# Spontaneous adaptation of ion selectivity in a bacterial flagellar motor

- 2 Pietro Ridone<sup>1</sup>, Tsubasa Ishida<sup>2</sup>, Yoshiyuki Sowa<sup>2,3</sup>, Matthew A. B. Baker<sup>1,4\*</sup>
- <sup>1</sup>School of Biotechnology and Biomolecular Sciences, University of New South
- 4 Wales, Sydney, Australia.

1

9

10

11

- <sup>2</sup>Department of Frontier Bioscience, Hosei University, Tokyo, Japan.
- <sup>3</sup>Research Center for Micro-Nano Technology, Hosei University, Tokyo, Japan.
- <sup>7</sup> CSIRO Synthetic Biology Future Science Platform, Brisbane, Australia.
- \*correspondence: matthew.baker@unsw.edu.au

## **ABSTRACT**

- Motility provides a selective advantage to many bacterial species and is often
- achieved by rotation of flagella that propel the cell towards more favourable
- conditions. In most species, the rotation of the flagellum, driven by the Bacterial
- Flagellar Motor (BFM), is powered by H<sup>+</sup> or Na<sup>+</sup> ion transit through the torque-
- generating stator subunits of the motor complex. The ionic requirements for motility
- appear to have adapted to environmental changes throughout history but the
- molecular basis of this adaptation, and the constraints which govern the evolution of
- the stator proteins are unknown. Here we use CRISPR-mediated genome
- 20 engineering to replace the native H<sup>+</sup>-powered stator genes of Escherichia coli with a
- compatible sodium-powered stator set from *Vibrio alginolyticus* and subsequently
- 22 direct the evolution of the stators to revert to H<sup>+</sup>-powered motility. Evidence from
- 23 whole genome seguencing indicates both flagellar- and non-flagellar-associated
- 24 genes that are involved in longer-term adaptation to new power sources. Overall,
- 25 transplanted Na<sup>+</sup>- powered stator genes can spontaneously incorporate novel
- 26 mutations that allow H<sup>+</sup>-motility when environmental Na<sup>+</sup> is lacking.

**INTRODUCTION** 

27

59

Bacterial motility via the flagellar motor represents one of the earliest forms of 28 locomotion. This rotary motility imparts such significant selective advantage<sup>1,2</sup> that 29 resources are allocated to chemotaxis even in the absence of nutrient gradients<sup>3,4</sup>. 30 Furthermore, the evolutionary origins, and subsequent adaptation of the motor are of 31 significant scientific and public interest<sup>5</sup>, since the BFM holds prominence as an 32 ancient, large, molecular complex of high sophistication. 33 34 The torque that drives the BFM is supplied by motor-associated transmembrane protein-complexes known as stators. The stator complex, an asymmetric 35 heteroheptamer (in E. coli: MotA<sub>5</sub>MotA<sub>2</sub>) most likely acts itself as a miniature rotating 36 nanomachine coupling ion transit to rotation<sup>6,7</sup>. The stators are essential for motility, 37 as they drive rotation, and are accessible for studies in experimental evolution due to 38 their unambiguous role in connecting a specific environmental cue (presence of the 39 coupling ion) to an easily discernible phenotype (cell swimming). Furthermore, the 40 stators have been subject to protein engineering approaches for many years, in 41 particular the synthesis of chimeric stator constructs that enable the motor of *E. coli*, 42 natively proton-driven, to be powered by sodium ion flow<sup>8</sup>. The majority of stators are 43 proton driven, but many that are sodium-driven can be found in nature, and this 44 divergence is presumed to have occurred in the distant past<sup>9-11</sup>. Past reports have 45 argued that H<sup>+</sup>-coupled motility diverged from Na<sup>+</sup>-coupled machinery in ancestral 46 times<sup>12</sup>, but the molecular basis for this adaptation, and the evolutionary landscape 47 that constrains stator adaptation remains unclear. 48 49 In order to simulate the effects of natural evolution on stator adaptation we designed an experiment where an E. coli strain, expressing only a sodium-powered stator, 50 would be introduced to a non-lethal environment (soft agar swim plate) which lacked 51 the power source for the stator (Na<sup>+</sup>). Our hypothesis was that the population would 52 undergo selection for upmotile variants, adapting its stators to function in the new 53 environment. 54 We used genomic editing techniques (no-SCAR CRISPR/Cas9<sup>13</sup> and λ-Red<sup>14</sup>) to 55 replace the native motA motB stator genes of the E. coli BFM with chimeric sodium-56 powered pomA potB (henceforth pots) stator genes derived from V. alginolyticus<sup>8</sup>. 57 We transplanted the pots stator genes at the same location and orientation of the 58 native motA motB locus to preserve the native genomic context of the motile RP437

- 60 E. coli strain. We then examined which genetic changes occurred during growth on
- soft-agar in depleted sodium, that is, under selective pressure for proton-driven
- 62 motility. We performed our directed evolution experiments of our pots E.coli strain in
- the absence of antibiotics to avoid additional, undesired selective pressures <sup>15</sup>.

