EXAMINING c-di-GMP AND POSSIBLE QUORUM SENSING REGULATION IN Pseudomonas fluorescens SBW25: LINKS BETWEEN INTRA- AND INTER-CELLULAR REGULATION BENEFITS COMMUNITY COOPERATIVE ACTIVITIES SUCH AS BIOFILM FORMATION


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Bacterial success in colonizing complex environments requires individual response to micro-scale conditions as well as community-level cooperation to produce large-scale structures such as biofilms. Connecting individual and community responses could be achieved by linking the intracellular sensory and regulatory systems mediated by bis-(3′-5′)-cyclic dimeric guanosine monophosphate (c-di-GMP) and other compounds of individuals with intercellular quorum sensing (QS) regulation controlling populations. There is growing evidence to suggest that biofilm formation by many pseudomonads is regulated by both intra and intercellular systems, though in the case of the model Pseudomonas fluorescens SBW25 Wrinkly Spreader in which mutations increasing c-di-GMP levels result in the production of a robust cellulose-based air-liquid interface biofilm, no evidence for the involvement of QS regulation has been reported. However, our recent review of the P. fluorescens SBW25 genome has identified a potential QS regulatory pathway and other QS–associated genes linked to c-di-GMP homeostasis, and QS signal molecules have also been identified in culture supernatants. These findings suggest a possible link between c-di-GMP and QS regulation in P. fluorescens SBW25 which might allow a more sophisticated and responsive control of cellulose production and biofilm formation when colonising the soil and plant-associated environments P. fluorescens SBW25 normally inhabits.

KEYWORDS: cyclic dimeric guanosine monophosphate (c-di-GMP), acyl homoserine lactone (AHL), bacterial cellulose, biofilms, Pseudomonas, quorum sensing (QS).

The formation of biofilms is often considered an exemplar of cooperation within bacterial communities, though cooperation is always balanced by competition within the developing community [1, 2]. Biofilm formation involves the coordinated attachment of founder cells to an appropriate...
Recently a linkage has been reported between c-di-GMP and QS regulation in other pseudomonads [12-14] and other bacteria (e.g. Burkholderia and Vibrio [15, 16]), and this has driven us to re-examine the *P. fluorescens* SBW25 genome for potential QS-related genes and culture supernatants for QS signals. Here we suggest how *P. fluorescens* SBW25 might use QS regulation to further refine cellulose production and biofilm formation in natural environments (as opposed to *in vitro* microcosms), and consider more generally the benefits of linking the sensory and regulatory systems of individuals with the QS-based cooperative activity of populations and communities.

**Pseudomonas fluorescens SBW25 and the rise of the Wrinkly Spreaders**

*P. fluorescens* SBW25 is a soil and plant-associated pseudomonad originally isolated from a sugar beet (*Beta vulgaris*) leaf [17], and is able to colonise the leaf and root surfaces of a range of different cultivated plants and weeds. The 6.7 Mbp genome was initially mapped by Southern analysis before the complete genome sequence was obtained [17-19]. *P. fluorescens* SBW25 has been used to investigate plant and soil-microbe interactions, regulation and metabolism, biofilm formation and experimental evolution studies [10, 11, 20-22]. In the latter, bacterial populations are grown in artificial environments known as microcosms and subject to varied selective pressures [20, 23-25]. One aim of this is to provide mechanistic explanations linking adaptive mutation and the underlying molecular biology of an organism with the fitness improvements of ‘evolved’ mutants compared to the ancestral strain.

