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Retraction of DNA-bound type IV competence pili initiates DNA uptake during natural transformation in *Vibrio cholerae*

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Natural transformation is a broadly conserved mechanism of horizontal gene transfer in bacterial species that can shape evolution and foster the spread of antibiotic resistance determinants, promote antigenic variation, and lead to the acquisition of novel virulence factors. Surface appendages called competence pili promote DNA uptake during the first step of natural transformation¹, however, their mechanism of action has remained unclear due to an absence of methods to visualize these structures in live cells. Here, using the model naturally transformable species Vibrio cholerae and a pilus labeling method, we define the mechanism for type IV competence pilus-mediated DNA uptake during natural transformation. First, we show that type IV competence pili bind to extracellular doublestranded DNA via their tip and demonstrate that this binding is critical for DNA uptake. Next, we show that type IV competence pili are dynamic structures and that pilus retraction brings tip-bound DNA to the cell surface. Finally, we show that pilus retraction is spatiotemporally coupled to DNA internalization and that sterically obstructing pilus retraction prevents DNA uptake. Together, these results indicate that type IV competence pili directly bind DNA via their tip and mediate DNA internalization through retraction during this conserved mechanism of horizontal gene transfer

It is widely believed that competence pili bind extracellular DNA and then actively translocate this DNA across the outer membrane via retraction^{2,3}. In the predominant alternate model, competence pili act as gatekeepers to open the outer membrane secretin

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CKE and ABD designed and coordinated the overall study. ABD, CKE, TND, JW, AVC, and NB performed the experiments. YVB, ABD, CKE, TND, and NB analyzed and interpreted data. CKE and ABD wrote the manuscript with help from YVB.

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Supplementary Discussion

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pore via their dynamic activity allowing for passive DNA diffusion across the outer membrane⁴. The main distinction between the proposed models for DNA uptake is the ability of pili to bind DNA. Previously, pilus-DNA binding was visually assessed by electron microscopy, which provides a static view and makes differentiating between coincidental colocalization and de facto binding difficult⁵. To test these models directly, we sought to observe both pili and DNA by fluorescence microscopy. Type IV competence pili are dynamic surface appendages that are polymerized and extended via the action of an ATPase, PilB, and retracted by depolymerization via the action of an antagonistic ATPase, PilT. We first labeled type IV competence pili in a hyperpiliated *pilT* mutant of V. cholerae^{6,7}, using a technique⁸ in which an amino acid residue of the major pilin subunit PilA is replaced by a cysteine for subsequent labeling with a thiol-reactive fluorescent maleimide dye (AF488mal) (Supplementary Fig. 1a-c). Neither the cysteine mutation nor labeling with AF488-mal substantially affected the transformation frequency either in wildtype cells or a constitutively competent strain where *pilT* is intact (Supplementary Fig. 1d-e). The constitutively competent strain maintained competence for an extended period of time (\mathfrak{B} hours) (Supplementary Fig. 2). Because we sought to characterize the function of type IV competence pili downstream of competence induction, we used constitutively competent strains throughout the remainder of the study.

To directly observe pilus-DNA binding, we fluorescently labeled both pili and DNA and tracked them by microscopy in real time. Imaging revealed several cells harboring DNAbound pili, and adherence was evident from the co-localized movement of DNA and pili (Fig. 1a, Supplementary Movie 1). We further confirmed DNA adherence using sheared pili bound to microfluidic channels by observing the correlated movement of DNA with single pilus fibers along with subsequent DNA detachment when flow was maintained (Fig. 1b, Supplementary Fig. 3, Supplementary Movies 2–3). We found that the majority of DNA interactions occurred at the pilus tip (Fig. 1c). DNA binding assays with fluorescently labeled DNA demonstrated that the piliated *pila-cys* and parent strains bound DNA equally, while the non-piliated *pilA* mutant was deficient in DNA binding (Fig. 1d). Furthermore, competition assays with unlabeled competitors indicated that this binding was specifically inhibited by nucleic acids with a strong preference for double-stranded DNA (dsDNA) (Fig. 1e). This result is consistent with dsDNA binding to Streptococcus pneumoniae competence pili⁵ and the preferential uptake of dsDNA in *Neisseria spp.*⁹, although ssDNA can be transformed with similar efficiency in the latter¹⁰. Importantly, we found that DNA binding was dependent on the pilus fiber and could not be attributed to the outer membrane secretin^{11,12}, which still localized properly in the *pilA* mutant (Supplementary Fig. 4a–c). Thus, type IV competence pili specifically bind to extracellular dsDNA.

Because the majority of DNA binding occurs at the pilus tip, we hypothesized that retraction helps bring tip-bound DNA to the cell. Indeed, we found that type IV competence pili are highly active surface structures, with 47% of cells grown under competence inducing conditions making at least one pilus of an average length of ~1.0 µm within a one-minute period (Fig. 2a–b, Supplementary Fig. 5a–c, and Supplementary Movie 4). Extension and retraction rates of type IV competence pili were significantly different, with retraction occurring twice as fast (Supplementary Fig. 5b). Pilus dynamic activity, however, was not altered by the presence of extracellular DNA (Supplementary Fig. 6). In some cases, cells

synthesized pili repeatedly at the same site, suggesting that the same assembly machinery may be reused (Fig. 2a, Supplementary Movie 4). These frequent extension and retraction events allow for repetitive interactions of type IV competence pili with the extracellular environment. Using fluorescently labeled DNA, we also found that retracting type IV competence pili brought tip-bound DNA to the cell surface (Fig. 2c, Supplementary Fig. 7, Supplementary Movies 5–6).

