1	ComM is a hexameric helicase that promotes branch migration during
2	natural transformation in diverse Gram-negative species
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13	ABSTRACT
14	Acquisition of foreign DNA by natural transformation is an important mechanism of adaptation
15	and evolution in diverse microbial species. Here, we characterize the mechanism of ComM, a
16	broadly conserved AAA+ protein previously implicated in homologous recombination of
17	transforming DNA (tDNA) in naturally competent Gram-negative bacterial species. In vivo, we
18	found that ComM was required for efficient comigration of linked genetic markers in Vibrio
19	cholerae and Acinetobacter baylyi, which is consistent with a role in branch migration. Also,
20	ComM was particularly important for integration of tDNA with increased sequence heterology,
21	suggesting that its activity promotes the acquisition of novel DNA sequences. In vitro, we
22	showed that purified ComM binds ssDNA, oligomerizes into a hexameric ring, and has
23	bidirectional helicase and branch migration activity. Based on these data, we propose a model
24	for tDNA integration during natural transformation. This study provides mechanistic insight into
25	the enigmatic steps involved in tDNA integration and uncovers the function of a protein
26	required for this conserved mechanism of horizontal gene transfer.
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29	

30 INTRODUCTION

31 Natural competence is a physiological state in which some bacterial species can take up free 32 DNA from the environment. Some competent species regulate the genes required for this 33 process and, depending on the organism, competence can be induced in response to the 34 availability of certain nutrients, quorum sensing pathways, or by DNA damage / stress (1). The 35 Gram-negative bacterium Vibrio cholerae is activated for competence during growth on chitin, 36 a polymer of β 1,4-linked N-acetyl glucosamine (2). Chitin is the primary constituent of 37 crustacean exoskeletons and is commonly found in the aquatic environment where this 38 facultative pathogen resides. Soluble chitin oligosaccharides indirectly induce expression of 39 TfoX, the master regulator of competence (3,4), which regulates expression of competence-40 related genes in concert with HapR, the master regulator of quorum sensing (5,6). 41 Acinetobacter baylyi ADP1, on the other hand, is a naturally competent Gram-negative microbe 42 that is constitutively active for competence throughout exponential growth (7). 43 44 During competence, dsDNA is bound extracellularly, however, only a single strand of this DNA is 45 transported into the cytoplasm. Competent bacterial species may use this ingested DNA as a 46 source of nutrients, however, if this DNA has sufficient homology to the host chromosome, the 47 incoming DNA can also be integrated into the bacterial genome by homologous recombination 48 (8). This process of DNA uptake and integration is referred to as natural transformation. As a 49 result, natural transformation is an important mechanism of horizontal gene transfer and can 50 lead to the repair of damaged DNA or facilitate acquisition of novel genetic information. 51 Homologous recombination of single-stranded transforming DNA (tDNA) with the host 52 chromosome requires the function of RecA, which facilitates homology searching and initiates 53 strand invasion of tDNA through the formation of a displacement loop (D-loop). Following RecA 54 mediated strand invasion, DNA junctions of this D-loop can then be moved in a process known 55 as branch migration to increase or decrease the amount of tDNA integrated. Then, by a 56 presently unresolved mechanism, this intermediate is resolved to stably integrate tDNA into the 57 host chromosome. The molecular details involved in the integration of tDNA downstream of

58 RecA strand invasion, however, are poorly understood.

59

One previously studied gene from the competent species *Haemophilus influenzae, comM*, is not required for DNA uptake but is required for the integration of tDNA into the host chromosome, a phenotype consistent with ComM playing a role in homologous recombination during natural transformation (9). The function of ComM, however, has remained unclear. Here, through both *in vivo* and *in vitro* characterization of ComM in *V. cholerae* and *A. baylyi*, we uncover that this protein functions as a hexameric helicase to aid in branch migration during this conserved mechanism of horizontal gene transfer.

67

68 MATERIALS AND METHODS

69 Bacterial strains and growth conditions

70 The parent V. cholerae strain used throughout this study is E7946 (10), while the A. baylyi strain

used is ADP1 (also known as BD413) (11). A description of all strains used in this study are listed

in Table S1. Strains were routinely grown in LB broth and plated on LB agar. When required,

73 media was supplemented with 200 μg/mL spectinomycin, 50 μg/mL kanamycin, 100 μg/mL

74 carbenicillin, $10 \,\mu\text{g/mL}$ trimethoprim, or $40 \,\mu\text{g/mL}$ X-Gal.

75

76 Construction of mutants and transforming DNA

- 27 Linear PCR product was constructed using splicing-by-overlap extension PCR exactly as
- 78 previously described (12). All primers used to construct and detect mutant alleles are described
- in **Table S2**. The pBAD18 Kan plasmid used as tDNA was purified from TG1, a *recA*+ *E. coli* host.

80 Mutants were made by cotransformation exactly as previously described (13).

81

82 <u>Transformation assays</u>

- 83 Transformation assays of *V. cholerae* on chitin were performed exactly as previously described
- 84 (3). Briefly, ~10⁸ CFU of mid-log *V. cholerae* were incubated statically in instant ocean (IO)
- 85 medium (7g/L; aquarium systems) containing chitin from shrimp shells (Sigma) for 16-24 hours
- 86 at 30°C. Then, tDNA was added (500 ng for linear products containing an antibiotic resistance
- 87 cassette inserted at the non-essential locus VC1807 and 1500 ng for plasmid DNA), and

reactions were incubated at 30°C for 5-16 hours. Reactions were outgrown with the addition of LB medium to each reaction by shaking at 37°C for 1-3 hours and then plated for quantitative culture onto medium to select for the tDNA (transformants) or onto plain LB (total viable counts). Transformation efficiency is shown as the number of transformants / total viable counts. In cases where no colonies were observed, efficiencies were denoted as below the limit of detection (LOD) for the assay.

- 95 Chitin-independent transformation assays were performed exactly as previously described
 96 using strains that contain an IPTG inducible P_{tac} promoter upstream of the native TfoX gene
 97 (14). Briefly, strains were grown overnight with 100 µM IPTG. Then, ~10⁸ cells were diluted into
 98 instant ocean medium, and tDNA was added. Reactions were incubated statically for 5 hours
 99 and then outgrown by adding LB and shaking at 37°C for 1-3 hours. Reactions were plated for
 100 quantitative culture as described above.
- 101

102 For transformation of *A. baylyi* ADP1, strains were grown overnight in LB media. Then, ~10⁸

103 cells were diluted into fresh LB medium, and tDNA was added (~100ng). Reactions were

104 incubated at 30°C with agitation for 5 hours and then plated for quantitative culture as

105 described above.

106

107 Protein expression and purification

ComM and Pif1 were cloned into StrepII expression vectors, expressed in Rosetta 2(DE3) pLysS
 cells using autoinduction medium (15), and purified using Strep-Tactin Sepharose (IBA). ComM
 protein preparations lacked detectable nuclease activity. For details please see Supplementary
 Methods.

112

113 *Electrophoretic mobility shift assay*

114 The ssDNA probe (BBC742) and dsDNA probe (annealed BBC742 and BBC743) were labeled

- using Cy5 dCTP (GE Healthcare) and Terminal deoxynucleotidyl Transferase (TdT; Promega).
- 116 Reactions were composed of 10 mM Tris-HCl pH 7.5, 20 mM KCl, 1 mM DTT, 10% Glycerol, 100

- 117 μg/ml BSA, 9 nM Cy5 labeled probe, purified ComM protein at the indicated concentration, and
- 118 0.1 mM ATP where indicated. Reactions were incubated at RT for 30 minutes and run on 8%
- 119 Tris-borate acrylamide gel at 150 V for 45 min. Probes were detected using a Chemidoc MP
- 120 imaging system (Biorad).
- 121
- 122 Blue native PAGE
- 123 Blue native PAGE was performed essentially as previously described (16). Purified ComM (2.5
- 124 μM) was incubated for 30 min at room temperature in reaction buffer [10 mM Tris-HCl pH 7.5,
- 125 20 mM KCl, 1 mM DTT, 10% Glycerol] with 5 mM ATP and/or 5 μ M ssDNA (oligo ABD363) where
- 126 indicated. For additional details see **Supplementary Methods**.
- 127

128 <u>Negative stain electron microscopy</u>

- 129 For negative stain electron microscopy, sample was prepared by applying 4 µL of ComM-ATP-
- 130 ssDNA solution onto a glow-discharged continuous carbon film coated copper grid (EMS) and
- 131 stained with 0.75% (w/v) uranyl formate. EM micrographs were acquired using a 300 kV JEM-
- 132 3200FS electron microscopy with 20-eV energy slit under low dose conditions ($\leq 20 \text{ e}^2/\text{Å}^2$) on a
- 133 Gatan UltraScan 4000 4k x 4k CCD camera. Additional details for EM image collection and
- analysis are provided in the **Supplementary Methods**. The EM electron density map has been
- 135 deposited to EMDataBank.org with the accession number EMD-8575.
- 136

137 *Phylogenetic Trees*

138 ComM homologs were identified using a protein BLAST search of diverse bacterial genomes

139 followed by phylogenetic analysis. Starting with a broadly representative set of genomes

- adapted from Wu and Eisen (17), an initial ComM candidate pool was generated from a
- 141 comprehensive protein BLAST search of all predicted CDS translations against the V. cholerae
- 142 ComM allele with a 0.001 e-value cutoff. Subsequent sequence alignment with MUSCLE (18)
- and maximum likelihood phylogenetic reconstruction with FastTree (19) identified a single
- 144 clade of alleles with high sequence similarity (generally well above 50%) to *V. cholerae* ComM
- 145 (data not shown), with the remaining alleles excluded from further analysis. After manually

pruning divergent alleles with alignments covering <70% of ComM, the retained sequences
comprised the set of true ComM orthologs used in the final sequence alignment and ComM
phylogeny reconstruction. For whole genome phylogenetic reconstruction, Phylosift (20)
identified and aligned a default set of 36 highly conserved marker genes. FastTree was used for
initial reconstruction, whereas RAxML (21) subsequently estimated the maximum likelihood
phylogeny for a reduced set of representative genomes under the LG (22) substitution model
with gamma-distributed rate variation.

153

154 *Helicase and branch migration assays*

Helicase assay substrates were 5' end-labeled with T4 polynucleotide kinase (T4 PNK; NEB) and $\gamma[^{32}P]$ -ATP. The 2-strand forked helicase substrates was annealed by combining equimolar amounts of a labeled oligonucleotide with the unlabeled complement and incubating at 37°C overnight.

