

Evolutionary resurrection of flagellar motility via rewiring of the nitrogen regulation system

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Title: Evolutionary resurrection of flagellar motility via rewiring of the nitrogen regulation system

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One Sentence Summary:

Rapid repeatable rewiring of regulatory networks: a nitrogen regulatory gene evolves a
new function, restoring flagella to immotile bacteria.

4

5 **Abstract**:

A central process in evolution is the recruitment of genes to regulatory networks. We 6 7 engineered immotile strains of the bacterium *Pseudomonas fluorescens* that lack flagella due to deletion of the regulatory gene *fleQ*. Under strong selection for motility, 8 these bacteria consistently regained flagella within 96 hours via a two-step evolutionary 9 pathway. Step 1 mutations increase intracellular levels of phosphorylated NtrC, a distant 10 11 homologue of FleQ, which begins to commandeer control of the *fleQ* regulon at the cost of disrupting nitrogen uptake and assimilation. Step 2 is a switch-of-function mutation 12 that redirects NtrC away from nitrogen uptake and towards its novel function as a 13 14 flagellar regulator. Our results demonstrate that natural selection can rapidly rewire regulatory networks in very few, repeatable mutational steps. 15

16

17 Main Text:

A longstanding evolutionary question concerns how the duplication and recruitment of genes to regulatory networks facilitates their expansion (*1*), and how networks gain mutational robustness and evolvability (*2*). Bacteria respond to diverse environments using a vast range of specialised regulatory pathways, predominantly two-component systems (*3*), which are the result of adaptive radiations within gene families. Due to past cycles of gene duplication, divergence and horizontal genetic
transfer, there is often extensive homology between the components of different
pathways (4), raising the possibility of cross-talk or redundancy between pathways (5).
Here we monitor the recovery of microbial populations from a catastrophic gene
deletion: bacteria engineered to lack a particular function are exposed to environments
that impose strong selection to re-evolve it, sometimes by recruitment of new genes to
regulatory networks (6, 7, 8, 9).

In the plant-associated soil bacterium P. fluorescens, the master regulator of 30 flagellar synthesis is FleQ (also called AdnA), a σ^{54} -dependent enhancer binding protein 31 (EBP) that activates transcription of genes required for flagellum biosynthesis (10, 11). 32 33 The starting *P. fluorescens* strain is AR2; this strain lacks flagella, due to deletion of fleQ, and is unable to move by spreading motility due to mutation of viscosin synthase 34 (viscB), resulting in a distinctive, point-like colony morphology on spreading motility 35 medium (SMM) (12) (Figure 1A). We grew replicate populations of AR2 on SMM; when 36 local nutrients became depleted, starvation imposed strong selection to re-evolve 37 motility. To demonstrate that this finding was not strain-specific, these experiments were 38 replicated in a different strain of *P. fluorescens*, Pf0-2x. This strain is a $\Delta fleQ$ variant of 39 Pf0-1, already viscosin-deficient, and is thus unable to move by spreading or swimming 40 motility. 41

After 96 hours incubation of AR2 and Pf0-2x at room temperature on SMM, two breakout mutations were visible conferring first slow (AR2S and Pf0-2xS) and then fast (AR2F and Pf0-2xF) spreading over the agar surface (Fig. 1A). The AR2F strain produces flagella, but we could not detect flagella in EM samples for AR2S (Fig. 1B).

Genome resequencing revealed a single nucleotide point mutation in *ntrB* in strain AR2S, causing an amino acid substitution within the PAS domain of the histidine kinase (HK) sensor NtrB (T97P). The fast-spreading strain AR2F had acquired an additional point mutation in the σ^{54} -dependent EBP gene *ntrC*, which alters an amino acid (R442C) within the DNA-binding domain (Table 1 & S2).

