Differential response to prey quorum signals indicates predatory specialization of myxobacteria and ability to predate *Pseudomonas aeruginosa*

Shukria Akbar,1 Kayleigh E. Phillips,1 Sandeep K. Misra,1 Joshua S. Sharp1,2 and D. Cole Stevens 1*

1Department of BioMolecular Sciences, University of Mississippi, University, MS.
2Department of Chemistry and Biochemistry, University of Mississippi, University, University, MS.

Summary

Multiomic analysis of transcriptional and metabolic responses from the predatory myxobacteria *Myxococcus xanthus* and *Cystobacter ferrugineus* exposed to prey signalling molecules of the acylhomoserine lactone and quinolone quorum signalling classes provided insight into predatory specialization. Acylhomoserine lactone quorum signals elicited a general response from both myxobacteria. We suggest that this is likely due to the generalist predator lifestyles of myxobacteria and ubiquity of acylhomoserine lactone signals. We also provide data that indicates the core homoserine lactone moiety included in all acylhomoserine lactone scaffolds to be sufficient to induce this general response. Comparing both myxobacteria, unique transcriptional and metabolic responses were observed from *Cystobacter ferrugineus* exposed to the quinolone signal 2-heptylquinolin-4(1H)-one (HHQ) natively produced by *Pseudomonas aeruginosa*. We suggest that this unique response and ability to metabolize quinolone signals contribute to the superior predation of *P. aeruginosa* observed from *C. ferrugineus*. These results further demonstrate myxobacterial eavesdropping on prey signalling molecules and provide insight into how responses to exogenous signals might correlate with prey range of myxobacteria.

Introduction

The uniquely multicellular lifestyles of myxobacteria have motivated continued efforts to explore the myxobacterium *Myxococcus xanthus* as a model organism for cooperative behaviours including development (Islam et al., 2020; Sharma et al., 2021), motility (Mercier et al., 2020; Rendueles and Velicer, 2020; Zhang et al., 2020b), and predation (Thiery and Kaimer, 2020; Zhang et al., 2020a; Sydney et al., 2021). Often attributed to their need to acquire nutrients as generalist predators (Nair et al., 2019) and capacity to prey upon clinical pathogens (Livingstone et al., 2017), myxobacteria have also been a valuable resource for the discovery of novel specialized metabolites as potential therapeutic lead compounds (Herrmann et al., 2017; Baltz, 2019; Perez et al., 2020). The diversities in structural scaffolds and observed activities as well as the unique chemical space associated with myxobacterial metabolites when compared with more thoroughly explored Actinobacteria make myxobacteria excellent sources for efforts focused on the discovery of therapeutics (Herrmann et al., 2017; Baltz, 2019). However, the connection between myxobacterial predation and production of these biologically active metabolites remains underexplored. Currently, only the metabolites myxovirescin (Xiao et al., 2011; Ellis et al., 2019; Wang et al., 2019) and myxoprincomide (Cortina et al., 2012; Muller et al., 2016) have been directly implicated to be involved during *Myxococcus xanthus* predation of *Escherichia coli* and *Bacillus subtilis*, respectively. In fact, the chemical ecology of predator–prey interactions between myxobacteria and prey remains underexplored (Findlay, 2016). The predatory capacity or prey range of myxobacteria cannot be directly correlated with phylogeny (Livingstone et al., 2017; Arend et al., 2020). Presently, the best determinants for broadly assessing prey ranges are genetic features that might provide specific traits to overcome predation resistances of individual prey. For example, myxobacteria possessing the formaldehyde dismutase gene *fdm* demonstrated comparatively better predation of toxic formaldehyde secreting *Pseudomonas aeruginosa* an opportunistic pathogen observed to be somewhat
recalcitrant to myxobacterial predation (Sutton et al., 2019).

The recent observation that acylhomoserine lactone (AHL) quorum signals from prey microbes impact the predatory capacity of *M. xanthus* suggests that quorum signals might influence predator–prey interactions (Lloyd and Whitworth, 2017). Although two orphaned, functional AHL synthases have been reported, no myxobacteria have been observed to produce AHLs (Albataineh et al., 2021). However, a recent survey of signalling systems within the family Myxococcaceae reported the presence of conserved AHL receptor (LuxR) homologues and inferred that many myxobacteria within the two genera *Myxococcus* and *Corallococcus* are capable of sensing AHL signalling molecules (Whitworth and Zwarycz, 2020). While this suggests that predatory myxobacteria might eavesdrop on prey quorum signalling, the observed reaction from *M. xanthus* might also simply be the result of exogenous AHLs as a nutrient source. Herein we utilize a combination of transcriptomics and metabolomics to determine how myxobacterial responses to quorum signals produced by *P. aeruginosa* might indicate predatory capacity.