#### RESULTS

64

65

## Preparation and Directed Evolution of a Na<sup>+</sup> powered E.coli strain

- The RP437 strain was edited to carry the chimeric *pomApotB* stator genes in place
- of the native E. coli motAmotB genes (Fig. 1A) via the no-SCAR Method<sup>13</sup> and
- traditional λ-Red recombineering respectively<sup>14</sup> (Supplementary Fig. 1&4). Following
- verification of successful editing by colony PCR and Sanger Sequencing
- 70 (Supplementary Fig. 2AB), a no-SCAR Pots clone was selected and tested on swim
- 71 plates (Fig. 1B). The edited strain was able to swim on sodium-rich (85 mM NaCl)
- soft agar plates but not on potassium-rich sodium-poor (67 mM KCl, ~8 mM [Na<sup>+</sup>])
- plates (Fig. 1B). This edited strain exhibited the same swimming behaviour as the
- control stator-less strain with motility restored via an inducible plasmid vector that
- could express the *Pots* construct (RP6894 Δ*motAmotB* + pSHU1234, hereby pPots).
- We next challenged this *pots* strains to survive on K<sup>+</sup> based soft agar for prolonged
- 77 periods (Fig. 1C). Motile subpopulations arose spontaneously from inoculated
- colonies within a few days. Cells from the edge of these motile flares were passaged
- onto fresh swim agar for up to 5 passages at 3-4 days intervals (Supplementary Fig.
- 80 5). When multiple flares occurred in a single swim ring, each was individually
- passaged (Fig. 1C), and could be recapitulated (Fig. 1D). Directed evolution
- consistently generated swimming flares when *pots* clones were cultured on agar
- containing yeast extract and tryptone (~8 mM [Na<sup>+</sup>]), but not on minimal media (~
- 1mM total [Na<sup>+</sup>]) or when the *pots* construct was expressed via a plasmid
- (Supplementary Fig. 6). One *pots* strain generated using  $\lambda$ -Red methods<sup>14</sup>, which
- carried the native V. alginolyticus pomA promoter, also successfully produced flares
- 87 (Supplementary Fig. 4).
- 88 Lineages were selected for whole-genome sequencing (WGS) after a preliminary
- screening for mutations in the stator genes by Sanger-sequencing PCR amplicons
- spanning the genomic *pomApotB* locus (Fig. 2A). Variant calling to the MG1655
- 91 reference genome was used to compare single nucleotide polymorphisms (SNPs)
- between members of the same lineage. Our intended *pomApotB* edit was the only

93 difference between the RP437 and *Pots* genomes, indicating that neither no-SCAR

94 nor λ-Red editing had resulted in off-target edits (Fig. 2A). 153 SNPs were called as

variants between our experimental parent RP437 and the MG1655 reference which

- were shared across all lineage members (Supplementary Table).
- 97 Several lineages whose descendants could swim in reduced sodium had mutations
- 98 at the pots locus (L3.3-4-5: PotB G20V; L6.4-5: PomA L183F; L8.3-4-5: PotB
- 99 G20W). In contrast, lineages passaged only on 85 mM Na<sup>+</sup> agar (L1 & L2)
- accumulated mutations not in stators but in the flagellar components. Lineages
- passaged on ~8 mM sodium-poor agar whose descendants could not swim (L4) had
- no mutations on any flagellar genes.
- To examine mutation reproducibility in the stators at higher throughput, we subjected
- 55 pots colonies to directed evolution in 8 mM [Na<sup>+</sup>] agar. These yielded a total of 42
- flares within the first 3 days of incubation, which were then passaged four more
- times at 3-day intervals. At the end of this experiment we selected the 20 terminal
- lineage members which produced the largest swim rings and Sanger-sequenced
- them at the *pots* locus (Supplementary Fig. 7). For these we observed a total of 5
- mutations in pomA (S25C, D31N, P177A, P177Q and L183F) and one more new
- mutation in *potB* (L36Q).
- Over the course of these experiments, we found that three stator residues underwent
- mutation at the same site twice (PomA L183F (2x), PomA P177A & P177Q, PotB
- G20V & G20W). WGS revealed that the *pitA* gene had mutated in three separate
- lineages (L2.5, L3.5, L8.5) with one of the mutations occurring twice (PitA
- 115 W112\*Stop).
- To test for the capacity for reversion, we took sequenced lineages that swam in 8
- mM [Na<sup>+</sup>] agar (L3.3-4-5, L6.4-5, L8.3-4-5) and reintroduced them to an environment
- with 85 mM [Na<sup>+</sup>] (Supplementary Fig. 7C). After 10 rounds of daily passaging, no
- reversion in the mutants that had enabled low-sodium swimming was observed, with
- only a single additional mutation gained: a *potB* T21A mutation in the terminal
- descendant of potB G20W pitA W112\*stop (L8.5).
- We further tested whether evolution could be more easily directed on minimal media
- when starting from a more favourable vantage. We examined all stator mutants that
- swam on 8 mM [Na<sup>+</sup>] plates in conditions of further sodium scarcity (minimal media: 1
- mM [Na<sup>+</sup>]). Initially, only the strains with *potB* G20V and *pomA* P177Q mutations