159

160 The short 3-strand branch migration substrates were constructed in a 2-step annealing process. 161 First, we incubated equimolar amounts of a labeled strand and unlabeled partial complement 162 at 95°C for 5 min. and then slow cooled to 25°C to create a forked substrate. An equimolar 163 amount of a third strand was then added to the resulting forked product and incubated 164 overnight at 37°C. Following annealing, the three-stranded products were PAGE purified. 165 Primers used for all probes and additional details can be found in the **Supplementary Methods**. 166 Helicase and branch migration activity was then assessed by incubating the indicated 167 concentrations of ComM, Pif1, or Hrg1 with 5 mM ATP and 0.1 nM DNA substrate in 168 resuspension buffer (25 mM Na-HEPES (pH 7.5), 5% (v/v) glycerol, 300 mM NaOAc, 5 mM 169 MgOAc, and 0.05% Tween-20). Reactions were incubated at 37°C for 30 min and stopped with 170 the addition of 1x Stop-Load dye (5% glycerol, 20 mM EDTA, 0.05% SDS, and 0.25% 171 bromophenol blue) supplemented with 400 µg/mL SDS-Proteinase K followed by a 10-min 172 incubation at 37°C. Reactions were then separated on 8% 19:1 acrylamide:bis-acrylamide gels 173 in TBE buffer at 10 V/cm. Gels were then dried between layers of Whatman filter paper under 174 vacuum at 55°C for 20 mins and then exposed to a phosphor imaging screen for 24-48 hours

prior to scanning on a Typhoon 9210 Variable Mode Imager to image radiolabeled DNA probes.

- 176 DNA unwinding and branch migration were quantified using ImageQuant 5.2 software.
- 177

178 The long 3-strand recombination intermediates were generated by RecA-mediated strand 179 exchange between circular single-stranded ϕ X174 virion DNA (NEB) and PstI-linearized double-180 stranded ϕ X174 DNA (NEB) essentially as previously described (23). Briefly, recombination 181 reactions were conducted in strand exchange buffer (25 mM Tris acetate pH 8.0, 1mM DTT, 1% 182 glycerol, 10 mM magnesium acetate, 3 mM potassium glutamate, 10 U/mL creatine 183 phosphokinase, 12.5 mM phosphocreatine, and 50 μ g/mL acetylated BSA). First, 7.1 μ M RecA 184 (NEB) was incubated with 44 μ M (in nucleotides) circular single-stranded ϕ X174 virion DNA 185 (CSS) at 37°C for 10 mins. Then, 2 mM ATP and ~0.84 µM SSB (Promega) were added to the 186 reaction and incubated at 37°C for 8 mins. Finally, the strand exchange reactions were started 187 by the addition of ~16.7 μ M (in nucleotides) of PstI-linearized ϕ X174 and incubated at 37°C for 188 an additional 20 mins to allow for the generation of recombination intermediates. Reactions 189 were then deproteinated and cleaned up using a PCR purification kit (Qiagen). To assess branch 190 migration of recombination intermediates, the deproteinated DNA was then incubated with 191 ComM in strand exchange buffer containing 5 mM ATP at 37°C for 10 mins. Reactions were 192 then stopped using 1X stop load dye and separated on a 0.8% agarose gel in TAE. Gels were 193 then stained with GelGreen (Biotium), and scanned on a Typhoon 9210 Variable Mode Imager. 194

195 DNA damage assay

196 For DNA damage assays, ~10⁸ cells of a midlog *Vibrio cholerae* culture in instant ocean medium

- 197 were treated with the indicated concentration of MMS or MMC for 1 hour at 30°C. To
- 198 determine viable counts, reactions were plated for quantitative culture on LB agar.
- 199
- 200 <u>GFP-ComM western blots</u>

201 The indicated strains of *V. cholerae* were grown with or without 100 µM IPTG. Cell lysates were

- run on 10% SDS PAGE gels and electrophoretically transferred to PVDF. This membrane was
- 203 then blotted with primary rabbit anti-GFP (Invitrogen) or mouse anti-RpoA (Biolegend)

- antibodies and a goat anti-rabbit or anti-mouse IRDye 800CW (LI-COR) secondary as
- appropriate. Bands were detected using a LI-COR Odyssey classic infrared imaging system.
- 206

207 **RESULTS**

208 <u>ComM is required for integration of tDNA during natural transformation</u>

209 Previously, we performed an unbiased transposon-sequencing screen (Tn-seq) in V. cholerae to 210 identify genes involved in natural transformation (3). One gene identified in that screen was 211 VC0032, which encodes a homolog of *comM* from *H. influenzae*. ComM was previously 212 implicated in the integration of tDNA during natural transformation in *H. influenzae* (9). To 213 determine if this was also the case in V. cholerae, we performed chitin-dependent natural 214 transformation assays using two distinct sources of tDNA. One was a linear PCR product that 215 inserts an antibiotic resistance cassette at a non-essential locus, while the other was a 216 replicating plasmid that lacks any homology to the host genome. We hypothesized that 217 transformation with linear product requires both DNA uptake and chromosomal integration, 218 while transformation with the plasmid only requires DNA uptake. To test this, we transformed a 219 recombination deficient $\Delta recA$ strain of V. cholerae, and found that, as expected, this strain 220 could not be transformed with linear product but could be transformed with a replicating 221 plasmid, consistent with plasmid transformation being recombination-independent (Fig. S1). 222 Additionally, plasmid transformation in this assay is dependent on natural competence, as 223 mutants in genes required for uptake ($\Delta pilA$) (5) and cytoplasmic protection ($\Delta dprA$) of tDNA 224 are not transformed (5,24,25) (Fig. S1). Using this assay, we find that a comM mutant 225 $(\Delta VC0032)$ in V. cholerae displays a ~100-fold reduction for transformation with linear product, 226 while rates of plasmid transformation were equal to the WT (Fig. 1A). This is consistent with 227 ComM playing a role downstream of DNA uptake, and potentially during recombination. These 228 assays were performed on chitin to induce the natural competence of this organism. To 229 determine if ComM is playing a role specifically downstream of competence induction, we 230 performed a chitin-independent transformation assay by overexpressing the competence 231 regulator TfoX. Under these conditions, a comM mutant still had reduced rates of 232 transformation when transformed with a linear PCR product (Fig. 1B). Cumulatively, these

233	results are consistent with ComM playing a role in the late steps of transformation downstream
234	of DNA uptake.
235	
236	To confirm that the phenotypes observed are due to mutation of <i>comM</i> , we complemented
237	strains by integrating an IPTG inducible P_{tac} -comM construct at a heterologous site on the
238	chromosome (at the <i>lacZ</i> locus). Our previous work has indicated that this expression construct
239	is leaky (14) (Fig. S2), and consistent with this, we observe complementation even in the
240	absence of inducer (Fig. 1C).
241	
242	ComM promotes branch migration through heterologous sequences in vivo
243	V. cholerae ComM is a predicted AAA+ ATPase, and members of this family have diverse
244	functions (26). To determine if ComM had structural similarity to any AAA+ ATPase of known
245	function, we submitted the primary sequence of this protein to the Phyre2 server (27). Despite
246	a lack of significant homology by BLAST, this analysis revealed structural similarity to MCM2-7,
247	the replicative helicase of eukaryotes (28,29). As a result, we explored whether ComM
248	functions as a helicase to promote branch migration during natural transformation.
249	
250	To test this in vivo, we assessed comigration of linked genetic markers on a linear tDNA product
251	(Fig. 2A). If branch migration during transformation is efficient, we hypothesized that both
252	markers would be integrated into the host chromosome. However, if branch migration is
253	inefficient, we hypothesized that we may observe integration of one marker but not the other.
254	For this assay, we generated two tDNA constructs that contained a Spec ^R marker upstream of
255	<i>lacZ</i> as well as a genetically linked point mutation in the <i>lacZ</i> gene that was either 820bp or
256	245bp downstream of the Spec ^R marker (Fig 2A). We selected for the Spec ^R marker and
257	screened for integration of the linked <i>lacZ</i> mutation as an indirect measure of branch migration.
258	To prevent post-recombination repair of the <i>lacZ</i> allele by the mismatch repair (MMR) system,
259	these experiments were performed in <i>mutS</i> mutant backgrounds. As with previous
260	experiments, a <i>comM</i> mutant is severely reduced for transformation efficiency for the Spec ^R
261	marker compared to the parent (Fig. 2A, left and 1A). Of those cells that integrated the Spec ^R

262 marker, the comigration efficiency of the *lacZ* mutation was higher than 90% for both products 263 (820bp and 245bp) in the parent strain background (Fig. 2A, right), which is consistent with 264 highly efficient branch migration in this background. When *comM* is deleted, however, the 265 comigration efficiency for the *lacZ* mutation drops significantly for both products compared to 266 the parent strain, and the reduction is more severe for the product where the *lacZ* mutation is 267 farther away (Fig. 2A, right). Cumulatively, these data suggest that comM may play a role in 268 branch migration during natural transformation to increase the amount of tDNA integrated into 269 the host chromosome.

270

271 To promote horizontal gene transfer, tDNA integrated during natural transformation must be 272 heterologous to the host chromosome. So next, we decided to test whether ComM promotes 273 integration of heterologous tDNA. To that end, we created strains that contain an inactivated 274 Tm^{κ} marker integrated in the chromosome. The marker was inactivated with either a 29bp 275 deletion or a nonsense point mutation. We then transformed these strains with tDNA that 276 would restore the Tm^R marker. Again, to eliminate any confounding effects of MMR, we 277 performed these experiments in *mutS* mutant backgrounds. First, we find that integration of a 278 point mutation is similar to a 29bp insertion in the parent strain background, indicating that in 279 the presence of ComM, tDNA is efficiently integrated regardless of sequence heterology. In the 280 *comM* mutant, however, we find that a point mutation is significantly easier to integrate 281 compared to the 29bp insertion (Fig. 2B). This finding is consistent with ComM promoting 282 branch migration through heterologous sequences during natural transformation; however, in 283 its absence, the integration of heterologous sequences is unfavored.

284

285 <u>ComM hexamerizes in the presence of ATP and ssDNA</u>

286 Because our *in vivo* data suggested that ComM acts as a branch migration factor, we next

- 287 decided to test the biochemical activity of this protein in vitro. First, we determined that N-
- terminally tagged ComM (GFP-ComM) was functional *in vivo* while a C-terminally tagged fusion
- 289 (ComM-GFP) was not (Fig. S2). Furthermore, recent studies indicate that *comM* is part of the
- 290 competence regulon (30). Using a strain where *comM* was N-terminally tagged with GFP at the

native locus, we found that *comM* protein levels are increased under competence inducing
 conditions (TfoX overexpression), indicating that native regulation of tagged *comM* is
 maintained (Fig. S2).