By exposing myxobacteria to structurally and functionally distinct classes of prey quorum signals comparing ubiquitous AHL signals and quinolone signals more unique to pseudomonads (Papenfort and Bassler, 2016), we anticipated that a differential response exclusive to a specific signal class would support predatory eavesdropping and perhaps correlate with improved predation of *P. aeruginosa*. For these experiments we exposed each myxobacterium to AHL signals (Galloway et al., 2011) as well as the quinolone signal 2-heptylquinolin-4(1H)-one (HHQ; Deziel et al., 2004; Dubern and Diggle, 2008; García-Reyes et al., 2020). Ubiquitous to Proteobacteria (notably excluding myxobacteria) and numerous other non-Proteobacteria genera, AHLs are the most common class of quorum signal autoinducers and are often implicated in interspecies communication within polymicrobial communities (Shiner et al., 2005; Mukherjee and Bassler, 2019). Also associated with the modulation of interspecies and interkingdom behaviours (Reen et al., 2011), the quinolone signal HHQ contributes to the pathogenicity of *P. aeruginosa* by participating in the regulation of various virulence factors (Dubern and Diggle, 2008; Reen et al., 2011). Exploration of the myxobacterial response to prey quorum signals not only provides insight into the impact of shared chemical signals might have on predator–prey interactions within bacterial communities but may also provide further genetic determinants that indicate predatory capacities of myxobacteria.

As a model organism for developmental studies, *M. xanthus* is the best characterized myxobacterium and has already demonstrated a behavioural response to exogenous AHLs (Lloyd and Whitworth, 2017). However, we suspected that routine use as a laboratory strain and well-explored specialized metabolism (Cortina et al., 2012; Hermann et al., 2017) of *M. xanthus* might diminish its viability as the sole myxobacterium for these experiments. Also of note, *M. xanthus* constitutively secretes lytic enzymes and specialized metabolites associated with prey lysis which may limit observable transcriptomic and metabolomic responses to exogenous quorum signals (Livingstone et al., 2018). Therefore, *Cystobacter ferrugineus* was also included as a more recent myxobacterial isolate with a less explored biosynthetic capacity and prey range (Akbar et al., 2017; Goes et al., 2020). Both *M. xanthus* and *C. ferrugineus* have an annotated solo LuxR-type AHL receptor present in their genomes (WP_011555271.1 and WP_071900454.1; Subramoni and Venturi, 2009; Tobias et al., 2020; Xu, 2020). However, homology-based annotation of these features only indicates the helix–turn–helix DNA-binding domain of LuxR receptors, and neither include an AHL-binding site motif (PF03472; Baikalov et al., 1996; Vannini et al., 2002; Mukherjee and Bassler, 2019).

Despite the absence of a canonical receptor, exogenous AHLs have been observed to stimulate the motility and predatory activity of *M. xanthus* (Lloyd and Whitworth, 2017). Also of note, neither *M. xanthus* nor *C. ferrugineus* possess a homologous PqsR-type HHQ receptor (Diggle et al., 2003; Wade et al., 2005). Utilizing a multilocus approach to assess the transcriptomic and metabolomic responses of *M. xanthus* and *C. ferrugineus* when exposed to AHL and quinolone signals, we sought to determine if structurally and functionally dissimilar quorum signals from prey elicit distinct responses from predatory myxobacteria that correlate with successful predation of *P. aeruginosa*.

**Results**

C6-AHL induces a general transcriptional response from both *M. xanthus* and *C. ferrugineus*

Exposure experiments utilizing a concentration of C6-AHL previously shown to elicit a predatory response from *M. xanthus* (9 μM) were conducted on plates of *M. xanthus* and *C. ferrugineus* (Lloyd and Whitworth, 2017). Exposure experiments were conducted in triplicate for both *M. xanthus* and *C. ferrugineus* with DMSO exposures serving as vehicle, negative controls for comparative analysis. Comparative transcriptomic analysis from RNAseq data revealed C6-AHL exposure impacted transcription of a total of 76 genes from *C. ferrugineus* experiments and just nine genes from *M. xanthus* experiments when only considering a ≥4-fold

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change in transcription at $p \leq 0.01$ (Fig. 1, Appendix S1). For this reason, our analysis of M. xanthus exposure experiments was expanded to include significant features at $p \leq 0.05$ resulting in an updated total of 56 impacted features from C6-AHL exposure (Fig. 1). While this indicates less variability across C. ferrugineus exposure experiments, we contend that this expansion provides a broader and more thorough analysis of statistically significant impacted features for our analysis. A similar consideration of C. ferrugineus genes with ≥4-fold change in transcription at $p \leq 0.05$ by C6-AHL exposure would provide an additional 119 impacted genes for consideration (Appendix S1). M. xanthus features included at the more stringent significance cutoff of $p \leq 0.01$ are indicated in Fig. 1. These data revealed that C6-AHL exposure elicited a general downregulation of genes with a total of 55 downregulated genes observed from M. xanthus and 51 genes from C. ferrugineus. Only one gene was observed to be upregulated by M. xanthus when exposed to C6-AHL, and 25 total upregulated genes were observed from C. ferrugineus during C6-AHL exposure.