NtrB and NtrC comprise a two-component system: under nitrogen limitation NtrB 51 phosphorylates NtrC, which activates transcription of genes required for nitrogen uptake 52 and metabolism. To determine how mutations in this separate regulatory pathway 53 restored motility in the absence of FleQ, we performed microarray and gRT-PCR 54 analyses of the ancestral and evolved strains (Fig. S1 & Table S1). The expression of 55 56 genes required for flagellum biosynthesis and chemotaxis was abolished in AR2 compared to wild-type SBW25 (Fig. 2A). The ntrB mutation in AR2S partially restores 57 the expression of flagellar genes, and over-activates the expression of genes involved 58 in nitrogen regulation, uptake and metabolism. The subsequent ntrC mutation in AR2F 59 reduces the expression of nitrogen uptake and metabolism genes, while further up-60 regulating flagellar and chemotaxis gene expression to wild-type levels (Fig. 2B). While 61 AR2S and AR2F showed higher growth rates than the ancestor in LB medium (the 62 medium on which the mutants arose; Tukey-Kramer HSD test, growth in LB compared 63 to AR2: AR2S, P < 0.001; AR2F, P < 0.001) (Fig. 1C), both mutants grew poorly in 64 minimal medium with ammonium as the sole nitrogen source (Tukey-Kramer HSD test, 65 growth in M9 + ammonium compared to AR2: AR2S, P < 0.001; AR2F, P = 0.001). This 66 67 is likely to be the result of ammonium toxicity due to the strong up-regulation of genes

involved in ammonium uptake and assimilation, indicating a pleiotropic cost of thisadaptation.

70 Sequencing of the *ntrBC* locus from independently evolved replicate strains showed that evolution often followed parallel trajectories in both AR2 and Pf0-2x: 71 mutation of *ntrB* gave a slow-spreading strain and this was followed by mutation of *ntrC* 72 yielding a fast-spreading strain (Table 1; Fig. 3A & C). While all Pf0-2xF replicates 73 carried mutations in ntrC, several Pf0-2xS strains were not mutated in ntrB, suggesting 74 an alternative evolutionary pathway. Genome resequencing of these strains revealed 75 mutations in glnA or glnK (Table 1; Fig. 3B) likely to result in loss of function leading to 76 abnormally high levels of phosphorylated NtrC: reduced ammonium assimilation by 77 78 glutamine synthetase (glnA) would impose severe nitrogen limitation in the cell irrespective of nitrogen availability, whereas GlnK is a PII-protein that regulates both 79 NtrB and glutamine synthetase activities. 80

These data suggest a predictable two-step evolutionary process: Step 1 81 increases levels of phosphorylated NtrC, via either (a) a direct regulatory route with 82 mutations in NtrB or GlnK, or (b) a physiological route with loss-of-function mutations 83 reducing glutamine synthetase activity and causing NtrB activation, partially or 84 intermittently reactivating the flagellar cascade. In Step 2, NtrC adapts to enhance 85 activation of the flagellar genes and in doing so, becomes a less potent activator of 86 nitrogen uptake genes. This model explains the microarray data and is consistent with 87 the predicted structural effects of the mutations (Figs. 2 and 3). Specifically, for NtrB the 88 structural changes are likely to either increase kinase or reduce phosphatase activity. In 89

support of this, the mutated NtrB(D228A) repeatedly emerging in Pf0-2xS resembles
NtrB(D227A) in *P. aeruginosa* which constitutively activates the Ntr system (*13*).

NtrC is a distant homologue of FleQ, sharing 30% amino acid identity and the 92 same three structural domains (TM-score > 0.7; P < 0.001; Fig. 3D) (14): an N-terminal 93 receiver domain, a conserved central σ^{54} -interacting domain, and a C-terminal DNA-94 binding domain containing a helix-turn-helix (HTH) motif flanked by highly disordered 95 regions. We posit that an overabundance of phosphorylated NtrC activates transcription 96 of flagellar genes through functional promiscuity (15). Consistent with this, the ntrC 97 mutations in fast-spreading strains are predominantly located within or adjacent to the 98 HTH domain and likely influence enhancer-binding specificity; one is a frameshift 99 abolishing the HTH altogether (Fig. 3C). The evolved NtrC' must be constitutively 100 phosphorylated by overactive NtrB to enable its new multipurpose role, with the result 101 that flagellum biosynthesis and nitrogen regulation are probably no longer responsive to 102 environmental stimuli. Consequently, there is a trade-off between nitrogen utilization 103 104 and motility (Fig. 1A & C).

