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Characterizing Genetic Circuit Components in E. coli towards a Campylobacter jejuni Biosensor.

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Abstract

Campylobacter jejuni is responsible for most cases of bacterial gastroenteritis (food poisoning) in the United Kingdom. The most common routes of transmission are by contact with raw poultry. Current detection systems for the pathogen are time-
consuming, expensive or inaccessible for everyday users. In this article we propose a cheaper and faster system for detection of *C. jejuni* using a synthetic biology approach. We aimed to detect *C. jejuni* by the presence of xylulose, an uncommon bacterial capsular saccharide. We characterized two sugar-based regulatory systems that displayed potential to act as tools for detection of xylulose. Using a two-plasmid reporter system in *Escherichia coli*, we investigated the regulatory protein component (MtlR) of the mannitol operon from *Pseudomonas fluorescens*. Our findings suggest that the promoter of *mtlE* is activated by MtlR in the presence of a variety of sugar inducer molecules, and may exhibit cross-activity with a native regulator of *E. coli*. Additionally, we engineered the L-arabinose transcriptional activator (AraC) of *E. coli* for altered ligand specificity. We performed site-specific saturation mutagenesis to generate AraC variants with altered effector specificity, with an aim to generate a mutant activated by xylulose. We characterized several mutant AraC variants which have lost the ability to respond specifically to the native L-arabinose effector. We promote this technique as a powerful tool for future iGEM teams to create regulatory circuits activated by novel small molecule ligands.

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**Competing Interests**

The authors have declared that no competing interests exist.

**Ethics Statement**

N/A

**Data Availability**
Introduction

*Campylobacter jejuni* is a Gram-negative, microaerophilic, corkscrew-shaped bacteria which has been implicated as being one of the most common causes of human gastroenteritis worldwide [1][2][3]. Infection with *C. jejuni* causes common symptoms such as diarrhoea, abdominal pain, fever, headache, nausea and vomiting [3]. *C. jejuni* is harboured by poultry, though it has been reported in other meat products, raw milk, and in untreated drinking water [3]. The high prevalence of *C. jejuni* makes it an interesting target for synthetic biology-based solutions.

Traditional methods for detection of *C. jejuni* include culture-based techniques, which are relatively cheap to perform and require less training than other methods [4]. However, they are incredibly time and labour intensive and therefore, in recent years, a move has been made towards use of rapid detection testing [5]. Such techniques include enzyme immunoassay and lateral flow systems, which require only one to two hours to give a result [6]. However, use of these methods requires highly trained employees, and so detection of *C. jejuni* in an industrial or agricultural setting would require outsourcing to specialists. In addition, a comparison of three rapid detection systems demonstrated a high number of false negative results, which is a drawback when considering detection of *C. jejuni* to reduce the incidence of disease outbreaks [7].

To reduce incidences of food poisoning by *C. jejuni*, and to improve upon current methods of detection, we decided to create a biosensor that was able to detect the presence of *C. jejuni* quickly and accurately. We envisaged a two-part biosensor that would specifically require two sensory inputs associated with *C. jejuni* to report a reliable result. The first molecule we identified as a marker for *C. jejuni* was autoinducer-2 (AI-2) [9]. AI-2 is a secreted quorum-sensing molecule. However, many varied gram-positive and gram-negative bacterial species sense their population density and surrounding bacterial environment using this molecule [10]. On the other hand, this ubiquity meant that AI-2 gene regulation was well characterised, with prior iGEM teams having worked on the *E. coli* AI-2 quorum sensing regulatory system.
Therefore, we searched for another marker with greater specificity to *C. jejuni* to use in tandem with AI-2.

Xylulose is a rare sugar found incorporated within the polysaccharide capsule of *C. jejuni* [8]. The presence of xylulose is uncommon in bacterial polysaccharide capsules [8]. Additionally, the glycosidic bonds which incorporate xylulose were found to be extremely acid-labile [8], providing a possibility to release the molecule from the capsule and allow for whole-cell based detection. For the detection of xylulose two possible avenues were explored. One exploits the mannitol metabolism operon of *Pseudomonas fluorescens* [13]. It has been previously reported that xylulose acts as a direct inducer of the regulatory protein MtlR, activating transcription from the p_mtie promoter [14]. To investigate its suitability for biosensor, we characterized the p_mtie/MtlR regulatory system in *E. coli*.