Comparing annotated features impacted by C6-AHL exposure across both datasets and their putative roles by general system, numerous features involved in signal transduction pathways and transcriptional regulation were included in both datasets with seven regulatory features downregulated by M. xanthus and six downregulated by C. ferrugineus (Fig. 2). Both myxobacteria also had a TetR family transcriptional regulator upregulated by C6-AHL exposure. Multiple features associated with primary and specialized metabolisms and cell wall biogenesis and maintenance were downregulated by C6-AHL exposure across both datasets. Impacted specialized metabolism features included in biosynthetic gene clusters (BGCs) provided by the BGC prediction platform antiSMASH were further analysed to determine putative role and associated metabolites. A total of six BGCs from M. xanthus contained genes impacted by C6-AHL exposure including one core biosynthetic gene cluster (WP_011554909.1) from an annotated thiopeptide/bacteriocin BGC and one regulatory gene (WP_011552861.1) from a putative nonribosomal peptide synthetase-polyketide synthase hybrid BGC (Table S3; Fig. 1.

Transcriptomic data from myxobacteria exposed to C6-AHL.
A. Differentially expressed genes and features from M. xanthus exposed to C6-AHL when compared with signal unexposed M. xanthus control ($p \leq 0.05$); * indicates features also impacted at $p \leq 0.01$.
B. Differentially expressed genes from C. ferrugineus exposed to C6-AHL when compared with signal unexposed C. ferrugineus control ($p \leq 0.01$). Data depicted as an average log2 fold change from three biological replicates. Impacted features annotated as hypothetical not included.

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Fig. S1; Blin et al., 2021a, 2021b). None of the BGC-associated genes from *M. xanthus* impacted by C6-AHL were from characterized gene clusters with assigned metabolites. A total of 16 BGCs from *C. ferrugineus* included features impacted by C6-AHL exposure, however, no features annotated as core biosynthetic genes, regulatory genes or transport genes were observed (Table S3, Fig. S1).

Considering previous reports that C6-AHL exposure suppresses *M. xanthus* sporulation (Lloyd and Whitworth, 2017), we sought to determine if C6-AHL exposure effected either of the transcriptional regulators associated with *M. xanthus* sporulation FruA or MrpC (Ogawa et al., 1996; Robinson et al., 2014; Marcos-Torres et al., 2020). While no significant change in FruA was observed, transcription of the gene product MrpC was downregulated 1.7-fold by *M. xanthus* exposure to C6-AHL. However, transcription of the FruA (WP_071904077.1) or MrpC (WP_071900118) homologues from *C. ferrugineus* was not significantly changed by C6-AHL exposure. While no obvious predatory features associated with motility or lytic enzymes were directly impacted in our C6-AHL exposed *M. xanthus* results, we suspect that this could be due to the previously reported constitutive toxicity of *M. xanthus* observed in both the presence and absence of prey (Livingstone et al., 2018). The increased transcription of lytic enzymes and mobile genetic elements observed from *C. ferrugineus* exposed to C6-AHL suggest a predatory response; however, these features could also be associated with a defence response akin to phage defence. Transcription of neither of the annotated LuxR-type receptors (*M. xanthus*, WP_011555271.1; *C. ferrugineus*, WP_071900454.1) was affected by C6-AHL exposure. Overall considering the most significantly impacted features across both datasets, C6-AHL exposure elicited somewhat similar responses from both myxobacteria including numerous features associated with transcriptional regulation and signal transduction, primary and specialized metabolisms, and cell wall maintenance.

HHQ elicits contrasting responses from *M. xanthus* and *C. ferrugineus*

Comparative transcriptomics from RNAseq data from exposure experiments with HHQ (9 μM) introduced to plates of *M. xanthus* and *C. ferrugineus* were also conducted in triplicate with DMSO exposures serving as HHQ unexposed, negative controls for comparative analysis. Comparative transcriptomic analysis from RNAseq data revealed HHQ exposure led to a ≥4-fold (*p* ≤ 0.05) change in transcription of a total of 186 genes from *C. ferrugineus* and 31 total genes from *M. xanthus* (Fig. 3). Unlike the similar responses elicited by C6-AHL exposure, contrasting responses were apparent when comparing data between the myxobacteria. Data from *M. xanthus* experiments revealed overlap between responses to C6-AHL and HHQ with a total of nine...
upregulated genes and 22 downregulated genes including five genes also downregulated by C6-AHL exposure. Overlapping annotated features impacted by both C6-AHL and HHQ included a NmrA/HSCARG family protein (WP_011556972.1), an immunity 49 family protein (WP_011550233.1), a CHASE2 domain-containing protein (WP_011554259.1), and two hypothetical proteins (WP_011555268.1 and WP_011552217.1). Comparing