To determine if pilus-DNA interactions were required for DNA uptake, we next sought to disrupt this interaction. Minor pilins can initiate pilus assembly and localize to the pilus tip^{13,14}, which we hypothesized could promote non-specific DNA binding¹⁵ by ionic interactions. Consistent with this, we made targeted mutations of several positively charged residues within the major and minor pilins and identified Arg/Lys mutations in two independent minor pilins (VC0858R165Q and VC0859K148Q) that additively reduced DNA binding (Fig. 2d and Supplementary Fig. 5f-g), but did not substantially alter pilus dynamics (Supplementary Fig. 5a-c, h and Supplementary Movie 7). These minor pilin mutants had significantly reduced rates of natural transformation (Fig. 2e) and DNA internalization (Fig. 2f), consistent with pilus-DNA interactions playing an important role in DNA uptake. Additionally, a distinct minor pilin point mutant (VC0858^{R168Q}) reduced DNA binding and natural transformation to the level of a *pilA* mutant (Supplementary Fig. 5d-e, g). While this mutant produced as many pili as the pilA-cys strain (Supplementary Fig. 5c), these pili displayed statistically significantly altered pilus dynamics (Supplementary Fig. 5a-b, h and Supplementary Movie 8). Despite this, many of the pilus events in the VC0858^{R168Q} mutant were comparable to the rates of extension and retraction in the *pilA-cys* strain (Supplementary Fig. 5a–b), which further supports that DNA binding is critical for DNA uptake.

Next, we assessed the spatiotemporal relationship between pilus dynamic activity and DNA internalization. As a marker for DNA uptake into the periplasm we used a fluorescent fusion to ComEA, a critical competence protein that binds to DNA in the periplasm and results in the formation of readily observed foci upon DNA uptake^{16–18}, a finding that we recapitulate here (Supplementary Fig. 8). We found that an average of 14% of cells formed new ComEA foci in the presence of DNA, and that of those events, $64 \pm 3\%$ were immediately preceded by a retracting pilus at the same location (Fig. 3a–b, Supplementary Movie 9). This value is likely an underestimate, as it does not account for short pili that cannot be resolved by light microscopy, pili occluded by the cell body, or pili that may form outside the plane of focus. For the events where readily observed pilus retraction precedes DNA uptake, ComEA focus formation only occurred after what appears to be the completion of pilus retraction, which is consistent with DNA binding to the tip of type IV competence pili (Fig. 1a–c). The non-piliated *pilA* mutant formed no ComEA foci either in the presence or absence of DNA (Fig. 3b). Thus, these data suggested that complete retraction of DNA bound pili is required for DNA uptake.

To test this model further, we assessed whether retraction was required for DNA uptake. Because deletion of the retraction ATPase *pilT* alters the activity of type IV MSHA pili¹⁹ and may also have pleiotropic effects on gene expression²⁰, we tested this hypothesis more directly by sterically blocking type IV competence pilus retraction. This was accomplished

by co-labeling *pilA-cys* cells with biotin-maleimide and AF488-mal before treating with neutravidin (a high-affinity biotin binding protein), which specifically blocks retraction of extended pili through the outer membrane secretin likely by thickening the pilus fiber (Fig. 3c-d, Supplementary Fig. 9, Supplementary Movies 10-11). Obstruction of pilus retraction resulted in a ~100-fold decrease in transformation frequency, indicating that at least ~99% of all DNA uptake events require retraction of an extended pilus (Fig. 3e). Furthermore, blocking pilus retraction resulted in very few DNA internalization events (Fig. 3b). Importantly, blocking pilus retraction using this approach did not inhibit binding to dsDNA (Supplementary Fig. 10). Recent work demonstrates that ComEA acts as a molecular ratchet to internalize DNA across the outer membrane, but it is currently unclear how DNA uptake is initiated because ComEA is localized to the periplasm^{2,16,17}. Our results support a previously proposed model²¹ whereby ComEA-dependent internalization is initiated by the dynamic activity of DNA-binding type IV competence pili. Consistent with this, we found that DNA was not internalized in a *comEA* mutant despite this strain maintaining normal type IV competence pilus dynamic activity (Supplementary Fig. 11). Together, these results indicate that retraction of DNA-bound type IV competence pili initiates the process of DNA uptake into the periplasm during natural transformation with concomitant uptake facilitated by ComEA-dependent molecular ratcheting.