Use of a gene regulatory system outside the context of the native organism may be problematic. For this reason, we aimed to construct an alternative xylulose sensor, by exploiting components of the L-arabinose operon, native to *E. coli* [15]. Several previous studies have shown that its regulatory protein, AraC, can be engineered to activate transcription in response to non-native small molecules [16][17][18]. Site-saturation mutagenesis of residues positioned within the ligand-binding pocket of AraC (Fig 1), coupled with fluorescence-based cell sorting, allowed isolation of AraC variants with altered effector specificity [16]. Based on these findings we aimed to use multiple site-saturation mutagenesis and fluorescence-based screening to generate mutant AraC responsive specifically to xylulose.
Fig 1. Crystal structure of AraC binding pocket with bound L-arabinose (Ara). The four key residues which are important for ligand binding pocket are indicated. Structure was generated using PyMOL.

Materials and Methods

Standard molecular biology techniques.

All protocols used during this work, including a standardised set for routine laboratory techniques, are detailed in S3 File. All ligation reactions described herein were transformed into DH5α commercial chemically competent E. coli cells, plated on L-agar containing the required antibiotics, and grown at 37°C overnight. All plasmid constructs herein were verified by diagnostic restriction digest and DNA sequencing prior to use or further subcloning.

Plasmid design for expression of p:\textit{mtlE}/MtIR regulatory system

All plasmids were constructed using BioBrick Standard Assembly. The biobrick p:\textit{tet} promoter (R0040) followed by a medium-strength ribosome binding site (RBS) B0032 was ordered for synthesis by Integrated DNA Technologies (IDT) as complementary oligos, that were annealed and ligated into a pSB1C3 plasmid backbone. \textit{mtlR} coding sequence was ordered as a gBlock Gene Fragment by IDT and inserted behind the R0040+B0032 promoter/RBS part in pSB1C3. This produced the regulatory plasmid BBa_K2442202 (Fig 2).

The sequence of the p:\textit{mtlE} promoter with its native RBS was obtained from Liu et al. (2015) [14] and supplied as oligo by IDT. A reporter plasmid was constructed by ligating GFP coding sequence (BBa_E0040; obtained from iGEM Distribution Kit) downstream of the p:\textit{mtlE} promoter into pSB1C3 backbone. The resulting plasmid BBa_K2442206 is shown in Fig 2.
Fig 2. Diagrams of the regulatory plasmid (left) and GFP reporter plasmid (right) for the expression of the MtlR/pmtE regulatory system.

Plasmid design for expression of pBAD/AraC regulatory system

Fragments containing parts R0011 (LacI-regulated promoter) upstream of ribosome-binding site (RBS) B0032 were synthesised by IDT as oligonucleotides. Wild type araC sequence as described by Miyada et al. (1980) [19] was amplified from BBa_I0500 using primers araC_BBPre_F and araC_BBSuf_R (Table 1) to introduce BioBrick prefix and suffix to both ends of WT araC. The PCR product was subsequently inserted downstream of R0011 promoter and B0032 RBS in pSB1C3 backbone to generate the regulatory plasmid BB_K2442104 (Fig 3).

Minimal pBAD promoter was synthesised by IDT as a gBlock (see S1 Fig for full sequence). The fragment was ligated into pSB3k3, upstream of part BBa_I13500 (containing B0034 RBS and GFP). This resulted in the final reporter plasmid BBa_K2442102 (Fig 3).
Fig 3. Diagrams of the regulatory plasmid (left) and GFP reporter plasmid (right) for the expression of the AraC/p_{BAD} regulatory system.