Wild-type *P. fluorescens* SBW25 produces smooth round colonies on agar plates, and will grow throughout the liquid column of static microcosms over a period of several days. However, during this time the expanding population will radiate and accumulate a variety of mutants including WS [10, 11]. These are a class of mutants characterised by a flattened, wrinkled colony morphology and the ability to produce a robust and well-attached biofilm at the A-L interface of static microcosms (these characteristics are collectively known as the WS phenotype) (Fig. 1). Although most biofilm research has focussed on liquid-solid surface (L-S) interface biofilms formed by bacteria in laminar flow cells or in microtitre plates [26-28], the ability to form biofilms at the A-L interface is common amongst...
Fig. 1. The Wrinkly Spreader produces a biofilm in static microcosms and wrinkled colonies on agar plates. Developing populations of P. fluorescens SBW25 in static microcosms will radiate and produce a number of mutants including the Wrinkly Spreaders (WS). (A) Wild-type P. fluorescens SBW25 grows in the liquid column of static microcosms (left) whereas Wrinkly Spreaders produce a robust biofilm at the air-liquid (AL) interface. The microcosms are 30 ml Universal glass vials with 6 ml liquid growth medium and are typically incubated for 3 days to allow biofilm-formation. (B) The Wrinkly Spreader forms wrinkled colonies readily distinguished from wild-type round and smooth colonies on agar plates. (C) Both the WS biofilm and colony contain extensive networks of extracellular cellulose fibres. Shown here is a confocal laser scanning microscopy (CLSM) image of a WS biofilm sample stained with Calcofluor to visualise cellulose (blue) surrounding green fluorescent protein (GFP)-expressing WS cells (green). These images were taken using a Carl Zeiss LSM-510 CLSM and the scale bar indicates 5 µm.

environmental pseudomonads [29-32]. Many abiotic and biotic factors influencing L-S and A-L interface biofilm-formation are the same and probably impact on other types of bacterial aggregation too [8].

Wrinkly Spreaders are adaptive mutants in static microcosms containing King’s B (KB) liquid growth medium [33], and show improved competitive fitness (W) compared to the wild-type P. fluorescens SBW25 ancestor and other non-biofilm-forming competitors (W = 1.5 – 2.5 [9, 34, 35]) (W is the ratio of Malthusian parameters for continuously growing populations and by definition, adaptive mutants will have a competitive fitness (W) advantage over the ancestor, i.e. W > 1; when W = 1, the two strains are neutral with respect to one another, and when W < 1 the mutant is at a disadvantage). However, in shaken microcosms where biofilms cannot form, or on agar plates where the WS phenotype is unstable, the fitness of Wrinkly Spreaders is much reduced and at times even lower than the ancestor (W = 0.1 – 1.0 [34-36]).

The evolutionary and ecological success of the Wrinkly Spreader is explained by the ability to rapidly colonise the A-L interface of static microcosms by biofilm-formation, allowing it to intercept oxygen (O₂) diffusing into the liquid from the air above [37, 38]. P. fluorescens SBW25 is an aerobe that grows more slowly under micro-aerobic conditions. The initial colonists of static microcosms rapidly establish a strong O₂ gradient through the up-take of O₂, effectively dividing their environment into an O₂-rich zone at the A-L interface and an O₂-depleted zone below (in this respect, wild-type P. fluorescens SBW25 is responsible for niche construction [11, 39]). Since growth is O₂-limited in these microcosms, bacteria located at the A-L interface will grow faster than those below. As Wrinkly Spreaders can colonise the A-L interface without the constant need for swimming by biofilm-formation, better access to O₂ and higher growth rates provide the fitness advantage, allowing them to outcompete the ancestor and other non-biofilm-forming competitors [37]. Growth at the A-L interface clearly impacts on metabolism, as bacteria recovered from within and underneath biofilms can be differentiated by Raman spectroscopy [40]. In ecological terms, the A-L interface can also be thought of as an unoccupied niche and colonisation of this space by biofilm-forming Wrinkly Spreaders would avoid the constant competition between the wild-type P. fluorescens SBW25 ancestor and other non-biofilm-forming competitors in the O₂-depleted niche below.