127

128

129

130

131

132

133

134

135

136

137

138

139

140

141

142

143

144

145

146

147

148

149

150

151

152

153

154

155

156

157

158

could swim, and this capacity was maintained following five passages over 12 days. Sanger sequencing revealed that the terminal descendant of potB G20V pitA G432R mutant (L3.5) gained a further mutation in pomA (M20L) (Supplementary Fig. 7B). A summary of all stator gene mutations obtained from all directed evolution experiments is provided in Fig. 2D. Finally, we characterized rotational phenotypes of the parent and evolved strains in the presence and absence of sodium using a tethered cell assay (Fig. 2B, Supplementary Fig. 8). The rotation speeds of single cells were measured under sodium-free 67mM KCl or 85mM NaCl buffer conditions and also tested in presence of 100 µM of the sodium-blocker phenamil<sup>16</sup>. The potB G20V (L3.3) actively rotated in sodium-free buffers and in presence of 100 µM Phenamil (Fig. 2B) – indicating sodium-free rotation and a disrupted phenamil binding site (Fig. 2B). We further confirmed this phenotype was reversible in single cells by tracking rotation of individual tethered cells as the buffers were sequentially exchanged (Supplementary Fig. 9). We confirmed the PotB G20V H<sup>+</sup>-powered phenotype by introducing the same point mutation (GGG to GTG) on our plasmid vector (pPots) and testing motility in tethered cell assay when the protein was expressed in the stator-less \( \Delta motAB \) RP6894 (Fig. 2C). Finally, we mapped mutants to their homologous position on the recently published high-resolution B. subtilis MotA<sub>5</sub>B<sub>2</sub> structure<sup>17</sup> (Fig. 2EF). All stator mutations accumulated at sites proximal to or within the predicted ion-transport pore, at the interface between the PotB transmembrane domain and the third and fourth transmembrane domains of PomA (Fig. 2F). **DISCUSSION** In our experiments we observed repeated mutation around the pore of the stator complex in response to ion-scarcity in the cell's surrounding environment. We confirmed phenotypic changes by measuring rotation in the absence and presence of the sodium channel blocker phenamil. Our strains adapted quickly to drive rotation in a sodium-poor environment within 2 weeks, indicating that the stators are highly adaptive. Previous reports have shown that bacterial motility can adapt<sup>18</sup> and be rescued<sup>19</sup> via remodelling of the flagellar regulatory network. Ni et al. observed that evolutionary

160

161

162

163

164

165

166

167

168

169

170

171

172

173

174

175

176

177

178

179

180

181

182

183

184

185

186

187

188

189

190

191

192

adaptation of motility occurs via remodeling of the checkpoint regulating flagellar gene expression<sup>18</sup>. Their experiments tracked adaptive changes in swim plates, matching our experiments, however their only selection criteria were for improved swimming in an unhindered swimming population. In agreement with our results (fliM A161V), they found fliM (M67I and T192N) to be the amongst the first genes to mutate in the improved swimmer population but they did not report any changes in flgL ( $\triangle$ A57-Q58), nor, significantly, did they see any mutations in any stator genes. Flagellum-mediated motility also appears to be naturally robust to the loss of regulatory factors, such as the enhancer-binding protein fleQ in *P. fluorescens*, which function can be substituted by distantly related homologous proteins<sup>19</sup>. In contrast, our *E. coli pots* strain faced selective pressure from ion scarcity. Our scenario is reminiscent of previous semi-solid agar experimental evolution studies on the adaptation of antibiotic resistance and recapitulates similar results. In the MEGA plate experiments of Baym et al., they similarly saw that the phosphate transporter pitA was repeatedly mutated, often to a frameshifted or nonsense variant<sup>15</sup>. In a similar experiment, the isocitrate dehydrogenase *icd* was also seen to mutate often<sup>1</sup>. Mutations in stators are known to affect ion usage and may confer dual-ion coupling capacity. For example, the substate preference of the *B. alcalophilus* MotPS stator (Na<sup>+</sup>/K<sup>+</sup> and Rb<sup>+</sup>) was changed with the single mutation M33L in MotS, causing the loss of both K<sup>+</sup>- and Rb<sup>+</sup>-coupling motility in E. coli<sup>20</sup>. Similarly a bi-functional B. clausii MotAB stator (Na<sup>+</sup>/H<sup>+</sup>) triple mutant (MotB V37L, A40S and G42S) was selective only for sodium ions while the combination of mutations G42S, Q43S and Q46A made MotB selective only for H<sup>+</sup> <sup>21</sup>. Except the previously reported variant pomA L183F<sup>22</sup>, none of the other mutations we report have been observed in previous studies involving random mutagenesis in motB using ethylmethane sulfonate<sup>23</sup>, or by mutagenesis in the pomA gene by using hydroxylamine<sup>24</sup>, or in *potB* using error-prone PCR<sup>25</sup>. Adaptation of plasmidencoded stator genes<sup>12,26</sup> has been previously reported. However, a distinct advantage of editing stator genes directly on the E. coli genome is that we can direct the evolution of sodium-stators in vivo and without antibiotics – something not possible in wild-type Vibrio sp. since the cells do not survive at low sodium. Conversely, it is difficult to direct evolution towards reversion because it is not possible, particularly in *E. coli*, to drastically reduce the proton concentration to incentivise the stators' use of sodium. Nevertheless, the fact that no revertants were

