The Genetics of Prey Susceptibility to Myxobacterial Predation: A Review, Including an Investigation into *Pseudomonas aeruginosa* Mutations Affecting Predation by *Myxococcus xanthus*

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Abstract

Bacterial predation is a ubiquitous and fundamental biological process, which influences the community composition of microbial ecosystems. Among the best characterised bacterial predators are the myxobacteria, which include the model organism *Myxococcus xanthus*. Predation by *M. xanthus* involves the secretion of antibiotic metabolites and hydrolytic enzymes, which results in the lysis of prey organisms and release of prey nutrients into the extracellular milieu. Due to the generalist nature of this predatory mechanism, *M. xanthus* has a broad prey range, being able to kill and consume Gram-negative/positive bacteria and fungi. Potential prey organisms have evolved a range of behaviours which protect themselves from attack by predators. In recent years, several investigations have studied the molecular responses of a broad variety of prey organisms to *M. xanthus* predation. It seems that the diverse mechanisms employed by prey belong to a much smaller number of general “predation resistance” strategies. In this mini-review, we present the current state of knowledge regarding *M. xanthus* predation, and how prey organisms resist predation. As previous molecular studies of prey susceptibility have focussed on individual genes/metabolites, we have also undertaken a genome-wide screen for genes of *Pseudomonas aeruginosa* which contribute to its ability to resist predation. *P. aeruginosa* is a World Health Organisation priority 1 antibiotic-resistant pathogen. It is metabolically versatile and has an array of pathogenic mechanisms, leading to its prevalence as an opportunistic pathogen. Using a library of nearly 5,500 defined transposon insertion mutants, we screened for “prey genes”, which when mutated allowed increased predation by a fluorescent strain of *M. xanthus*. A set of candidate “prey proteins” were identified, which shared common functional roles and whose nature suggested that predation resistance by *P. aeruginosa* requires an effective metal/oxidative stress system, an intact motility system, and mechanisms for detoxifying antimicrobial peptides.
Introduction

Predation is a fundamental and ubiquitous biological process, shaping the biotic composition of ecosystems. Microbial predation dictates the flow of nutrients between members of the microbial food web, with predators incorporating biomass from members of lower trophic levels [Zhang and Leuders, 2017]. Predators are defined as organisms that can kill and consume other “prey” organisms, and in microbes, predation can be facultative or obligate [Pérez et al., 2016]. Nevertheless, many microbes possess “anti-competitor” mechanisms that kill competing cells (e.g., antibiotic-producing Streptomyces spp.), releasing nutrients for general consumption. Consequently, the line between anti-competitor mechanisms and predatory activity is a fine one, and many organisms traditionally not viewed as predators have subsequently been shown to be true predators [Kumbhar et al., 2014].

Microbial predation has been investigated for more than 75 years, and research into the predatory mechanisms of predators has been reviewed and compiled recently [Pérez et al., 2016; Findlay, 2016; Jurkevitch and Mitchell, 2020; Whitworth et al., 2020]. Some microbial predators are considered specialists, while others are more generalist, with predatory activity being dependent on the availability of particular prey organisms. The two best understood microbial predators are the BALOs (Bdellovibrio and like organisms) and the myxobacteria, which possess very different predatory mechanisms [Pérez et al., 2016].

BALOs are typically endobiotic predators, attaching to the outer membrane of Gram-negative prey bacteria and entering their periplasm to form a bdelloplast. Within the bdelloplast, the contents of the prey cell are consumed and the BALO cell replicates before progeny are released from the lysing bdelloplast [Laloux, 2020]. Due to the requirement for periplasmic invasion and residency, the endobiotic mode of predation requires a Gram-negative prey organism.

In contrast, “public goods” predators exemplified by the myxobacteria, but also including Herpetosiphon spp. and Streptomyces spp., secrete antibiotics and/or lytic enzymes which kill prey organisms from the outside [Pérez et al., 2016]. The undirected nature of such predation is associated with broad prey ranges, with prey including Gram-negative bacteria, Gram-positive bacteria and fungi [Morgan et al., 2010; Livingstone et al., 2018a]. Myxobacteria are widely known to produce large numbers of secondary metabolites, many of which have antibiotic activity [Korp et al., 2016; Findlay, 2016]. This, coupled with their broad prey range, which includes important human pathogens, has stimulated many studies into various aspects of myxobacterial predation [reviewed by Berleman and Kirby 2009; Muñoz-Dorado et al., 2016; Furness et al., 2020; Thiery and Kaimer, 2020].