Wrinkly Spreader lineages (i.e. subpopulations) arise in communities, founded by wild-type P. fluo-
rescens SBW25 and containing numerous other mutant lineages, in which there is both inter-lineage and intra-lineage completion. This type of complex interaction is described by the Red Queen hypothesis in which all genotypes are locked into constant competition against one another [41]. In this type of community, the Wrinkly Spreader fitness advantage is negative-frequency dependent: at low relative numbers they have a significant advantage and are largely competing against non-biofilm–forming lineages. In this situation the cooperative activity within the WS lineage that forms the biofilm might be better explained by the Black Queen hypothesis where genotypes are mutually dependent [42]. However, as Wrinkly Spreader numbers grow, completion within the WS lineage increases, reducing the fitness advantage for all Wrinkly Spreaders.

In bacterial social evolution theory, biofilms are considered common or public goods in which individual bacteria are expected contribute to the costly EPS matrix through cooperative social behaviour [1, 2, 43, 44]. However, this apparent cooperation might actually result from one or more of three mechanisms, including the response of individual bacteria to environmental stimuli, a community response to QS regulation, and mutation overriding normal regulatory systems (Fig. 2). This cooperation is explained by Hamilton’s inclusive fitness (or kin selection theory) which states that cooperation evolves between genetically related individuals [44], and could be facilitated by commonality of sensory systems and QS regulation in which all sufficiently-related individuals respond to the same signals, or to the clonal expansion of a single mutant lineage.

Unlike many other pseudomonads, P. fluorescens SBW25 has not been shown to use QS regulation and therefore appears inquorate. Perhaps uniquely, biofilm-formation by Wrinkly Spreaders has been shown to be the result of genetic mutation and clonal expansion, rather than by the cooperative effort of a community.

Unravelling the underlying molecular biology of the Wrinkly Spreader

Mini-transposon mutagenesis of the archetypal Wrinkly Spreader, a particular WS isolate also known as the Large Spreading Wrinkly Spreader (LWS) and strain PR1200, was used to identify the key genes and regulatory pathways responsible for the WS phenotype [10, 34, 35]. Two key operons, referred to as wss (WS structural operon) and wsp (WS phenotype regulatory operon), were found to be essential for the WS phenotype.

The ten gene wss operon (wssA–J) encodes a bacterial cellulose synthase, with the core subunits encoded by WssB, WssC, WssD, and WssE [34]. Cellulose was first reported to be produced by Acetobacter xylinus (now known as Gluconacetobacter hansenii) in a thick, buoyant biofilm, and it is also

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**Fig. 2.** Three routes leading to the formation of biofilms. Biofilms might be the result of communities of bacteria all responding to the same environmental stimuli (top), quorum sensing (QS) regulation (middle), or mutation (bottom). The initiation of biofilm-formation might be the result of the detection of a signal by all individuals indicating that conditions are appropriate for settlement and growth. Alternatively, QS signals produced by a few individuals might then commit the entire community to biofilm-formation, or mutation in one individual could induce the appropriate behaviour. In the first two situations, variation within the community (from minor genotype differences between lineages or subpopulations to species differences, indicated by white, grey and black fillings) will result in a heterogeneous biofilm-founding population. However, in the case of mutation, the biofilm will be produced by the clonal expansion of the mutant lineage with no heterogeneity in the early stages of biofilm-formation.
produced in substantially lower amounts by enteric bacteria such as *Escherichia coli* K-12 and *Salmonella enteritidis* 3934 [32, 46, 47]. In *Escherichia, Gluconacetobacter* and *Salmonella* spp., the genes required for cellulose production are contained within bacterial cellulose synthase (*bcs*) operons similar to the *P. fluorescens* SBW25 *wss* operon [34], though *bcs* operon duplications and other rearrangements are common [48]. Unlike most *bcs* operons however, the *P. fluorescens* SBW25 *wss* operon also encodes proteins involved in the acetylation of cellulose (*WssF–I*) [34, 49, 50] which are also found in several closely-related *Pseudomonas* plant pathogens as well as in the bird pathogen *Bordetella avium* 197N [49, 51, 52]. This unusual modification of the polymer by *P. fluorescens* SBW25 was proven by purifying EPS from WS biofilms and subsequent chemical composition and linkage analysis [49].