Previous studies have shown that type IV pili require a PilT retraction ATPase to retract^{22,23}. Contrary to this, we found that *pi/T* mutants were still capable of retraction as observed by correlated changes in cell body fluorescence that occur upon extension and retraction of labeled pili during time-lapse experiments (Fig. 4a, Supplementary Fig. 12, Supplementary Movie 12). The *pilT* mutant, however, exhibited a 10,000-fold reduction in transformation frequency and very little DNA internalization, despite exhibiting pilus retraction (Fig. 3b, Fig. 4a–d). Transformation of the $\Delta pilT$ mutant above the levels observed in the $\Delta pilA$ strain contrasts with a prior study⁷, which may be due to technical or strain differences (see Supplementary Discussion). Importantly, sterically blocking pili with biotin-mal and neutravidin as discussed above inhibited residual retraction events and transformation of the *pilT* mutant, indicating that retraction is still required for natural transformation in this background (Supplementary Fig. 13). We also found that PilT-independent natural transformation and retraction is not mediated by the PilT homolog PilU (Supplementary Fig. 14). Our data thus far indicate that tip bound DNA is internalized via retraction through the outer membrane secretin pore. Therefore, one possibility is that incomplete retraction in the pilT mutant may partially account for the highly reduced rate of transformation and DNA uptake observed.

DNA could be thread through the PilQ outer membrane secretin pore from one end, or may undergo extreme bending to fit through this pore. We found that cells could internalize linear DNA and circular plasmids (i.e. DNA lacking a free end) equally well (Supplementary Fig. 15), suggesting that DNA is likely bent when threaded across the membrane through PilQ. Because dsDNA has a persistence length of ~50 nm (or 150 bp), the extreme bending required to bring DNA through the relatively small PilQ pore (diameter of 7–8 nm based on closely related systems)^{24,25} would require a high force of retraction. Accordingly, another explanation for lack of DNA uptake in *pilT* mutants may be a requirement for a high retraction force and speed to pull DNA through PilQ. To test whether the force and speed of

retraction are altered in *pilT* mutants, we employed a previously described micropillar assay in which cells use pili to bind micropillars that bend as pilus retraction occurs, resulting in measurable retraction rates and forces²⁶. The *pilT* mutant exhibited a significant reduction in both the speed and force of retraction (Fig. 4c–d). Consequently, reduced transformation in the *pilT* mutant may be explained by either a lack of full retraction and/or a reduction in the force or speed of retraction, which prevents translocation of DNA across the outer membrane. Coupled with recent reports that divergent pili lacking a *pilT* homologue can retract^{8,13}, these data support a model whereby *pilT*-independent retraction may be conserved across diverse systems including competence pili from species that lack defined pilus retraction machinery (e.g. *S. pneumoniae, B. subtilis*, and *H. influenzae*). Whether additional canonical type IV pili also exhibit PilT-independent retraction, however, remains to be seen.

Binding and transport of large DNA molecules across the cell envelope is critical for horizontal gene transfer in diverse microbial species, and DNA taken up during this process can also be used as a nutrient. Here, we clarify the mechanism of action for type IV competence pili, the complex nanomachines that promote DNA uptake during natural transformation (Fig. 4e). DNA uptake in *V. cholerae* is sequence independent¹⁵, and we show that minor pilins likely promote DNA binding at the pilus tip. DNA binding to pili in *Neisseria* is also mediated by a minor pilin, ComP, which binds to a species-specific DNA uptake sequence (DUS)^{27,28}. Despite minor pilins playing a role in DNA binding in both of these systems, the proteins involved lack homology, suggesting that distinct mechanisms for pilus DNA binding likely exist among diverse competence pilus systems.

Binding of DNA to the pilus tip as observed here is also compelling because the secretin pore through which the pilus extends and retracts is only wide enough to accommodate the pilus fiber and would likely exclude DNA bound along the pilus length^{24,25}. Thus, retraction of tip bound DNA could allow for threading DNA through the secretin pore in the wake of a retracting pilus fiber. Our data indicate that pilus retraction, however, is not sufficient for DNA uptake into the periplasm, but likely works in conjunction with the molecular ratchet ComEA^{2,17} to mediate efficient DNA uptake.

We show that type IV competence pili in *V. cholerae* exhibit different rates of extension and retraction, which contrasts with other type IV pilus systems that have been characterized where rates of extension and retraction are equal^{8,29}. Notably, the speed of extension and/or retraction were markedly slower than the rates observed for other canonical type IV pili as in *Neisseria gonorrhoeae* (retraction: ~1.2 µm/s³⁰) and *Pseudomonas aeruginosa* (extension and retraction = ~0.5 µm/s²⁹). The mechanism underlying this difference is unclear, however, the pilus from *V. cholerae* studied here is largely dedicated to competence, while the pili in *N. gonorrhoeae* and *P. aeruginosa* can mediate twitching motility, which could account for this difference.

While the type IV competence pili of *V. cholerae* studied here are long micrometer-scale surface appendages, many naturally competent species (such as *Bacillus subtilis* and *Haemophilus influenzae*) possess short "pseudopili" that are thought to simply span the cell envelope. Both long competence pili and short pseudopili, however, could engage in DNA

uptake through a similar mechanism by binding DNA via their exposed tip and using retraction to pull bound DNA through the outer membrane in Gram-negative species or to pull this DNA through the thick peptidoglycan layer in Gram-positive species. Long extended pili involved in competence (as in *V. cholerae, Neisseria spp., S. pneumoniae*, and *Acinetobacter baylyi*) may not be critical for recruitment of freely diffusible DNA (since short pseudopili would suffice), but instead, these structures may be important for binding and uptake of DNA bound to surfaces as would likely be the case in bacterial biofilms.