Myxobacteria secrete diffusible antibiotics, which result in zones of growth inhibition of prey lawns, whether surrounding myxobacterial colonies [Xiao et al., 2011] or around paper disks infused with extracted antibiotics [Schneiker et al., 2007]. However, experiments into the predation of Escherichia coli by the myxobacterium Myxococcus xanthus suggest that the antibiotic responsible for growth inhibition (myxovirescin) does not directly kill prey but rather impedes its growth. This suggests that myxobacteria also secrete non-antibiotic substances, which are responsible for prey killing.

Gram-negative bacteria, such as the myxobacteria, secrete outer membrane vesicles (OMVs), which are pinched off portions of the outer membrane enclosing an aqueous volume derived from contents of the periplasm [Whitworth, 2011]. Both proteins and metabolites can be packaged into OMVs, and purified OMVs are themselves predatory, even those purified from non-predatory bacteria [Kadurugamuwa and Beveridge, 1996; Li et al., 1998]. Myxobacterial OMVs were first characterised by Kahnt et al. [2010] and shortly thereafter shown to be able to kill both E. coli and a Pseudomonas sp. strain [Evans et al., 2012]. Several studies have characterised the contents of OMVs from M. xanthus [Kahnt et al., 2010; Berleman et al., 2014; Whitworth et al., 2015; Zwarycz et al., 2020], finding an abundance of hydrolytic enzymes, proteases and secondary metabolites, consistent with a role in predation. A survey of OMV proteomes from different strains of M. xanthus found that despite substantial commonality, there was also considerable variability and individuality in OMV proteome composition even within that single myxobacterial species [Zwarycz et al., 2020].

There are several benefits to predators for secreting hydrolytic enzymes within OMVs rather than directly into the extracellular milieu. The delivery of OMV contents is targetable – OMVs can fuse with specific target membranes, delivering cargo into the prey cytoplasm/periplasm, or they can be triggered to lyse at the surface of prey cells [Kadurugamuwa and Beveridge, 1996]. However, there is currently little understanding of how cargo proteins are targeted for packaging into OMVs, or how OMVs can be targeted to particular destinations. Enclosing cargo proteins protects them from environmental stresses and allows them to maintain their concentrations and potency at a distance from the producing cell. At the
same time, packaging cargo within OMVs reduces their rate of transport away from the producing cell [Whitworth, 2011]. This may appear to be a counter-productive behaviour for predatory cells; however, reducing the migration of predatory secretions away from the producer ensures that prey killing only happens proximal to the predatory cell. Such behaviour may be important in reducing competition from kin and other competitors for the nutrients released by lysing prey in the public commons [Marshall and Whitworth, 2019].

It is believed that the broad prey range of myxobacteria and other “public goods” predators is a consequence of their indiscriminate secretion of a multi-valent cocktail of OMVs, hydrolases and antibiotics. However, while myxobacteria are all able to kill a wide range of prey organisms, individual strains show dramatic differences in their profiles of predatory activity against specific prey [Morgan et al., 2010; Livingstone et al., 2017]. It is thought that the patchy mosaic of predatory activity and prey susceptibility observed is a consequence of individualised differences in the composition of each strains’ multi-valent predatory cocktail. Such individuality is observed in the OMV cargo proteins of different strains and also in the genes within their genomes. Pan-genome analysis of M. xanthus, Myxococcus spp. and Coralloccoccus spp. demonstrates that myxobacterial genomes contain large numbers of accessory genes, which are typically unique to those individual strains [Livingstone et al., 2018b; Zwarycz et al., 2020; Chambers et al., 2020].

The lack of correlation between predatory activity, predator phylogeny and prey susceptibility [Livingstone et al., 2017] suggests that individual strains of predator possess unique complements of predatory metabolites and hydrolases, and that what we deduce about the predatory mechanisms of one strain may not be relevant to another predatory strain, even within the same species. Nevertheless, the genomic and proteomic diversity of predatory strains presents opportunities. For instance, genome-wide association studies have been used to correlate myxobacterial predatory efficiency with the presence/absence of specific genes, allowing the identification of enzymes which assist in effective predation of specific prey [Sutton et al., 2019].