194

195

196

197

198

199

200

201

202

203

204

205

206

207

208

209

210

211

212

213

214

215

216

217

218

219

220

221

222

223

224

225

observed agrees with previous work suggesting that requirements for Na<sup>+</sup> binding are more strict than for H<sup>+</sup> binding, and that mutations that convert a sodium powered motor to a proton powered motor are more accessible than the reverse<sup>27,28</sup>. We observed a convergence of mutations on the stator genes and even to the very same nucleotide (GAG (L) to GAA (F) in two separate pomA L183F lineages). Since this was from a clonal population under identical environmental constraints, it suggests that adaptation of the stators is prioritized in changing environments. Mutation of pore-proximal residues into hydrophobic residues (eg. G20V) might hint at a mechanism for varying constrictions in the pore to alter the efficiency of ion binding. Motility confers a fitness advantage that is worth significant energetic investment despite the high cost of synthesizing the flagellar machinery<sup>4,29</sup>. This advantage can only be seized if the correct ions are available for stator-conversion into torque. CRISPR/Cas9 has become a widespread method used for precise genomic edits, yet the reversion of such edits appears rapid and gene targeted. Here, ion scarcity supplied a strong selective pressure that allowed us to identify novel mutations correlated with altered ion-specificity. Our transplant of an unfit protein and the cells' rapid reversion of this edit demonstrates tight evolutionary regulation of the stator subunit in an ancient molecular complex. **MATERIALS AND METHODS** Starting strains and plasmids. E. coli strain RP437 was used as the parent strain for genomic editing experiments<sup>30</sup>. The pSHU1234 (pPots) plasmid encoding pomA and potB25 was used as the template to generate the double stranded donor DNA. This was used to replace the motA and motB gene on the RP437 chromosome. Liquid cell culturing was done using LB broth (NaCl or KCl, Yeast Extract, Bacto Tryptone). Cells were cultured on agar plates composed of LB Broth and 1% Bacto Agar (BD, U.S.A.). Swim plate cultures were performed on the same substrates adjusted for agar content (0.3% Bacto Agar). Editing E. coli with Cas9-assisted Recombineering. This procedure was adapted from the no-SCAR method<sup>13</sup>. The target strain to be edited (E.coli RP437) was sequentially transformed first with the pKD-sgRNA-3'MotA (Sm<sup>+</sup>) plasmid, encoding a sgRNA sequence directed at the 3' end of *motA*, and then

with the pCas9cr4 (Cm<sup>+</sup>) plasmid to yield a parent strain harboring both plasmids. 226 The overlap extension PCR technique<sup>31</sup> was employed to assemble linear double 227 stranded DNA molecules (dsDNA) using 3 starting dsDNA fragments. The resulting 228 donor DNA was electroporated in the plasmid-bearing host and the successfully 229 edited clones were selected via colony PCR and Sanger sequencing of the motAB 230 locus. A list of primers and PCR protocols used in this work is provided in 231 Supplementary Fig. 10. 232 Construction of Pots by λ-Red Recombineering 233 Chromosomal replacement from motAmotB to pomApotB was achieved by using a λ 234 235 Red recombination system, with plasmid pKD46 encoding the Red system and positive selection for the recovery of swimming ability<sup>32</sup>. Motile clones were selected 236 by isolating motile flares on swim plates (Supplementary Fig. 4A). 237 Tethered cell assay preparation and analysis 238 The tethered cell assay was performed as previously described<sup>33</sup>. The tethered cells 239 time lapse videos were recorded at 40x magnification on a phase contrast 240 microscope (Nikon). Time lapse videos were collected using a camera (Chameleon3 241 CM3, Point Grey Research) recording 20 s-long videos at 20 frames per second. 242 Time lapse videos were collected using a camera (Chameleon3 CM3, Point 243 Grey Research) recording 20s-long videos at 20 frames per second. Single cell 244 tracking experiments were collected using a camera (DMK21AU618, Imaging 245 Source) recording 10s-long videos recorded at 60 frames per second. Custom 246 LabView software 10,25 was employed as previously reported to estimate specific 247 rotational parameters of the tethered cells such as rotation frequency (speed), 248 clockwise and counterclockwise bias and switching frequency. Visualization of the 249 250 data was performed using Graph Pad Prism 8. Single Nucleotide Polymorphism (SNP) analysis 251 Whole genome sequencing of 22 E. coli strains was performed using a MiSeq 2x 252 150bp chip on an Illumina sequencing platform. Sequencing was carried out at the 253 Ramaciotti Centre for Genomics, Kensington and delivered as demultiplexed fastQ 254 files (Quality Control: >80% bases higher than Q30 at 2×150 bp). The SNP calling 255 and analysis was performed using Snippy<sup>34,35</sup>. The short reads from sequencing 256 were aligned to the MG1655 reference E. coli genome (GenBank: U00096.2) and to 257 258 a synthetic genome based on MG1655, edited to contain the Pots stator sequences from pPots (pomA/potB) at the motAB locus. 259