Methods

Bacterial strains and culture conditions

All *V. cholerae* strains used throughout this study are derivatives of the El Tor isolate E7946³¹. Competence induction in *V. cholerae* requires two cues – chitin oligosaccharides and quorum sensing which activate the expression of TfoX and HapR, respectively^{6,32–34}. Thus, to constitutively activate competence, we ectopically expressed TfoX (via P_{tac} -tfoX) and constitutively activated quorum sensing via inactivation of $luxO^{35,36}$. For a list of all strains used throughout this study, see Supplementary Table 1. Bacteria were routinely cultivated on LB Miller broth and agar supplemented with kanamycin (50 µg/mL), spectinomycin (200 µg/mL), trimethoprim (10 µg/mL), chloramphenicol (1 µg/mL), erythromycin (10 µg/mL), carbenicillin (100 µg/mL), and/or zeocin (50 µg/mL) as appropriate. Instant ocean medium (7 g/L – Aquarium Systems) was used throughout the study where indicated.

Construction of mutant strains

Mutants were constructed by MuGENT, Exo-MuGENT, and/or natural transformation exactly as previously described^{35,37}. Briefly, mutant constructs for deletions or fluorescent fusions were generated via splicing-by-overlap (SOE) PCR to stitch (1) the upstream region of homology (aka the UP arm), (2) the mutation (aka the MIDDLE arm, which could represent an antibiotic resistance cassette or a fluorescent gene), and (3) the downstream region of homology (aka the DOWN arm). For a list of primers used to generate all mutant constructs, see Supplementary Table 2. The UP arm was amplified via an F1/R1 primer pair, while the DOWN arm was amplified with F2/R2 primer pair. All antibiotic resistance cassettes were amplified with ABD123 (ATTCCGGGGGATCCGTCGAC) and ABD124 (TGTAGGCTGGAGCTGCTTC). Fluorescent genes were amplified with the primers indicated in Supplementary Table 2. SOE PCR reactions were carried out using a mixture of the UP, MIDDLE, and DOWN arms in a PCR reaction as template with the F1 and R2 primers. These constructs were then introduced into naturally competent strains as described below in natural transformation assays (for products with an antibiotic resistance marker) or by co-transformation (for constructs that lack a resistance marker – i.e. fluorescent fusions) as previously described³⁵.

For point mutations, mutant constructs were generated via a single PCR reaction using an allele specific F primer that contained >35bp of homology upstream of the point mutation and an R2 primer to generate the downstream region of homology. This product was introduced into a *recJ exoVII* mutant of *V. cholerae* via co-transformation as previously

described³⁷. Subsequently, point mutations were amplified off of the gDNA of these strains with F1/R2 primers and then transferred to strains of interest by co-transformation³⁵.

Mutants were confirmed by PCR and/or sequencing. The P_{tac} -tfoX and $\Delta lacZ$::lacIq mutant constructs are from a previously published study³⁵. To design cysteine replacement mutants and minor pilin lysine/arginine mutations, amino acid residues from pilin sequences were analyzed for solvent accessibility as described previously⁸ using solvent accessibility prediction software³⁸.

Natural transformation assays

Chitin-dependent and chitin-independent transformation assays were conducted as previously described^{35,37}. Unless otherwise noted, natural transformation assays were chitinindependent using strains containing IPTG-inducible P_{tac} -tfoX and $\Delta luxO$ mutations to induce competence. For chitin-independent transformation assays, the indicated strains were grown to late-log in LB + 100 μ M IPTG (to induce TfoX expression) + 20 mM MgCl₂ + 10 mM CaCl₂. Then, ~10⁸ CFUs of this culture were diluted into instant ocean medium. Next, ~500 ng of a transforming DNA was added to each reaction and allowed to incubate for an additional 5 hours at 30°C to allow for natural transformation. For each strain tested, negative controls were performed where no DNA was added. VC1807 is a frame-shifted transposase and its mutation does not impact the growth of V. cholerae³⁵, thus, PCR products for $\Delta VC1807$:: Ab^R (Ab^R = Cm^R, Zeo^R, Tm^R, or Erm^R resistance cassettes depending on the resistance profiles of strains tested) containing 3 kb homology arms (required for maximum rates of transformation 35) were used to test transformation frequency throughout this manuscript. Next, reactions were outgrown by addition of 1 mL of LB and shaking at 37°C for 2 hours followed by plating on media to select for transformants (i.e. media to select for integration of $\Delta VC1807::Ab^{R}$) and on nonselective media to assess the total viable counts. Transformation frequency was defined as the number of transformants divided by the total viable counts. Where indicated, 25 µg/mL of Alexa Fluor 488 C₅ Maleimide (AF488-mal)(Thermo Fisher) was added to the chitin-independent transformation assays 15 min before the addition of transforming DNA.

For chitin-dependent natural transformation assays (only used in Supplementary Fig. 1c), $\sim 10^8$ CFUs of a mid-log culture of the indicated strain were incubated on chitin powder from shrimp cells (Alfa Aesar) in instant ocean medium for 24 hours to allow for competence induction. Then, 500 ng of Δ VC1807::Ab^R transforming DNA was added, and reactions were incubated, outgrown, and plated exactly as described above for chitin-independent transformation assays to attain the transformation frequency.