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Fig. 3. Transcriptomic data from myxobacteria exposed to HHQ.
A. Differentially expressed genes and features from *M. xanthus* exposed to HHQ when compared with signal unexposed *M. xanthus* control (p ≤ 0.05).
B. Differentially expressed genes from *C. ferrugineus* exposed to HHQ when compared with signal unexposed *C. ferrugineus* control (p ≤ 0.05). Data depicted as an average log2 fold change from three biological replicates. Impacted features annotated as hypothetical not included.
impacted genes from AHL and HHQ exposure experiments, further overlap between putative roles of annotated genes was also observed from *M. xanthus* with multiple impacted genes predicted to be involved in transcriptional regulation and signal transduction and cell wall biogenesis and maintenance (Figs. 2 and 4). Of note, the pleiotropic regulator MrpC was also downregulated 2.2-fold in *M. xanthus* exposed to HHQ which is comparable to 1.7-fold downregulation of MrpC observed with C6-AHL exposure. Only one gene (WP_011554915.1) included in a putative thiopeptide/bacteriocin BGC was significantly impacted by HHQ exposure (Table S3, Fig. S1).

Unlike the overlap in responses to both signals observed from *M. xanthus*, none of the 186 genes effected by HHQ exposure overlapped with the 76 genes impacted by C6-AHL exposure. Considering annotated genes by functional category, *C. ferrugineus* genes upregulated by HHQ exposure (156 total) were largely associated with signal transduction and transcriptional regulation, various metabolic pathways, and multiple genes associated with protein translation and turnover, cell wall biogenesis and maintenance and specialized metabolism were downregulated (29 total; Fig. 4). A total of six BGCs from *C. ferrugineus* contained genes impacted by HHQ exposure. These included three regulatory genes from a nonribosomal peptide synthetase (WP_071896979.1), a type III polyketide synthase (WP_071896419.1), and a linear azol(in)e-containing peptide pathway (WP_071903679.1) as well as a transport-related gene (WP_071904585.1) from a type I polyketide synthase pathway (Table S3, Fig. S1). Interestingly, an annotated FAD-dependent oxidoreductase (WP_071901324.1) homologous to the monoxygenase PsqH from *P. aeruginosa* (91% coverage; 38% identity) which hydroxylates HHQ to yield 2-heptyl-3-hydroxyquinolin-4(1H)-one or pseudomonas quinolone signal (PQS) was upregulated 31-fold (Diggle et al., 2003; Ritzmann et al., 2021). An outlier to the contrasting responses to HHQ, an annotated DUF2378 family protein (*M. xanthus*, WP_011553830.1; *C. ferrugineus*, WP_084736518.1) was significantly upregulated in both myxobacteria; DUF2378 family proteins are ~200 amino acid proteins with no known function that are exclusive to myxobacteria. Overall, these results indicate that *M. xanthus* exhibits a similar transcriptional response to both C6-AHL and HHQ whereas HHQ elicits a distinct response from *C. ferrugineus* dissimilar from the more general response observed from both myxobacteria when exposed to C6-AHL.

**Differential metabolic impact of AHL and HHQ signals**

Subsequent exposure experiments were conducted with *M. xanthus* and *C. ferrugineus* exactly as done for our RNAseq experiments with an additional AHL signal,
3-oxo-C6-AHL, also included. Crude, organic phase extracts generated from these experiments were subjected to untargeted mass spectrometry and the XCMS-MRM (v3.7.1) platform (Domingo-Almenara et al., 2018; Forsberg et al., 2018) was utilized for comparative analysis and determination of statistical significance for all detected features. Comparing features with significantly impacted intensities ($p \leq 0.02$) during these signal exposure experiments, all three signals elicited a more apparent response from *C. ferrugineus* (Fig. 5, Fig. S2). Despite the comparatively diminished response from *M. xanthus*, two general trends were apparent when comparing the signals responses between both myxobacteria. First, C6-AHL and 3-oxo-C6-AHL exposure resulted in highly similar responses from both myxobacteria with few to no uniquely impacted features specific to either AHL signal (Fig. 5). Second, HHQ exposure induced a dramatic change in the metabolic profile of *C. ferrugineus* that was not observed from HHQ-exposed *M. xanthus*. A total of 47 features from *C. ferrugineus* were impacted by both AHL signals while 133 features were affected by HHQ exposure. Intrigued by the difference in responses, additional experiments where both myxobacteria were exposed to exogenous C6-AHL and HHQ simultaneously were done. Comparative analysis of results revealed that the addition of C6-AHL did not dramatically impact the change in metabolic profile observed from either myxobacteria when exposed to HHQ (Fig. 6). Conversely, impacted features observed in our previous AHL exposure experiments were not observed in our C6-AHL + HHQ experiments. For example, of the 47 total overlapping *C. ferrugineus* features with significantly changed intensities during AHL exposure conditions, 36 were not observed to change during C6-AHL + HHQ exposure experiments. The Global Natural Products Social Molecular Networking (GNPS) platform (Wang et al., 2016) was used to determine if any impacted features could be associated with characterized metabolites. Although networks included detected features with exact masses for myxovirescin