The Genetic Basis of Prey Susceptibility/Resistance to Predatory Attack

As described above, the effectiveness with which a prey organism is consumed by a predatory microbe is highly strain specific. But it is equally clear that the peculiarities of the prey organism also have a profound effect on the outcomes of predatory encounters. There is no correlation between predatory activity and prey phylogeny, and closely-related prey organisms can be preyed upon with very different efficiencies by the same predator [Livingstone et al., 2017]. In general, the role of the prey in the predator-prey relationship has received less attention than that of the predator; however, there are growing numbers of studies which have investigated prey determinants of susceptibility and/or resistance to myxobacterial predation.

Fungi

While myxobacteria can effectively prey upon fungi, very few studies have investigated the mechanisms involved. Recently, Li et al. [2019] demonstrated that an outer membrane β-barrel glucanase (GluM) was able to degrade fungal cell walls and impede infection by plant pathogens. GluM showed specificity towards β-1,6-glucans, and fungi lacking such linkages were immune to myxobacterial predation, implying GluM is important in dictating predatory specificity towards fungal prey.

Sinorhizobium meliloti

M. xanthus can attack the Gram-negative rhizobium Sinorhizobium meliloti via two distinct predatory modes. In “frontal attack” colonies expand towards each other and encounter each other as dense populations of cells, while during “wolf-pack attack” small groups of cells operate independently to infiltrate prey populations. The effectiveness of frontal attack has been found to be reduced by S. meliloti through the production of the exopolysaccharide galactoglucan EPS II [Pérez et al., 2014]. During frontal attack, the area between S. meliloti and M. xanthus accumulates copper. This induces the predator to express copper-resistance mechanisms, while the prey produces the antioxidant melanin. Prey strains incapable of producing melanin were found to be more sensitive to predation, suggesting that copper is used by the predator to induce metal/oxidative stress in the prey [Contreras-Moreno et al., 2020].

Streptomyces coelicolor

Recently elevated to the status of a predator in its own right, Streptomyces coelicolor is a model Gram-positive organism for studies into secondary metabolism and sporulation. When colonies of M. xanthus and S. coelicolor expand towards each other, the vegetative hyphae of S. coelicolor are lysed, and it responds by producing an aerial mycelium and synthesising the antibiotic actinorhodin [Pérez et al., 2011]. Thus, S. coelicolor resists predation through two strategies: aerial hyphae undergo...
differentiation into chains of spores, which are typically predation-resistant cell types, and antibiotic production represents a “counter-attack” response to predation.

**Bacillus subtilis**

*Bacillus subtilis* is another model Gram-positive organism for studies on sporulation, and it is perhaps not surprizing that *B. subtilis* is induced to sporulate by myxobacterial predation, forming predation-resistant spores [Müller et al., 2014]. Predatory attack by *M. xanthus* also prompted *B. subtilis* to produce specialised structures called megastructures to house sporulating cells, which prevented *M. xanthus* from acquiring nutrients from the prey colony [Müller et al., 2015]. As well as differentiating into predation-resistant forms, *B. subtilis* also employs a counter-attack mechanism, producing the polyketide antibiotic bacillena, which protects the colony from predation [Müller et al., 2014].

**Escherichia coli**

The model Gram-negative bacterium *E. coli* is one of the most commonly studied prey organisms killed by myxobacteria. Biofilm formation, particularly production of the curli and cellulose matrix components, has been found to provide *E. coli* with protection from predation by *M. xanthus* [DePas et al., 2014]. Similarly, in an artificial evolution experiment, Nair et al. [2019] showed that predator-prey co-cultures of *M. xanthus* and *E. coli* drove adaptive evolution of mucoidy in *E. coli* in addition to changes in its outer membrane protease OmpT (both features involved with *E. coli* pathogenesis).

Transcriptome profiling of *E. coli* revealed widespread changes in gene expression when co-incubated with *M. xanthus*, with 40% of *E. coli* genes being significantly differentially expressed [Livingstone et al., 2018c]. Addition of purified *M. xanthus* OMVs or culture supernatant also induced expression of hundreds of *E. coli* genes. Genes for LPS synthesis, ribosome function and the electron transport chain/oxidative phosphorylation were particularly and consistently up-regulated during exposure to *M. xanthus* and/or its secretions, suggesting they are the primary molecular targets of attack [Livingstone et al., 2018c]. Unfortunately, it is not clear whether the genes induced by *E. coli* while under attack from *M. xanthus* actually make a difference to predatory outcomes (e.g., prey longevity or survival) or merely reflect the prey trying (unsuccessfully) to maintain homeostasis.

