the amino acid sequence level (see Fig. S5A in the supplemental material). Both of these species’ Cff1p homologs have high conservation in the putative ligand binding domain, and they each possess a residue equivalent to E44 in *S. cerevisiae* Cff1p (Fig. S5A, arrow). To test for function, we cloned these two genes and introduced them into our cff1Δ *S. cerevisiae* mutant under the control of the native *S. cerevisiae* CFF1 promoter. Unlike the cff1Δ *S. cerevisiae* mutant that produced no MHF, the mutant carrying each homolog produced activity sufficient to induce maximal light production in the *V. harveyi* TL-26 reporter strain (Fig. 4A). The E38A and E30A substitutions in the *B. cinerea* and *T. versicolor* Cff1p proteins (equivalent to E44A in *S. cerevisiae* Cff1p), respectively, eliminated production of the activity (Fig. 4B). In both cases, the WT and mutant proteins were made at approximately the same levels and were equally stable (Fig. S5B). HPLC fractionation confirmed that MHF was indeed produced in cff1Δ *S. cerevisiae* carrying the WT CFF1 homologs, and no MHF could be detected in the cases in which mutant CFF1 alleles were present (see Fig. S6 in the supplemental material). Thus, both the *T. versicolor* and *B. cinerea* CFF1 genes can complement the cff1Δ *S. cerevisiae* defect and restore MHF production, and a glutamate at a position equivalent to 44 in *S. cerevisiae* Cff1p is required. In the cases of the other 10 fungi that Tourneroche et al. reported to possess AI-2 activity (49), we do not know whether they make a different active molecule or, alternatively, if they possess Cff1p proteins that are unrecognizable through the database search we performed.

A search of the nonredundant protein sequence database (50) for proteins with homology to *S. cerevisiae*, *T. versicolor*, and *B. cinerea* Cff1p uncovered three additional *Saccharomyces* species possessing proteins with an average identity of 90% to Cff1p. More broadly, we identified 410 non-*Saccharomyces* fungal species possessing proteins harboring 26% to 71% identity to Cff1p. Included were members of the Ascomycota phylum, such as *Aspergillus fumigatus* and *Neurospora crassa*, and the Basidiomycota phyla, such as *Cryptococcus neoformans*. We also identified >350 prokaryotes possessing putative proteins with 25% to 45% identity to Cff1p. These species exist in 11 phyla and include multiple pseudomonads, staphylococci, and bacilli. Our search also uncovered 25 other organisms, spanning all domains, with potential Cff1p homologs. The
majority of these organisms exist in the marine environment, for example, the acorn worm *Saccoglossus kowalevskii*, the green alga *Ostreococcus lucimarinus*, and the archaeon *Methanohalophilus halophilus* (Fig. 4C; see Fig. S7 in the supplemental material). At present, we do not know whether this representation indicates a dominant biological function in the marine niche or if it stems from biased sampling. Notably, E44 is conserved in 97% of these potential homologs (Fig. 4C, arrow).
It is intriguing that bacterial species may possess cff genes (note that we designate the bacterial genes and proteins as cff and Cff, respectively, and the fungal genes and proteins CFF1 and Cff1p, respectively). Thousands of bacterial species are known to synthesize the interspecies quorum-sensing AI AI-2 via possession of luxS encoding the AI-2 synthase (10). To investigate whether bacteria could potentially make both MHF and AI-2, we performed database analyses. According to the nonredundant protein sequences database (50), only ~20% of bacterial species possessing a cff homolog also harbor luxS (Fig. 5A). The majority of bacteria that possess both luxS and cff genes belong to the Firmicutes phylum (Fig. 5B), a phylum considered ancestral to other bacterial phyla. This pattern suggests that a shared ancestor possessed both genes and that, generally, the species maintained only one of the two genes as they diverged.