Cellulose fibres and clumps could also be visualised in WS biofilm samples by fluorescent or confocal laser scanning microscopy [32, 49, 53] (Fig. 1). Mini-transposon insertions in *wssA–E* abolished cellulose production and the WS phenotype, whereas insertions in *wssF–I* resulted in the production of un-acetylated cellulose and a partial WS phenotype [34, 45, 49, 54]. In contrast, mutation of *wssJ* had no impact, suggesting that the last gene in the *wss* operon may be redundant [45]. Furthermore, like several other pseudomonad biofilms, the WS biofilm contains extracellular DNA which may also contribute to biofilm strength and resilience (O. Moshynets et al. unpublished observations; to be published elsewhere).

The seven gene *wsp* operon (*wspA–F, R*) encodes a chemosensory-like complex in which the receipt of an unknown environmental signal results in the activation of WspR, a GGDEF-domain containing diguanylate cyclase (DGC) response regulator [34, 55, 56]. Activated WspR synthesises c-di-GMP from GTP [57]. C-di-GMP had been earlier identified as a regulator of cellulose synthase activity in *Glucgonacetobacter* spp. [58] and is an important intracellular regulator found in many bacterial species. Intracellular concentrations of c-di-GMP is mediated by the opposing actions of two types of enzymes: DGCs containing GGDEF-domains that synthesize c-di-GMP, and phosphodiesterases (PDEs) containing EAL or HD-GYP domains that degrade c-di-GMP. Levels of c-di-GMP influence a wide range of phenotypes including cellular responses that effect pathogenesis, motility, secretion, cytotoxicity, synthesis of secondary metabolites and environmental stress adaptation as well as QS [59-62]. Many of these phenotypes are also regulated by the Gac/Rsm master regulon, and recently links between the c-di-GMP and Gac/Rsm networks have been identified in *P. aeruginosa* PAK [63] (the Gac/Rsm regulon is also present in *P. fluorescens* SBW25 [64]). Modulation of c-di-GMP is also associated with control of motility, biofilm formation through adhesion and EPS production, including cellulose, and other social behaviours.

The nature of the Wsp complex and the link to cellulose production through c-di-GMP suggested a model explaining the WS phenotype [56]. In this, mutations in Wsp subunits that increased WspR activation in the absence of the normal environmental signal were predicted to result in the expression of the WS phenotype (Fig. 3). The model was confirmed by the identification of a *wspF* (S301R) mutation in the archetypal Wrinkly Spreader, close to the active site and predicted to reduce the methyl esterase activity of the subunit required to limit WspR activation. Allelic replacement experiments in which the *wspF* mutation was changed to the wild-type sequence and vice versa confirmed that this mutation was sufficient to explain the WS phenotype [56]. Further analysis of other independently-isolated Wrinkly Spreaders identified additional mutations in *wspF* and *wspE*, as well as in several other DGCs or DGC-associated loci (*aws* and *mws*), all of which are predicted to result in increased levels of c-di-GMP and the expression of the WS phenotype [56, 65, 66] (exogenous Fe can non-specifically induce cellulose production by *P. fluorescens* SBW25 and the formation of a weak, poorly attached biofilm [67]). A total of 39 GGDEF-domain containing proteins have been identified in *P. fluorescens* SBW25 genome [18], and the activation of some of these by intra- and inter-genic mutations in negative regulators of the DGC or the DGCs themselves can also activate the WS phenotype in a Δ*aws* Δ*mws* Δ*wsp* triple-deletion mutant [68] (loss of function mutations often lead to bacterial adaptation to new conditions [69]). This strongly suggests that the c-di-GMP connection between the Wsp chemosensory-like complex and the WS cellulose synthase is not unique, and any mutation occurring in radiating populations of *P. fluorescens* SBW25 in static microcosms that up-regulates c-di-GMP levels might result in Wrinkly Spreaders (mutations will arise and fix with a frequency corresponding to their ease of occurrence and fitness relative to other Wrinkly Spreader competitors).
The WS model is not appropriate for the regulation of cellulose production by *P. fluorescens* SBW25 in natural environments

