CRISPR/Cas9-based Genome Editing in *Pseudomonas aeruginosa* and Cytidine Deaminase-Mediated Base Editing in *Pseudomonas* Species

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**HIGHLIGHTS**

The pCasPA/pACRISPR system enables efficient genetic manipulation in *P. aeruginosa*

The pnCasPA-EBC system allows rapid C→T base editing in *Pseudomonas* species

The two tools will accelerate a variety of investigations in *Pseudomonas* species

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CRISPR/Cas9-based Genome Editing in Pseudomonas aeruginosa and Cytidine Deaminase-Mediated Base Editing in Pseudomonas Species

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SUMMARY

Pseudomonas species are a large class of gram-negative bacteria that exhibit significant biomedical, ecological, and industrial importance. Despite the extensive research and wide applications, genetic manipulation in Pseudomonas species, in particular in the major human pathogen Pseudomonas aeruginosa, remains a laborious endeavor. Here we report the development of a genome editing method pCasPA/pACRISPR by harnessing the CRISPR/Cas9 and the phage λ-Red recombination systems. The method allows for efficient and scarless genetic manipulation in P. aeruginosa. By engineering the fusion of the cytidine deaminase APOBEC1 and the Cas9 nickase, we further develop a base editing system pnCasPA-BEC, which enables highly efficient gene inactivation and point mutations in a variety of Pseudomonas species, such as P. aeruginosa, Pseudomonas putida, Pseudomonas fluorescens, and Pseudomonas syringae. Application of the two genome editing methods will dramatically accelerate a wide variety of investigations, such as bacterial physiology study, drug target exploration, and metabolic engineering.

INTRODUCTION

Pseudomonas is a genus of gram-negative Gammaproteobacteria existing in diverse environments, such as soil, water, plant, and human body (Palleroni, 1993; Peix et al., 2009). The significant biomedical, ecological, and industrial importance of these bacteria has rendered them as an excellent focus for scientific research (Gross and Loper, 2009; Peix et al., 2009). For instance, Pseudomonas aeruginosa is a major human pathogen and is capable of causing severe infections in vulnerable patients hospitalized with cystic fibrosis, burns, acquired immunodeficiency syndrome, or cancer. Extensive research has been focused to dissect the molecular basis of the infection mechanisms and to develop novel therapeutic means against P. aeruginosa infections (Hauser et al., 2011; Sadikot et al., 2012). Pseudomonas putida is a common soil bacterium and has been widely utilized for bioremediation and high-value chemical production (Moreno and Rojo, 2013; Nikel et al., 2016). Great efforts of metabolic engineering have been made to boost the capacity of P. putida in harsh environment and chemical production (Loeschcke and Thies, 2015; Wu et al., 2011).

Basic physiology studies and applied investigations in Pseudomonas species would benefit greatly from rapid and efficient genome editing tools. Despite the recent progress in the development of CRISPR-based genome engineering methods in Pseudomonas species, including the CRISPR-based systems for genome engineering in P. putida (Aparicio et al., 2017; Sun et al., 2018) and the dCas9-based transcription inhibition system in Pseudomonas (Tan et al., 2018), the genetic manipulation methods in most Pseudomonas species, such as the major human pathogen P. aeruginosa, are still time consuming and laborious (Hmelo et al., 2015; Hoang et al., 1998; Martinezgarcía and Lorenzo, 2011; Thomas et al., 2009). For instance, to construct a clean deletion mutant in P. aeruginosa, a two-step selection process is often required. First, a target gene is replaced by an antibiotic marker via homologous recombination. Second, the antibiotic marker is eliminated with the help of the FLP recombinase, leaving a scar sequence in place of the deleted gene (Hoang et al., 1998).