294

295 To characterize ComM in vitro, we expressed StrepII-ComM (N-terminal tag) in E. coli and 296 purified it to homogeneity. The peak of recombinant ComM eluting from preparative gel 297 filtration chromatography had a calculated molecular weight of 57 kDa (Fig. S3). As the 298 predicted mass of ComM is 61 kDa, this suggests that ComM exists as a monomer in solution. 299 Many helicases oligomerize in their active state, including all known MCM family helicases (31). 300 So, we next tested whether purified ComM oligomerizes *in vitro*. Because ComM is a predicted 301 ATPase and may interact with DNA, we hypothesized that these factors may be required for its 302 oligomerization and activity. To assess oligomerization, we performed blue-native PAGE (16). In 303 this assay, ComM appears to oligomerize robustly in the presence of ATP and ssDNA, with some 304 oligomerization also observed in the presence of ATP alone (Fig. 3A). This latter observation, 305 however, may be due to a small amount of contaminating ssDNA that remains bound to ComM 306 during purification. Defining the number of subunits in this oligomer was unreliable by blue-307 native PAGE due to lack of resolving power by the gel. However, the higher molecular weight 308 species generated in the presence of ATP and ssDNA was likely larger than a dimer. Therefore, 309 we attempted to observe ComM oligomers by negative stain transmission electron microscopy 310 (TEM). In the absence of ligands, ComM particles were small and uniform, consistent with our 311 gel filtration results and demonstrated the purity of our protein preparations. In the presence 312 of ATP and ssDNA, we observed ring-like densities for ComM that upon 2-D averaging revealed 313 that this protein forms a hexameric ring (Fig. 3B and 3C). Furthermore, we generated a 3-D 314 reconstruction from the negative stain TEM images of ComM in the presence of ATP and ssDNA 315 (~13.8 Å resolution, see **Supplementary Methods**), which revealed that ComM forms a three-316 tiered barrel-like structure with a large opening on both ends (Fig. 3D). The pore on the bottom 317 of this barrel is ~18Å, which can accommodate ssDNA but not dsDNA, while the 26 Å pore is 318 able to accommodate dsDNA. The size of these pores, however, may be an underestimate due 319 to the stain used during EM and/or ssDNA that may be bound and averaged into the 3-D

- 320 construction. Regardless, these data suggest that ComM forms a hexameric ring, consistent
- 321 with structures adopted by many AAA+ helicases (26). ComM also oligomerized in the presence
- 322 of ADP and the non-hydrolysable ATP analog AMP-PNP, indicating that ATP is not a strict
- 323 requirement for hexamer formation (Fig. S4).
- 324

325 <u>ComM binds ssDNA and dsDNA in the presence of ATP</u>

- 326 Because ATP was required for oligomerization in the presence of ssDNA, we hypothesized that 327 it would also be required for DNA binding. To test this, we mutated the conserved lysine in the 328 Walker A motif of ComM, which in other AAA+ ATPases, promotes ATP binding (26). In vivo, we 329 find that this mutation abrogates ComM function (Fig. S5A), while protein stability is 330 maintained (Fig. S2). To determine if ATP binding was required for ComM to bind ssDNA, we performed electrophoretic mobility shift assays (EMSAs) using purified ComM and ComM^{K224A} in 331 332 vitro. We find that WT ComM binds both ssDNA and dsDNA in an ATP-dependent manner (Fig. **S5B-D**). Consistent with this, ComM^{K224A} displays greatly reduced ssDNA binding even in the 333 334 presence of ATP (Fig. S5B). Also, using non-labeled competitor DNA of differing lengths in 335 EMSAs, we found that ComM preferentially bound ssDNA >60bp in length. Taken together, 336 these results indicate that ATP is important for ComM to bind DNA and function during natural
- 337 transformation.
- 338

339 *ComM has helicase activity* in vitro

340 Thus far, our data suggest that ComM may play a role in branch migration *in vivo*. Some branch 341 migration factors (e.g., RuvAB and RecG (32,33)) display helicase activity. So next, we tested the 342 helicase activity of purified ComM in vitro. We observed enzymatic unwinding of a forked DNA 343 substrate with increasing concentrations of ComM. Assuming that ComM hexamers are the 344 active oligomeric form, the helicase activity had an apparent K_M of 50.8 nM (Fig. 4A). As expected, the purified ATP binding mutant ComM^{K224A} did not display helicase activity in this 345 346 assay compared to WT ComM and Pif1, a previously characterized helicase (34) that served as a 347 positive control in these assays (**Fig. 4B**). Furthermore, the non-hydrolysable ATP analog ATPyS

inhibited helicase activity, which is consistent with ATP hydrolysis being required for ComMfunction (Fig. 4B).

350

351 Most helicases exhibit a preferred directionality (either 5' to 3' or 3' to 5'). A forked DNA 352 substrate, however, does not distinguish between these activities. To determine whether 353 ComM had a preferred directionality, we tested helicase activity on forked substrates where 354 one of the two tails is inverted (by a 3'-3' linkage) relative to the remainder of the oligo (35). 355 Thus, directional substrates either have two 5' ends (to assess 5' to 3' directionality) or two 3' 356 ends (to assess 3' to 5' directionality). As controls in these assays, we used the unidirectional 357 helicases Pif1 (a 5' to 3' helicase (34)) and SftH (a 3' to 5' helicase (36)). While Pif1 and SftH 358 exhibited unidirectional helicase activity as expected, ComM unwound all of the substrates 359 tested (Fig. 4C). Taken together, these data suggest that ComM is an ATP-dependent 360 bidirectional helicase. Bidirectional activity of a single motor protein like ComM, while 361 uncommon, is not unprecedented (37-39).

362

363 <u>ComM exhibits branch migration in vitro</u>

364 While our data above clearly indicate that ComM is a hexameric helicase, not all helicases can 365 promote branch migration. To more formally test whether ComM can promote branch 366 migration, we performed *in vitro* branch migration assays using short 3-stranded substrates 367 that more closely resemble the junctions of the D-loop that form during natural transformation 368 (Fig. S6A and S6B)(40). These substrates contained a small region of heterology (indicated by 369 the grey box), which prevents spontaneous branch migration as previously described (40). 370 Using these substrates, we observed resolution of both the 5' to 3' and 3' to 5' substrates (Fig. 371 **S6C** and **D**), which is consistent with the bidirectional helicase activity of ComM. This activity 372 was inhibited when ComM was incubated with AMP-PNP, a nonhydrolyzable analog of ATP (Fig. **S6C** and **D**). And no activity was observed with ComM^{K224A} even if incubated with ATP (Fig. **S6C** 373 374 and **D**). Thus, the branch migration activity observed is ATPase-dependent. It is formally 375 possible that resolution of these short 3-stranded products was the result of only helicase 376 activity (via the sequential removal of the unlabeled strand followed by the labeled strand or by

377 unwinding of just the labeled strand). To address this, we also performed helicase assays using forked substrates that are derived from the oligos used to make the three-stranded branch 378 379 migration substrates. This analysis indicated that ComM helicase activity on the forked 380 substrates was markedly less efficient than its activity on the 3-stranded substrates (compare 381 Fig. S6C-D to S6E-F), which suggests that the activity observed on the 3-stranded substrates is 382 the result of *bong fide* branch migration and not simply helicase activity. The forked substrates 383 used in this assay have 60 bp of annealed sequence (Fig. S6E-F), while the substrates previously 384 used to test helicase activity only had 20 bp of annealed sequence (Fig. 4A). Thus, reduced 385 activity on the forked DNA substrates in this assay is likely attributed to poor processivity for 386 ComM helicase activity.

387

388 While the short 3-stranded substrates used above suggest that ComM possesses branch 389 migration activity, we further tested 3-stranded branch migration using long DNA substrates. 390 Long 3-stranded substrates were generated by RecA-mediated strand exchange between 391 circular single-stranded (CSS) and linear double-stranded (LDS) ΦX174 DNA (23,41). Strand 392 exchange reactions were allowed to proceed until the majority of the LDS was reacted with the 393 CSS to form intermediates (INT = joint molecules that have not completed strand exchange) or 394 nicked product (NP = the product formed upon complete strand exchange). Reactions were 395 then deproteinated, and the resulting DNA was used to assess resolution of the long 3-stranded 396 INT via branch migration (23,41). We found that ComM could efficiently drive the reaction in 397 both directions, resolving the INT structures into both NP and LDS (Fig. 5), while reactions where ComM^{K224A} was added showed no activity. These results are consistent with ComM 398 399 promoting branch migration of these long DNA substrates. Together, these results indicate that 400 ComM exhibits branch migration on 3-stranded junctions in vitro on substrates that are likely 401 similar to the structures formed during natural transformation *in vivo*.

402

403 *ComM is broadly conserved*

404 Next, we assessed how broadly conserved ComM was among bacterial species. Homologs of

405 ComM are found almost ubiquitously among Gram-negative species, including all known Gram-

negative naturally competent microbes (Fig. 6 and Fig. S7). Among this group, only select
species lacked a ComM homolog, suggesting that loss of ComM occurred relatively recently
(Fig. 6 and Fig. S7). By contrast, we see a pervasive lack of ComM homologs among species
within the Bacilli and Mollicutes, (Fig. 6 and Fig. S7), suggesting that ComM may have been lost
in a common ancestor for these two Classes. Interestingly, all known naturally competent
Gram-positive species fall within the Bacilli group, indicating that branch migration during
natural transformation must occur via a ComM-independent mechanism in these microbes.

413

414 *ComM promotes branch migration in* Acinetobacter baylyi

Because ComM is broadly conserved among Gram-negative naturally competent microbes,
next, we tested its role during natural transformation in the model competent species *A. baylyi*

417 ADP1. As in *V. cholerae*, a *comM* mutant of *A. baylyi* displayed greatly reduced rates of natural

418 transformation when using linear tDNA (**Fig. 7A, left**). Also, this mutant displayed reduced

419 comigration of linked genetic markers, consistent with *comM* playing a role in branch migration

420 (Fig. 7A, right). Additionally, we observed that ComM is required for integration of tDNA

421 containing larger regions of heterologous sequence (Fig. 7B). These data are consistent with

422 what was observed in *V. cholerae* (Fig. 2) and suggests that ComM is a conserved branch

423 migration factor important for natural transformation in diverse Gram-negative bacterial424 species.

425

426 *RadA is not required for natural transformation in* V. cholerae.

427 While ComM is broadly conserved, it is absent in all of the known Gram-positive naturally 428 competent species (Fig. 6). In the Gram-positive Streptococcus pneumoniae, mutants of radA 429 (also known as *sms*) display reduced rates of natural transformation but are not affected at the 430 level of tDNA uptake, similar to what is observed for V. cholerae comM mutants in this study 431 (42) and in *H. influenzae* (9). Similar results are also seen in *radA* mutants in *Bacillus subtilis* 432 (43). E. coli RadA has recently been shown to promote branch migration during RecA-mediated 433 strand exchange (23). Also, it was very recently demonstrated that RadA is a hexameric helicase 434 that promotes bidirectional D-loop extension during natural transformation in S. pneumoniae

(38). Interestingly, *radA* is broadly conserved and both *V. cholerae* and *A. baylyi* contain *radA*homologs (VC2343 and ACIAD2664, respectively). RadA, however, is not required for natural
transformation in *V. cholerae* (Fig. S8). Also, preliminary Tn-seq data from our lab indicates that
RadA is not important for natural transformation in *A. baylyi*. Thus, it is tempting to speculate
that in Gram-positive competent species, RadA carries out the same function that ComM plays
during natural transformation in Gram-negative species.