**Bacillus licheniformis**

Gram-positive *Bacillus licheniformis* is able to resist *M. xanthus* predation by inducing expression of an enzyme catalysing the glycosylation of the antibiotic myxovirescin A [Wang et al., 2019]. Glycosylation of myxovirescin is thought to reduce its affinity for its target protein LspA (signal peptidase II), thereby attenuating its antibiotic activity. Accordingly, transplantation of the *yjiC* gene encoding the glycosylase was able to make recipient *E. coli* resistant to predation by *M. xanthus*. In *B. licheniformis*, predation by *M. xanthus* induced expression of both *yjiC* and *lspA*, suggesting a two-pronged response to the presence of myxovirescin [Wang et al., 2019].

**Unifying Principles**

The molecular details of the responses of specific prey organisms to *M. xanthus* attack are very different from one another, and yet there are commonalities in the general features of the prey responses. We would broadly divide the observed prey strategies for resisting *M. xanthus* predation into 5 categories:

- **Being innately resistant.** Some prey organisms are intrinsically resistant to the predator’s molecules of attack. For instance, fungi lacking β-1,6-glucans resist *M. xanthus* predation.
- **Producing specific counter-measures.** The prey specifically neutralise harmful molecules produced by the attacker. Examples above include the production of melanin by *S. meliloti* and glycosylation of myxovirescin by *B. licheniformis*.
- **Reinforcing the cell exterior.** Production of a thickened outer layer can be achieved in several ways, including sporulation (as exemplified by *S. coelicolor* and *B. subtilis*), or production of an outer sugary coating (e.g., production of EPS/LPS and mucoidy by *E. coli*).
- **Building external protective structures.** Usually a communal response to predation, prey cells cooperate by secreting an engulfing matrix, physically separating themselves from the predators. This category includes formation of megastructures by *B. subtilis* and biofilm formation by *E. coli*.
- **Counter-attacking.** By producing its own toxic molecules, the prey pre-emptively disables *M. xanthus*, impeding its predation. Examples of this category include production of actinorhodin by *S. coelicolor* and bacillena production by *B. subtilis*.

Some prey organisms appear to employ just one of these strategies (or one is sufficient to become resistant), but others use multiple strategies, and it is likely that further strategies for the avoidance of *M. xanthus* predation are yet to be discovered. While some predation resistance
strategies are adopted by multiple prey, the molecular mechanisms by which they implement each strategy are very different. The genes responsible for some mechanisms could conceivably be transferred between prey while maintaining their function (for instance, transformation with yjiC or acquisition of the bacillaene biosynthetic gene cluster), but other more complex strategies are less likely to be transferable (e.g., megastructure formation and the ability to sporulate).

At the moment, we cannot predict from their genomes whether any particular combination of prey and myxobacterial predator will demonstrate predation, or whether the prey will manifest any particular survival strategy. Perhaps attempting to acquire such understanding is a fruitful task given the complexity of the phenomena involved; however, it is tempting to try. If we can discover at the genomic level how prey and predator genes interact to manifest observed predator-prey phenotypes, we might be able to elevate our understanding of predation resistance mechanisms from anecdotal to universal – a foundational goal of comparative genomics [Whitworth, 2008].

To date, most studies of prey susceptibility have been descriptive or have investigated individual genes that confer the particularly strong selective advantages during M. xanthus predation. We believe it is desirable to take a holistic approach to screen entire genomes for genes that make a quantifiable contribution to predation resistance. While several experimental approaches to achieve this are widely used, we took advantage of a pre-existing library of defined transposon insertion mutants of Pseudomonas aeruginosa, which includes strains carrying disruptions of nearly 4,600 genes [Liberati et al., 2006].

P. aeruginosa is a Gram-negative flagellated rod-shaped bacterium that is ubiquitous in the environment and commonly found in soil, in water and on plant and animal tissues. It is a facultative anaerobe capable of respiring nitrate and can grow both planktonically or attached to a surface, forming a biofilm at high cell densities [Sønderholm et al., 2017]. It is also an opportunistic human pathogen, causing infections when the host’s normal immune defences are disrupted. With the rise in antibiotic-resistant strains, it has been defined by the World Health Organisation as a highest-priority pathogen.