Mutant AraC Library Construction

Site directed mutagenesis by PCR was performed according to the protocol in S3 File. QIAQuick PCR purification of the product was performed according to the Qiagen protocol (S3 File). The purified PCR product was ligated into pSB1C3 backbone, downstream of R0011 promoter and B0032 RBS. Ligation reaction was ethanol-precipitated, and transformed into electrocompetent DH5α by electroporation. This resulted in the araC mutant library. Colonies carrying the mutant library were washed off transformation plates with 1.5ml double distilled H_2O, (ddH_2O) then transferred to a 1.5ml tube. The cell pellet was centrifuged at room temperature, the supernatant was then discarded and pellet resuspended in 1.5ml ddH_2O. This centrifugation-resuspension step was repeated 3 times to remove agar plate debris. Plasmid DNA was extracted using the Qiagen Plasmid MiniPrep Kit with 2 of each PB buffer and PE buffer wash steps to ensure maximum extract purity. Sequencing primer VF2 was then added and sample sent for sequencing.

Characterization of the p_{mtlE}/MtlR system activity in E. coli

To characterize activity of the p_{mtlE}/MtlR in E. coli, we studied the levels of GFP fluorescence using a 96 well plate in the FLUOstar Omega plate reader (BMG Labtech). Each strain was grown to saturation in an overnight culture of LB with
appropriate antibiotics for the relevant plasmids (chloramphenicol and kanamycin). Then the culture was diluted 1:100 with fresh LB+antibiotic and placed into a black bottomed 96 well plate. All readings were performed in the plate reader at 37°C shaking at 200 RPM. GFP fluorescence was measured (excitation at 485nm and emission at 530nm) every hour for 8 hours.

**Mutant Screening**

Plasmid DNA containing the araC mutant libraries were transformed into DS941 strain *E. coli* carrying the reporter plasmid K2442102 (in pSB3K3 vector). Transformants were plated on LB agar medium containing chloramphenicol plus kanamycin (to select for K2442102 and K2442104) plus one of the tested inducers: arabinose, xylose or decanal. Concentrations were 40mM for arabinose and xylose and 2mM for decanal. As a control test, transformants were also plated on LB medium containing chloramphenicol and kanamycin only. Fluorescence images of the conditional transformation plates were obtained using a GE-Healthcare Typhoon FLA-9500 laser scanner. Excitation was recorded at 473nm, emission was recorded using 520-540nm filter. Colonies which exhibited fluorescence were observed as dark colonies on the scan. Out of those which appeared dark on xylose or decanal plates, 200 were replica short-streaked onto plates containing xylose, arabinose, decanal, or no additive. Each colony of interest was picked and then immediately short-streaked onto each new condition plate using the same toothpick, into the same position using grids. This method of replica plating ensures the plates can later be aligned and short streaks of the same origin directly compared between the plates. After overnight incubation at 37°C, fluorescence scans of each plate were again obtained using the laser scanner.

**Liquid culture fluorescence assay for AraC mutants**

Colonies of interest were inoculated into L-broth containing chloramphenicol and kanamycin. Each liquid culture was grown overnight at 37°C, shaking at 225 rpm. The following day the culture was diluted 1:100 into fresh L-broth containing the above antibiotics plus one of the following inducer conditions:

- 40mM xylose
- 40mM arabinose
200µl of each culture to be tested was placed in a well of a clear-based 6-well plate, then incubated at 37°C shaking at 300 rpm for 12 hours in a BMG FLUOstar Omega fluorescence plate reader. GFP fluorescence of each culture was measured at 1 hour time intervals for the duration of the culture experiment, using excitation wavelength 485nm and emission wavelength 530nm. Cell growth was simultaneously tracked by measuring optical density at 600nm.