To test whether additional Cff and Cff1p homologs are functional, we selected 10 additional organisms, namely, four prokaryotes (Pseudomonas chlororaphis, Umezawaea tangerina, Staphylococcus aureus, and Algoriphagus boritolerans), two fungi (A. fumigatus and C. neoformans), and four organisms from other kingdoms (O. lucimarinus, S. kowalevskii, M. euhalobius, and Achlya hypogyna) that possess putative Cff or Cff1p proteins (Table 1). Our choices included clinically relevant organisms (i.e., S. aureus and C. neoformans) and marine organisms (i.e., O. lucimarinus and S. kowalevskii) that may coexist with bacteria that use AI-2-LuxP-mediated quorum sensing. Additionally, we tested proteins from organisms representing unique phyla and proteins with various levels of amino acid identity relative to S. cerevisiae Cff1p (Table 1). To assay activity, we expressed...
the candidate genes in our cff1Δ S. cerevisiae strain under the control of the endogenous S. cerevisiae CFF1 promoter. All of the homologs except those from U. tangerina and O. lucimarinus complemented the loss of Cff1p in S. cerevisiae, driving sufficient MHF production to induce light production in the V. harveyi TL-26 reporter strain (Fig. 5C). This result suggests that these, and likely many other, organisms harbor the potential to synthesize MHF.

Compared to BLASTp, Interpro (51) provides a dramatically larger set of proteins under the class “uncharacterized protein, YML079W-like” (i.e., the S. cerevisiae systematic name for Cff1p). This group contains proteins from 3,000 different organisms, including 2,400 species of bacteria. Notably, 95% of these putative Cff homologs have the conserved glutamate at the position corresponding to 44 in the S. cerevisiae Cff1p. Thus, the Interpro data suggest that, potentially, the number of bacterial species containing cff is on the same scale as those possessing luxS, as Interpro lists ~2,500 different species (primarily bacterial) that contain luxS. In this data set, ~15% of the bacterial species contain both luxS and cff.

We were especially intrigued that the Interpro database search revealed a virus with a potential Cff homolog, Pandoravirus salinus. Using the above strategy, we tested the functionality of the P. salinus Cff homolog and found that, indeed, the viral cff gene complements the loss of CFF1 in S. cerevisiae (Fig. 5C). This finding provides initial validation for the output of the larger Interpro data set and presages the existence of many additional functional Cff1p homologs. Moreover, this final result, coupled with the other findings here, shows that MHF production could occur across the archaeal, bacterial, and eukaryotic domains, as well as among viruses.

**DISCUSSION**

Quorum sensing is the process by which bacteria monitor their local cell population density and determine when it is appropriate to engage in collective behaviors (1–3). Bacteria often employ multiple AIs, encoding distinct information about species relatedness, which presumably enables them to take a census of “self” and “other.” The AI-2-LuxP quorum-sensing pathway is proposed to be used for the latter, to monitor the total cell density of the vicinal community. Previously, we showed that mammalian cells can make a mimic of AI-2 that activates quorum sensing via LuxP, providing a founding example of interdomain quorum-sensing-mediated communication (13). Here, we show that S. cerevisiae, another eukaryotic organism, makes MHF, which can
induce quorum-sensing behavior through the canonical AI-2 pathway. Our preliminary studies suggest that MHF production may be widespread across domains. Presumably, the ability of MHF to substitute for AI-2 enables bacteria that possess LuxP to detect the presence of other bacterial cells, higher organisms, and possibly viruses in the environment.