The large genome of P. aeruginosa provides it with metabolic versatility and an array of pathogenic mechanisms. It causes infection of burn wounds, often resulting in invasive infections sepsis and death, while respiratory infection is the leading reason of morbidity in cystic fibrosis patients [Dolan, 2020]. P. aeruginosa PA14 is a clinical isolate obtained originally from a burn patient, which displays pathogenicity in a variety of genetically tractable model hosts and mice [Mahajan-Miklos et al., 2000]. Myxobacteria are generally able to prey upon P. aeruginosa [Livingstone et al., 2017], and understanding how myxobacterial predators kill this prey (and how the prey resists predation) might allow the development of predation-inspired therapies for treating P. aeruginosa infections [Pérez et al., 2020]. We therefore screened mutants of P. aeruginosa strain PA14 for genes which on disruption affected predatory efficiency during predation by M. xanthus.

**High-Throughput Screening of Prey Susceptibility Genes**

To identify candidate “prey” genes, we screened a library of P. aeruginosa strain PA14 transposon insertion mutants (e.g., mutated “predation resistance” genes) for mutations that affected susceptibility to predation by M. xanthus. In a 96-well plate format, defined mutations of strain PA14 were co-incubated with a fluorescent strain of M. xanthus expressing mCherry (strain EH715).

**Bacterial Strains**

Primers mCherry_F_NdeI (ATTCACATATGGTT-GAGCAAGGGCGAG) and mCherry_R_EcoRI (TAC-TATGAAATTCTACTTGTACAGCTGTCGCTCCATG) were used to amplify mCherry from pMKK227 [Koch et al., 2011]. The product was inserted into pMR3679 under the control of a vanillate-inducible promoter [Ineista et al., 2012], producing pJS007. To obtain strain EH715, around 1 µg of pJS007 was used to transform M. xanthus strain DK1622 [Kaiser, 1979] by electroporation. This strain was maintained in the presence of kanamycin at 100 µg/mL. Incubation of EH715 in 50 µM vanillate for 18 h gave suitable induction of red fluorescence.

The library of P. aeruginosa strain PA14 transposon insertion mutants used here was the PA14NR Set, described by Liberati et al. [2006]. Most of the mutants contained insertions of MAR2xT7 (a derivative of Himar1 – a mariner family transposon), while a minority contained TnphoA insertions. The PA14NR Set comprises a total of 5,459 mutants, corresponding to 4,596 predicted strain PA14 genes (77% of predicted genes).

**Predator-Prey Assays in 96-Well Plates**

M. xanthus EH715 was grown to late exponential phase in DCY (20 g casitone, 2 g yeast extract, 10 mL 1 M...

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**Tris pH 7.4, 8 mL 1 M MgSO₄ per L) supplemented with 50 µM kanamycin and 50 µM vanillate. Cells were harvested by centrifugation and resuspended in TM buffer (10 mL 1 M Tris pH 7.4, 8 mL 1 M MgSO₄ per L). *P. aeruginosa* strains were propagated by inoculating into 96-well plates containing LB (10 g tryptone, 10 g NaCl, 5 g yeast extract per L) and incubated overnight at 30°C. Pre-dation assays were set up in singlicate 96-well plates with each well containing an equal volume of *P. aeruginosa* culture and strain EH715 in buffer. Assay plates were incubated at 30°C, and OD₆₀₀ (optical density at 600 nm) and fluorescence (excitation at 587 nm, emission at 610 nm) were measured after 0 and 24 h. Controls included “cell-free” (just buffer and LB) and “predator-only” wells.