For cotransformation assays, cells were incubated with two genetically unlinked markers. One marker was a Δ VC1807::Erm^R mutant construct, while the other marker was a previously described Δ VCA1045 mutant construct³⁹, which inactivates the sole mannitol transporter in *V. cholerae*. In these assays, 10ng of the Δ VC1807::Erm^R PCR product was added to transformation assays and then ~3 µg of Δ VCA1045 PCR product was either added to cells immediately or after a 3 hour delay. Cell were then outgrown and plated onto Erm containing media to select for integration of the Δ VC1807::Erm^R product. 24 colonies were then patched into M9 minimal medium containing mannitol as a sole carbon source to

screen for integration of the Δ VCA1045 mutant construct. The cotransformation frequency was defined as the percent of the Δ VC1807::Erm^R mutants that incorporated the Δ VCA1045 product.

To physically obstruct pilus retraction, $\sim 10^8$ CFUs of the indicated strains grown to late-log under competence inducing conditions were washed in instant ocean medium and incubated with 50 µg/mL EZ link Biotin-PEG11-maleimide (biotin-mal)(Thermo Fisher) for 45 min at room temperature. Then, cells were pelleted and washed twice in instant ocean medium. Next, $\sim 10^7$ CFUs were diluted into fresh instant ocean medium. Where indicated, neutravidin (Thermo Fisher) was then added to reactions at a final concentration of 1.32 mg/mL and incubated at room temperature for 30 min. Then, 500 ng of $\Delta VC1807::Ab^{R}$ transforming DNA was added and reactions were incubated at 30°C. For strains where *pilT* was intact, reactions were allowed to proceed for 10 min before the addition of 10 units of DNAse I (NEB) to prevent additional DNA uptake. This was performed because production of new pilins during the normal 5-hour DNA incubation would circumvent our method to block pilus retraction. For transformation assays in the $\Delta pilT$ mutant background, a 10 min incubation with DNA was insufficient to obtain any transformants. Thus, reactions were allowed to proceed for 1 hour at 30°C prior to DNAse I addition as described above. Following DNA addition, all reactions were incubated at 30°C for a total of 5 hours (regardless of the time of DNAse I addition) to allow for DNA integration. Reactions were then outgrown and plated exactly as described above to attain the transformation frequency.

For natural transformation assays of minor pilin DNA binding mutants, strains were incubated with 5 ng of Δ VC1807::Ab^R transforming DNA. After 5 mins, DNAse I was added to prevent additional DNA uptake, and reactions were then outgrown and plated as described above. This was performed to sensitize the assay to the effects of DNA binding on natural transformation.

Pilin labeling, imaging, and quantification

Pilin labeling was achieved using AF488-mal. Cultures were grown to late-log phase in LB supplemented with 20 mM MgCl₂, 10 mM CaCl₂, and 100 µM IPTG to induce competence. ~10⁸ CFUs were then centrifuged at 16,000 × g for 1 min and then resuspended in instant ocean medium supplemented with 20 mM MgCl₂ and 10 mM CaCl₂ before labeling with 25 µg/ml of AF488-mal for 30 min. Labeled cells were centrifuged, washed once, and resuspended in instant ocean medium. Cell bodies were imaged using phase contrast microscopy while labeled pili were imaged using fluorescence microscopy on a Nikon Ti-2 microscope using a Plan Apo 60X objective, a GFP or DsRed filter cube, a Hamamatsu ORCAFlash4.0 camera and Nikon NIS Elements Imaging Software. Cell numbers were quantified using MicrobeJ⁴⁰. To determine pilus length and rates of extension and retraction, labeled cells were imaged by time-lapse microscopy every second for 1 min. For pilus length measurements, pili that were already retracting when imaging began were excluded from the analysis. For extension and retraction rate calculations, only cells that made a single pilus that began extension and fully retracted within the 1 min window were analyzed. To determine the number of pili made per cell, the percent of cells that make pili, and the percent of cells that that form a ComEA-mCherry focus, labeled cells were imaged in the

presence or absence of 1 μ g DNA by time-lapse microscopy every 10 s for 5 min. Pilus length, extension and retraction rates, number of pili made per cell, the percent of cells that make pili, and the percent of cells that form a ComEA-mCherry focus were manually calculated using measurement tools of the NIS Elements analysis software. To block pilus retraction for fluorescence microscopy, cells were co-labeled with 25 μ g/ml of AF488-mal and 25 μ g/ml biotin-mal for 30 min before washing. Where indicated, 1.32 mg/mL neutravidin was then added to washed cells and incubated for 30 min at room temperature before imaging. All imaging was performed under 0.2% Gelzan (Sigma) pads made with instant ocean medium.