The mammalian AI-2 mimic is distinct from MHF, suggesting that multiple compounds might be exploited for AI-2-like interactions between eukaryotes and bacteria. Plants, insects, and fungi have previously been shown to produce MHF (42–44, 52–54), and based on the findings presented here, it is possible that MHF could be widely used for cross-domain interactions between bacteria and fungi. While future studies are necessary to understand the ecological significance of MHF in such presumptive interdomain interactions and, more specifically, what, if any, traits are controlled by MHF in organisms that produce MHF, it is already known that cross-communication between eukaryotes and prokaryotes can shape each participant’s biology. For example, quorum-sensing pathways drive behavior across domains in both mutualistic and parasitic relationships. Examples include communication between the gut and the microbiome (55), competition between C. albicans and P. aeruginosa during infection of the lung (56), and the ability of Legionella pneumophila to alter eukaryotic host cell migration (57). Beyond bacteria and eukaryotes, recent work demonstrates that quorum-sensing-mediated cross-communication occurs between bacteria and phages (58–60). Specifically, an AI called 3,5-dimethylpyrazin-2-ol (DPO) that is produced by many species of bacteria is detected by a phage that uses the information encoded in DPO to determine whether to enter the lytic or the lysogenic state (59). Our present work provides evidence for a virus with the potential to make MHF. Presumably, interdomain chemical interactions could enable sharing of and cheating on public goods, could enhance symbiosis or predation, or could allow particular organisms to coestablish niches in otherwise inhospitable environments. Notably, bacteria that synthesize AI-2 generally use it to regulate their own quorum-sensing-controlled behaviors (61). We do not yet know if S. cerevisiae and other eukaryotes use MHF as a quorum-sensing signal. Our intention now is to perform transcriptome sequencing (RNA-seq) on cff1Δ S. cerevisiae in the presence and absence of exogenously supplied MHF to discover the endogenous response.

MHF is a volatile compound that is used as a flavorant. Food scientists have previously shown that low levels of MHF exist in fermented foods, such as soy sauce and malt (62–64). As alluded to above, MHF is hypothesized to form spontaneously from pentose sugars that undergo the Maillard reaction (i.e., during cooking) or as a byproduct (in fungi). Notably, MHF is also a breakdown product of DPD; however, S. cerevisiae lacks LuxS, so DPD is an unlikely source of MHF in fungi (44, 65). In the fungus Zygosaccharomyces rouxii, it is proposed that, following production of ribulose-5-phosphate by the pentose phosphate pathway, MHF forms spontaneously as a breakdown product (42). Here, we show that Cff1p, which is presumably an enzyme, is required for MHF production in S. cerevisiae. Z. rouxii contains a CFF1 homolog, which suggested to us that the pathway we discovered here could be relevant in this organism. Indeed, cell-free culture fluids from WT Z. rouxii possess activity capable of inducing light production in V. harveyi TL-26, while reporter fluids prepared from a Z. rouxii cff1D strain are devoid of that activity (see Fig. S8 in the supplemental material).

S. cerevisiae CFF1 is a gene of unknown function. The crystal structure of S. cerevisiae Cff1p has been solved and shows similarity to sugar epimerases and isomerases (26). The Cff1p protein, like other cupins, contains a jelly-roll fold similar to those in germin-and auxin-binding proteins. In Cff1p, the regions adjacent to the cupin motif are distinct from previously studied members of this protein superfamily. Cff1p has notable similarity to the Salmonella enterica serovar Typhimurium protein RmlC, which catalyzes the conversion of dTDP-6-deoxy-o-xylo-4-hexulose to dTDP-6-deoxy-L-lyxo-hexulose (66). The crystal structure of RmlC bound to a substrate analog has been compared with that of S. cerevisiae Cff1p and shows that Cff1p possesses a binding pocket that could accommodate both the nucleotide with which Cff1p was cocrystallized and a sugar moiety.
However, no sugar was present in the Cff1p crystal. The authors hypothesized that the pocket may bind a sugar nucleotide. Accordingly, the jelly-roll fold motif is shared with enzymes, such as phosphoglucose isomerase and dTDP-4-keto-6-deoxy-o-hexulose-3,5-epimerase (26). Given this relatedness and the fact that DPD, the nonborated precursor to AI-2, is a sugar, we suspect that Cff1p could be the synthase for MHF and that MHF is made from a sugar substrate.