441

442 <u>ComM is not required for DNA repair</u>

443 ComM homologs are also found among diverse non-competent species (Fig. 6). Moreover, 444 many bacterial helicases are implicated in promoting branch migration during other types of 445 homologous recombination, including during DNA repair. So next, we wanted to determine if 446 ComM also plays a role in DNA repair independent of its role during natural transformation. 447 ComM is poorly expressed in the absence of competence induction in V. cholerae (Fig. S2). 448 Thus, to test the role of ComM in DNA damage, we tested survival of P_{tac} -tfoX and P_{tac} -tfoX 449 Δ*comM* strains under competence inducing conditions (i.e. ectopic expression of TfoX). A WT 450 strain and a recA mutant were also included in these assays as controls. DNA damage was 451 tested using methyl methanesulfonate (MMS - methylates DNA / stalls replication forks) and 452 mitomycin C (MMC – alkylates DNA / generates interstrand DNA crosslinks)(44,45). As 453 expected, the recA mutant was more sensitive to these treatments compared to the WT, 454 consistent with a critical role for homologous recombination in DNA repair (Fig. S9) (46,47). The 455 P_{tac} -tfoX $\Delta comM$ mutant, however, was as resistant to these DNA damaging agents as the P_{tac} -456 tfoX strain, indicating that this branch migration factor either does not play a role during DNA 457 repair or that the activity of this protein is redundant with other branch migration factors in the 458 context of repair (Fig. S9). Furthermore, induction of competence (via ectopic expression of 459 TfoX) showed little to no difference in DNA damage repair compared to the WT, indicating that 460 natural competence plays a limited role in DNA repair in V. cholerae under the condition tested 461 (Fig. S9).

462

463 **DISCUSSION**

464 Natural transformation is an important mechanism of horizontal gene transfer in bacterial 465 species. It is dependent upon activation of bacterial competence or the ability to bind and take 466 up exogenous DNA. Altogether, our in vivo and in vitro data elucidate a role for ComM as a 467 helicase / branch migration factor that promotes the integration of tDNA during natural 468 transformation. In our model, integration is initiated by RecA-mediated strand invasion and 469 formation of a D-loop that generates a three-stranded intermediate structure (Fig. 8). Through 470 its bidirectional helicase and/or branch migration activity, ComM then likely promotes 471 expansion of the D-loop, which enhances the integration of tDNA into the genome (Fig. 8). 472 Following branch migration, the junctions are resolved to mediate stable integration of tDNA by 473 an unresolved mechanism.

474

475 Our data suggest that ComM is important for the incorporation of heterologous sequences. The 476 main drivers for evolution and maintenance of natural transformation in bacterial species are 477 heavily debated. One model suggests that this process is largely for enhancing genetic diversity, 478 while another hypothesis is that natural transformation evolved as a mechanism for acquisition 479 of DNA as a nutrient (48,49). These processes are not mutually exclusive. Because ComM 480 affects only the integration of heterologous tDNA (and not its uptake), however, the activity of 481 this protein supports a role for natural transformation in adaptation and evolution through the 482 acquisition of novel genetic material. Other competence genes that are involved specifically in 483 homologous recombination (e.g., *dprA*) also support this hypothesis.

484

485 Our in vitro data suggest that ComM forms a hexameric ring structure in its active state similar 486 to that of eukaryotic MCM2-7 and bacterial DnaB (31,50). Our 3D reconstruction reveals a 487 three-tiered barrel-like complex with a ~26-Å pore and ~18-Å pore on the top and bottom, 488 respectively. We propose that ComM either forms around, or is loaded onto the displaced 489 single strand of the native genomic DNA and acts as a wedge to dissociate these strands. 490 Alternatively, dsDNA may enter the 26-Å pore, be unwound in the central channel of the 491 complex with single strands bring extruded through the side channels that are evident between 492 the tiers of the barrel structure. Such side channels are not uncommon among hexameric AAA+

helicases (*e.g.*, SV40 T-antigen (51,52) and the archaeal MCM (28,53)) and have been
hypothesized to be exit channels for extruded ssDNA. The pore sizes observed in our
reconstruction appear to be consistent with those found in other ring helicases, however, it has
been shown that the size of the opening and channel can change depending on the nucleotide
bound (ATP vs. ADP) (51,54). Future work will focus on characterizing the ComM structure
bound to ADP and AMP-PNP, which may help inform the structural changes associated with the
catalytic cycle of this hexameric helicase.

500

501 Our phylogenetic analysis indicates that ComM is broadly conserved in bacterial species, and is 502 largely, only excluded from the Bacilli and Mollicutes. There is also, however, evidence for 503 isolated examples for ComM loss in species that fall outside of these two groups, which 504 suggests that these have occurred relatively recently. Some of these represent obligate 505 intracellular pathogens or endosymbiots, which commonly contain highly reduced genomes 506 (e.g. Buchnera and Chlaymdia) (55,56). Another example of recent ComM loss is among 507 *Prochlorococcus* species, which are related to other naturally competent cyanobacteria (e.g. 508 Synechococcus and Synechocystis spp.). Interestingly, we found that Prochlorococcus lacked 509 many of the genes required for natural transformation (e.g. dprA, comEA, comEC, etc.) (57), 510 indicating that competence may have been lost in this lineage of cyanobacteria. Another 511 notable example of ComM loss is in Acinetobacter baumanii, which is an opportunistic 512 pathogen that is closely related to A. baylyi. Many strains of A. baumanii, including the one 513 analyzed here (ATCC 17978), contain an AbaR-type genomic island integrated into comM 514 (58,59). Strains that contain this horizontally transferred genomic island display low rates of 515 natural transformation while those that lack it display higher rates of natural transformation 516 (60), indicating that AbaR island-dependent inactivation of ComM may inhibit this mechanism 517 of horizontal gene transfer in A. baumanii. This represents another example in a growing list of 518 horizontally acquired genomic elements that inhibit natural transformation (61-63). Our 519 phylogenetic analysis also indicates that ComM is highly conserved among non-competent 520 bacterial species. This suggests that ComM may have a function outside of natural 521 transformation. Alternatively, it is possible that many of these species are capable of natural

transformation, however, the conditions required for competence induction have not yet been
identified. Indeed, the inducing cue for natural transformation in *V. cholerae* was only
discovered in 2006 (2).

525

526 Our data suggest that branch migration in Gram-positive and Gram-negative species has 527 diverged in their dependence on distinct factors (RadA and ComM, respectively). The tract 528 length of DNA recombined into S. pneumoniae during natural transformation is ~2.5 kb on 529 average (64). In *H. influenza*e, the mean recombination tract length is ~14 kb (65). Thus, 530 compared to RadA-dependent branch migration in Gram-positive species, ComM may facilitate 531 the integration of more tDNA in Gram-negative species. Other proteins that impact DNA 532 integration during natural transformation, however, may confound this overly simplified 533 comparison.

534

535 ComM is not essential for natural transformation because transformants are still observed in a 536 comM mutant. This suggests that other proteins may be involved in promoting the integration 537 of tDNA in the absence of this branch migration factor. Also, our data suggest that these 538 alternative branch migration factors are less efficient at incorporating tDNA with sequence 539 heterology compared to ComM in both V. cholerae and A. baylyi. This role could be carried out 540 by another helicase. Candidates include RuvAB, RecG, and PriA, which have all previously been 541 implicated in branch migration (32,33,66). In fact, the primasome helicase PriA is essential for 542 natural transformation in *Neisseria gonorrhoeae*. This may be due to a role in branch migration, 543 however, PriA is also essential for restarting stalled replication forks and priA mutants have a 544 severe growth defect (67). Also, it has been proposed that PriA helicase activity may facilitate 545 the uptake of tDNA through the inner membrane (68). A recG homologue in Streptococcus 546 pneumoniae (mmsA) was shown to have a mild effect on transformation when deleted (69). 547 RecG reverses stalled replication forks and promotes branch migration opposite to the direction 548 of RecA-mediated strand exchange in E. coli (70,71). Also, RecG acts on three-strand 549 intermediates in vitro, which could give credence to involvement in natural transformation 550 where such a structure is formed (41). Because RecG works counter to the direction of RecA-

mediated strand exchange, however, it is currently unclear how RecG may play a role in natural
transformation in the presence of ComM. Another possibility is that RecA, which has inherent
ATP-dependent unidirectional branch migration activity (72,73), could promote branch
migration independent of other canonical branch migration factors. Future work will focus on
identifying genetically interacting partners of ComM to uncover the role of additional factors
required for efficient integration of tDNA during natural transformation.

557

In addition to a AAA+ ATPase domain, ComM also contains two magnesium (Mg) chelatase domains. Mg chelatase domain-containing proteins have only previously been implicated in inserting Mg into protoporphyrin rings in photosynthetic organisms (74). While it is currently unclear how these domains participate in ComM function, to our knowledge, this study is the first to indicate a function for a Mg-chelatase domain-containing protein in a nonphotosynthetic organism.

564

565 In conclusion, the results from this study strongly support that ComM enhances natural

566 transformation by promoting ATP-dependent bidirectional helicase activity and/or branch

567 migration activity, which allows for the efficient integration of heterologous tDNA.

568 Furthermore, our data in V. cholerae and A. baylyi as well as previous work from H. influenzae

indicate that this branch migration factor is a broadly conserved mechanism for integration of

570 tDNA in diverse naturally transformable Gram-negative species.

571

572 **ACKNOWLEDGEMENTS**

This work was supported by US National Institutes of Health Grant Al118863 to A.B.D., an
Indiana University Collaborative Research Grant to M.L.B., an American Cancer Society
Research Scholar grant (RSG-16-180-01-DMC) to M.L.B., startup funds from the Indiana
University College of Arts and Sciences to ABD, and DTK was supported by NIH grant R35
GM122556 to Yves Brun.

578

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783		

784 **FIGURE LEGENDS**

Fig. 1 – *ComM is required for integration of DNA during natural transformation.* (A) Chitin-

- 786 dependent natural transformation assays in the indicated strains using a linear PCR product or a
- 787 replicating plasmid as tDNA. (B) Chitin-independent transformation assays of the indicated
- strains with linear PCR product as tDNA. (C) Complementation of *comM in trans* tested in chitin-
- dependent transformation assays using a linear PCR product as tDNA. All data are shown as the

790 mean \pm SD and are the result of at least three independent biological replicates. ** = p<0.01, 791 *** = p<0.001, NS = not significant

792

793 Fig. 2 – ComM promotes branch migration through heterologous sequences in vivo. (A) Chitin-794 dependent transformation assay performed using tDNA that contained linked genetic markers 795 separated by 820 bp or 245 bp. (B) Chitin-dependent transformation assay performed in Tm^s 796 strains (Tm^R marker inactivated by a nonsense point mutation or 29-bp deletion) using tDNA 797 that would revert the integrated marker to Tm^R. All strains in **A** and **B** contained a mutation in 798 *mutS* to prevent MMR activity. In schematics above bar graphs, X's denote possible crossover 799 points for homologous recombination. All data are shown as the mean ± SD and are the result 800 of at least three independent biological replicates. *** = p < 0.001

801

Fig. 3 – ComM hexamerizes in the presence of ATP and ssDNA. (A) Blue-native PAGE assay of
purified ComM in the indicated conditions. (B) Negative stain EM of purified ComM under the
indicated conditions. Representative ring-like densities observed in the presence of 5 mM ATP
and 5 μM ssDNA are indicated by black arrows. Scale bar = 50 nm. (C) A representative 2-D class
average of the ring-like densities observed by negative stain EM reveals a hexameric complex.
(D) 3-D reconstruction (~13.8 Å resolution) of the ring complex imposing C6 symmetry.