Fluorescent DNA binding and localization assays

Strains used for DNA binding and localization assays contained $\Delta pilT$ mutations to ensure the presence of extended pili for DNA binding and to diminish DNA uptake. Fluorescently labeled Cy3- and MFP488-DNA was generated using a 6 kb ΔVC1807::Ab^R PCR product (the same product used for transformation efficiency assays throughout this manuscript) and the LabelIT kit (Mirus Biosciences), which covalently adds a fluorescent label onto ~1 in every 20-60 bp. For DNA binding assays, the indicated strains were grown to late log in LB + 100 μ M IPTG + 20 mM MgCl₂ + 10 mM CaCl₂. Then, ~10⁸ CFUs were centrifuged and washed in instant ocean medium. Cells were then incubated with 100 ng of Cy3- or MFP488-labeled DNA at room temperature for 15 min (MFP488-labeled DNA was used for DNA pulldowns of minor pilin mutant strains because they contained *comEA*-mCherry, which has significant spectral overlap with Cy3). Cells were then pelleted and washed twice with fresh instant ocean to remove unbound DNA. Reactions were then transferred to a 96well plate, and Cy3 or MFP488 fluorescence was determined on a Biotek H1M plate reader. Where indicated, 5 µg ssDNA (PhiX virion DNA, NEB), 5 µg dsDNA (PhiX RFII DNA, NEB), or 10 µg acetylated BSA (Promega) were added to DNA binding reactions as nonlabeled competitors. Where indicated, strains were labeled with biotin-mal and/or neutravidin exactly as described above for natural transformation assays.

For microscopy localization assays on cell-associated pili, cells were labeled as described above before they were added to a coverslip with BSA (0.2 mg/mL) and 4 ng Cy3-labeled DNA. After incubation for 5–15 min, cells were imaged using a Cy3 filter cube every 3 s for 2 min. During imaging, we noticed the majority of DNA binding events occurred at the ends of pilus fibers. However, we experienced several limitations using this setup that prevented quantitative analysis. For example, the large number of pili in $\Delta pi/T$ mutants obscured how many fibers were binding DNA puncta, the cell body fluorescence prevented observation of DNA binding to pili shorter than 1 µm, and Brownian motion in this setup resulted in pili and DNA movement outside of the focal plane.

For robust quantification of DNA localization along pilus filaments, we used sheared pili bound to surfaces within microfluidic channels to alleviate the limitations of using cellassociated pili and to further demonstrate that DNA binding was pilus-dependent. Microfluidic channel devices were constructed exactly as described previously⁸. Channels were pre-treated with BSA (0.2 mg/mL) in instant ocean medium to reduce nonspecific adherence. Then, ~ 10^8 AF488-mal labeled cells were vortexed to shear pili, and then cells

and sheared pili were added to microchannels and incubated for 30 min at room temperature to allow for pilus attachment to the microfluidic channel surfaces. Next, 11 ng of Cy3labeled DNA in a total volume of 400 μ l instant ocean medium was added to the channel to both flush away unattached cells and pili and to introduce DNA to the microchannel. Then, the chamber was incubated at room temperature for 5–15 min with DNA to allow for DNApilus association before imaging. To confirm DNA-pilus binding, gravity flow was applied to microfluidic channels by altering the heights of the input and output tubing to bring both pili and bound DNA into the same field of view for imaging. Only DNA molecules that exhibited colocalization with pili during flow directional changes were analyzed, allowing for exclusion of glass-bound DNA from the analysis. Pili shorter than 0.5 μ m were excluded from the analysis since their movement could not be resolved with directional flow changes. Pilus length and DNA localization was determined using MicrobeJ³². Pilus lengths were normalized where the glass bound end was fixed at 0 and the free end was 1.0, and relative DNA binding was then plotted along the normalized length.

To observe type IV competence pili pulling fluorescent DNA, *pilA*-cys cells harboring native *comEA* (TND0651) were grown to late-log in LB + 100 μ M IPTG + 20 mM MgCl₂ + 10 mM CaCl₂, washed in instant ocean and then mixed with ~5 ng of Cy3-labeled DNA and acetylated BSA was added to 100 μ g/mL. Next, a wet mount was prepared for microscopy on an inverted scope by placing 5–10 μ L of sample onto a 22 × 50 mm coverslip and then covering the sample with a 22 × 22 mm coverslip. Time-lapse imaging was then performed as described above.

DNA internalization assay

~10⁸ CFUs of cells grown to late-log in LB + 100 μ M IPTG + 20 mM MgCl₂ + 10 mM CaCl₂ were pelleted and resuspended in instant ocean medium. Cells were then incubated with 10 ng of either an MFP488-labeled 6kb Δ VC1807::Ab^R PCR product or an ~6kb supercoiled plasmid (pBAD18Kan) at room temperature for 1 hour. Next, 10 units of DNase I was added to reactions and incubated at room temperature for 1 min to degrade any remaining extracellular DNA. Cells were then washed once with instant ocean medium to remove excess dye and imaged. The percent of cells with internalized DNA was quantified using MicrobeJ.

mCherry-PilQ localization

Cells were grown to late-log in LB + 100 μ M IPTG + 20 mM MgCl₂ + 10 mM CaCl₂ and imaged. Localization of mCherry-PilQ foci was determined and plotted using MicrobeJ.

Cell body fluorescence quantification

Strains were labeled with AF488-mal as indicated above and imaged for 10 s intervals over a 20 min window. Integrated fluorescence intensity was measured for both the cell body and the pilus for each cell for each time point using ImageJ⁴¹. Fluorescence intensities were normalized for photobleaching over the time-lapse experiment by multiplying the measured fluorescence intensity by the percentage of fluorescence lost from the total fluorescence intensity (the sum of the cell body and pilus intensities) for each time point. After adjustment for photobleaching, the cell body fluorescence for each time point was divided

by the cell body fluorescence measured at time 0. The relative cell body intensity changes were plotted with the measured pilus length for each time point.