Although the WS phenotype, including both competitive fitness and wrinkleality, is variable amongst independently-isolated Wrinkly Spreaders [70], it appears that WS activating mutations are binary in nature and result in the unregulated over-production of cellulose and biofilm formation. In the soil and plant-associated environments colonised by *P. fluorescens* SBW25, greater responsiveness to environmental stimuli and flexibility in behaviour is required, and both cellulose production and biofilm formation are expected to be controlled by a complex network of regulatory systems. Some of these have been identified with wss operon transcription regulated by the DGCs AwsR and WspR, as well as by the master regulators of alginate (AlgR) and flagella (FleQ), during the colonisation of the Sugar beet rhizosphere where the ability to produce cellulose provides a fitness advantage for *P. fluorescens* SBW25 [71, 72]. However, the environmental stimuli these systems respond to have yet to be identified, and it is notable that QS regulation, involved in the regulation of EPS production and biofilm formation in many pseudomonads, has not been reported in *P. fluorescens* SBW25, nor have QS genes been identified in the analysis of Wrinkly Spreaders or *P. fluorescens* SBW25 rhizosphere colonisation.

Challenging the view that *P. fluorescens* SBW25 is inquorate

Recently, a link between QS regulation and c-di-GMP has been identified in pseudomonads [12-14]. It is generally assumed that the primary quorum signalling molecules or autoinducers that bind their cognate receptors to directly or indirectly activate gene expression used by Gram-negative bacteria are N-acyl-homoserine lactones (AHLs) [3-6]. Pseudomonads have been reported to produce a variety of different AHLs controlling the production of virulence factors, EPS, and motility that are important for bacterial adaptation and colonisation in a range of environments [73-76]. In *P. aeruginosa* PA14, the QS regulator LasR induces the expression of TpbA which in turn represses the DGC activity of TpbB, resulting in lowered c-di-GMP levels [77] (a slightly different TpbB analogue known as YfiR has also been investigated in *P. aeruginosa* PA01 [78]). This link drove us to examine the *P. fluorescens* SBW25...
genome for QS components which might also be linked to c-di-GMP homeostasis and therefore to cellulose synthesis and biofilm–formation (this analysis will be published elsewhere).

We have identified a potential AHL synthase homologue (PFLU0008), as well as homologues of LasR (PFLU1386) and TpbA (PFLU0050) in *P. fluorescens* SBW25, and hypothesize that they may form a QS regulatory pathway acting through WspR or other DGCs to repress cellulose production (Fig. 4). First, the presence of a HdtS-like AHL synthase homologue [73] suggests that *P. fluorescens* SBW25 might produce AHLs under the right conditions, and in preliminary experiments, we have detected at least two different AHL molecules in culture extracts using a standard AHL bio-reporter assay [79, 80]. Secondly, although *P. fluorescens* SBW25 was not originally annotated as having a LasR homologue [19], PFLU1386 is a putative regulator belonging to the same LuxR family of transcriptional regulators. Despite a poor level of homology (32% ID, amino-acid level sequence identity), it has been possible to model the 3D structure of PFLU1386 using a LasR–AHL co-crystal template and dock several AHLs including DHL (N-dodeconyl-L-homoserine lactone, C12-HSL) and 3-O-DHL (N-(3-oxo-dodecanoyl)-L-homoserine lactone, 3-O-C12-HSL) (Fig. 5). This suggests that PFLU1386 might bind AHLs and induce the transcription of as-yet unidentified QS-dependent genes. Finally, the 3D structure of PFLU0050 can also be modelled using a TpbA crystal template [81], suggesting that these two proteins share a high degree of structural similarity, and that PFLU0050 may also show AHL-dependent tyrosine phosphatase activity and regulate a cognate TpbB-like DGC. Using the *P. aeruginosa* PA14 TpbB amino-acid sequence, we identified a possible TpbB homologue, PFLU5210 (62% ID), which had previously been identified as AwsR which regulates wss operon transcription [65, 72]. PFLU0050 may also regulate the activity of AwsR as well as more distantly-related DGCs (e.g. WspR which shares 35% ID with TpbB) to control c-di-GMP levels.