### **AUTHOR CONTRIBUTIONS**

- 261 PR and MABB designed and executed experiments in strain editing, molecular
- biology, microbiology and rotational measurement. TS and YS executed experiments
- in strain editing and rotational measurement. MABB executed bioinformatics
- surrounding variant calling. MB supervised the design, execution and writing of the
- project. All authors contributed to writing and revision of the manuscript.

### 266 **ACKNOWLEDGEMENTS**

We would like to acknowledge Myu Yoshida and Rie Ito for technical assistance.

### 268 **FUNDING**

260

- YS was supported by JSPS KAKENHI (JP18H02475 and JP20K06564), MEXT
- 270 KAKENHI (JP19H05404) and Takeda Science Foundation. MABB was supported by
- a UNSW Scientia Research Fellowship, a CSIRO Synthetic Biology Future Science
- 272 Platform 2018 Project Grant, and ARC Discovery Project DP190100497.

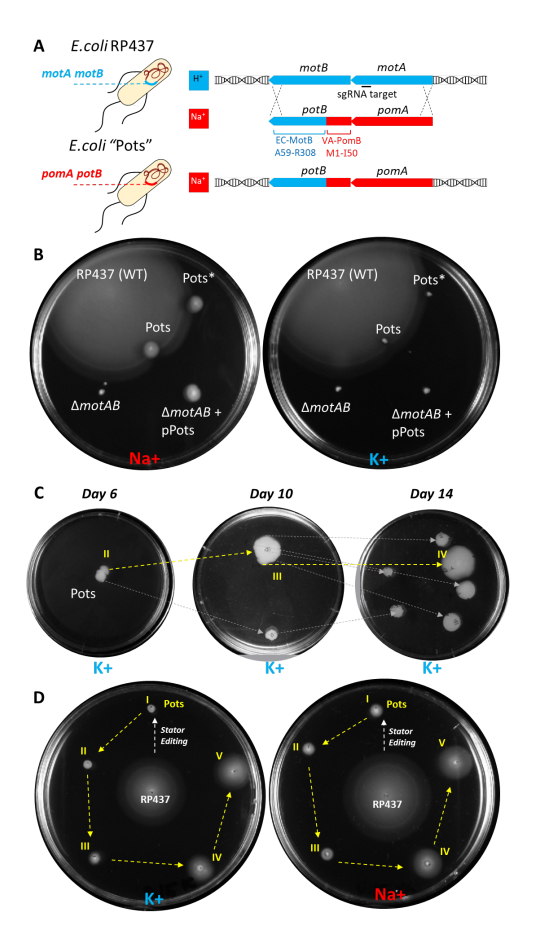
#### REFERENCES

- Fraebel, D. T. *et al.* Environment determines evolutionary trajectory in a constrained phenotypic space. *Elife* **6**, doi:10.7554/eLife.24669 (2017).
- 276 2 Roszak, D. B. & Colwell, R. R. Survival strategies of bacteria in the natural environment.
  277 *Microbiol Rev* **51**, 365-379 (1987).
- 278 3 Gude, S. *et al.* Bacterial coexistence driven by motility and spatial competition. *Nature* **578**, 588-592, doi:10.1038/s41586-020-2033-2 (2020).
- Ni, B., Colin, R., Link, H., Endres, R. G. & Sourjik, V. Growth-rate dependent resource
   investment in bacterial motile behavior quantitatively follows potential benefit of
   chemotaxis. *Proc Natl Acad Sci U S A* 117, 595-601, doi:10.1073/pnas.1910849117 (2020).
- Pallen, M. J. & Matzke, N. J. From The Origin of Species to the origin of bacterial flagella. *Nat Rev Microbiol* **4**, 784-790, doi:10.1038/nrmicro1493 (2006).
- Deme, J. C. *et al.* Structures of the stator complex that drives rotation of the bacterial flagellum. *Nat Microbiol* **5**, 1553-1564, doi:10.1038/s41564-020-0788-8 (2020).
- Santiveri, M. *et al.* Structure and function of stator units of the bacterial flagellar motor.
   2020.2005.2015.096610, doi:10.1101/2020.05.15.096610 %J bioRxiv (2020).
- 289 Asai, Y., Yakushi, T., Kawagishi, I. & Homma, M. Ion-coupling determinants of Na+-driven and H+-driven flagellar motors. *J Mol Biol* **327**, 453-463, doi:10.1016/s0022-2836(03)00096-2 (2003).
- Takekawa, N. *et al.* Structure of Vibrio FliL, a New Stomatin-like Protein That Assists the Bacterial Flagellar Motor Function. *mBio* **10**, doi:10.1128/mBio.00292-19 (2019).
- Islam, M. I., Lin, A., Lai, Y. W., Matzke, N. J. & Baker, M. A. B. Ancestral Sequence
   Reconstructions of MotB Are Proton-Motile and Require MotA for Motility. *Front Microbiol* 11, 625837, doi:10.3389/fmicb.2020.625837 (2020).
- Lai, Y. W., Ridone, P., Peralta, G., Tanaka, M. M. & Baker, M. A. B. Evolution of the Stator Elements of Rotary Prokaryote Motors. *J Bacteriol* **202**, doi:10.1128/JB.00557-19 (2020).
- Takekawa, N. *et al.* Sodium-driven energy conversion for flagellar rotation of the earliest divergent hyperthermophilic bacterium. *Sci Rep* **5**, 12711, doi:10.1038/srep12711 (2015).
- Reisch, C. R. & Prather, K. L. The no-SCAR (Scarless Cas9 Assisted Recombineering) system for genome editing in Escherichia coli. *Sci Rep* **5**, 15096, doi:10.1038/srep15096 (2015).

303	14	Datsenko, K. A. & Wanner, B. L. One-step inactivation of chromosomal genes in Escherichia
304		coli K-12 using PCR products. Proc Natl Acad Sci U S A 97, 6640-6645,
305		doi:10.1073/pnas.120163297 (2000).

- Baym, M. *et al.* Spatiotemporal microbial evolution on antibiotic landscapes. *Science* **353**, 1147-1151, doi:10.1126/science.aag0822 (2016).