CRISPR/Cas9, obtained from bacterial adaptive immune systems (Horvath and Barrangou, 2010), has been engineered for genome editing in a variety of organisms, such as mammalian cells (Cong et al., 2013; Mali et al., 2013), Saccharomyces cerevisiae (Bao et al., 2015; Ryan and Cate, 2014), Escherichia coli (Jiang et al., 2013), and Staphylococcus aureus (Chen et al., 2017). In this system, the Cas9 DNA nuclease forms a
complex with a single guide RNA (sgRNA). The complex can be precisely guided to any genomic locus via base pairing of the programmable 20-nucleotide (nt) sequence of sgRNA with the genomic DNA when an adjacent protospacer-adjacent motif (PAM, e.g., 5′-NGG-3′ for *Streptococcus pyogenes* Cas9 [Jinek et al., 2012]) is present in the target locus (Anders et al., 2014; Nishimasu et al., 2014; Wang et al., 2016; Wiedenheft et al., 2012). After binding, the Cas9 DNA nuclease cleaves the target locus, generating a double-stranded DNA break in the genome. Given that bacterial cells do not possess the non-homologous end-joining repair pathway, only the cells that have undergone homologous recombination can survive after the double-stranded DNA break of the genome. Thereby, it is possible to achieve a one-step seamless genome editing in *Pseudomonas* species with the utilization of the CRISPR/Cas9 system.

The recent development of “base editors” opens a new avenue for genome editing in biological systems. The systems directly catalyze the conversion of bases via a deamination reaction without generating a double-stranded DNA break or utilizing a donor repair template (Gaudelli et al., 2017; Komor et al., 2016). Until now, two kinds of base editors have been developed. The cytidine editor mediates the conversion of C to T (or G to A) (Bannos et al., 2018; Gu et al., 2018; Komor et al., 2016), whereas the adenosine editor effects an A to G (or T to C) substitution (Gaudelli et al., 2017). The base editors are typically composed of a defective Cas9 protein (Cas9D10A or Cas9D10AH840A) and a deaminase fused to the Cas9 protein. Guided by the Cas9/sgRNA complex, the deaminase can be directed to any genomic locus to perform base editing in the single-stranded DNA (ssDNA) generated upon Cas9/sgRNA binding. By catalyzing the conversion of CAA, CAG, CGA, or TGG to TAA, TAG, or TGA codons, the cytidine base editor is capable of inactivating a target gene by generating a premature stop codon.

In this study, we developed a CRISPR/Cas9-mediated genome editing method, allowing for efficient and scarless genetic manipulation in *P. aeruginosa*. Moreover, we developed a base editing system, enabling highly efficient C to T (or G to A) conversion in *Pseudomonas* species. The two genome editing methods would greatly simplify the genetic manipulation in *Pseudomonas* species and accelerate a wide variety of investigations.

**RESULTS**

**Construction of CRISPR/Cas9-based Genome Editing System in *P. aeruginosa***

To develop an efficient and convenient genetic manipulation method in *Pseudomonas* species, we first sought to harness the CRISPR/Cas9 system for genome editing in these bacteria. We constructed a single-plasmid system pCasPAIgm that possessed both the *Streptococcus pyogenes* Cas9 (spCas9) protein and the corresponding sgRNA expression cassettes (Figure S1A) [Jinek et al., 2012]. We then assembled a 20-nt spacer and repair arms (~1 kb each) into the plasmid to assess the editing efficiency of this system in a model *P. aeruginosa* strain PAO1. This system succeeded in deleting the wspF gene (GenBank: PA3703) with an efficiency of 9/11 (Figure S1B). However, it failed to delete the nalD gene (GenBank: PA3574) or rhlR (GenBank: PA3477) gene (Figures S1C and S1D). The discrepancy in the efficiency of the system in deleting different genes is likely ascribed to the weak intrinsic homologous recombination capacity of *P. aeruginosa*, which was further confirmed by the observation that only a small amount of colonies (10–20 colonies) could be obtained after editing even in the successful case of the deletion of the wspF gene.