Negative stain electron microscopy

To ensure that the pili imaged via EM were type IV competence pili, we inactivated the two other type IV pilus systems (MSHA and TCP) in *V. cholerae* yielding strain SAD2094 (see Supplementary Table 1 for additional strain details). We also generated a *pilA*-cys mutation in this background (SAD2093). Strains were gown to late-log in LB + 100 μ M IPTG + 20 mM MgCl₂ + 10 mM CaCl₂ and then treated with biotin-mal and/or neutravidin exactly as described above for natural transformation assays.

To prepare the negative stain specimen, 4 μ L of sample solution was applied on a glowdischarged, carbon-coated, 300-mesh copper grid for 30 s. The excess solution was removed by a piece of filter paper. The grid was washed with 4 μ L of Milli-Q water, stained with 4 μ L of 0.75% (w/v) uranyl formate for 25 s and blotted to dry. The images were acquired at a nominal magnification of 10,000x, 20,000x or 40,000x using JEOL JEM-1400 Plus transmission electron microscope operated at 120 kV and a Gatan 4k × 4k OneView Camera.

Micropillar retraction assay

To ensure that all measurements recorded were due to type IV competence pili, all the experiments were performed with strains lacking all external appendages other than type IV competence pili as well as exopolysaccharide production (i.e. mutants lacking flagella, MSHA pili, TCP pili, and VPS production). See Supplementary Table 1 for additional strain details. Strains were grown overnight in LB either directly from frozen stocks or from a single colony off of an LB agar plate. 50 µL of the overnight liquid culture was subcultured into 3 ml of LB + 100 μ M IPTG + 20 mM MgCl₂ + 10 mM CaCl₂ and allowed to grow at 30 °C for 5 hours. Then, 100 μ L of this culture was centrifuge for 5 min at 20,000 × g and the bacteria were resuspended in instant ocean medium. At different times, 10 µL of the resuspension was added to micropillars in an observation chamber as previously reported²⁶. Briefly, silica molds were inverted on activated coverslips with polyacrylamide gels in between. The result is an array of flexible micropillars in a hexagonal array of $3 \mu m \times 3 \mu m$. Once the bacteria are in contact with the micropillars, 10Hz movies of the top of the pillars are recorded. The motion of the tips of the pillars is tracked using a cross-correlation algorithm in ImageJ⁴¹. The amplitude and speeds of the pillars' motions were then analyzed using Matlab. Finally, we calibrated the pillars' stiffness constant using optical tweezers as previously described²⁶. The pillars used in this study have a stiffness constant of 17±4 pN/ μm.

Statistics

Significance was calculated using statistical tests on the GraphPad (Prism) 5.0 software. Statistical differences between two groups were analyzed using two-tailed Student's t-tests. Complete statistics can be found in Supplementary Table 3. Sample sizes were chosen based on historical data and no statistical methods were used to predetermine sample size.

Data availability statement

The data that support the findings of this study are available from the corresponding author upon request.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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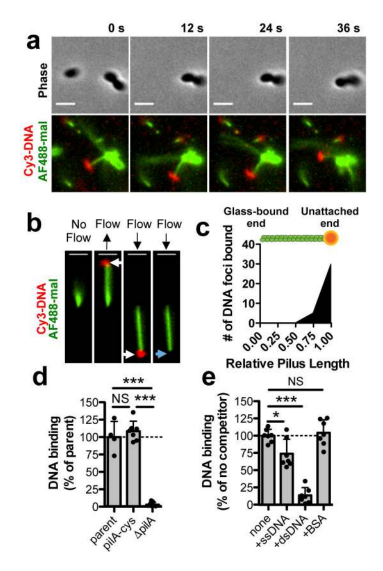


Fig. 1. The tips of type IV competence pili directly bind DNA.

Because type IV competence pili are dynamically active and relatively few pili are extended at any given time, all pilus-DNA binding studies were performed in $\Delta pilT$ backgrounds to ensure that cells contained numerous extended pili as shown in Supplementary Fig. 1c. (**a**) Montage of time-lapse imaging of a *pilA*-cys $\Delta pilT$ cell labeled with AF488-mal after mixing with Cy3-labeled DNA in a wet mount. Scale bar, 2 µm. (**b**) Example of a sheared pilus labeled with AF488-mal bound to glass within a microfluidic device after the addition of fluorescently labeled DNA. Before flow is applied, the relatively long pilus fiber is out of the field of view, but after flow is applied, bound DNA becomes apparent (white arrows) and moves with the direction of flow. When flow is maintained, the DNA detaches from the pilus fiber (blue arrow). Black arrows above images represent the direction of flow. Scale bar, 2 µm. Data in **a-b** are representative of 3 independent experiments. (**c**) Localization of bound DNA along sheared pili in microfluidic devices. Pili were normalized in length to 1.00 (arbitrary units), and the position of DNA foci bound along the length for each fiber was plotted, n = 51 independent DNA-bound pili analyzed. Green rod above plot is a visual representation of a labeled pilus fiber bound to orange DNA. (**d**) Relative DNA binding

assay using a fluorescently labeled 6 kb PCR product. parent n = 4, pilA-cys n = 7, Δ pilA, n = 7. (e) Relative DNA binding as in **d** except with the addition of the indicated non-labeled competitors at 50-fold excess. Competitors were ssDNA (PhiX174 virion), dsDNA (PhiX174 RFII), or bovine serum albumin (BSA). n = 7 for all samples. Each data point in **d-e** represents an independent biological replicate and bar graphs indicate the mean ± SD. Statistical comparisons were made by two-tailed Student's *t*-test. NS = not significant, **P* < 0.05, ****P* < 0.001.