- Atsumi, T., Sugiyama, S., Cragoe, E. J., Jr. & Imae, Y. Specific inhibition of the Na(+)-driven flagellar motors of alkalophilic Bacillus strains by the amiloride analog phenamil. *J Bacteriol* **172**, 1634-1639, doi:10.1128/jb.172.3.1634-1639.1990 (1990).
- Deme, J. C. *et al.* Structures of the stator complex that drives rotation of the bacterial flagellum. *Nat Microbiol* **5**, 1553-+, doi:10.1038/s41564-020-0788-8 (2020).
- Ni, B. *et al.* Evolutionary Remodeling of Bacterial Motility Checkpoint Control. *Cell Rep* **18**, 866-877, doi:10.1016/j.celrep.2016.12.088 (2017).
- Taylor, T. B. *et al.* Evolution. Evolutionary resurrection of flagellar motility via rewiring of the nitrogen regulation system. *Science* **347**, 1014-1017, doi:10.1126/science.1259145 (2015).
- Terahara, N., Sano, M. & Ito, M. A Bacillus Flagellar Motor That Can Use Both Na+ and K+ as a Coupling Ion Is Converted by a Single Mutation to Use Only Na+. *Plos One* **7**, doi:ARTN e4624810.1371/journal.pone.0046248 (2012).
- Terahara, N., Krulwich, T. A. & Ito, M. Mutations alter the sodium versus proton use of a Bacillus clausii flagellar motor and confer dual ion use on Bacillus subtilis motors. *P Natl* Acad Sci USA **105**, 14359-14364, doi:10.1073/pnas.0802106105 (2008).
- Sudo, Y., Terashima, H., Abe-Yoshizumi, R., Kojima, S. & Homma, M. Comparative study of the ion flux pathway in stator units of proton- and sodium-driven flagellar motors. *Biophysics* (*Nagoya-shi*) **5**, 45-52, doi:10.2142/biophysics.5.45 (2009).
- Garza, A. G., Harris-Haller, L. W., Stoebner, R. A. & Manson, M. D. Motility protein interactions in the bacterial flagellar motor. *Proc Natl Acad Sci U S A* **92**, 1970-1974, doi:10.1073/pnas.92.6.1970 (1995).
- Kojima, S., Kuroda, M., Kawagishi, I. & Homma, M. Random mutagenesis of the pomA gene encoding a putative channel component of the Na(+)-driven polar flagellar motor of Vibrio alginolyticus. *Microbiology (Reading)* **145 ( Pt 7)**, 1759-1767, doi:10.1099/13500872-145-7-1759 (1999).
- Ishida, T. *et al.* Sodium-powered stators of the bacterial flagellar motor can generate torque
   in the presence of phenamil with mutations near the peptidoglycan-binding region. *Mol Microbiol* 111, 1689-1699, doi:10.1111/mmi.14246 (2019).
- Nishino, Y., Onoue, Y., Kojima, S. & Homma, M. Functional chimeras of flagellar stator proteins between E-coli MotB and Vibrio PomB at the periplasmic region in Vibrio or E-coli. *Microbiologyopen* **4**, 323-331, doi:10.1002/mbo3.240 (2015).
- Mulkidjanian, A. Y., Galperin, M. Y., Makarova, K. S., Wolf, Y. I. & Koonin, E. V. Evolutionary primacy of sodium bioenergetics. *Biol Direct* **3**, 13, doi:10.1186/1745-6150-3-13 (2008).
- Hase, C. C., Fedorova, N. D., Galperin, M. Y. & Dibrov, P. A. Sodium ion cycle in bacterial pathogens: evidence from cross-genome comparisons. *Microbiol Mol Biol Rev* **65**, 353-370, table of contents, doi:10.1128/MMBR.65.3.353-370.2001 (2001).
- Colin, R. & Sourjik, V. Emergent properties of bacterial chemotaxis pathway. *Curr Opin Microbiol* **39**, 24-33, doi:10.1016/j.mib.2017.07.004 (2017).
- Parkinson, J. S. Complementation analysis and deletion mapping of Escherichia coli mutants defective in chemotaxis. *Journal of Bacteriology* **135**, 45-53 (1978).
- Higuchi, R., Krummel, B. & Saiki, R. K. A general method of in vitro preparation and specific mutagenesis of DNA fragments: study of protein and DNA interactions. *Nucleic Acids Res* **16**, 7351-7367, doi:10.1093/nar/16.15.7351 (1988).
- 351 32 Kinosita, Y. *et al.* Distinct chemotactic behavior in the original Escherichia coli K-12 352 depending on forward-and-backward swimming, not on run-tumble movements. *Sci Rep* **10**, 353 15887, doi:10.1038/s41598-020-72429-1 (2020).
- Nishiyama, M. & Kojima, S. Bacterial motility measured by a miniature chamber for highpressure microscopy. *Int J Mol Sci* **13**, 9225-9239, doi:10.3390/ijms13079225 (2012).