**Data Interpretation**

We expected that after a period of incubation, increased susceptibility to predation of strain PA14 mutants by EH715 would be manifested as greater fluorescence compared to controls. This is because the suscep-
Susceptible strain PA14 mutants would be relatively less abundant, while the population of fluorescent strain EH715 would be more abundant, as compared to the controls. As strain PA14 grows much faster than EH715, we also expected that after incubation, a lower OD$_{600}$ than controls might indicate greater predatory activity. From amongst the 63 plates, 192 mutants were selected which exhibited particularly high fluorescence and/or low optical density measurements. Nucleotide sequences of the mutated genes were translated and annotated with Prokka [Seemann, 2014] and then the annotation was enriched with EggNOG 5.0 [Huerta-Cepas et al., 2019]. This approach yielded 85 candidate “prey proteins,” and encouragingly, some of those proteins were identified from multiple plates (AstB, TrpE and YhhJ). The COG (clusters of orthologous groups) categories [Galperin et al., 2019] to

| Function                          | Identifier | Description                                                                 |
|-----------------------------------|------------|-----------------------------------------------------------------------------|
| Amino acid metabolism             | AspH       | Aspartyl/asparaginyl beta-hydroxylase                                       |
| Amino acid metabolism             | AstB       | N-succinylarginine dihydrofase                                              |
| Amino acid metabolism             | Ivd        | Isovaleryl-CoA dehydrogenase 1                                              |
| Amino acid metabolism             | PhzB       | Chorismate binding enzyme                                                    |
| Amino acid metabolism             | PuuA       | Glutamine synthetase, catalytic domain                                       |
| Amino acid metabolism             | TrpE       | Anthranilate synthase component 1                                            |
| Chemotaxis                        | CheA       | Chemotaxis protein histidine kinase                                          |
| Chemotaxis                        | FliH       | Flagellar assembly protein                                                   |
| Chemotaxis                        | FliP       | Flagellar biosynthetic protein                                               |
| GGDEF protein                     | DR97_5,071 | GGDEF diguanylate cyclase domain protein                                    |
| GGDEF protein                     | PleD       | Response regulator with GGDEF diguanylate cyclase domain                    |
| Ion transport                     | FilE       | Cation-efflux pump                                                          |
| Ion transport                     | HisJ       | Ligated ion channel L-glutamate- and glycine-binding site                   |
| Ion transport, redox              | KefB       | Glutathione-regulated potassium-efflux system protein                       |
| LPS production                    | AraT       | Transfers L-Ara4N to lipid A                                                 |
| LPS production                    | LptC       | Lipopolysaccharide export system protein                                     |
| Protease                          | ApeB       | Aminopeptidase I zinc metalloprotease (M18)                                 |
| Protease                          | PilP       | Df-1/Pilp family intracellular protease                                     |
| Protein secretion                 | BabA       | Part of the outer membrane protein assembly complex                         |
| Protein secretion                 | FhaC       | Member of a two-partner secretion pathway (TPS)                             |
| Protein secretion                 | Hcp        | Type VI secretion system effector                                           |
| Protein secretion                 | TadB       | Type II secretion system (T2SS), protein F                                 |
| Protein secretion                 | YscK       | YOP proteins translocation protein K (type III secretion)                   |
| Protein secretion                 | YscW       | Type III secretion system lipoprotein chaperone                             |
| Quorum quenching                  | PvdQ       | Acyl homoserine lactone acylase                                             |
| Redox processes                   | Gor        | Glutathione reductase                                                       |
| Redox processes                   | GrxD       | Glutaredoxin                                                                |
| Redox processes                   | NuoC       | NADH-quinone oxidoreductase subunit C/D                                    |
| Redox processes                   | NuoE       | Thioredoxin-like [2Fe-2S] ferredoxin                                        |
| Transcription factor              | ArsR       | Helix_turn_helix domain protein, arsenical resistance operon repressor      |
| Transcription factor              | HcrR       | Helix-turn-helix domain protein, rpiR family                                |
| Transcription factor              | OhrR       | Helix_turn_helix domain protein, multiple antibiotic resistance protein      |
| Transcription factor              | PilR       | Transcription regulatory protein, Fis family                                |
| Transport                         | CcmA       | ABC transporter, ATP-binding protein                                         |
| Transport                         | DctA       | C4-dicarboxylate transport protein                                          |
| Transport                         | DctQ       | Tripartite ATP-independent periplasmatic transporter component              |
| Transport                         | OpuCC      | Substrate binding domain of ABC-type glycine betaine transport system       |
| Transport                         | RbsC       | Binding-protein-dependent transport system permease family protein          |
| Transport                         | YdhC       | Sugar (and other) transporter                                               |
| Transport                         | YehB       | Binding-protein-dependent transport system inner membrane component         |
| Transport                         | YhhJ       | ABC-2 family transporter protein                                            |
which “prey proteins” were assigned are indicated in Figure 1. “Prey protein” membership of COG categories was significantly different from that of strain PA14 genome-encoded proteins ($\chi^2$ test, $p < 0.01$).