Our study suggests that the scope of organisms that can participate in quorum sensing through AI-2-type pathways continues to increase, hinting that AI-2 quorum sensing mediates interspecies bacterial communication and interdomain communication between bacteria and eukaryotes and possibly viruses. Bacteria can distinguish among closely related quorum-sensing AIs, and they are capable of decoding and integrating the information contained in blends of AIs to drive appropriate behaviors based on the cell density and the species identities of neighboring bacteria. Cooccurrences of organisms from different domains have important ecological and medical implications (67–70). We highlight a few examples involving S. cerevisiae, the main focus of the present work; the presence of S. cerevisiae improves the ability of Pseudomonas putida to grow in glucose-containing medium (71), enhances the growth of lactic acid bacteria in nitrogen-rich environments (72), and stimulates the growth of multiple Acinetobacter species through the secretion of ethanol (73). Our future work will focus on how domain-spanning quorum-sensing cross-communication influences the behavior of the various participants and affects global community structures and their functioning.

MATERIALS AND METHODS

Strains, plasmids, and media. Strains and plasmids used in this work are provided in Table S1 and S2, respectively, in the supplemental material. YML079W (CFF1) and YML063W (RPS1B) were deleted from S. cerevisiae using the standard kanMX insertion technique (38, 74). Geneteticin (G418 sulfate) (Thermo Fisher Scientific) was added at 200 μg/ml for KanMX selection. S. cerevisiae was grown in yeast extract-peptone-dextrose (YPD) medium, unless specified otherwise (Thermo Fisher Scientific). V. harveyi was grown in Luria-Marine (LM) medium, and AI-2 reporter assays were performed in autoinducer bioassay (AB) medium (29). To generate pRS416-CFF1, CFF1 and ~150 bp upstream and downstream were cloned into the XhoI and XbaI (New England BioLabs) sites of pRS416 using standard cloning procedures. We tested two constructs, one containing the CFF1 gene and 500 bp of upstream and downstream DNA and one containing the CFF1 gene with upstream and downstream regions of ~150 bp and encompassing only intergenic sequences between CFF1 and its neighboring genes. Both constructs drove production of the same amount of MHF, suggesting that all of the necessary promoter and terminator elements for CFF1 are in the intergenic regions. Therefore, we used the construct containing CFF1 and only the intergenic regions for the work reported here. DNA encoding the HALO sequence was fused to that encoding the 3′-terminating CFF1 using Gibson assembly (New England BioLabs) (75). DNA encoding CFF1 homologs was codon optimized and synthesized (Integrated DNA Technologies) before being inserted between the upstream and downstream regions of S. cerevisiae CFF1, cloned into pRS416 (76), and HALO tagged using the strategy outlined above or using Gibson assembly. Mutations in the constructs were generated by standard mutagenic PCR using PfuUltra polymerase (QuikChange II; Agilent Technologies).

Yeast, mammalian, and Z. rouxii Al-2 mimic production. S. cerevisiae cells were grown overnight in YPD or SD-ura (as needed for plasmid maintenance) with shaking at 30°C. The cells were pelleted by centrifugation for 10 min at 4,000 rpm. The pellet was washed twice with sterile water, followed by centrifugation as above. The cells were resuspended in water and incubated for 24 h at 30°C with shaking. For making yeast Al-2 mimic for molecule identification, the cells were resuspended at OD600 of 100 and incubated overnight at 30°C. When making yeast AI-2 mimic for all other assays, unless otherwise stated, the cells were resuspended at an OD600 of 10 and incubated overnight at 30°C. The cells were removed by centrifugation as above, and the resulting cell-free fluids were filtered through 0.22-μm polyethersulfone (PES) membranes (Millipore Sigma). Such preparations were used as the sources of yeast Al-2 mimic for activity assays and for further purification. The mammalian Al-2 mimic produced by Caco-2 cells was prepared as described previously (13). CFF1 was replaced with a KanMX cassette in Z. rouxii as previously described (77). Z. rouxii cell-free culture fluids were generated in water, as described above for S. cerevisiae.