The flagellar regulatory network may have an unusually dynamic evolutionary 105 history. Flagella are expensive to make, and not always advantageous. Pathogens 106 107 expressing flagella can trigger strong immune responses in the host, so rapid transitions are seen over short timescales between uniflagellate, multiflagellate and aflagellate 108 states (16). This volatility is reflected in the structure of regulatory networks: in close 109 relatives of *Pseudomonas*, *fleQ* appears not to be involved in flagellar gene expression 110 (17), and in Helicobacter pylori, a gene of unknown function can be used as a "spare 111 part" to permit motility in *flhB* mutants (18). Our results illustrate that trans-acting 112

mutations can contribute to gene network evolution (19), but that as predicted, such 113 mutations bear severe pleiotropic costs (20, 21). Genes can retain the potential to take 114 on the functions of long-diverged homologues, suggesting that some degree of 115 evolutionary resilience is a consequence of regulatory pathways that evolve via gene 116 duplication. While *de novo* origination of new functions in nature is likely to take longer 117 and involve more mutational steps, this system enables us to understand the adaptive 118 process in detail at the genetic and phenotypic level. Here we identified a novel, 119 tractable model for gene network evolution and observed, in real time, the rewiring of 120 gene networks to enable the incorporation of a modified component (NtrC') creating a 121 122 novel regulatory function, by a highly repeatable two-step evolutionary pathway with the same point mutations often recurring in independent lineages. 123

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TBT, LJJ, RWJ and MAB conceived and designed the study. TBT, GM and AA performed experiments. MWS and AHD performed independent lines of enquiry on Pf0-2x. DS conducted bioinformatics analysis of genome resequencing data, identified mutated genes and handled sequencing data. LM conducted the protein structure prediction and analysis. This work was supported by a Leverhulme grant to LJJ, MAB and RWJ, BBSRC grant BB/J015350/1 to RWJ, start-up funding from the University of York to MAB, Qassim University to AA, and Agriculture and Food Research Initiative Competitive Grant 2010-65110-20392 from the USDA's National Institute of Food and Agriculture, Microbial Functional Genomics Program to MWS. TBT, GM, LJJ, RWJ, MWS, DJS and MAB wrote the paper. We thank Graham Bell, Mark Pagel, Angus Buckling and James Moir for useful discussions; Peter Ashton for processing of microarray data; and Konrad Paszkiewicz and Exeter Sequencing Service facility and support from the following: Wellcome Trust Institutional Strategic Support Fund (WT097835MF), Wellcome Trust Multi User Equipment Award (WT101650MA) and BBSRC LOLA award (BB/K003240/1). Sequence data for genomic resequencing of AR2S and AR2F have been submitted to the SRA under accession numbers SRR1510202 and SRR1510203, respectively. The eArray design ID for the microarray is 045642. Microarray data have been submitted to the ArrayExpress database under accession number E-MTAB-2788 (www.ebi.ac.uk/arrayexpress).

Figures 1 – 3:

Fig. 1. Phenotypic assays of motility variants.

A Surface spreading motility assays of ancestral (AR2) and evolved slow spreading (AR2S) and fast spreading (AR2F) mutants, after 27 hours. **B** Electron microscopy confirms the presence of a flagellum in AR2F, but fails to confirm presence in AR2S. **C** Mean (N=4) cell doublings per hour (+/- 1 SEM). Strains were grown in differing nitrogen environments: 10 mM glutamine (GIn), glutamate (GIu) and ammonium (NH4) as the sole nitrogen source, or in high nutrient lysogeny broth (LB): AR2, $F_{3, 12} = 13.460$, *P* < 0.001; AR2S, $F_{3, 12} = 72.674$, *P* < 0.001; AR2F, $F_{3, 12} = 52.538$, *P* < 0.001. There were also differences between doubling rate of strains within each nitrogen medium (Glutamate (Glu), $F_{2, 9} = 12.654$, *P* = 0.002; Ammonium (NH4), $F_{2, 9} = 40.529$, *P* < 0.001), with the exception of Glutamine (Gln) ($F_{2, 9} = 3.703$, *P* = 0.067).

Fig. 2. Heat map of microarray expression profiles for all evolved and ancestral motility variants.

Heat maps show where there is significant ($P \le 0.05$) fold-change of ≥ 2 in genes selected based on GO-terms for **A** Bacterial-type flagellum (24 genes) and **B** Nitrogen compound transport (146 genes) for all strains. The gradation of colors reflects normalized raw signal values across the entire array. Genes are ordered according to chromosomal position to enable clearer visualization of coregulated gene clusters. Full transcriptome analysis is reported in Supplementary File "Microarray dataset.xlsx".

Fig. 3. Full chain multi-template 3D models of protein structures of slow and fast spreading motility variants.