Results

\( P_{m\text{tl}E} \) is regulated by MtlR and other native regulatory proteins in \( E. \text{coli} \)

From the literature, we identified a regulatory system that responds to xylulose - the mannitol-inducible promoter from \( P. \text{fluorescens} \) and its regulatory protein MtlR [14]. To utilize these parts in our dual-input biosensor, their activity needed to be characterized in \( E. \text{coli} \). Two constructs were assembled: the regulatory plasmid with a constitutively active \( p_{\text{tet}} \) promoter driving expression of the MtlR protein, and the reporter plasmid containing \( p_{m\text{tl}E} \) promoter regulating expression of GFP (Fig 1).

To test whether MtlR can activate \( p_{m\text{tl}E} \) in \( E. \text{coli} \), we measured GFP fluorescence in \( E. \text{coli} \) carrying the reporter plasmid alone, or both the regulatory and reporter plasmids. The experiment was done in presence of 6 structurally similar sugars (ribose, fructose, xylose, mannitol, arabinose and sorbitol) [14], to investigate substrate specificity of MtlR. Xylulose itself was not tested due to budgetary constraints. Fluorescence levels were compared to basal fluorescence levels of DH5α cells, not expressing either plasmid. Cells expressing the reporter plasmid alone showed higher levels of fluorescence than empty cells, in presence of all sugars tested (Fig 4 and S1 File). This suggests that \( P_{m\text{tl}E} \) may interact with native \( E. \text{coli} \) proteins. GFP fluorescence levels further increased in cells expressing both MtlR and the reporter plasmid (Fig 4 and S1 File). This supports previous findings that MtlR functions as an activator of \( p_{m\text{tl}E} \) in \( E. \text{coli} \). Although we were unable to test xylulose, we found that \( p_{m\text{tl}E} \) was induced in presence of a number of structurally similar sugars,
showing highest response to ribose and sorbitol (Fig 4 and S1 File). We conclude that $p_{mtlE}$ promoter functions in *E. coli*, and MtlR acts as its activator. However, the *P. fluorescens* $p_{mtlE}$ promoter is not strictly regulated by MtlR when expressed in *E. coli*.

**Figure 4. Activity of $p_{mtlE}/MtlR$ reporter circuit in *E. coli*.** Average relative fluorescence over optical density at hour 8. Shows fluorescence levels in presence of each sugar tested (as specified). The cells were expressing either reporter plasmid BBa_K2442206 alone, both reporter and regulatory plasmid BBa_K2442202, or neither (as specified). Low copy; PSB1C3 plasmid vector. High copy; PSB3k3 plasmid vector. *E. coli* strain is DH5α. n=1.

Split regulatory components of the L-arabinose operon from *E. coli* are functional and can be used for construction of L-arabinose-inducible systems

As the *P. fluorescens* MtlR regulatory system lacked specificity in *E. coli*, we aimed to utilize components of the L-arabinose operon from *E. coli* to generate a new tightly controlled xylulose-regulatory system. We chose to mutagenize the AraC protein to change its effector specificity, based on previous reports of successful engineering of AraC to respond to non-native inducers [16][17][18]. The $p_{BAD}$ promoter is regulated by the AraC transcriptional regulator, which drives expression from $p_{BAD}$ only in presence of L-arabinose [15]. In nature the $p_{BAD}$ promoter overlaps with the *araC*
coding region [20]. To allow for mutagenesis, we split araC from pBAD. The minimal pBAD was designed to retain all the sites required for AraC binding. The start codon of araC within pBAD has been changed from ATG→AGT (S1 Fig).

To test activity of minimal pBAD, we expressed regulatory and reporter plasmids in AraC-negative E. coli strain DS941 and plated the cells on LB medium. GFP fluorescence measurements demonstrated that AraC expressed from a separate plasmid can induce expression from minimal pBAD upon binding of arabinose, as cells exhibited fluorescence only in presence of arabinose (Fig 5 and S2 File). GFP fluorescence measurements revealed that minimal pBAD is inducible by L-arabinose 300-fold (Fig 5 and S2 File), showing improvement over previously characterized pBAD parts. The minimal pBAD promoter is tightly regulated by AraC expressed either from our regulatory plasmid, or from bacterial chromosome.
Figure 5. Activity of split araC and \( P_{BAD} \) fluorescence circuits in E. coli. Average relative fluorescence over optical density at hour 8. Shows fluorescence levels under no additive and 40mM Arabinose. E. coli strains which the plasmids were transformed into are DS941(araC-) or DH5a (araC+). Cells were expressing either regulatory construct BBa_K2442104, reporter construct BBa_K2442102, both, or neither (as specified). Low copy; PSB1C3 plasmid vector. High copy; PSB3k3 plasmid vector. Error bars are standard deviation; \( n=3 \)