808

809 Fig. 4 – ComM exhibits helicase activity in vitro. (A) A representative forked substrate helicase 810 assay with increasing concentrations of purified ComM. This forked substrate has 20 bp of 811 annealed sequence and 25 bp tails. Concentrations of ComM used (in hexamer) were 0, 10, 25, 812 37.5, 50, 75, 100, 150, 200, and 250 nM. Images were quantified and plotted as indicated. (B) 813 Helicase assay using forked DNA substrate with the indicated purified protein (100 nM Pif1 and 250 nM ComM / ComM^{K224A} hexamer) in the presence of 5 mM ATP (Columns 1, 2, and 3) or 814 815 ATPvS (Column 4). (C) Helicase assays using forked DNA substrates that accommodate enzymes 816 of either directionality or that can only be unwound by 5' to 3' or 3' to 5' activity. Directional 817 substrates contained one ssDNA tail that is inverted relative to the remainder of the strand 818 (inverted portion indicated in gray on the schematic above bars), thus, preventing helicase

activity in one direction. Substrates were incubated with 100 nM purified ComM (hexamer), 10 nM Pif1, or 50 nM SftH. All data are shown as the mean \pm SD and are the result of at least three independent replicates. *** = p<0.001.

822

823 Fig 5 – ComM exhibits 3-stranded branch migration activity on long DNA substrates in vitro. (A) 824 Schematic for RecA-mediated strand exchange between linear double stranded PhiX174 (LDS) 825 and circular single-stranded PhiX174 (CSS), which results in the formation of intermediates (INT) 826 that can be resolved to nicked product (NP) if strand exchange commences to completion. 827 Strand exchange reactions were deproteinated prior to complete strand exchange, and the 828 resulting DNA was used to assess branch migration-dependent resolution of intermediate 829 structures (INT). (B) Representative gel where deproteinated intermediates were incubated 830 with the proteins indicated. (C) Three independent replicates of the assay described in B were 831 quantified, and the relative abundance of the INT, NP, and LDS are shown as the mean ± SD. 832

Fig. 6 – *ComM is broadly conserved*. Estimated maximum likelihood phylogeny of diverse
species based on a concatenated alignment of 36 conserved proteins identified from whole
genome sequences. Species with an identified ComM homolog are highlighted in gray.
Competent species are designated by a star next to the species name. Major taxa are labeled
along their nodes. Pro: Proteobacteria (Greek letters indicate subdivisions); Bac: Bacilli; Mol:
Mollicutes; Cya: Cyanobacteria; Arc: Archaea. Scale bar indicates distance.

839

Fig. 7 – *ComM promotes branch migration in* Acinetobacter baylyi. (A) Transformation assay of *A. baylyi* using a linear tDNA product with linked genetic markers. (B) Transformation assay performed in Spec^S strains with tDNA that would revert the strain to Spec^R. Integration of the marker would either repair a point mutation or delete 180 bp of genomic sequence. All strains in A and B contain *mutS* mutations to prevent MMR activity. In schematics above bar graphs, X's denote possible crossover points for homologous recombination. All data are shown as the mean ± SD and are the result of at least three independent biological replicates. *** = p<0.001.

- 848 **Fig. 8** Proposed model for the role of ComM during natural transformation. ComM is shown as
- 849 a hexameric ring that promotes integration of tDNA via its bidirectional helicase and/or branch
- 850 migration activity. This can support integration of tDNA with a heterologous region, which is
- indicated by a gray box.

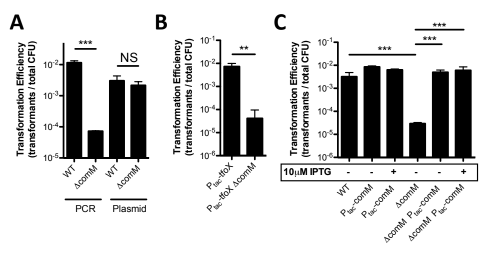


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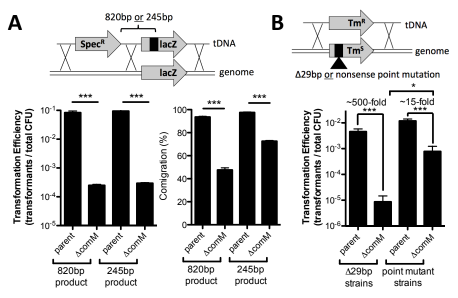


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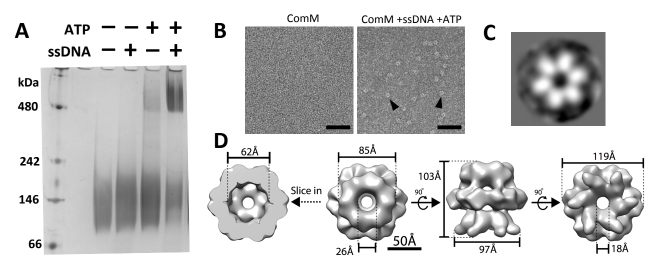


Fig. 3 – *ComM hexamerizes in the presence of ATP and ssDNA.* (**A**) Blue-native PAGE assay of purified ComM in the indicated conditions. (**B**) Negative stain EM of purified ComM under the indicated conditions. Representative ring-like densities observed in the presence of 5 mM ATP and 5 μ M ssDNA are indicated by black arrows. Scale bar = 50 nm. (**C**) A representative 2-D class average of the ring-like densities observed by negative stain EM reveals a hexameric complex. (**D**) 3-D reconstruction (~13.8 Å resolution) of the ring complex imposing C6 symmetry.

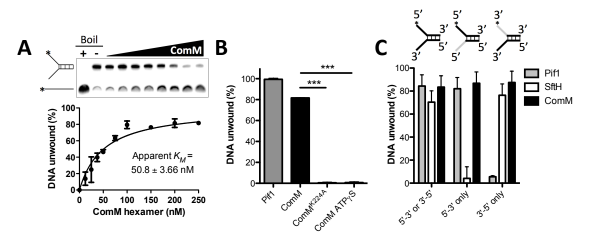


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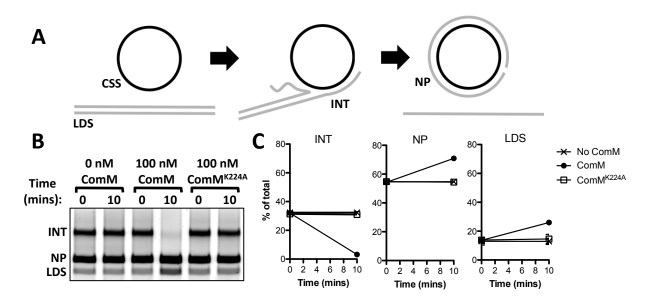


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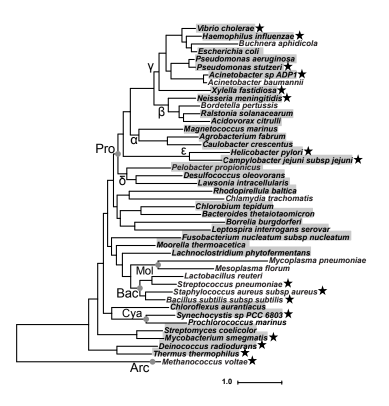


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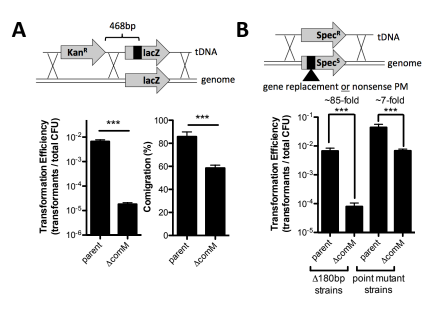


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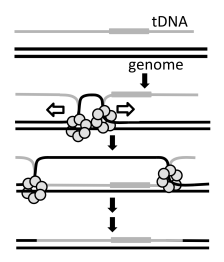


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SUPPLEMENTARY FIGURES

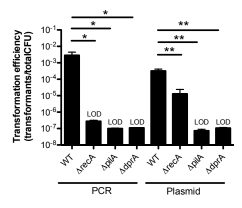


Fig. S1 – *Uptake of plasmid DNA is independent of recombination*. Chitin-dependent transformation assay with the indicated strains using either linear PCR product or plasmid as tDNA. All data are shown as the mean \pm SD and are the result of at least three independent biological replicates. * = p<0.05, ** = p<0.01, and LOD = limit of detection.

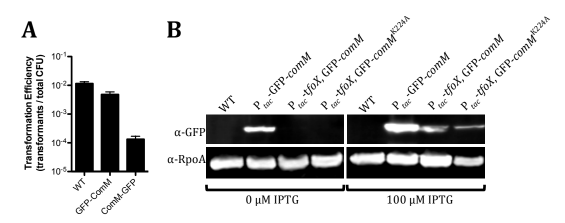


Fig. S2 – *N*-terminal ComM fusions are functional. (**A**) Chitin-dependent transformation assay with the indicated strains using linear PCR product as the tDNA. (**B**) Representative western blot to detect GFP-ComM and RpoA (loading control) in the indicated strains grown in the presence or absence of 100 μ M IPTG. Blot indicates that GFP-*comM* and GFP-*comM*^{K224A} at the native locus are induced when TfoX is ectopically expressed to induce competence. Also, this blot indicates that the P_{tac}-GFP-*comM* construct is leaky and expressed in the absence of inducer. Data in **A** are shown as the mean ± SD and are the result of at least three independent biological replicates.

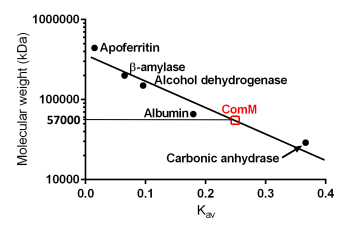


Fig. S3 – *ComM* is monomeric in soluble form. Purified StrepII-ComM was analyzed by gel filtration and compared to a set of protein standards to determine size.

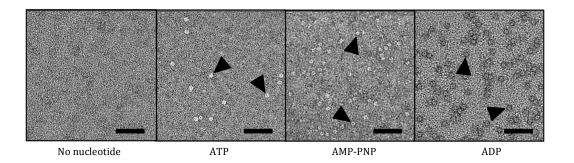


Fig. S4 – *ComM oligomerizes in the presence of ADP and AMP-PNP.* Negative stain EM of purified ComM incubated with 5 μ M ssDNA and 5mM of the indicated nucleotide or nucleotide analog. Representative ring-like densities in these samples are indicated by black arrows. Scale bar = 50 nm.