Slow spreading variants can either follow the direct regulatory route through mutation of NtrB or GlnK (**A**), or the physiological route through mutation of GlnA causing overactivation of NtrB (**B**); both routes are predicted to lead to hyperphosphorylation of NtrC. Fast spreading variants all show mutational changes to NtrC (**C**), which has a similar global structure to FleQ (**D**). The colour scheme represents the variation in models, which correlates with local (per-residue) quality and disorder. Regions coloured in blue and green represent low local variability in structure, while those in red show high local variability (see Table 1 and Table S2 for mutation details). § = All mutations mapped onto SBW25 wildtype protein structures for illustrative purposes; * = Truncated domain.

	Slow Spreaders (AR2/Pf0-2xS)	Fast Spreaders (AR2/Pf0-2xF)		
	Hyperphosphorylation of NtrC	Switch of NtrC-P specificity		
R2	ntrB T97P*	ntrC R442C*		
	ntrB V185K	ntrC K342E		
A	ntrB D179N	ntrC G452R		
	<i>ntrB</i> L184Q / V185I	ntrC K342V / Frameshift: V342		
		ntrC N454S		
	ntrB D228A [§]	ntrC R441S		
		ntrC N454S		
		ntrC P424L		
		ntrC N454S		
)-2x		ntrC L418R		
PfC		ntrC G414D		
	ntrB D228A3	ntrC N454S		
		ntrC F426V		
	gInK Frameshift: I86*	ntrC R442H*		
	gInA T237P*	ntrC A445V*		
	gInA Frameshift: T205*	ntrC R442C*		

* = Identified by genome resequencing

§ = Independent *ntrB* mutant strains, parent to multiple *ntrC* mutant strains

Table 1. Mutational trajectory towards slow and fast spreading phenotypes.

Mutations confirmed in slow spreading motility variants are predicted to result in hyperphosphorylation of NtrC; mutations in fast spreading variants lead to predicted switched specificity of NtrC-P towards FleQ targets. Slow and fast spreading variants share the same ancestry.



Nitrogen Environment

Figure 1



A Bacterial-type Flagellum

Figure 2



Figure 3

Supplementary Materials:

Title: Evolutionary resurrection of flagellar motility via rewiring of the nitrogen regulation system

Authors: Tiffany B. Taylor1⁺, Geraldine Mulley1⁺, Alexander H. Dills2, Abdullah S. Alsohim1,3, Liam J. McGuffin1, David J. Studholme4, Mark W. Silby2, Michael A. Brockhurst5, Louise J. Johnson1,^{*}, Robert W. Jackson1,6

Materials and Methods

Fig. S1.

Table S1.

Table S2.

198 Supplementary Materials:

199

200 Materials and Methods:

201

202 Microbiological methods

- 203 *P. fluorescens* strains used in this study: SBW25, SBW25Δ*fleQ*, AR2 (SBW25Δ*fleQ* IS-
- 204 Ω Km-hah: PFLU2552), and Pf0-2x (Pf0-1 Δ *fleQ*). Motility assays are as described in
- (12). All starting populations were from a single AR2 colony grown on LB agar (1.5%),

and stab inoculated into the center of a SMM plate using a sterile wire. The initial 206 observed motility mutants (AR2S and AR2F) were cultured immediately, cryopreserved 207 and used for subsequent genome resequencing and microarray analysis. Independently 208evolved motility mutants were isolated from AR2 and Pf0-2x and cryopreserved. 209 Mutation rates of all lineages were checked by plating an overnight culture onto LB agar 210 supplemented with 100 µg ml⁻¹ rifampicin. All strains were found to have a similar 211 mutation frequency: SBW25 = 6.98×10^{-9} cfu ml⁻¹; Δ fleQSBW25 = 3.72×10^{-8} cfu ml⁻¹; 212 $AR2 = 2.50 \times 10^{-8}$ cfu ml⁻¹; $AR2S = 2.3 \times 10^{-8}$ cfu ml⁻¹; $AR2F = 1.4 \times 10^{-8}$ cfu ml⁻¹. 213 M9 nitrogen modified medium (lacking ammonium) was supplemented with 10 mM 214 glutamate, glutamine or ammonium solutions. Strains were grown for 16 hours in LB at 215 216 27°C and diluted to OD 0.001 in M9 nitrogen modified media, and in LB. Optical density 595 nm was measured every 20 minutes, under continuous shaking and at an 217

incubation of 27°C, for 24 hours (Tecan, GENios).