Site-directed mutagenesis of the AraC protein provides a tool to develop biosensors responsive to non-native molecules

We performed site-directed saturation mutagenesis of the araC gene, targeting four amino acids within the L-arabinose binding pocket of AraC. The protocol successfully generated a mutant library of \(~24,000\) AraC variants. Sequencing confirmed that NNS mutations were introduced at correct codon positions (residues 8, 24, 80 and 82; Fig 6). E. coli DS941 transformed with the mutant library and reporter plasmids were screened for GFP fluorescence in presence of 3 different inducer molecules that were available to us, to identify colonies with altered AraC/\( P_{BAD} \) activity. Due to prohibitive cost, we were unable to test xylulose. Nevertheless, 4 colonies displaying altered expression patterns were identified and subsequently characterized (Fig 7 and S2 File). Two AraC mutants constitutively activated \( P_{BAD} \), without the requirement for L-arabinose. The other two had lost the ability to respond to arabinose. Potentially, the latter two variants could be responsive to a yet unidentified compound.
Figure 6. Sequencing trace from mutant library MiniPrep. Figure shows NNS mutations introduced at residues 8, 24, 80 and 82. N; any base. S; strong base (C or G).
Figure 7. Activity of AraC mutants compared to WT AraC. Average relative fluorescence over optical density at hour 8. Shows fluorescence levels under no additive, 40mM Arabinose, 40mM xylose and 2mM decanal. E. coli strain which the plasmids were transformed into is DS941 (araC-). Cells were expressing the reporter plasmid and regulatory plasmid with wild type (WT) or mutant (mut1-mut4) variant of AraC (as specified). Empty cells served as control. Low copy; PSB1C3 plasmid vector. High copy; PSB3k3 plasmid vector. For WT AraC (lane 1) error bars are standard deviation; n=3. For all other samples (lanes 2-6): n=1.

The 4 mutant plasmids were sequenced. Wild-type AraC possesses proline at residue 8, threonine at residue 24, histidine at 80 and tyrosine at 82. Mutant 1 (BBa_K2442105) has the following mutations: proline-8→glycine (P8G), histidine-80→proline (H80P) and tyrosine-82→tryptophan (Y82W). Mutant 2 (BBa_K2442106)
and 3 (BBa_K2442107) both had the same mutations as follows: proline-8→serine (P8S), threonine-24→aspartic acid (T24D), histidine-80→isoleucine (H80I) and tyrosine-82→alanine (Y82A). Sequencing of mutant 4 (BBa_K2442108) revealed a deletion within the araC coding region, which explains lack of AraC activity in the corresponding colonies.

**Discussion**

Due to lack of cheap and rapid detection methods for *C. jejuni*, we designed a biosensor responsive to a biomarker specific to the pathogen - xylulose. In our study we characterized activity of a mannitol-responsive regulator MtlR in *E. coli*. Liu et al. (2015) [14] previously reported that mannitol and xylulose act as direct inducers of MtlR to activate transcription from the p<sub>mtlE</sub> promoter. Our results suggest that this system is functional when expressed in *E. coli*, with MtlR activating transcription from p<sub>mtlE</sub>. However, reporter expression was also induced independently of MtlR. This was achieved in presence of a variety of sugars structurally similar to mannitol and xylulose, including arabinose, fructose, xylose, ribose and sorbitol. Contradictory to how the system works in *P. fluorescens* [14], the strongest reporter expression was achieved in presence of sorbitol and ribose, rather than mannitol. Although the p<sub>mtlE</sub> was previously found to be activated by sorbitol independently of MtlR [14], no response to ribose has yet been observed. We suggest that other, yet unidentified proteins naturally found in *E. coli* act as transcriptional activators of p<sub>mtlE</sub>.