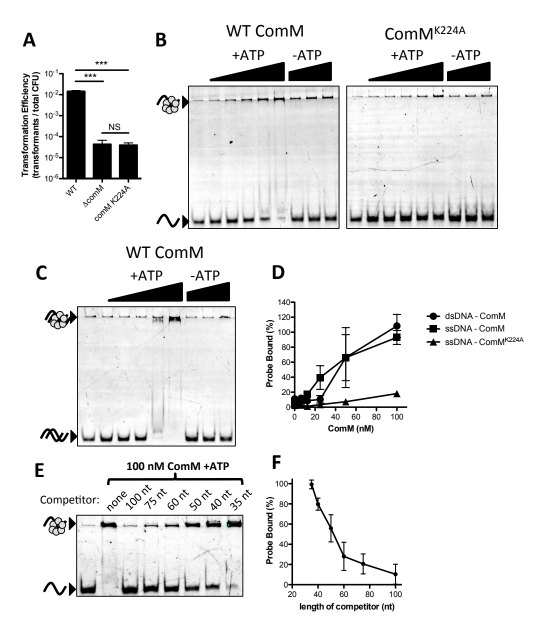


Fig. S5 – *ComM binds ssDNA in the presence of ATP.* (**A**) Chitin-dependent transformation assay of the indicated strains using a linear PCR product as tDNA. Data are shown as the mean ± SD and are the result of at least three independent biological replicates. *** = *p*<0.001, NS = not significant. (**B**) EMSA with purified ComM and ComM^{K224A} and a ssDNA probe. Protein concentrations (of the hexamer) used in the presence of ATP (+ATP) were 0, 6.25, 12.5, 25, 50, 100 nM, and in the absence of ATP (-ATP) were 25, 50, and 100 nM. Bound probe is retained in the well due to the large size of the DNA-bound oligomeric complex. (**C**) Representative EMSA with purified ComM and dsDNA probe. All reactions were performed in the presence of ATP and the protein concentrations (of hexamer) used were the same as in **B**. (**D**) Replicate EMSAs from **B** and **C** were quantified and plotted as indicated. (**E**) A representative EMSA where binding was competed with cold competitor DNA of the indicated length. The labeled probe was a 100 nt poly-dT oligo. The cold competitor was of the length indicated (also poly-dT) and was added at 100X molar excess to the labeled probe. (**F**) Replicate EMSAs from **E** were quantified and plotted as indicated. Data in **B**, **C**, and **E** are representative of at least three independent experiments.

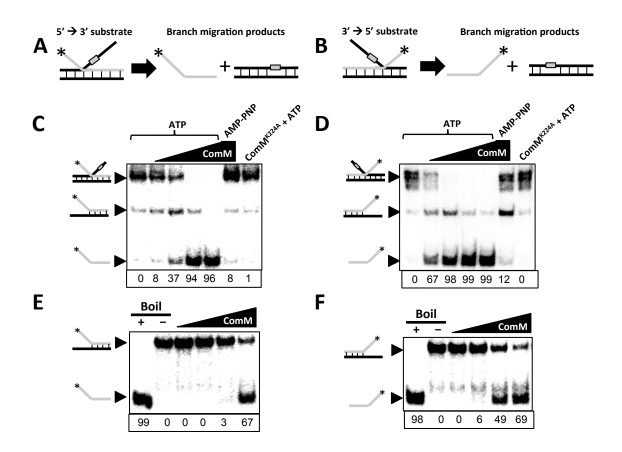


Fig. S6 – *ComM exhibits branch migration activity on short substrates* in vitro. Schematics of the substrates used to test (**A**) $5' \rightarrow 3'$ and (**B**) $3' \rightarrow 5'$ branch migration activity. The gray box indicates a region on the substrate that is not homologous to the complementary strand. This was introduced to prevent spontaneous branch migration. The labeled strand has 60 bp of annealed sequence and a 30 nt tail (**C** and **D**) Representative branch migration assays using the substrates described in **A** and **B**, respectively. Substrates were incubated with 0, 25, 50, 100, or 200 nM WT ComM hexamer or 200 nM ComM^{K224A} as indicated. Reactions were incubated with ATP or AMP-PNP as indicated. The % of final branch migration product generated at each concentration of ComM is indicated below each lane. (**E** and **F**) Representative helicase assays using forked substrates derived from the same oligos used to generate the three-stranded branch described in **A** and **B**, respectively. Substrates were incubated with 0, 25, 50, 100, or 200 nM to generate the three-stranded branch described in **A** and **B**, respectively. Substrates were incubated with 0, 25, 50, 100, or 200 nM to generate the three-stranded branch described in **A** and **B**, respectively. Substrates were incubated with 0, 25, 50, 100, or 200 nM wt ComM hexamer in the presence of ATP. The % of unwound product generated at each concentration of ComM is indicated below each lane. All data are representative of at least three independent experiments.



Fig. S7 – *ComM is broadly conserved.* (**A**) Phylogenetic trees of species based on a concatenated alignment of 36 conserved protein sequences. Green text indicates species with an identifiable ComM homolog. (**B**) Phylogenetic tree of ComM alleles.

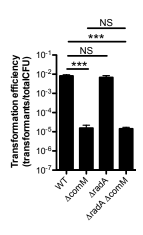


Fig. S8 – *RadA is not required for natural transformation in* V. cholerae. All strains contain P_{tac} -*tfoX* mutations and were transformed via chitin-independent transformation assays using a linear PCR product as the tDNA. All data are shown as the mean ± SD and the result of 6 independent biological replicates. *** = p<0.001, NS = not significant.

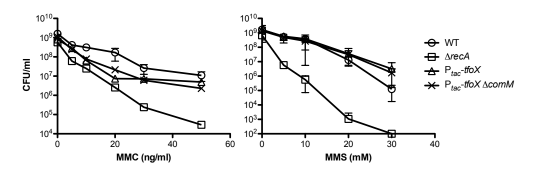


Fig. S9 – *ComM is not required for DNA repair.* Strains were treated with increasing doses of the DNA damaging agent indicated on the X-axis and then plated for viability. All data are shown as the mean ± SD and are the result of at least three independent biological replicates.

SUPPLEMENTARY TABLES

Table S1 – Strains used in this stud	Table S1	- Strains	used in	this stud
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Strain Name in Manuscript	Genotype and Antibiotic Resistances	Description	Reference / strain#
WT	Sm ^R	Wildtype <i>V. cholerae</i> O1 El Tor strain used throughout this study	(1) / (SAD030)
	Strains Used for T	ransformation Assays	
ΔcomM	ΔVC0032::Spec ^R	Replacement of VC0032 with a spectinomycin resistance cassette	This Study (SAD083)
P _{tac} - <i>tfoX</i>	VC1153 OE Kan ^R	OE KanR = a fragment of the Tn10 transposon from pDL1093, including the rrnB antiterminator, Ptac, LacI, and KanR. This fragment, and its use in overexpression of VC1153 (tfoX) is described in	(2) / (SAD061)
Δ <i>comM</i> Ptac- <i>tfoX</i>	VC0032::Spec ^R , VC1153 OE Kan ^R	Replacement of VC0032 with Spec ^R cassette in a VC1153 OE Kan ^R parent strain	This Study (SAD066)
P _{tac-} comM	P _{tac} - <i>comM</i> at <i>lacZ</i> locus Spec ^R	VC0032 fused with an IPTG inducible promoter at the lacZ locus	This Study (SAD1065)
ΔrecA	Δ <i>recA::</i> Spec ^R	Replacement of VC0543 with Spec ^R cassette	This Study (SAD081)
ΔpilA	Δ <i>pilA</i> ::Spec ^R	Replacement of VC2423 with Spec ^R cassette	This Study (SAD780)
∆dprA	∆ <i>dprA::</i> Spec ^R	Replacement of VC0048 with $Spec^{R}$	This Study (SAD079)
P _{tac-} comM, ΔcomM	P _{tac} - <i>comM</i> at <i>lacZ</i> locus Spec ^R , ΔVC0032::Amp ^R	VC0032 fused with an IPTG inducible promoter at the lacZ locus in a $\Delta comM$ parent strain	This Study (SAD1066)
Δ29bp (Tm ^s)	ΔVC1807::TmR*Δ29bp, ΔmutS MuGENT edit, LPQEN KanR	ΔVC1807::TmR*Δ29bp, ΔmutS MuGENT edit, LPQEN KanR; NOT TmR resistant	This Study (TND0226/ SAD1321)
Δ29bp (Tm ^s) ΔcomM	ΔcomM::CarbR, (ΔVC1807::TmR*Δ29bp, ΔmutS MuGENT edit, LPQEN KanR)	VC0032 deletion in a Tm ^s parent strain	This Study (TND0229/ SAD1322)
Point mutant (Tm ^s)	ΔmutS MuGENT edit, LPQEN::SpecR, ΔVC1807::TmR*TI	Δ mutS MuGENT edit, LPQEN::SpecR, Δ VC1807::TmR*TI; Δ VC1807::TmR*TI is the TmR cassette at VC1807, except it has a transition point mutation that introduces a premature stop codon.	This Study (TND0220/ SAD1323)
Point mutant (Tm ^s), ΔcomM	ΔmutS MUGENT edit, ΔcomM::CarbR, LPQEN::KanR, ΔVC1807::TmR*TI	ΔmutS MUGENT edit, ΔcomM::CarbR,LPQEN::KanR, ΔVC1807::TmR*TI;ΔVC1807::TmR*TI is the TmRcassette at VC1807, except it has atransition point mutation thatintroduces a premature stop codon.	This Study (TND0221/ SAD1324)
<i>comM</i> ^{K224A}	LPQEN::Kan ^R , <i>comM</i> ^{K224A}	K to A residue substitution disrupts ATP binding. FRT Kan cassette following the LPQEN amino acid sequence in <i>lacZ</i> (VC2338) used for	This Study (SAD1026)

		selection during co-transformation.	
gfp-comM	gfp-comM	N-terminal GFP-comM at native locus	This Study (SAD924)
comM-gfp	comM-gfp	C-terminal comM-GFP at native locus	This Study (SAD925)
parent (ADP1)	ΔACIAD1551::P _{tac} -lacZ, ΔmutS::SpecR	lacZ introduced into a defunct transposase (neutral gene = ACIAD1551), mutS deleted and replaced with Spec ^R (ACIAD1500)	This Study (TND0137/ SAD1325)
∆comM (ADP1)	ΔACIAD1551::P _{tac} -lacZ, ΔmutS::SpecR, ΔcomM	lacZ introduced into a defunct transposase (neutral gene = ACIAD1551), mutS deleted and replaced with Spec ^R (ACIAD1500), comM in-frame mutation (ACIAD0242)	This Study (TND0149/ SAD1326)
Δ180 (ADP1)	ΔmutS::Kan ^R	MutS deleted and replaced with a Kan ^R cassette	This Study (SAD742)
Δ180 ΔcomM (ADP1)	ΔmutS::Kan ^R , ΔcomM	MutS deleted and replaced with a Kan ^R cassette, Δ comM in-frame	This Study (TND0144/ SAD1327)
Point mutant (ADP1)	ACIAD1551:: <i>Spec^R</i> (point mutant), ΔmutS::Kan ^R	SpecR cassette in ACIAD1551 is inactivated with a point mutation. MutS deleted and replaced with Kan ^R	This Study (TND0150/ SAD1328)
Point mutant, ΔcomM (ADP1)	ACIAD1551:: <i>Spec^R</i> (point mutant), ΔmutS::Kan ^R , ΔcomM	SpecR cassette in ACIAD1551 is inactivated with a point mutation. mutS was deleted and replaced with Kan ^R and comM was deleted in-frame	This Study (TND0164/ SAD1329)
pBAD18 Kan	pBAD18 Kan	TG1 E. coli strain used to purifiy a replicating plasmid with a Kan ^R cassette	This Study (SAD233)
P _{tac} -tfoX, ΔradA	VC1153 OE Kan ^R , <i>radA::Spec^R</i>	<i>radA</i> (VC2343) deleted and replaced with Spec ^R in a TfoX overexpressing background.	This Study (TMN0135/ SAD1813)
P _{tac} -tfoX, ΔradA, ΔcomM	VC1153 OE Kan ^R , radA::Spec ^R , comM::Carb ^R	<i>comM</i> (VC0032) deleted and replaced with Carb ^R in a TfoX overexpressing, <i>radA</i> deletion background.	This Study (TMN0136/ SAD1814)
	Strains used in	n ComM Induction	
P _{tac} -gfp	Ptac-GFPmut3 <i>Spec</i> ^{<i>R</i>} at the <i>lacZ</i> locus	Replaced lacZ gene in SAD030 with a fragment from the transposon vector pDL1098 which encodes LacI, SpecR and has a Ptac promoter. Cloned GFPmut3 downstream of Ptac promoter	This Study (SAD559)
P _{tac} -gfp-comM	Ptac N-terminal <i>gfp-comM</i> at <i>lacZ Spec^R, ΔcomM Kan</i>	VC0032 deletion strain containing an N-terminally GFP tagged comM at the lacZ locus under the control of Ptac	This Study (SAD921)
gfp-comM	N-terminal GFP-comM at the native locus	N-terminally GFP tagged comM was cloned into SAD030 at the native locus	This Study (SAD924)
P _{tac} -tfoX, gfp-comM	VC1153 OE Kan ^R , N- terminally GFP-comM at the native locus	Amplified VC1153 OE Kan from SAD061 and cloned fragment into SAD924	This Study (TMN0140 / SAD1320)