356	34	Bush, S. J. et al. Genomic diversity affects the accuracy of bacterial single-nucleotide
357		polymorphism-calling pipelines. Gigascience 9, doi:10.1093/gigascience/giaa007 (2020).
358 359	35	Seemann, T. Snippy: fast bacterial variant calling from NGS reads. <i>Internet</i> https://github.com/tseemann/snippy (2015).
360		



**Figure 1. BFM stator editing and recapitulated directed evolution.** A) Schematic of E. coli RP437, carrying a proton-powered stator set (*motA motB*), undergoing an

362

homologous recombination-catalysed editing process to replace motAB with the sodium-powered chimeric stator pomA potB ('pots') via a dsDNA donor. The chimeric Pots construct consists of DNA that encodes pomA from V. alginolyticus (VA) and a spliced B subunit encoding for Vibrio p omB amino acids M1-I50 fused in frame to E. coli (EC) motB A59 - R308 (now potB A51 - R300). B) Soft agar swim plate assay to assess bacterial motility. The strains indicated above were inoculated on 0.3% Agar prepared with either NaCl LB (left) or KCl LB (right) media and incubated at 30°C for 24 hrs. The plates contain no antibiotics and 0.4% arabinose to induce expression from pPots. Strains: RP437 (WT, parent strain), Pots\* (not-fully cured, carries plasmid pCas9cr4), Pots (cured of all plasmid), \( \Delta motAB \) (E.coli RP6894) with or without pPots (pSHU1234, Cm<sup>+</sup>, encoding the pomApotB construct under arabinose induction). C) Directed evolution experiment plates. A single Pots colony was innoculated on a K<sup>+</sup> soft agar plate and incubated at 30°C until flares developed. The edge of the flare was then transferred onto a fresh plate and allowed to spread radially until it was again transferred at 4-day intervals. Yellow arrows indicate an improved swimmer subpopulation being propagated. Grey arrows indicate other portions of the colony being propagated. D) Full recapitulation of the evolved lineage shown in (C). Lineage members were inoculated on soft LB agar with or without sodium from glycerol stocks and incubated for 24 hrs at 30°C. Parent strain RP437 is also included in the centre of the plate as control. The phenotypic effect of stator replacement is highlighted by the white arrows. Passages 1 to 5 of the directed evolution experiment are indicated in roman numerals and each passage highlighted by yellow arrows. These passages correspond to: I = Pots, II = L4.2, III = L5.3, IV = L6.4 and V = L6.5 in Fig. 2, respectively.