![Fig. 5. Comparison of metabolomic response to C6-AHL, 3-oxo-C6-AHL and HHQ exposure experiments with *M. xanthus* (A and B) and *C. ferrugineus* (C and D). Numbers included in each Venn diagram account for a unique detected feature with a significantly impacted intensity upon exposure to the indicated signalling molecule provided by XCMS-multigroup analysis ($n = 3, p \leq 0.02$).](image)

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and degraded myxovirescin metabolites were observed in *M. xanthus* extracts and detected features with exact masses for various tubulysins were observed in *C. ferrugineus* extracts (Fig. S3), detected quantities were not impacted by AHL or HHQ exposure. From these results, we determined that both myxobacteria demonstrate a metabolic response unique to either HHQ or AHL-type chemical signals with a conserved response to both C6-AHL and 3-oxo-C6-AHL. These data also revealed a unique metabolic response from *C. ferrugineus* when exposed to HHQ similar to our previous transcriptomic observation.

**Conserved metabolomic response to AHLs and determination of core L-homoserine lactone elicitor**

Intrigued by the overlap in metabolic responses observed from AHL signal exposure, we were curious if the core homoserine lactone moiety present in all AHL-type quorum signals was sufficient to elicit a similar response. Untargeted mass spectrometry and XCMS-MRM analysis of additional exposure experiments with *C. ferrugineus* including either L-homoserine lactone (L-HSL) the stereoisomer present in natural AHL-type quorum signals (Papenfort and Bassler, 2016; Mukherjee and Bassler, 2019), D-homoserine lactone (D-HSL) the enantiomer of L-HSL, or boiled C6-AHL were completed to determine any overlap with previously observed responses to C6-AHL and 3-oxo-C6-AHL exposure. Comparative analysis of statistically impacted features by signal intensity (\(p \leq 0.02\)) revealed 20 overlapping features from the L-HSL, C6-AHL, and 3-oxo-C6-AHL exposure experiments and just three overlapping features from the L-HSL, D-HSL, C6-AHL, and 3-oxo-C6-AHL exposure experiments (Fig. 7, Fig. S4). Hierarchical clustering of detected feature intensities from L-HSL, D-HSL, C6-AHL, and control datasets also revealed clustering between L-HSL and C6-AHL datasets (Fig. S5). These
results suggest that the core homoserine lactone core present in all AHLs is sufficient for predatory eavesdropping by myxobacteria.

Oxidative detoxification of HHQ observed from C. ferrugineus

Comparing metabolomic datasets from HHQ exposure experiments, oxidized analogs of HHQ detected at 260.164 m/z were exclusive to the C. ferrugineus dataset. Authentic standards for the oxidized HHQ quinolone signals PQS and 2-heptyl-1-hydroxyquinolin-4(1H)-one (HQNO) were used to determine that both oxidized signals were present in HHQ-exposed C. ferrugineus extracts (Fig. 8; Fig. S6; Cao et al., 2020). Oxidative detoxification of quinolone signals with HHQ observed to be oxidized to either HQNO or PQS. The presence of a metabolite with an exact mass and similar MS2 fragmentation pattern matching PQS-N-oxide (PQS-NO) an oxidation product reported by Thierbach et al. was also observed in C. ferrugineus extracts from HHQ, PQS, and HQNO exposure experiments suggesting subsequent oxidation of both PQS and HQNO (Fig. 8; Figs. S7 and S8; Thierbach et al., 2017). These results suggest that C. ferrugineus possesses a detoxification route for quinolone signals not observed from M. xanthus and oxidizes the quinolone signals HHQ, PQS, and HQNO.

C. ferrugineus response to HHQ correlates with superior predation of P. aeruginosa

Predation assays using the lawn culture method were conducted in triplicate on lawns of P. aeruginosa with both M. xanthus and C. ferrugineus (Morgan et al., 2010). These assays confirmed that P. aeruginosa was comparatively more susceptible to predation by C. ferrugineus (Fig. 9A). Secretion of toxic quinolone signals in soil is thought to provide P. aeruginosa an advantage, and HHQ inhibition of soil-dwelling Bacillus atrophaeus swarming motility has been observed (Reen et al., 2015; Ritzmann et al., 2021). Subsequent swarming expansion assays with M. xanthus and C. ferrugineus on medias with and without supplemented HHQ were done to determine if similar antiswarming activities were observed. From these assays, we observed HHQ (9 μM) significantly inhibited M. xanthus swarming but not C. ferrugineus swarming (Fig. 9B and C). We suggest oxidative degradation of quinolone signals observed from C. ferrugineus contributes to predation of P. aeruginosa. Altogether these results indicate the unique response to exogenous HHQ observed from C. ferrugineus to be an evolved trait associated with exposure to quinolone signals that correlates with predation of quinolone signal-producing pseudomonads.
Discussion

Although the predatory lifestyles of myxobacteria have long been associated with their capacity as a resource for natural products discovery, the chemical ecology of predator–prey interactions remains underexplored (Findlay, 2016; Munoz-Dorado et al., 2016; Herrmann et al., 2017). The recent discovery that exogenous AHL quorum signals associated with Gram-negative prey bacteria increase the predatory capacity of \textit{M. xanthus} provides an excellent example of shared chemical space within microbial communities influencing predation (Lloyd and Whitworth, 2017). Utilizing comparative transcriptomics and metabolomics, we sought to determine the generality of predatory eavesdropping and how the phenomenon might correlate with prey range by comparing responses from \textit{M. xanthus} and \textit{C. ferrugineus} when exposed to structurally distinct quorum signals associated with the opportunistic pathogen \textit{P. aeruginosa}.