Our analysis of the *P. fluorescens* SBW25 genome has identified several other proteins that might play a role in QS regulation including some associated with alpha-hydroxy ketones (AHKs) which we have detected *P. fluorescens* SBW25 culture extracts using the Janthinobacterium sp. bio-reporter assay [82]. We have identified two potential AHK synthases (PFLU5220 and PFLU5614) and an AHK sensory protein (PFLU1736) in the *P. fluorescens* SBW25 genome, but have not identified a potential AHK-dependent regulator. This suggests that the potential *P. fluorescens* SBW25 AHK pathway is more similar to the Vibrio pathway which also lacks a regulator, than for example, the Legionella pathway which includes one [83].

**Fig. 4.** *P. fluorescens* SBW25 may utilise QS to regulate cellulose production. A bioinformatics analyses of *P. fluorescens* SBW25 proteins has identified potential homologues which may act together to produce a QS regulatory pathway that might reduce the activity of the Wss cellulose synthase by modulating c-di-GMP levels. The homologue of HdtS (PFLU0008) is predicted to synthesise AHLs which then bind to the LasR homologue (PFLU1386) in a QS manner. Bound PFLU1386 is predicted to induce transcription of the PFLU0050 gene resulting in the expression of a DGC regulator homologous to TpbA. PFLU0050 may then regulate the activity of the TpbB homologue (AwsR, PFLU5210). PFLU0050 might also repress the activity of other DGCs, possibly even WspR, to reduce c-di-GMP levels and repress cellulose production.

**Linking individual behaviour with community action**

Although biofilm–formation can result from mutation and the clonal expansion of adaptive mutant lineages, or through QS regulation, there are benefits in linking individual (intracellular) regulatory networks with community-based decision and
enforcement (Table). Bacteria have multiple and interacting regulatory networks which allow the appropriate response to environmental factors. For example, in pseudomonads, Fe-repressible and anaerobically (low \(O_2\))-induced small RNA (sRNA) repressors interact with QS regulation, whilst other sRNAs sequester RNA-binding proteins that regulate global translational repressors, including central elements controlling motility, biofilm-formation, QS regulation, the expression of virulence factors, and the utilisation of carbon sources, some of which are also regulated by c-di-GMP levels [4, 13, 84]. Although bacteria within a fairly uniform environment might be expected to show the same behaviour, micro-heterogeneity in local conditions, variation in cellular status and stochastic responses would lead to phenotypic heterogeneity within populations [85] and mean that not all individuals would be in an appropriate condition to invest in community effort or to benefit from the results of such endeavours. Indeed, if all individuals do not benefit, cooperation can breakdown leading to a public goods dilemma: in both \(P. aeruginosa\) PA01 and Wrinkly Spreader populations, the presence of cheaters reduces overall fitness, growth and biofilm formation [86, 87] (though not in all cases, e.g. [7]).

However, links between intracellular regulators such as c-di-GMP with community-based (intercellular) QS regulation provide a mechanism allowing individuals to opt-in and opt-out of cooperative activity. In particular, individual sensing and decision allows opting in to the early stages of biofilm formation when benefits might be expected in the future (i.e. providing a fitness benefit to that particular lineage), and opting out when biofilms mature or are damaged, when further benefits to cooperative activity are no longer likely. One example of the need for an opt-out might be found in Ancestor’s inhibition, where the constant production of EPS and growth at the surface of a biofilm pushes earlier generations into the middle of the structure where conditions may no longer be adequate [88,89]. In \(Escherichia coli\) K12, an oxygen-sensing phosphodiesterase and associated diguanylate cyclase have been described [81], suggesting a mechanism allowing \(O_2\)-stressed individuals in biofilms to stop EPS production and de-repress mobility by altering c-di-GMP levels; in \(P. aeruginosa\) PA01, the anaerobic regulator Anr may also interact with QS regulation in response to altering \(O_2\) levels [91]. Finally, QS regulation may impose a direct cost under some conditions, as many environmental and clinical isolates of \(P. aeruginosa\) have been found to have defective QS regulatory systems [92].