Phage recombination systems, such as λ-Red and RecET, have a higher homologous recombination capacity and have been engineered alone or in a couple of CRISPR systems for genetic manipulation in a variety of organisms (Datsenko and Wanner, 2000; Jiang et al., 2013; Jiang et al., 2015; Jiang et al., 2017; Murphy, 2012; Oh and van Pijkeren, 2014; Zhang et al., 1998). Thereby, we sought to increase the efficiency of homologous-recombination-mediated double-stranded DNA break repair in *P. aeruginosa* by introducing the λ-Red recombination system into the organism. Direct installation of the λ-Red system into the one-plasmid system pCasPAIgm did not improve the efficiency of deleting the nalD gene (Figures S1E and S1F). Inspired by the successes of using a two-plasmid system for CRISPR/Cas9-mediated genome editing in *E. coli* and other bacteria (Jiang et al., 2013, 2015), we designed and constructed a two-plasmid system pCasPA/pACRISPR. The pCasPA plasmid was capable of expressing the Cas9 nuclease and the λ-Red system proteins, Exo, Gam, and Bet. The expression of both the Cas9 nuclease and the λ-Red system was driven by the L-arabinose-inducible promoter P araB (Narayanan et al., 2006) (Figure 1A). The pACRISPR plasmid was used to express the sgRNA. The expression of the sgRNA was driven by the well-studied strong promoter trc (Brosius et al., 1985) (Figure 1B). The pACRISPR plasmid contained two seamless cloning sites (Figure 1C). The Bsal sites were used for the insertion of the 20-nt spacer by Golden Gate assembly.
Engler et al., 2009), and the XbaI and XhoI sites were utilized for one-step cloning of repair arms by Gibson assembly (Gibson et al., 2009). The counter-selectable sacB that worked as a lethal gene in the presence of sucrose was introduced into both plasmids to facilitate rapid plasmid curing after editing (Schweizer, 1992).

Assessment of the Editing Efficiency of the pCasPA/pACRISPR System

To test the functionality of the pCasPA/pACRISPR system in genome editing in P. aeruginosa, we first transformed the pCasPA plasmid into the PAO1 strain. After the induction by L-arabinose for 2 hr, the cells containing the pCasPA plasmid were collected and prepared as the electrocompetent cells. Next, the empty pACRISPR plasmid, pACRISPR assembled with a 20-nt rhlR spacer (pACRISPR-rhlR_spacer), pACRISPR assembled with the rhlR repair arms (pACRISPR-rhlR_repair), as well as pACRISPR assembled with both spacer and repair arms (pACRISPR-rhlR) were electroporated into the cells for genome editing (Figures 2 and 3A). As shown in Figure S2A, more than 10^3 colonies were observed for a single transformation with the pACRISPR and pACRISPR-rhlR_repair plasmid, whereas fewer than 10 colonies could be obtained for the same transformation with the pACRISPR-rhlRSpacer plasmid. The introduction of the rhlR spacer will produce an intact sgRNA that directs the Cas9 endonuclease to the rhlR gene locus to create a double-stranded break, leading to the death of cells. Around 100 colonies could be recovered when the cells were transformed with the pACRISPR-rhlR plasmid containing both the spacer and the repair arms. The rhlR gene in the colonies that was electroporated with the pACRISPR-rhlR plasmid was successfully deleted with an efficiency of 12/12, confirmed by PCR, sequencing, and the pigment production assay (loss of pigment production is a major phenotype for rhlR disruption [Brint and Ohman, 1995; Cao et al., 2014]) (Figure 3B). In addition, we also transformed the pACRISPR-rhlR plasmid into the pCasPA-containing...
PAO1 strain without the L-arabinose induction. As shown in Figure S2A, only several colonies were recovered on the plate, confirming that the \( \lambda \)-Red system is essential for the recombination and the leakage expression of the Cas9 protein is sufficient for the double-stranded DNA break.