Initial transcriptomic data comparing \textit{M. xanthus} and \textit{C. ferrugineus} exposed to C6-AHL revealed overlapping transcriptional responses from both myxobacteria. Originally referenced as predatory eavesdropping, we sought to determine the impact of C6-AHL on genes with annotations affiliated with predation and predatory responses such as motility features, lytic enzymes and specialized metabolism (Munoz-Dorado et al., 2016). Transcription of multiple genes associated with transcriptional regulation, metabolism and cell wall maintenance was influenced by exogenous C6-AHL across both myxobacteria. Although we observed numerous genes embedded in putative specialized metabolite biosynthetic pathways impacted by C6-AHL from both myxobacteria, none were included in the annotated BGCs for myxovirescin or myxoprincomide. Since these are the only two myxobacterial metabolites currently implicated in predation, further investigation is required to determine if C6-AHL induces production of biologically active, specialized metabolites (Xiao et al., 2011; Cortina et al., 2012; Muller et al., 2016; Ellis et al., 2019; Wang et al., 2019). The only other potential predatory features with a transcriptional response to C6-AHL exposure were putative lytic enzymes from \textit{C. ferrugineus}, and no annotated genes predicted to be involved in motility were affected by C6-AHL in either myxobacteria. Considering the original observation that AHLs stimulate predation by increasing the vegetative population of \textit{M. xanthus}, we suggest

Fig. 8. A. Extracted ion chromatograph (EIC) depicting presence of HQNO and PQS in HHQ exposed extracts from \textit{C. ferrugineus} and not observed in HHQ exposed extracts from \textit{M. xanthus}. B. EIC depicting presence of PQS-NO in HHQ exposed extracts of \textit{C. ferrugineus}, also not present in \textit{M. xanthus} extracts. Chromatographs rendered with MZmine v2.37. C. Oxidative detoxification of HHQ by \textit{C. ferrugineus} including exact mass values from ChemDraw Professional v17.1. D. Detected ion intensities for PQS-NO comparing crude extracts of \textit{C. ferrugineus} exposed to HHQ, PQS and HQNO; detected intensity data provided by XCMS-multigroup analysis (n = 3; p ≤ 0.02).
that the observed change in transcription of genes associated with metabolism and signal transduction from both myxobacteria may correspond with a similar population-based response and shift in vegetative state (Lloyd and Whitworth, 2017). The decreased transcription of the gene encoding for MrpC, a developmental regulator involved in sporulation (Robinson et al., 2014), observed in our M. xanthus dataset also supports a population-based response to C6-AHL.

Subsequent comparative metabolomics experiments indicated that C6-AHL and 3-oxo-C6-AHL elicit overlapping responses from both myxobacteria and that the core AHL moiety L-HSL also elicits a similar response from C. ferrugineus. We conclude that the overlap between C6-AHL, 3-oxo-C6-AHL and L-HSL indicates an evolved recognition of the homoserine lactone unit present in all AHL quorum signals (Papenfort and Bassler, 2016; Mukherjee and Bassler, 2019). As generalist predators, a more general process for AHL perception that responds to a core moiety in the scaffold of AHLs might be preferred to a specialized process associated with the variable N-acylamides of AHLs. We also suspect this centralized response to L-HSL may relate to the absence of a LuxR-type AHL receptor that includes a conserved AHL-binding domain. The ubiquity of AHL quorum signals amongst Gram-negative bacteria combined with the overlap in observed responses from M. xanthus and C. ferrugineus suggest that AHL-based eavesdropping by myxobacteria could be a general trait that benefits predation.

Unlike the overlap in responses to AHL exposure, the quinolone signal HHQ elicited a contrasting transcriptomic from M. xanthus and C. ferrugineus. Contrary to M. xanthus, C. ferrugineus upregulated genes associated with signal transduction and transcriptional regulation, various metabolic pathways, and multiple genes associated with protein translation and turnover, cell wall biogenesis and maintenance, and specialized metabolism when exposed to HHQ. Interestingly, an annotated FAD-dependent oxidoreductase homologous to the
monooxygenase PqsH from *P. aeruginosa*, which hydroxylates HHQ to yield PQS was upregulated 31-fold in *C. ferrugineus* exposed to HHQ. Subsequent metabolomic experiments confirmed the presence of two oxidized analogs of HHQ, PQS and HQNO (Dubern and Diggle, 2008; Thierbach et al., 2017). The detection of these oxidized quinolones as well as an additional feature, PQS-NO, previously associated with the oxidative detoxification of quinolone signals in HHQ exposed *C. ferrugineus* samples and absence in *M. xanthus* samples suggests that *M. xanthus* is unable to similarly metabolize HHQ (Dubern and Diggle, 2008; Thierbach et al., 2017; Ritzbach et al., 2021). Oxidative detoxification of quinolone signals produced by pseudomonads has previously been reported from strains of *Arthrobacter*, *Rhodococcus* and *Staphylococcus aureus* (Thierbach et al., 2017). Subsequent swarming expansion assays revealed HHQ inhibits *M. xanthus* swarming but not *C. ferrugineus* swarming. We suggest that this oxidative detoxification process contributes to the superior predation of *P. aeruginosa* observed from *C. ferrugineus* in our predation assays comparing both myxobacteria.