V. harveyi TL-25/TL-26 reporter assays. Bioluminescence reporter assays using V. harveyi strains TL-25 and TL-26 were performed as previously reported (13, 78). Briefly, vibrio strains were grown overnight at 30°C in LM medium with shaking. V. harveyi cultures were diluted 1:1,000 in AB medium containing 100 μM boric acid. In all cases, cultures were aliquoted into wells of black-sided, clear-bottom 96-well plates (Corning). DPD, AI-1, MHF, or AI-2 mimic preparation was added at the indicated amounts, and the mixtures were serial diluted. Following incubation at 30°C with shaking for 6 h, bioluminescence and cell density (OD600) were measured using an Envision plate reader (PerkinElmer). Data are presented as relative light units (RLUs), which are bioluminescence per OD600. *Max AI-2* shown in figures indicates
bioluminescence output following addition of 1.25 nM AI-2 (i.e., 3-THMF-borate). ‘Max Al-1’ depicted in figures indicates bioluminescence output following addition of 500 nM Al-1 (i.e., 3-hydroxy C4-homoserine lactone).

Yeast growth curve and survival assays. S. cerevisiae strains were grown overnight at 30°C in YPD medium with shaking to saturation. The cultures were diluted to OD600 of 0.1 in a black-sided, clear-bottom 96-well plate. Cells were incubated at 30°C with shaking, and time points were taken every 20 min. Growth was measured in a Synergy plate reader (BioTek). For survival analyses, S. cerevisiae strains were diluted 1:100,000 and plated onto YPD agar plates. Colonies were allowed to grow for 48 h and then counted manually.

Screen for S. cerevisiae genes required for MHF production. The S. cerevisiae Yeast Knockout (YKO) Collection (Dharmacon) is a 96-well plate arrayed library containing about 5,000 S. cerevisiae strains with single gene deletions spanning the yeast genome (37–39). After thawing, 5 μl of each strain in the library was transferred into 96-well plates containing 150 μl YPD+G418 sulfate, and the plates were incubated at 30°C with shaking overnight. A total of 10 μl of each strain was diluted into 150-μl fresh YPD+G418 sulfate medium and allowed to grow for 5 h at 30°C with shaking. The cells were pelleted at 4,000 rpm for 10 min and then resuspended in water. The wash and centrifugation steps were repeated two more times. The cells were pelleted at 4,000 rpm for a final 10 min and resuspended in AB medium that had been supplemented with 100 μM boric acid. The initial OD600 of each culture was measured to identify any yeast mutants that had grown poorly. Yeast deletion strains that exhibited slowed growth were eliminated from analysis. The plates were incubated overnight at 30°C with shaking. The following morning, the plates were subjected to centrifugation at 4,000 rpm for 10 min. A total of 75 μl of culture fluid from each well was combined with 75 μl of fresh AB medium that had been inoculated with a 1:1,000 dilution of an overnight culture of the V. harveyi TL-26 reporter strain, and the mixture was incubated at 30°C with shaking overnight. The following 8 h of incubation at 30°C with aeration, bioluminescence and OD600 were measured (Envision Plate Reader). Normalized RLUs were calculated by dividing the bioluminescence by the OD600 and then dividing by the average RLUs from all plates from which measurements were made on a single day. Mutants from the screen that appeared to be defective in production of activity were retested individually, as above. Mutants of interest were reconstructed using the standard KanMX deletion method (38, 74) and again examined for the ability to produce activity.

Cff1p alignment and phylogenetic tree production. Potential Cff1p homologs were identified using BLASTp (50) and searching against S. cerevisiae Cff1p with an E value cutoff of 1e-5. This analysis returned 1,975 sequences. The list of sequences was culled to remove duplicate species resulting in 744 sequences. Rough phylogenetic analyses demonstrated that sequences from species within a genus clustered; thus, the list of sequences was further trimmed to include only the top hit in each genus, delivering 367 total sequences. These sequences were aligned using Clustal Omega (79) in SnapGene software (GSL Biotech). The phylogenetic tree in Fig. S7 was constructed in MEGA-X using the maximum likelihood method and Jones-Taylor-Thornton (JTT) matrix-based model with 500 bootstrap replications as described previously (80). The alignment was visualized using ggmsa in R, pruning the ends of the alignment to the first and last amino acids of S. cerevisiae Cff1p.