**Concluding statement**

Our desire to understand the underlying molecular biology of the novel biofilm-forming Wrinkly Spreader, an adaptive mutant isolated from radiating populations of \(P. fluorescens\) SBW25 in static microcosms, uncovered a pathway linking mutations affecting c-di-GMP levels to the production of cellulose which forms the main matrix component of the biofilm. Although this analysis was aided by availability of the genome sequence, our current review of the coding potential of the genome suggests that \(P. fluorescens\) SBW25 may also have QS regulation capability. The interaction between intracellular regulatory networks involving key regulators such as...
c-di-GMP with intracellular QS regulation provides additional benefits to both individual bacteria and communities during biofilm formation. The acquisition of genome sequences is commonplace today and is often the starting point in investigating complex bacterial regulatory networks and behaviours. However, retrospective inspection of genomes can reveal additional elements which were not considered earlier, adding to our understanding of how bacteria interact with their environment and with each other.

The new challenge in understanding biofilm formation by *P. fluorescens* SBW25 and related pseudomonads is to determine the relative importance of environmental signals, QS regulation and mutation in the colonisation of new environments and adaptation over evolutionarily-relevant time scales. A better understanding of this will be of particular relevance to medical microbiology, where bacteria are able to colonise tissues in chronic infections by overcoming the immune system and avoiding antibiotics through biofilm formation and mutation.

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**Table. Benefits of linking individual regulatory networks with community-based decision and enforcement for the production of biofilms**

| Individual sensing and decision | To integrate various environmental chemical and physical signals to confirm that individual bacteria are in the appropriate circumstances and position for attachment and biofilm formation. C-di-GMP homeostasis linked to other networks to integrate oxygen sensing, metabolic status, energy reserves, nutrient availability, etc. Although individual bacteria can express biofilm matrix components, there is no defence against competitors (cheaters) using these and avoiding the cost of production. |
| Community decision and control | To implement the cooperative investment of all appropriately-positioned bacteria in the formation of a biofilm for mutual benefit. As the initial expression of quorum signals within a population of bacterial cells is stochastic, and if the population is also heterogeneous in terms of cell position and status, there is the chance that QS will force some members to cooperate when they might otherwise decide not to do so. |
| Individual and Community interaction | Regulatory interactions allow for the fine-tuning of individual and community decisions. Interactions allow individual bacteria to opt-in to community activity when they are ready and can benefit from this investment. Poorly-positioned or stressed bacteria can also opt-out from the inappropriate demands of the comfortable majority during the initiation of biofilm formation and in developed structures when individuals may no longer occupy optimal positions. Depending on the relative strengths of the individual and community regulatory systems, the interaction might also allow the adjustment of individual investment resulting from altered conditions, new construction costs resulting from damage, and prevent the development of cheaters. |
АНАЛИЗ c-di-GMP И МОЖЛИВОГО ПОЧУТТЯ КВОРУМУ В Pseudomonas fluorescens SBW25: ЗВ’ЯЗОК МІЖ ВНУТРІШНЬО-І МІЖКЛІТИВНОЮ РЕГУЛЯЦІЄЮ СПРИЯЄ КООПЕРАТИВНІЙ ПОВЕДІНЦІ В УГРУПОВАНИІ І ФОРМУванНІ БІОПЛІВКИ