219

220 Molecular methods

The mutations present in the original evolved mutants AR2S and AR2F, plus 3 Pf0-2xS

and 3 descended Pf0-2xF strains were identified by genome resequencing.

223 Genomic DNA was isolated using Puregene DNA extraction (Qiagen) or Wizard®

224 Genomic DNA Purification (Promega) kits. We resequenced the complete genomes of

mutants AR2S and AR2F (original) using the Illumina HiSeq and identified single-

- nucleotide variants (SNV) by alignment to the SBW25 reference as described previously
- 227 (22, 23). Further mutants were analyzed by targeted PCR amplification and sequencing

of *ntrBC* to determine whether the same two-step process had occurred; primers were

designed to flank the region where SNVs had been identified by whole-genome

230 resequencing (NtrC, F: CTTCATCCCCAACTCCTTGA, R:

231 AAGCTGCTGAAAAGCGAGAC; NtrB, F: CTTGCGCCTTGAGTACATGA, R:

232 ATGCGGTCTACCAGGTTACG).

233

234 AR2S and AR2F Isolate Genome Resequencing Details

Whole-genome resequencing was performed using the Illumina GA2x. We generated 235 12,065,035 pairs of 36-bp reads for AR2S (SRA accession SRR1510202) and 236 24,075,130 pairs of 36-bp reads for AR2F (SRR1510203). To identify mutations, we 237 aligned the genomic sequence reads against the Pseudomonas fluorescens SBW25 238 genome sequence (NCBI RefSeq accession NC_012660) using BWA-mem version 239 0.7.5a-r405 (http://www.ncbi.nlm.nih.gov/pubmed/19451168). We discarded reads that 240 did not uniquely map to a single location on the SBW25 genome in order to avoid 241 artifacts due to misalignment of sequence reads arising from repeat sequences. The 242 resulting average depths of coverage over the SBW25 genome were 124x and 144x for 243 AR2S and AR2F, respectively. For 99.86% of the SBW25 reference genome sequence 244 (6,713,197 out of 6,722,539 bp) we were able to unambiguously determine the 245 nucleotide sequence in both mutant genomes (on the basis that at these genomic 246 positions there was at least 95% consensus among all aligned reads at those positions 247 for each of the two resequenced genomes and depth of at least 5x). Over the 6,713,197 248 bp of the genome for which we could unambiguously determine the DNA sequence for 249 both mutant genomes, we found only three SNVs with respect to the reference genome 250

sequence. Two of these three variants were present in both resequenced genomes: at 251 position 376,439 both resequenced strains had a G whereas in the SBW25 reference 252 genome the base is T. This corresponds to a non-silent change from codon acc to 253 codon Ccc in gene PFLU0344 (*glnL/ntrB*). The second variant was at position 1,786,536 254 where SBW25 has A but the two resequenced mutants have G; this variant falls in an 255 intergenic region and we have no reason to suspect it has an effect on the phenotypes 256 observed. The third SNV is private to AR2F and falls within the gene PFLU0343 257 (glnG/ntrC). In SBW25 and AR2S the base is G but in AR2F it is A; this results in a non-258 silent change from codon cgt to codon Tgt. Aside from these three variant sites, the 259 remainder of the 6,713,197 bp of unambiguously resolved genome sequence contained 260 no variation from the SBW25 reference genome. 261

262

263 Pf0-2xS and Pf0-2xF Isolate Genome Resequencing Details

Library preparation was performed using Illumina Nextera XT kit with Nextera XT
 Indexes and sequenced as 250PE reads from MiSeq.

²⁶⁶ Whole-genome resequencing of Pf0-2x strains was performed using the Illumina MiSeq.