Although our results show an increase in fluorescence upon expression of the p<sub>mtlE</sub>/MtlR components in presence of sugars, it is important to note that our testing lacked a no-inducer control. As reporter expression from p<sub>mtlE</sub> was observed in absence of MtlR expression, it may be that this system is constitutively activated independently of any inducer sugar. Inclusion of such control would allow for better understanding of the system’s activity in *E. coli*. Moreover, due to time constraints the experiment was only performed on only one occasion, which could impact reliability of the results. This would be improved in future experiments by increasing the number of repetitions and samples tested, to determine statistical significance of the results obtained.
As we didn’t identify a natural regulatory system responsive specifically to xylulose, we performed mutagenesis of the L-arabinose-responsive protein AraC as an alternative route to detect our target sugar. To achieve this, we split the araC coding region from the overlapping sequence of the pBAD promoter. Our results suggest that the modified pBAD is functional and can be activated by AraC expressed from a separate construct. Most importantly, we demonstrated successful mutagenesis protocol of the AraC protein to generate variants with altered effector substrate specificity. However, due to time constraints, we were only able to screen 200 mutant colonies out of 24,000 colonies obtained. This is an extremely low number considering our protocol was expected to generate 160,000 mutant AraC variants with different amino acid combinations at the 4 randomised residue positions. Moreover, limited resources enabled us to test for response to a narrow range of inducer molecules. In the future, optimization of the transformation protocol to obtain enough colonies to cover the entire library, along with a larger scale mutant screening would potentially allow identification of an AraC variant responsive to xylulose. Such mutant could be utilized in a xylulose-sensing component in a C. jejuni biosensor. Moreover, a catalogue of AraC mutants could be utilized as a component of a biosensor toolbox, allowing for generation of genetic circuits regulated by small molecules of choice.

Although we aimed to generate a xylulose-regulated system, we were unable to test our constructs in presence of xylulose due to its prohibitive cost. Nevertheless, the sugar would be required to screen for AraC mutants responsive to xylulose. We found that xylulose isomerase enzyme is capable of converting the cheaper sugar, xylose, into xylulose [21][22]. We have considered development of a suitable expression plasmid, which in the future could be used to overexpress and purify the enzyme for production of xylulose in subsequent experiments (for full description see [http://2017.igem.org/Team:Glasgow/XyluloseBiosynthesis](http://2017.igem.org/Team:Glasgow/XyluloseBiosynthesis)).

Although xylulose is rarely found in bacterial capsules [8], potential contamination of the tested area by xylulose from other sources could lead to false positive results from our detector. For this reason, we designed our biosensor to detect two sensory inputs. Apart from xylulose, we identified autoinducer-2 (AI-2) as another marker for C. jejuni [9]. AI-2 is a secreted quorum sensing molecule. In future development of the biosensor components, the detectors for both xylulose and autoinducer-2 would form
two components of an AND gate that will ensure a positive result is given only when both xylulose and autoinducer-2 are present, improving specificity and accuracy of the detector (for full description see http://2017.igem.org/Team:Glasgow/ANDGate).

To increase efficiency of the whole-cell based biosensor, we went through several design iterations to create a device that would house the engineered bacteria to make the use of the biosensor simple and easy (for full description see http://2017.igem.org/Team:Glasgow/Applied_Design; http://2017.igem.org/Team:Glasgow/Hardware).

Conclusions

We conclude that the AraC mutagenesis protocol was successful at generating AraC variants with altered effector specificity. A larger scale mutant screen could result in identification of a xylulose-inducible AraC variant. For increased specificity, the xylulose-regulated $p_{BAD}/AraC$ system could be combined with an AI-2-sensing construct within an AND gate, and transformed into a host bacterium to produce a dual-input biosensor tool for rapid detection of C. jejuni.

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