364

365

366

367

368

369

370

371

372

373

374

375

376

377

378

379

380

381

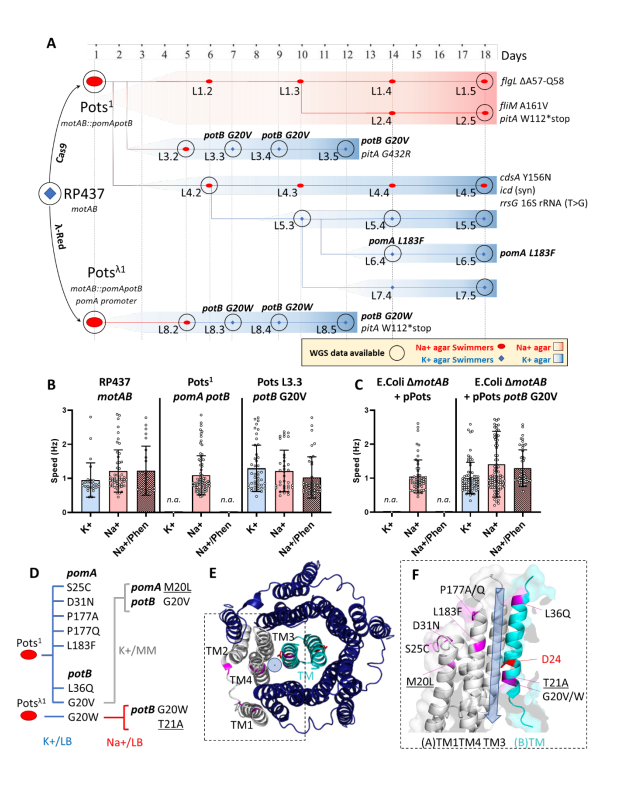
382

383

384

385

386



**Figure 2. Sequencing and phenotyping of evolved lineages.** A) Schematic of directed evolution experiments. The lineage members in the diagram are color-coded based on their ability to swim on either a Na<sup>+</sup> -based substrate (red ellipse) or K<sup>+</sup>-based swim agar (blue diamond). All lineage members labelled in blue retained their ability to swim on Na<sup>+</sup>-rich soft agar. Lineages passaged on K<sup>+</sup> agar are highlighted by a blue gradient bar, while lineages passaged on Na<sup>+</sup> agar are highlighted by a red gradient bar. All strains were locally Sanger sequenced at the *pomApotB* locus, but 21

399

400

401

402

403

404

405

406

407

408

409

410

411

412

413

414

415

416

417

418

419

420

421

422

423

424

425

426

427

428

429

430

strains, indicated by black circles, underwent whole genome sequencing (WGS). Lineage number and passage are indicated: ie L1.3 indicates the first lineage and the third passage of a motile flare from an initial inoculation site. SNPs identified via variant calling relative to reference genome of MG1655 are annotated next to each respective lineage member. Highlighted genes other than pomA and potB: pitA (metal phosphate:H<sup>+</sup> symporter), flgL (flagellar hook-filament junction protein 2), fliM (flagellar motor switch protein), cdsA (cardiolipin-diglyceride synthase), icd (isocitrate dehydrogenase), rrsG (16S ribosomal RNA). B) Single cell speed measurements using the tethered cell assay measured in Hz (revolutions/s). Blue bar indicates speed in 67 mM KCl motility Buffer, red bar: 85 mM NaCl motility Buffer; red patterned bar: 85 mM NaCL + 100 µM phenamil motility buffer. Number of cells analysed per condition (from left to right): RP437: 27, 51, 25; pots: n.a, 78, n.a; pots potB G20V 45, 36, 39 (n.a. indicates no visible rotating cell). Error bars indicate Standard Deviation (S.D.). C) Single cell speed measurements using the tethered cell assay in RP6894 ΔmotAB strain co-expressing pomA and potB G20V from pPots plasmid. Blue bar indicates speed in 67 mM KCl motility buffer, red bar: 85 mM NaCl motility buffer; red patterned bar: 85 mM NaCl + 100 µM phenamil motility buffer. Number of cells analysed per condition (from left to right): ( $\Delta motAB + pPots: n.a., 32, n.a; \Delta motAB +$ pPots potB G20V: 40, 63, 48). Error bars indicate S.D. D) Graphical summary of stator gene mutations detected across all directed evolution experiments and the growth conditions under which these mutations arose. LB indicates agar containing Yeast extract and Tryptone. MM indicates agar in minimal media. Mutations in a subsequent generation are underlined. E) View from the extracellular side of the transmembrane portion of B. subtilis MotA<sub>5</sub>B<sub>2</sub> stator complex (PDB: 6YSL). One monomer of subunit A is coloured in white and the TM domains of the B subunits are coloured in cyan. Mutant sites obtained in the directed evolution experiments are labelled in magenta, the catalytic aspartate residue essential for function is highlighted in red. The light blue circle indicates the predicted location of the ion transport pore (inward conduction). F) side view of the area highlighted by the dashed box in (E). Homologous residue location derived from protein sequence alignments. Primary mutation sites are indicated in black, while secondary mutation sites are underscored. The arrow at the interface between (A)TM3-4 and (B)TM indicates the predicted location of the ion transport pore.