We generated 1,145,122 forward and 1,141,017 reverse reads of 251-bp for Pf0-2xS_1,

1,279,360 forward and 1,272,990 reverse reads of 251-bp for Pf0-2xF_1.1, 1,962,339

forward and 1,941,676 reverse reads of 251-bp for Pf0-2xS_2, 907,328 forward and

270 906,083 reverse reads of 251-bp for Pf0-2xF_2.1, 935,862 forward and 932,415 reverse

reads of 251-bp for Pf0-2xS_3, and 1,194,665 forward and 1,190,975 reverse reads of

272 251-bp for Pf0-2xF_3.1. These genomic sequence reads were mapped against the

273 *Pseudomonas fluorescens* Pf0-1 genome sequence (NCBI RefSeq accession

NC_007492.2) using CLC Genomics Workbench 6.0; unmapped reads were not 274 included in further analysis. The average coverage depths for each library were: 44.6 for 275 Pf0-2xS_1, 49.8 for Pf0-2xF_1.1, 76.1 for Pf0-2xS_2, 35.3 for Pf0-2xF_2.1, 36.4 for Pf0-276 2xS 3, and 46.5 for Pf0-2xF 3.1. Out of the 6,438,405 bp Pf0-1 genome, 97.06-99.35% 277 of nucleotides (depending on sample) had a minimum 95% consensus among all 278 mapped reads. Using probabilistic variant detection to investigate unambiguous 279 nucleotide positions in each sequenced strain, we identified two SNVs and a 3bp 280 deletion common to all mutants (Table S2). These three common changes were also 281 present in the parental strain (Pf0-2x) and were thus not considered further. A 15bp 282 deletion was identified in both the Pf0-2xS1 and F1.1 strains with a unique SNV 283 occurring in the latter. An additional SNV was common to Pf0-2xS2 and Pf0-2xF2.1, yet 284 another to both Pf0-2xS3 and Pf0-2xF3.1, and a single unique SNV was discovered in 285 Pf0-2xF2.1 and another in Pf0-2xF3.1 with respect to the reference genome sequence. 286

287

The presence of *glnA*, *glnK* and *ntrC* mutations in resequenced strains was confirmed 288 by PCR of candidate regions from parental and evolved strains followed by Sanger 289 sequencing of the amplicons. Mutations in *ntrB* were detected by targeted PCR 290 amplification and sequencing, and *ntrC* mutations in derivatives of *ntrB* mutants were 291 detected the same way. In some cases, multiple primers were used to cover the full 292 length of the gene. This enabled us to be certain about the mutations, and rule out any 293 others. Primers used for Pf0-1 were: ntrB, F: ATTGCGCCTCGAGTACATGA and 294 TCGACACGGTTTCACTACGG, R: ATGCGGTCGACCAGATTGCG and 295 TCGGAGCCTTGGTTTGGTTT; ntrC, F: AAGCTGCTGAAAAGCGAGAC and 296

GATTAAGGGTCACGGTGCCT, R: CTTCATGCCGAGTTCCTTGA and
 CACTGGAACAAGGAGCCACA; glnA, F: GGAGGCCTTTCTTTGTCACG and
 ACGCTTGTAGGAGTTGGTGG and CGGGAAGTAGCCACCTTTCA, R:
 CCACCAACTCCTACAAGCGT and CAAGTCCGACATCTCCGGTT and
 CTACCCGCCCTAATTCACCC; glnK, F: GCCGGGCATTACGATAGACA, R:
 TGCGCTTAGACTTGAGTCGG.

303

304 Microarray design

305 For microarray analysis, RNA was extracted from SBW25, SBW25*A*fleQ, AR2, AR2S and AR2F using an RNeasy kit (Qiagen) and assayed for quality [2100 Bioanalyzer 306 (Agilent), and Nanodrop 1000 Spectrophotometer (Thermo Fisher Scientific)]. RNA was 307 labelled, hybridised to custom Agilent arrays and scanned according to the 308 manufacturer's instructions (Technology Facility, Dept. of Biology, University of York, 309 UK). Differentially expressed genes were identified by ANOVA using the Benjamini-310 Hochberg FDR correction. No commercially available microarrays were available for P. 311 fluorescens, so a custom design was created using the Agilent eArray system 312 (http://earray.chem.agilent.com/earray). The P. fluorescens SBW25 genome was 313 loaded into Artemis (24) (http://www.sanger.ac.uk/resources/software/artemis), and all 314 open reading frames marked as CDSs and these putative CDS written to a FASTA file 315 which was uploaded to eArray. A probeset was created containing five unique probes 316 per CDS, plus the appropriate Agilent control sequences. The probes were then laid out 317 onto a standard Agilent 8x60K format slide. Microarrays were validated using gRT-PCR 318 (MyiQTM, Bio-Rad). 319

321 RNA labelling and hybridisation

RNA isolated from each sample was labelled with Cy-3 using the Agilent Low Input
Quick Amp WT Labelling Kit (one color) according to the manufacturer's instructions.
Briefly, the RNA is reverse transcribed using a primer mix containing oligo-dT and
random nucleotide primers containing T7 promoters, then the cRNA amplified using the
T7 polymerase in the presence of Cy-3 labelled dCTP. The labelled cDNA was then
quality controlled and only samples with a specific activity of > 6 were used hybridised
to the arrays.