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Успішність бактеріальної колонізації складних еконіш вимагає індивідуальної відповіді на зміни умов на мікрорівні так само як і кооперацію на рівні угруповування для продукції таких великомасштабних структур як біоплівки. Координація індивідуальних відповідей і відповідей угруповування може бути досягнута шляхом зв’язування внутрішньоклітинних сенсорних і регуляторних систем, опосередкованих біс-(3′,5′)-циклічним димерним гуанозинмонофосфатом (c-di-GMP) та іншими сполуками індивідіумів із міжклітинною регуляцією – почутием кворumu (ПК), що контролює популяцію. Накопичується все більше доказів того, що формування біоплівки багатьма псевдомонадами регулюється як внутрішньоклітинними, так і міжклітинними регуляторними системами, хоча в разі моделю Pseudomonas fluorescens SBW25 Wrinkly Spreader, у котрій мутація що підвищує рівні c-di-GMP, призводять до створення міцної целюлозної біоплівки на кордоні розділу фаз повітря–рідина, не було виявлено ніякого свідчення залучення кворумзалежної регуляції. Проте наш недавній огляд геному P. fluorescens SBW25 виявив потенційний ПК-залежний регуляторній шлях та інші ПК-залежні гени, пов’язані з гомеостазом c-di-GMP, а молекули ПК-сигналінгу було ідентифіковано в культурі. Це дані свідчать про можливий зв’язок між c-di-GMP-регуляцією і ПК у P. fluorescens SBW25, що дозволяє проводити більш складний і гнучкий контроль продукції целюлози і утворення біоплівки за колонізації грунтів і еконіш, асоційованих з рослинами, – природними середовищами існування P. fluorescens SBW25.

Ключові слова: циклічний димерний гуанозинмонофосфат (c-di-GMP), ацилгомосеринлактон (AHL), бактеріальна целюлоза, біоплівки, Pseudomonas, почутие кворуму.

АНАЛИЗ c-di-GMP И ВОЗМОЖНОГО ЧУВСТВА КВОРУМУ У Pseudomonas fluorescens SBW25: СВЯЗЬ МЕЖДУ ВНУТРІ- И МЕЖКЛЕТОЧНОЙ РЕГУЛЯЦИЕЙ СПОСОБСТВУЕ КООПЕРАТИВНОМУ ПОВЕДЕНИЮ В СООБЩЕСТВЕ И ФОРМировАНИЮ БИОПЛЁНКИ

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Успешность бактериальной колонизации сложных экосистем требует индивидуального от- вета на изменения условий на микромасштабном уровне.
но как и кооперацию на уровне сообщества для продукции таких крупномасштабных структур как биопленки. Координация индивидуальных ответов и ответов сообщества может быть достигнута путем связывания внутриклеточных сенсорных и регуляторных систем, опосредованных бис-(3',5')-циклическим димерным гуанозинофосфатом (c-di-GMP) и другими соединениями индивидумов с межклеточной регуляцией—чувством кворума (ЧК), контролирующим популяцию. Накапливается всё больше доказательств того, что формирование биопленки многими псевдомонадами регулируется как внутриклеточными, так и межклеточными регуляторными системами, хотя в случае модельной Pseudomonas fluorescens SBW25 Wrinkly Spreader, у которой мутации, повышающие уровни c-di-GMP, приводят к созданию прочной целлюлозной биоплёнки на границе раздела фаз воздух–жидкость, не было обнаружено никакого свидетельства вовлечения кворумзависимой регуляции. Однако наш недавний обзор генома P. fluorescens SBW25 выявил потенциальный ЧК-зависимый регуляторный путь и другие ЧК-зависимые гены, связанные с гомеостазом c-di-GMP, а молекулы ЧК-сигналинга были идентифицированы в культуре. Эти данные свидетельствуют о возможной связи между c-di-GMP-регуляцией и ЧК у P. fluorescens SBW25, что позволяет проводить более сложный и гибкий контроль продукции целлюлозы и образования биопленки при колонизации почв и экосистем, ассоциированных с растениями, – естественными средами обитания P. fluorescens SBW25.

Ключевые слова: циклический димерный гуанозимнофосфат (c-di-GMP) ацилгомосеринлактон (AHL), бактериальная целлюлоза, биопленки, Pseudomonas, чувство кворума.

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