Because the \( \lambda \)-Red recombination system is capable of promoting homologous recombination with the utilization of short (~50 bp each) repair templates (Liang and Liu, 2010), we systematically investigated the editing efficiencies of the two-plasmid system when different lengths of repair templates were utilized. First, we tested the editing efficiencies of using the repair templates cloned into the pACRISPR plasmid. As shown in Figure S2B, the efficiencies of deleting the \( \text{nalD} \) gene were 100% when 500 bp or longer repair arms were utilized. However, the editing efficiency reduced significantly when shorter repair templates were utilized. No desired mutants could be obtained when a 45-bp repair arm was utilized (Figure S2B). In addition to the circular double-stranded DNA (cloned into the pACRISPR plasmid in this case), the \( \lambda \)-Red recombination system is also capable of using linear ssDNA or double-stranded DNA for recombination (Datsenko and Wanner, 2000; Ellis et al., 2001). Thereby, we tested the editing efficiency of the two-plasmid system when different lengths of linear ssDNA or double-stranded DNA were utilized as the repair templates. We co-transformed the linear repair templates and the pACRISPR plasmid into the cells that contained the pCasPA plasmid. The editing efficiencies of using the linear 45-nt single-strand DNA, 100-bp double-stranded DNA, 200-bp double-stranded DNA, 500-bp double-stranded DNA, and 1-kb double-stranded DNA were 2/12, 3/12, 9/12, 5/12, and 1/12, respectively (Figure S2C). The reduced editing efficiency of using 500-bp or longer linear repair templates is likely ascribed to the decreased transformation efficiency of long linear DNAs (Ghanta et al., 2018).

Given the differential editing efficiencies of using different types and lengths of the repair templates, we used 500-bp circular repair templates (assembled into the pACRISPR plasmid) for the subsequent genome editing experiments. We tested the efficiencies of the two-plasmid system in the deletions of four other genes in the PAO1 strain. The efficiencies of deleting \( \text{lasR} \) (GenBank: PA1430), \( \text{rsaL} \) (GenBank: PA1431), \( \text{algR} \) (GenBank: PA5261), and \( \text{rhlB} \) (GenBank: PA3478) genes were 11/12, 12/12, 5/12, and 8/12, respectively (Figures 3C and S3A).

To assess the capacity of the pCasPA/pACRISPR system in large-fragment deletion, we constructed three plasmids pACRISPR-\( \text{rhlR} \)3k, pACRISPR-\( \text{rhlR} \)5k, and pACRISPR-\( \text{rhlR} \)10k that were used to delete a 3-, 5-
and 10-kb DNA fragment, respectively, in the rhlR gene locus. As shown in Figure S3B, the 3-kb DNA fragment could be deleted with an efficiency of 12/12. However, the attempts to delete both the 5- and 10-kb DNA fragments failed. In addition to large-fragment deletion, we applied the pCasPA/pACRISPR system for multiplex gene editing. We cloned the spacers and repair arms of both the rhlR and lasR genes into a single pACRISPR plasmid and transformed it into the bacterial cells for editing. As shown in Figure S3C, the rhlR and lasR genes could be deleted simultaneously. However, the transformation colony-forming units (CFUs) decreased dramatically. Only 10–20 colonies were obtained from a single transformation (Figure S3D).