Samples were hybridised to the arrays, incubated and washed in accordance with the
manufacturer's instructions, and the slides scanned using an Agilent Microarray
Scanner and the fluorescence quantified with the Feature Extraction software. The
resulting files were then loaded into GeneSpring for further analysis.

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334 Initial Data Analysis

GeneSpring software version 12.6 was used for all data analysis. Initial analysis was performed to identify those genes that were differentially expressed between any conditions in the experiment, followed by Gene Set Expression Analysis (GSEA) on the basis of Gene Ontology terms. Briefly, the sample replicates were grouped, and then ANOVA performed to identify differentially expressed genes, using the Benjamini-Hochberg FDR p-value correction (with a FDR of 5%), and a Tukey HSD post-hoc test used for each gene to identify which samples were significantly different from each

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other for that gene. Those GO categories that were over-represented in the set of
 differentially expressed genes were then identified.

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345 **3D protein models**

The IntFOLD server (*25*) version 2 was used to build multi-template 3D models from the wild type and mutant sequences of AdnA, NtrB, NtrC, FleQ, GlnA and GlnK. The 3D models were quality checked using the ModFOLD4 protocol (*26*). Intrinsic protein disorder was predicted using the DISOclust method (*27*). Binding sites were predicted using the FunFOLD2 method (*28*). Models were structurally aligned and scored using TM-align (*29*).

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353 Further Discussion of Microarray Results

Full transcriptome analysis is reported separately in Supplementary File "Microarray dataset.xlsx". When genes were filtered for significant fold changes ($P \le 0.05$) of ≥ 2 , and sorted by descending fold changes, the genes that showed the greatest changes in expression are those highlighted in this study. Specifically, when comparing AR2 and AR2S, the large majority of genes with greatest up regulation are related to nitrogen transport, whereas when comparing AR2S to AR2F, the large majority of genes are related to flagellar motility and chemotaxis.

Our microarray data show that the average level of flagella gene expression in AR2S is between 1.5-2-fold lower than in SBW25, but considerably higher than AR2 (between 3 and 75 fold). There are at least two possible models to explain why we were unable to 364 observe flagella in AR2S: (i) Phosphorylated NtrC (NtrC-P) only weakly interacts with flagella gene promoters, but the over-abundance of NtrC-P in AR2S saturates NtrC-365 dependent promoters and the excess NtrC-P drives expression of flagella genes. The 366 level of transcription is suboptimal, thus the flagella may be unstable and support limited 367 or transient motility; (ii) Promiscuous activity of over-abundant NtrC-P (e.g. interacting 368 with RpoN from solution) leads to expression from flagella promoters, but this is not 369 equivalent in all cells and only a proportion of the population express sufficient levels to 370 produce a functional flagellum at any one time. 371



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Fig. S1: Validation of gene expression changes detected by microarray using qRT-PCR.

A – **E** Box plots show gene expression, relative to wild-type SBW25, in genes related to nitrogen utilization and flagellar function. Note, data presented on log scales.

A. SBW25∆*fleQ*

	GENE	GENE CHIP ARRAY		REAL TIME PCR	
	SYMBOL	Fold Change	p-value	Fold Change	p-value
Bootorial type	flgF	-79.06	1.52E-07	-8.4	0.196
Flagellum	Pflu4448 (Flagellin)	-158.33	2.09E-07	-22.59	0.002
Regulation of	gInA	1.01	2.24E-06	9.14	0.187
Nitrogen	ntrB	-1.54	1.37E-07	-4.5	0.41
Utilization	ntrC	-1.74	1.58E-07	-1.39	0.136

B. AR2

	GENE	GENE CHIP ARRAY		REAL TIME PCR		
	SYMBOL	Fold Change	p-value	Fold Change	p-value	
Bootorial type	flgF	-76.48	1.52E-07	-2.84	0.265	
Flagellum	Pflu4448 (Flagellin)	-128.22	2.09E-07	-6.61	0.004	
Regulation of	gInA	1.03	2.24E-06	3.27	0.393	
Nitrogen	ntrB	-1	1.37E-07	-2.24	0.424	
Utilization	ntrC	-1.41	1.58E-07	-1.31	0.231	