$D = tf_0 V$	VC1153 OE <i>Kan^R</i> , N-	Amplified <i>comM</i> ^{K224A} mutation from	This Study
P _{tac} -tfoX, gfp-comM ^{K224A}	terminal <i>gfp-comM</i> ^{K224A} at	SAD1026 and cloned fragment into	(TMN0148 /
gjp-comm ⁿ²² m	the native locus, <i>Spec</i> ^{<i>R</i>}	SAD1320	SAD1545)
Strains Used for Protein Purification			
ComM	ComM cloned into <i>Amp^R</i> StrepII expression vector	ComM N terminally tagged with 4x StrepII	This Study (pMB486 /
			SAD1330)
ComM ^{K224A}	ComM ^{K224A} cloned into Amp ^R StrepII expression vector	ComM ^{K224A} N terminally tagged with 4x StrepII	This Study (pMB488 / SAD1331)
Rosetta 2 (DE3)		E. coli expression strain	(pMB131/ SAD1332)

Table S2- Primers used in this study-

Primer	Primer Sequence (5' to 3')*	Description
Name		_
Mutant co	onstructs	
ABD855	CATGAATCACTTTGGCATGAGG	$\Delta comM$ F1
ABD856	gtcgacggatccccggaatCATTGCTTCCCTTAGTATTTGATC	$\Delta com M R1$
ABD857	gaagcagctccagcctacaTAGTACTCTGACCTGCAGAGTTC	$\Delta comM$ F2
ABD858	AAATTCCAGAAAAACCACGTC	$\Delta com M R2$
BBC749	CCGTGAAGCGAGCATGGTcgcACCCGTCCCCGGAG	<i>comM</i> ^{K224A} R1
BBC750	CTCCGGGGACGGGTgcgACCATGCTCGCTTCACGG	<i>comM</i> ^{K224A} F2
ABD812	AAATGGAGTTTGATCGCATTGGC	Δ <i>recA</i> F1
ABD921	gtcgacggatccccggaatCATTACTCTCTCCGGATAGTCACTC	ΔrecA R1
ABD922	gaagcagctccagcctacaTAATCGGCAGGCTGAATGCAAAG	ΔrecA F2
ABD815	TGATCAGCGTTTGGAATACGTCG	ΔrecA R2
BBC401	ACCAGCAAAGCTAATAAAATCGAG	Δ <i>pilA</i> F1
BBC402	gtcgacggatccccggaatGAGCATATGCCTTGCTACACAAG	$\Delta pilA R1$
BBC403	gaagcagctccagcctacaACTGCAGGTGCAACAATTAACTAA	$\Delta pilA$ F2
BBC404	CGCCATACTAACCCAATACACTC	$\Delta pilA R2$
ABD820	CGCTCTTATCTGCTTGGATAATGG	$\Delta dprA$ F1
ABD998	gtcgacggatccccggaatCATTAACTGGCATCATCAACC	$\Delta dprA R1$
ABD999	gaagcagctccagcctacaTAGCTATGATGATGGATATTTTGATG	$\Delta dprA$ F2
ABD823	TGAAGTACAAGGCCAGTTACTGG	$\Delta dprA R2$
BBC907	AAAGAGCAGTTGTCGCTAGAC	Δ <i>radA</i> F1
BBC908	gtcgacggatccccggaatCAATCCTCGAACTTGCTCTCAC	Δ <i>radA</i> R1
BBC909	gaagcagctccagcctacaTAATGGGTAGTTGGTTTTGAAC	Δ <i>radA</i> F2
BBC910	ATGAAGAAATCTTAGTCCGCAG	Δ <i>radA</i> R2
ABD824	TTTAGCCCCATTGGCGAACTGGG	ΔmutS F1
ABD825	GAGTATCTTTGACGTATTGGATCtcatattatactaCATAATCTTATGTC	ΔmutS R1
	GCTGCTTATC	
ABD826	GATAAGCAGCGACATAAGATTATGtagtataatatgaGATCCAATACGT	ΔmutS F2
	CAAAGATACTC	
ABD360	AGATCTTGCCTGATGACGCTTTACTC	ΔmutS R2
BBC717	AAATAGATTTGGTGACTTTACCTCC	VC1807::Ab ^R F1
ABD340	gtcgacggatccccggaatACGTTTCATTAGTCACCTCTATTGTTAACTTG TTC	VC1807::Ab ^R R1
ABD341	gaagcagctccagcctacaTAGTCGAAAATAAAAAAAAGAGGCTCGCCTC	VC1807::Ab ^R F2
BBC718	CTTTACGCCTGATTGTCTACAC	VC1807::Ab ^R R2

ABD332	GGCTGAACGTGGTTGTCGAAAATGAC	lacZ F1
ABD263	gtcgacggatccccggaatAACTGATCCAATTTTTCAGCGCATATTTTGG	lacZ LPQEN::Ab ^R R1
ABD262	gaagcagctccagcctacaTGCCGCAGGAAAACCGCCCCCTaATC	lacZ LPQEN::Ab ^R F2
ABD256	CCCAAATACGGCAACTTGGCG	lacZ R2
ABD269	gaagcagctccagcctacaAATTGTGTAAACGTTTCCACAATTTAAATAG AGG	Spec ^R upstream of lacZ R1
ABD268	gtcgacggatccccggaatGGTGAGTGGTTCACAGAATCGGTG	Spec ^R upstream of lacZ F2
ABD495	AAAAAAATCTTCAATCGCGAGTATCGGCTaGCGGTAGAGATACACA TCGCGAAAGATGCC	lacZ 820bp linked R1
ABD494	TCTTTCGCGATGTGTATCTCTACCGCtAGCCGATACTCGCGATTGAA GATTTTTTTATCC	lacZ 820bp linked F2
ABD329	GAACATGGGGTGTACGGCAGTGCCATTaAACGATGTGCGGGTTTTG CCAATCTTG	lacZ 245bp linked R1
ABD328	CAAGATTGGCAAAACCCGCACATCGTTtAATGGCACTGCCGTACACC CCATGTTC	lacZ 245bp linked F2
BBC1157	GTAAAACTTGAACGTGTTACGAATTGATTCAAAAGTCTTGCGTC	Tm ^R Δ29bp R1
BBC1158	GACGCAAGACTTTTGAATCAATTCGTAACACGTTCAAGTTTTAC	Tm ^R Δ29bp F2
BBC747	GTCCAACCAACAGCCATTGGTTtTAGGTAATAGCTTTAAACAGGAG C	Tm ^R Point mutation
BBC748	GCTCCTGTTTAAAGCTATTACCTAaAACCAATGGCTGTTGGTTGGA C	Tm ^R Point mutation
BBC498	GGGTAACGCCAGGGTTTTCtCAGTCACGACGTTGTAAAAC	SpecR point mutant R1
BBC499	GTTTTACAACGTCGTGACTGGtAAAACCCTGGCGTTACCC	SpecR point mutant F2
BBC280	TCCACCACTTCCACCtGCGACGTTCTGCGCACTGAGC	GFP-ComM R1
BBC351	GTTCTTCTCCTTTACGCATTACGTTACCTCCTTTTGATCAAAAAGCC TTCAGC	ComM-GFP R1
BBC352	GCaGGTGGAGCAGGTGGAGGACTTGCGATCATTCATAGC	ComM-GFP F2
ABD688	CCACTGTTGCGCAGTTGAATACC	tfoX OE Kan F1
ABD691	ATGATGTCAAACCATGAACCCGG	tfoX OE Kan R2
BBC331	CAATTTCACACAGGATCCCGGGAGGAGGTAACGTAATGGGACTTGC GATCATTC	P _{tac} -comM F
BBC365	tgtaggctggagctgcttcCTAGACGTTCTGCGCACTGAG	P _{tac} -comM R
DOG0140	GTTGCTGCATTTGTTCGATCTG	ΔcomM (ADP1) F1
DOG0141	gtcgacggatccccggaatCATACTATTATTGTTCCATTATGGTGC	ΔcomM (ADP1) R1
DOG0142	gaagcagctccagcctacaTATCGCAGTGAACATAGCTAAAA	ΔcomM (ADP1) F2
DOG0143	ATCAGTGGTTGGGAAGGTG	ΔcomM (ADP1) R2
Inserts fo	r cloning	
MB1225	CGGGATCCATGGGACTTGCGATCATTCATAGCCG	comM ORF F1
MB1214	CGATCGATCTCGAGCTAGACGTTCTGCGCACTGAGC	comM ORF R1
MB1215	TATTGTTTCTCGGCCCTCCGGGGACGGGTGCGACCATGCTCGCTTCACGG CTGTGCGATT	<i>comM^{K224A}</i> F1
MB1216	AATCGCACAGCCGTGAAGCGAGCATGGTCGCACCCGTCCCCGGAGGGC CGAGAAACAATA	<i>comM</i> ^{K224A} R1
Oligomer	ization assays/Negative stain EM	1
ABD363	CGTTAAATGAAATTAATACGACTCACTATAGGGAGAGAGGGTTTGCTCTGT	ssDNA substrate for oligomerization
EMCA	TTGAGAAGCC	ongomenzation
EMSA pro		
BBC742	ATTCCGGGGATCCGTCGACCTGCAGTTCAGAAGCAGCTCCAGCCTACA	EMSA binding probe (ssDNA, dsDNA)
BBC743	TGTAGGCTGGAGCTGCTTCTGAACTGCAGGTCGACGGATCCCCGGAAT	EMSA binding probe (dsDNA)