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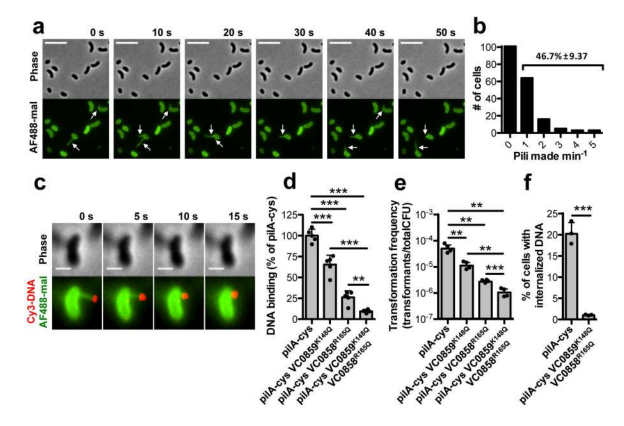


Fig. 2. Type IV competence pilus dynamic activity and DNA binding are critical for DNA internalization.

(a) Montage of time-lapse imaging of *pilA*-cys cells after labeling with AF488-mal dye. The white arrows indicate pili. Scale bar, 5 µm. (b) Number of cells making 0–5 pili within a one-minute period. Data are from three independent, biological replicates, n = 192 total pili observed. Percentage is the percent of cells within the population that make pili within the one-minute time-frame \pm SD. (c) Montage of time-lapse imaging of a retracting *pilA*-cys strain after labeling with AF488-mal and incubation with fluorescently labeled DNA in a wet mount. Scale bar, 1 µm. Similar events were captured in three independent experiments. (d) Relative DNA binding assay of the indicated strains in $\Delta pi/T$ mutant backgrounds using a fluorescently labeled 6 kb PCR product. (e) Natural transformation assays of the indicated strains. Cells were incubated with 5 ng of transforming DNA (tDNA) and DNase I was added to reactions after 10 mins to prevent additional DNA uptake. (f) DNA internalization assays of the indicated strains using a fluorescently labeled 6 kb PCR product. VC0858 and VC0859 encode minor pilins. Each data point in **d-f** represents an independent biological replicate (d-e, n = 4 for all samples; f, n = 3 for all samples) and bar graphs indicate the mean \pm SD. All statistical comparisons were made by two-tailed Student's *t*-test. ***P* < 0.01, ****P* < 0.001.

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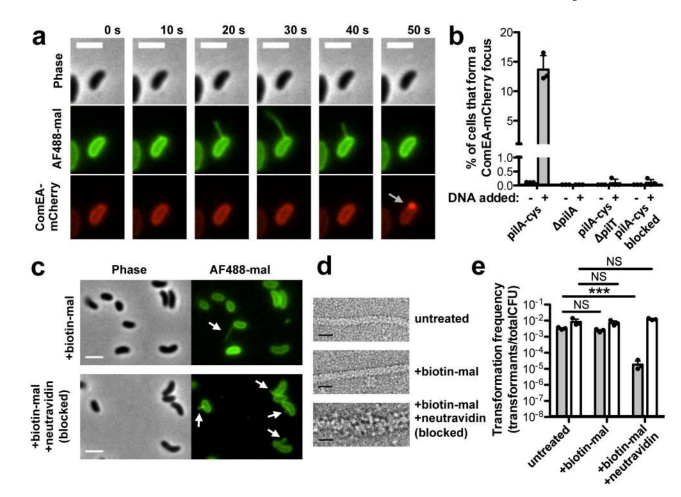


Fig 3. Pilus retraction is required for DNA uptake.

(a) Montage of time-lapse imaging of *pilA*-cys ComEA-mCherry strain labeled with AF488mal showing ComEA focus formation after pilus retraction (gray arrow). Scale bar, 2 µm. (b) Percent of cells that formed a ComEA-mCherry focus within a five min window in the presence or absence of DNA. Cells that had already formed ComEA-mCherry foci at the start of imaging were excluded from the analysis. Data are from three independent experiments (n = 3 for each condition) and shown as the mean \pm SD. (c) Static images of *pilA*-cys cells labeled with a 1:1 ratio of biotin-mal:AF488-mal with or without added neutravidin. White arrows indicate pili. Scale bar, 2 µm. (d) Transmission electron micrographs of *pilA*-cys pili with the indicated treatments. While untreated and biotin-mal treated pili exhibit a similar thickness, the biotin-mal + neutravidin treated pilus is approximately twice as thick. Scale bar, 10 nm. Images in **c-d** are representative of two independent experiments. (e) Natural transformation assays of *pilA*-cys (black bars) or parent (white bars) strains after the indicated treatment. Cells were incubated with 500 ng of tDNA and DNase I was added to reactions after 10 mins to prevent additional DNA uptake. Data are from three independent, biological replicates and shown as the mean \pm SD. Statistical comparisons were made by two-tailed Student's *t*-test. NS = not significant. ***P < 0.001.

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