C. AR2S

	GENE	GENE CHIP ARRAY		REAL TIME PCR		
	SYMBOL	Fold Change	p-value	Fold Change	p-value	
Bootorial type	flgF	-2.53	1.52E-07	3.13	0.192	
Flagellum	Pflu4448 (Flagellin)	-1.71	2.09E-07	4.34	0.042	
Regulation of	gInA	6.08	2.24E-06	25.49	0.008	
Nitrogen	ntrB	28.18	1.37E-07	26.45	0.017	
Utilization	ntrC	27.14	1.58E-07	49.14	0.059	

D. AR2F

GENE		GENE CHIP ARRAY		REAL TIME PCR	
	SYMBOL	Fold Change	p-value	Fold Change	p-value
Bootorial type	flgF	1.1	1.52E-07	4.49	0.1
Flagellum	Pflu4448 (Flagellin)	3.6	2.09E-07	16.57	0.065
Regulation of	gInA	11.34	2.24E-06	175.63	0.005
Nitrogen	ntrB	58.27	1.37E-07	96.09	0.006
Utilization	ntrC	63.72	1.58E-07	49.73	0.03

Table S1: Mean fold change (relative to SBW25) from qRT-PCR and microarrays with p-values (A - D).

Colours correspond to genes with significant (≥ 2 fold change) up regulated (red) and down regulated (blue) changes.

Strain	Nucleotide change	AA change	Gene	Domain/Function
AR2S_1	T376439G*	T97P	ntrB	PAS domain
AR2F_1.1	G374322A*	R442C	ntrC	HTH domain
AR2S_2	G376175A/ T376174A	V185K	ntrB	HK domain
AR2F_2.1	T374622C	K342E	ntrC	HTH domain
AR2S_3	G376193A	D179N	ntrB	HK domain
AR2F_3.1	C374291G	G452R	ntrC	Miscellaneous function
AR2S_4	G376176A T376174A	L184Q V185I	ntrB	HK domain
AR2F_4.1	T374622C/ / A374625	K342V <i>FS</i> from V342	ntrC	HTH domain
Pf0-2xS_1	∆6431771-3*	ΔG85	gidB	Methyltransferase domain
	Δ61/14/2-8/*	FS from 186	ginK	PII regulator
Pf0-2xF_1.1	C3828791*	R442H	ntrC	HIH domain
Pf0-2xS 2	∆6431771-3*	ΔG85	gidB	Methyltransferase domain
	T388861G* ∆902225*	T237P <i>FS</i> from S544	gInA carB	Catalytic domain Miscellaneous function
Pf0-2xF_2.1	G382870A*	A445V	ntrC	HTH domain
Pf0-2xS_3	∆6431771-3*	ΔG85	gidB	Methyltransferase domain
	Δ388958*	<i>FS</i> from T205	glnA	Catalytic domain
Pf0-2xF_3.1	G382880A*	R442C	ntrC	HTH domain
Pf0-2xS_4	A384603C	D228A	ntrB	Miscellaneous function
Pf0-2xF_4.1	A382843G	N454S	ntrC	HTH domain
Pf0-2xF_4.2	C382883A	R441S	ntrC	HTH domain
Pf0-2xF_4.3	A382843G	N454S	ntrC	HTH domain
Pf0-2xF_4.4	C382933T	P424L	ntrC	HTH domain
Pf0-2xF_4.5	A382843G	N454S	ntrC	HTH domain
Pf0-2xS_5	A384603C	D228A	ntrB	Miscellaneous function
Pf0-2xF_5.1	T382951G	L418R	ntrC	Miscellaneous function
Pf0-2xF_5.2	G382963A	G414D	ntrC	Miscellaneous function
Pf0-2xF_5.3	A382843G	N454S	ntrC	HTH domain
Pf0-2xF_5.4	T382928G	F426V	ntrC	HTH domain

* = Identified by genome resequencing.

Table S2: Full details of mutations identified in strains with parental AR2 and Pf0-2x origin.

Numbers before decimal points represent independent lineages, and numbers after represent derived strains. Note fast spreading mutants contain all nucleotide changes within a lineage.