Despite being considered keystone taxa within microbial communities (Petters et al., 2021), the extent of myxobacterial bacterivory and its contribution to nutrient cycling within microbial food webs remains unknown. These results further demonstrate myxobacterial eavesdropping on prey signalling molecules and provide insight into how responses to exogenous signals might correlate with prey range of myxobacteria. Although broadly considered generalist predators, predatory specialization has been observed from myxobacteria. Oxidation of quinolone signals and superior predation of *P. aeruginosa* observed from *C. ferrugineus* provides an example of how prey signalling molecules and the shared chemical ecology of microbial communities influence myxobacterial predation.

**Experimental procedures**

**Cultivation of *M. xanthus* and *C. ferrugineus***

*Cystobacter ferrugineus* strain Cbfe23, DSM 52764, initially obtained from German Collection of Microorganisms (DSMZ) in Braunschweig, and *Myxococcus xanthus* strain GJLV were employed in this study. *Cystobacter ferrugineus* was grown on VY/2 agar (5 g l⁻¹ baker's yeast, 1.36 g l⁻¹ CaCl₂, 0.5 mg l⁻¹ vitamin B₁₂, 15 g l⁻¹ agar, pH 7.2). Whereas, CTTYE agar (1.4% w/v agar, 1% Casitone, 10 mM Tris–HCl (pH 7.6), 1 mM potassium phosphate (pH 7.6), 8 mM MgSO₄, 0.5% yeast extract) was utilized to culture *M. xanthus*.

**Quorum signal exposure experiments**

For signal exposure conditions, required volumes for 9 μM of filter sterilized, HHQ (Sigma), C₆-AHL (Cayman Chemical), 3-oxo-C₆-AHL (Cayman Chemical), l-HSL (Cayman Chemical), d-HSL (Cayman Chemical), PQS (Sigma) and HQNO (Sigma) from a 150 mM stock prepared in DMSO were added to autoclaved medium at 55°C. Boiled AHL samples were prepared according to established methodology (Lloyd and Whitworth, 2017). For RNA-seq and LC–MS/MS analysis, *C. ferrugineus* was cultivated on VY/2 agar medium, and *M. xanthus* was cultured on CTTYE agar medium. For all signal exposure experiments both myxobacteria were grown at 30°C with *C. ferrugineus* grown for 10 days and *M. xanthus* grown 14 days.

**RNA sequencing experiments and analysis**

Myxobacterial cells were scrapped from the agar plates and stored in RNA-ladder. Total RNA was isolated from the samples using the RNeasy PowerSoil Total RNA Kit (Qiagen) following the manufacturer's instructions. Consistent aliquots of biomass (500 mg) from each myxobacteria were used for RNA extractions. The concentration of total RNA was determined using the Qubit RNA Assay Kit (Life Technologies). For rRNA depletion, first, 1000 ng of total RNA was used to remove the DNA contamination using Baseline-ZERO DNase (Epicentre) following the manufacturer's instructions followed by purification using the RNA Clean and Concentrator-5 columns (Zymo Research). DNA free RNA samples were used for rRNA removal by using RiboMinus rRNA Removal Kit (Bacteria; Thermo Fisher Scientific) and final purification was performed using the RNA Clean and Concentrator-5 columns (Zymo Research). rRNA depleted samples were used for library preparation using the KAPA mRNA HyperPrep Kits (Roche) by following the manufacturer's instructions. Following the library preparation, the final concentration of each library was measured using the Qubit dsDNA HS Assay Kit (Life Technologies), and average library size for each was determined using the Agilent 2100 Bioanalyzer (Agilent Technologies; Tables S1 and S2). The libraries were then pooled in equimolar ratios of 0.75 nM, and sequenced paired end for 300 cycles using the NovaSeq 6000 system (Illumina). RNA sequencing was conducted by MR DNA (Molecular Research LP). RNAseq analysis was performed using ArrayStar V15 and the R-package DESeq2 for differential expression data. Raw data from RNAseq analysis publicly available at the National Centre for Biotechnology Information Sequence Read Archive under the following BioProjects PRJNA555507, PRJNA730806 and PRJNA730808.
AntiSMASH analysis

Biosynthetic gene clusters from *M. xanthus* DK1622 deposited in the antiSMASH database version 3 (Blin et al., 2021b) were used in our analysis of specialized metabolite genes included in BGCs. The .gbk file for the *C. ferrugineus* genome sequence deposited at NCBI (NZ_MPIN00000000.1) was analysed with antiSMASH version 6 (Blin et al., 2021a) to determine BGCs and identify included specialized metabolite genes. All BGC-associated genes resulting from these analyses are provided in Table S3.