Yeast Al-2 mimic purification and identification. Approximately 250 ml of concentrated crude yeast Al-2 mimic preparation was filtered through a 0.45-μm polyvinylidene difluoride (PVDF) filter (Millipore Sigma) and separated on a 2- by 23-cm Luna C18 column (Phenomenex) using a mobile phase consisting of 5% water in methanol at a flow rate of 10 ml/min. The component of interest was eluted in 8 min in approximately 6 ml of mobile phase. The fraction was dried by rotary evaporation to remove methanol. To enable further concentration, the aqueous solution was saturated with sodium chloride and extracted with dichloromethane. The dichloromethane layer containing the product of interest was dried, and the sample was refrigerated. Fifty collections were processed in this manner. The products were combined and dissolved in 5 μl of deuterated water to a concentration of 0.003 mg/ml (determined subsequently by HPLC analysis using a standard curve). A “background” sample was generated by collecting the same volume of mobile phase in an area containing no UV-visible eluting components. Both samples were analyzed by 1H and 13C NMR spectroscopy, and although the data were inconclusive, several key signals were detected in the active sample that were not present in and not obscured by the background sample. For example, in the 13C NMR, signals at δ 194, 172, and 134 ppm were observed reproducibly. These data combined with mass spectrometry data led to a collection of possible candidates for the active component (Fig. S2A, Fig. S2A). The structure was confirmed by GC-MS analysis, including matching of the fragmentation pattern of the active component against a database of known structures. Analysis was carried out on an RTX-1ms (25 μm) 30-m column with 0.32-mm internal diameter (ID) (Restek). The samples were incubated at 40°C for 2 min and then subjected to a temperature ramp up of 10°C/min to 300°C, followed by a 5-min hold at 300°C. Mammalian Al-2 mimic activity was separated on a 25- by 0.46-cm Synergy 10-μm Polar-RP column using a mobile phase of 5% water in methanol at a flow rate of 0.5 ml/min.

Protein gels. To assess the levels of Cff1p, the Cff1p and Cff homologs, and the Cff1p and Cff variants, yeast cells producing the protein of interest were grown to exponential phase in SD-Ura. In each case, cells equivalent to 10 OD600 units were pelleted for 10 min at 4,000 rpm and resuspended in 50 μl yeast protein extraction reagent (Y-PER; Thermo Fisher Scientific) supplemented with 1× Halt protease inhibitor cocktail (Thermo Fisher Scientific). These mixtures were incubated at room temperature for 20 min with agitation and then subjected to centrifugation at 13,000 rpm in a tabletop centrifuge. The supernatants were collected and incubated with 1 μM HaloTag TMR ligand (Promega) at room temperature for 20 min. Next, 4× Laemmli sample buffer was added, and the mixtures were incubated at 70°C March/April 2021 Volume 12 Issue 2 e03303-20 mbio.asm.org

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for 10 min. Samples were loaded onto 4% to 20% gradient mini-protein precast gels (Bio-Rad). Following electrophoresis, proteins were visualized using the Cy3 filter set on an ImageQuant LAS 4000 instrument. To assess total protein, gels were stained with Coomassie brilliant blue R-250 staining solution (Bio-Rad) and visualized using an ImageQuant LAS 4000 instrument.

Data availability. All data and materials published in the manuscript are available upon request.

SUPPLEMENTAL MATERIAL
Supplemental material is available online only.
FIG S1, TIF file, 1.8 MB.
FIG S2, PDF file, 0.1 MB.
FIG S3, TIF file, 1.2 MB.
FIG S4, TIF file, 1 MB.
FIG S5, TIF file, 2.1 MB.
FIG S6, TIF file, 1.4 MB.
FIG S7, TIF file, 1.4 MB.
FIG S8, TIF file, 0.3 MB.
TABLE S1, DOCX file, 0.01 MB.
TABLE S2, DOCX file, 0.01 MB.

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