Individual COG categories enriched (more than 3-fold) among “prey proteins” were A (RNA processing and modification), N (cell motility), O (post-translation modification, protein turnover, and chaperones), and V (defence mechanisms), while COG category J (translation, ribosomal structure and biogenesis) was more than 3-fold depleted compared to the genome-encoded protein set. As apparent in Figure 1 and as might be expected, prey proteins were generally depleted (>2-fold) in the COG super-category “information storage and processing” (which comprises individual COG categories A, B, J, K and L).

**Functional Roles of “Prey Proteins”**

Many of the candidate “prey proteins” had overlapping functional roles or common mechanisms, and/or had roles which would be expected to impact on predation resistance (Table 1). For instance, 5 proteins were motility related, including 2 flagella proteins, 2 flagella assembly proteins, and the chemotaxis regulator CheA – presumably being able to swim away from predators reduces the impact of predation. Also present in the list of “prey proteins” were several proteins involved in protein secretion systems (6 proteins), amino acid metabolism (6 proteins), transport (12 proteins), proteases (2 proteins), and cyclic-di-GMP signalling (2 proteins), which might also hint at general aspects of predator resistance. For instance, cyclic-di-GMP signalling promotes biofilm formation [Güvener and Harwood, 2007].

Other “prey proteins” suggest that *M. xanthus* might attack *P. aeruginosa* with cationic antimicrobial peptides and/or metal/oxidative stress. Five “prey proteins” are involved in processes affecting redox homeostasis, 4 proteins are ion transporters (including a redox-regulated transporter and a cation efflux pump), while a further 4 are transcription factors (including a likely arsenate resistance regulator and a multi-drug resistance regulator). Two prey proteins are involved in LPS (lipo polysaccharide) production, including a glycosyl transferase, which modifies lipid A conferring resistance to cationic antimicrobial peptides and polymyxin. Metal/oxidative stress is known to mediate myxobacterial predation of *S. meliloti* [Contreras-Moreno et al., 2020] and many peptides produced by myxobacteria have potent antimicrobial activity [Wenzel and Müller, 2009]. The electron transport chain is a known target of myxobacterial predation, and of ubiquitous myxobacterial antibiotics such as the myxalamides [Gerth et al., 1983; Livingstone et al., 2018c].

Also within the list of “prey proteins” was PvdQ, an acylase involved in both siderophore production and acyl homoserine lactone (AHL) degradation in *P. aeruginosa*. AHLs are widely used bacterial quorum signalling molecules, which promote biofilm formation in *Pseudomonas*. While biofilm formation is thought to provide protection from predation, *M. xanthus* is able to sense the AHLs produced by prey, stimulating its predatory activity [Lloyd and Whitworth, 2017]. Potentially, PvdQ might increase resistance to predation by preventing AHL-mediated eavesdropping by predators, as well as by stimulating antibiotic resistance [Wang et al., 2011].

It would now be interesting to characterise the “prey proteins” in more detail, ideally using engineered non-polar mutations. Studies could then be undertaken on those mutants to assess their phenotype, gene expression patterns and the mechanism by which they conferred any selective advantage when subjected to predatory attack.

**Conclusion**

The bacterial predator-prey relationship is multi-facetted, seemingly unique to each combination of predator and prey organism. However, it remains likely that identification of specific prey responses to predation will illuminate general features of predation resistance responses. We screened nearly 5,500 mutants of *P. aeruginosa* (covering 77% of genes in the genome) to identify the genes for proteins conferring resistance to predation by *M. xanthus*. The candidate “prey proteins” identified were enriched for particular COGs and many had overlapping functional roles. The nature of the “prey proteins” identified suggests that effective resistance of predation by *P. aeruginosa* requires systems for resisting metal/oxidative stress and for detoxifying antibiotics and antimicrobial peptides, as well as a functional motility system.

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Author Contributions

D.E.W., M.T.S. and N.P.T. devised the project, D.E.W., M.T.S. and N.P.T. supervised the screening experiment, J.S. generated EH715 under the supervision of E.H., N.S. performed the screening experiments and data analysis, all authors edited and approved the final version of the manuscript.

Conflict of Interest Statement

There are no conflicts of interest to declare.

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