Organic phase extraction of metabolites

After cultivation, myxobacterial plates were manually diced and extracted with excess EtOAc. Pooled EtOAc was filtered and dried in vacuo to provide crude extracts for LC–MS/MS analysis. LC–MS/MS analysis of the extracted samples was performed on an Orbitrap Fusion instrument (Thermo Scientific, San Jose, California) controlled with Xcalibur version 2.0.7 and coupled to a Dionex Ultimate 3000 nanoUHPLC system. Samples were loaded onto a PepMap 100 C18 column (0.3 mm × 150 mm, 2 μm, Thermo Fisher Scientific). Separation of the samples was performed using mobile phase A (0.1% formic acid in water) and mobile phase B (0.1% formic acid in acetonitrile) at a rate of 6 μl min⁻¹. The samples were eluted with a gradient consisting of 5% to 60% solvent B over 15 min, ramped to 95% B over 2 min, held for 3 min, and then returned to 5% B over 3 min and held for 8 min. All data were acquired in positive ion mode. Collision induced dissociation (CID) was used to fragment molecules, with an isolation width of 3 m z⁻¹ units. The spray voltage was set to 3600 V, and the temperature of the heated capillary was set to 300°C. In CID mode, full MS scans were acquired from m/z 150 to 1200 followed by eight subsequent MS² scans on the top eight most abundant peaks. The nominal orbitrap resolution for both the MS¹ and MS² scans was 60,000. The expected mass accuracy based on external calibration was <3 ppm. MZmine 2.53 was used to generate extracted ion chromatograms (Pluskal et al., 2010). Organic phase extraction was utilized in combination with a lower mass range cutoff of 150 m z⁻¹ to minimize detected primary metabolites and effectively prioritize analysis of less polar, higher mass specialized metabolites.

XCMS analysis

Generated data were converted to .mzXML files using MS-Convert (Adusumilli and Mallick, 2017). Multigroup analysis of converted .mzXML files was done using XCMS-MRM and the default HPLC/Orbitrap parameters (Domingo-Almenara et al., 2018; Forsberg et al., 2018). Within the XCMS-MRM result tables, determination of signal-impacted detected features was afforded by filtering results for those with a p ≤ 0.02.

GNPS analysis

Converted .mzXML files were used to generate mass spectrometry molecular networks using the GNPS platform release 28.2 (Wang et al., 2016). All included LC–MS/MS data were deposited as a publicly available MassIVE dataset (MSV000087999).

Lawn culture predation assays

*Pseudomonas aeruginosa* ATCC 10145ᵀ was purchased from the American Type Culture Collection (ATCC). The predation experiment was performed according to (Pham et al., 2005; Morgan et al., 2010). Briefly, an overnight grown culture of *P. aeruginosa* was pelleted at 5000 x g. The cell pellet was washed with TM buffer and pelleted again. The pelleted cells were resuspended in TM buffer to an OD₆₀₀ 0.5. A 250 μl volume of resuspended cell suspension was utilized to make a uniform bacterial lawn on a WAT agar plate. Myxobacterium *M. xanthus* GJV1 was grown on CTTYE agar, and *C. ferrugineus* was grown on VY/2 agar for 7 days. A 600 mm² agar block of each myxobacteria was excised and placed at the centre of the *P. aeruginosa* cell lawn. Assays were incubated at 30°C and swarm diameters measured after 4 days.

Swarming expansion assays

Swarming expansion assays were performed using modified established methods (Lloyd and Whitworth, 2017). Myxobacteria were grown on either VY/2 or CTTYE agar media as previously described for 4 days at 30°C. A homogenous bacterial suspension was created using sterile distilled water, vortexed, and equal to an OD₆₀₀ 0.5 for *C. ferrugineus* and an OD₆₀₀ 1.0 for *M. xanthus*. Then 5 μl of myxobacterial suspension was aliquoted within the centre of 100 mm × 15 mm plates supplemented within respective medias with either 9 μM filter sterilized HHQ (Sigma), without HHQ, or supplemented with 34 μM filter sterilized tetracycline (Thermo Fisher Scientific). Conditions without HHQ and tetracycline were used as controls. Swarming diameters (cm) were assessed and recorded daily for 10 days.

PRISM v7.0d was used to measure the statistical significance of changes in swarm expansion rates across the strains using two-way ANOVA and the Tukey’s multiple comparisons test.
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**Supporting Information**

Additional Supporting Information may be found in the online version of this article at the publisher's web-site.

**Appendix S1**: Supporting Information

**Appendix S2**: Supporting Information