MB1040		100 bp poly-dT ssDNA
1121010	<u> </u>	probe
	ттт т	
MB1039	<u> </u>	75 bp poly-dT ssDNA
		probe
MB1038	<u> </u>	60 bp poly-dT ssDNA
		probe
M551	דדד דדד דדד דדד דדד דדד דדד דדד דדד דד	50 bp poly-dT ssDNA
	TT	probe
MB1037	<u> </u>	40 bp poly-dT ssDNA
		probe
MB1140		35 bp poly-dT ssDNA probe
Helicase	substrates	·
MB1167	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	Poly-dT 5' tail
MB1168	GAACGCTATGTGAGTGACACTTTTTTTTTTTTTTTTTTT	Poly-dT 3' tail
MB1510	TTTTTTTTTTTTTTTTTTTTTT/ilnvdT/GTGTCACTCACATAGCGT	Poly-dT inverted 5' tail
	TC	
MB1511	GAACGCTATGTGAGTGACAC/ilnvdT/TTTTTTTTTTTTTTTTTTTTTT	Poly-dT inverted 3' tail
	TTT	
Short thr	ee-stranded branch migration substrates	
BBC1916	TTTGATAAGAGGTCATTTTTGCGGATGGCTTAGAGCTTAATTGCT	5' to 3' bottom strand
	GAATCTGGTGCTGTAGGTCAACATGTTGTAAATATGCAGCTAAAG	
BBC1915	TTTTTTTTTTTTTTTTTTTTTTTTTTCACAGCACCAGATTCAGCA	5' to 3' top strand with
	ATTAAGCTCTAAGCCATCCGCAAAAATGACCTCTTATCAAA	polyDT tail (also for forked helicase
		substrate)
BBC1917	CTTTAGCTGCATATTTACAACATGTTGACCTACAGCAAAGAATTCA	5' to 3' top strand for 3'
	GCAATTAAGCTCTAAGCCATCCGCAAAAATGACCTCTTATCAAA	branch
BBC1913	CTTTAGCTGCATATTTACAACATGTTGACCTACAGCACCAGATTCA	3' to 5' bottom strand
	GCAATTAAGCTCTAAGCCATCCGCAAAAATGACCTCTTATCAAA	
BBC1912	TTTGATAAGAGGTCATTTTTGCGGATGGCTTAGAGCTTAATTGCT	3' to 5' top strand with
	GAATCTGGTGCTGTTTTTTTTTTTTTTTTTTTTTTTTTT	polyDT tail (also used for forked helicase substrate
BBC1914	TTTGATAAGAGGTCATTTTTGCGGATGGCTTAGAGCTTAATTATA	3' to 5' top strand for 5'
	AAATCTGGTGCTGTAGGTCAACATGTTGTAAATATGCAGCTAAAG	branch

*Lowercase letters indicate overlap sequences for SOE PCRs or mutated nucleotides when generating point mutations

SUPPLEMENTARY METHODS

Protein expression and purification

The *comM* open reading frame was PCR-amplified from *V. cholerae* genomic DNA using oligonucleotides MB1225 (CGGGATCCATGGGACTTGCGATCATTCATAGCCG) and MB1214 (CGATCGATCTCGAGCTAGACGTTCTGCGCACTGAGC), digested with *Bam*HI and *Xho*I, and ligated into the same sites in plasmid pMB131 to generate pMB486. This cloning added an N-terminal 4x Strep-tag II to the translated protein. The expression plasmid encoding the ATPase- and helicase-dead *comM-K224A* allele (pMB488) was created site-directed mutagenesis of pMB486 with oligonucleotides MB1215

(TATTGTTTCTCGGCCCTCCGGGGACGGGTGCGACCATGCTCGCTTCACGGCTGTGCGATT) and MB1216 (AATCGCACAGCCGTGAAGCGAGCATGGTCGCACCCGTCCCCGGAGGGCCGAGAAACAATA) and verified by DNA sequencing (ACGT, Inc.). Expression plasmids were transformed into Rosetta 2(DE3) pLysS cells and selected for at 37°C on LB medium supplemented with 100 μ g/mL ampicillin and 34 μ g/mL chloramphenicol. Fresh transformants were used to inoculate one or more 5-mL LB cultures supplemented with antibiotics and incubated at 30°C for ~6 h with aeration. These starter cultures were then diluted 1:100 in ZYP-5052 autoinduction medium containing 1x trace metals mix (3), 100 μ g/mL ampicillin, and 34 μ g/mL chloramphenicol and incubated at 22°C with agitation to OD₆₀₀ >3 (15-18 h). Cells were harvested by centrifugation for 10 min at 5,500 x g and 4°C. Cell pellets were weighed and frozen at -80°C prior to lysis or for long-term storage.

Frozen cell pellets were thawed at room temperature by stirring in 4 mL/g cell pellet resuspension buffer (25 mM Na-HEPES (pH 7.5), 5% (v/v) glycerol, 300 mM NaOAc, 5 mM MgOAc, and 0.05% Tween-20) supplemented with 1x protease inhibitor cocktail (Sigma), and 20 µg/mL DNase I. Cells were lysed by six passed through a Cell Cracker operated at >1000 psi. All subsequent steps were performed at 4°C. The soluble fraction was clarified by centrifugation for 30 min at 33,000 x g followed by filtering the supernatant through a 0.22-µm membrane. This mixture was then applied to a Strep-Tactin Sepharose column (IBA) pre-equilibrated in resuspension buffer using an ÄKTA Pure (GE Healthcare Life Sciences). The column was washed with 20 column volumes (CVs) of resuspension buffer, 10 CVs of resuspension buffer supplemented with 5 mM ATP, and 10 CVs of resuspension buffer. Protein was eluted with 15 CVs of resuspension buffer supplemented with 2.5 mM desthiobiotin (IBA). Column fractions were examined on 8% SDS-PAGE gels run at 20 V/cm and stained with Coomassie Brilliant Blue R-250 (BioRad). Peak fractions were pooled, concentrated with Amicon Ultra-4 30K centrifugal filters, and loaded onto a HiPrep 16/60 Sephacryl S-200 HR column (GE Healthcare Life Sciences) pre-equilibrated in gel filtration buffer (25 mM Na-HEPES (pH 7.5), 5% glycerol, 300 mM NaCl, 5 mM MgCl₂, and 0.05% Tween-20). The protein was eluted with 1.5 CVs gel filtration buffer, and fractions were analyzed by SDS-PAGE as above. Peak fractions were pooled, snap-frozen with liquid nitrogen, and stored at -80°C.

The *Saccharomyces cerevisiae* Pif1 helicase was overexpressed In Rosetta cells from plasmid pMB330 as described for ComM above. Pif1 purification was likewise identical, except the protein from the Strep-Tactin column was polished by Ni-affinity chromatography instead of size exclusion. Briefly, the pooled peak fractions were applied to a His60 Ni Superflow (Clontech) gravity column, washed with 10 CVs resuspension buffer supplemented with 25 mM imidazole (pH 8), and eluted with 4.5 CVs of a step gradient of resuspension buffer containing 100 mM, 250, and 500 mM imidazole (pH 8). Peak fractions were pooled, buffer exchanged into storage buffer (4), snap-frozen

with liquid nitrogen, and stored at -80°C. The *Mycobacterium smegmatis* SftH was purified exactly as previously described (5).

ComM preps were tested for nuclease activity by incubating a labeled ssDNA probe with 100 nM of each protein prep in resuspension buffer for 1 hour at 37°C. Samples were then deproteinated with 1X stop load buffer and separated by native PAGE to assess degradation of the probe. All ComM protein preps used lacked detectable nuclease activity.

Blue native PAGE

Oligomerization of ComM protein was assayed using Blue Native PAGE electrophoresis. 2.5 μ M purified ComM was incubated for 30 min at room temperature in reaction buffer [10 mM Tris-HCl pH7.5, 20mM KCl, 1mM DTT, 10% Glycerol] with 5 mM ATP and/or 5 nM ssDNA (oligo ABD363) where indicated. 1 μ L 20x sample buffer [5% Coomassie G-250, 0.5 M aminocaproic acid pH 7] was added to each reaction and samples were run on 4-16% Native PAGE gels [gel buffer = 0.5 M aminocaproic acid pH 7.0, 0.05 M Bis-Tris pH 7.0]. The cathode buffer was composed of 50 mM Tricine, 15 mM Bis-Tris pH 7.0, 0.02% Coomassie G-250, while the anode buffer was composed of 50 mM Bis-Tris pH 7.0. Samples were run at 150 V for 30 min, then 200 V for 45 min.

Negative stain electron microscopy

The nominal magnification for the images is 60,000x, which is equivalent to 1.8 Å per pixel at the final image. Initial image processing, particle boxing, and CTF determination were performed using EMAN2 (6). A phase-flipped particle dataset was then imported into Relion (7) for 2D classification. Classes showing noisy images were discarded at this stage. As we observed clear six-fold symmetry from the classes, the subsequent processing imposed C6 symmetry. The remaining "good" classes were used to generate the initial models using e2initialmodel.py. The 3D classification was carried out using the initial model that was low-pass filtered to 40 Å to eliminate the possible effect from the model bias. Three 3D classes were obtained; the highest population (46%) of the classes was subjected to further structure refinement in Relion. Approximately 32,958 particles were used to generate the final 3D reconstruction. The reported resolution is ~13.8 Å using gold-standard Fourier shell correlation at a 0.143 cutoff; however, it is an over-estimated value because of the use of negative stain. The structure is rendered using UCSF Chimera (8).

Helicase Assays

Fork substrates for helicase assays were made by 5'-end labelling oligonucleotides (**Table S2**) with T4 polynucleotide kinase (T4 PNK; NEB) and γ [³²P]-ATP. Labelled oligonucleotides were separated from free label using illustra ProbeQuant G-50 micro columns (GE Healthcare) following the manufacturer's instructions. Oligonucleotides were annealed by incubation with an equimolar amount of partially complementary oligonucleotides overnight at 37°C in annealing buffer (20 mM Tris-HCl [pH 8], 4% glycerol, 0.1 mM EDTA, 40 µg/mL BSA, 10 mM DTT, and 10 mM MgOAc) (9).

The DNA fork that allows for 5'-3' and 3'-5' activity was made by annealing oligonucelotides MB1167 with MB1168. The DNA fork that only allows for 5'-3' helicase activity was made by

annealing MB1167 / MB1511. The DNA fork that only allows for 3'-5' helicase activity was made by annealing MB1168 / MB1510. DNA unwinding was assessed by incubating the indicated concentrations of helicase with 5 mM ATP and 0.1 nM radiolabelled fork in resuspension buffer. Reactions were incubated at 37°C for 30 min and stopped with the addition of 1x Stop-Load dye (5% glycerol, 20 mM EDTA, 0.05% SDS, and 0.25% bromophenol blue) supplemented with 400 µg/mL Proteinase K followed by a 10-min incubation at 37°C. Unwound DNA was then separated on 8% 19:1 acrylamide:bis-acrylamide gels in TBE buffer at 10 V/cm. Gels were dried under vacuum and imaged using a Typhoon 9210 Variable Mode Imager. DNA binding was quantified using ImageQuant 5.2 software.

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