We noticed that some colonies could escape the CRISPR-induced death in the editing experiments. To probe the possible mechanism, we picked 4 escaped transformants from the PAO1ΔnalD plate. First, we sequenced the target gene (nalD gene locus) and no mutations were observed. Second, we tried to amplify the sgRNA fragment (~1 kb) in the editing plasmid of the escaped transformants. As shown in Figure S3E, no band of ~1 kb was observed for the 4 escaped transformants, whereas a clear band of ~1 kb could be observed for the transformant that was successfully edited (ck). The results revealed that the editing plasmids from the escaped transformants contained mutations or deletions in DNA sequences coding for Cas9/sgRNA.
In addition to the genome editing in the *P. aeruginosa* PAO1 strain, we investigated the editing efficiency of the pCasPA/pACRISPR system in another widely utilized *P. aeruginosa* strain PAK. The efficiencies of deleting *rscA*, *algR*, and *lasR* genes in the PAK strain were 10/12, 8/12, and 6/12, respectively (Figure S3F). In addition to gene deletion, we assessed the capacity of the two-plasmid system in gene insertion in *P. aeruginosa*. The native *rhlA* (GenBank: PA3479) promoter was successfully replaced by two foreign *trc* and *rpsL* promoters, both with efficiencies of 11/12 (Figures 3D and 3F). Together, these experiments demonstrated that the two-plasmid system pCasPA/pACRISPR possessed a great capacity for genome editing in *P. aeruginosa*.

**Plasmid Curing after Editing**

To cure the plasmids after editing, one colony from the PAO1 strain containing the desired *naID*-gene deletion was cultured in fresh Luria-Bertani (LB) medium until growth was evident. The culture was diluted for $10^6$ folds with fresh LB medium, and 100 mL diluted culture was plated onto the LB plates in the presence or absence of 5% w/v sucrose. Notably, much fewer colonies grew on the plate that had sucrose than on the plate that did not have sucrose (Figure S3G). Six individual colonies from the plate containing sucrose were randomly picked and streaked onto three different LB agar plates (no antibiotics, 100 μg/mL tetracycline, and 150 μg/mL carbenicillin). All the six colonies grew normally on the plate without antibiotics, whereas no growth of colonies was observed on the plates containing tetracycline or carbenicillin (Figure S3H), thus confirming that both the pCasPA and the pACRISPR plasmids could be easily cured after editing.

**Construction of the Base-Editing System pnCasPA-BEC**

To expand the utility of the pCasPA/pACRISPR system, we assessed the capacity of this system for genome editing in other widely studied microbes, including *P. putida*, *Pseudomonas fluorescens* (a plant growth-promoting bacterium), and *Pseudomonas syringae* (a major plant pathogen). However, the extremely low efficiency for the transformation of the pCasPA plasmid into these microbes prevents its applications in genome editing in these bacteria. The large size of the pCasPA plasmid (17653 bp) and the toxicity of the Cas9 protein and the λ-Red system may be the possible reasons for the low transformation efficiency (Jiang et al., 2014; Sun et al., 2018). Thereby, we sought to develop a base editing system as an alternative way for genome editing in these bacteria, because base editing systems have been demonstrated to be emerging tools for genetic manipulation in microbes without generating double-stranded DNA break or sacrificing transformation CFUs (Gu et al., 2018). We designed and constructed a base editing system pnCasPA-BEC (Figure 5A). In this plasmid, the cytidine deaminase (rat APOBEC1) was fused to the N terminus of the Cas9 nickase (SpCas9D10A) via an XTEN linker (Komor et al., 2016) (Figures 4A and 4B). The expression of the fusion protein and the sgRNA were driven by the *trc* promoter (SpCas9D10A) via an XTEN linker (Komor et al., 2016) (Figures 4A and 4B). The expression of the fusion protein and the sgRNA were driven by the *rpsL* promoter (from PAO1) and the *trc* promoter, respectively. Two *BsaI* sites were engineered into the plasmid for seamless cloning of the 20-bp spacer using Golden Gate assembly. The broad-host-range replicon mSF was introduced in this system for plasmid replication in *Pseudomonas* species. In addition, we utilized the *sacB* gene for plasmid curing after editing.

**Assessment of the Editing Efficiency of the pnCasPA-BEC System in Pseudomonas Species**

We first assessed the capacity of the pnCasPA-BEC system in base editing in the PAO1 strain. We designed spacers of *rhlR* and *rhlB* genes containing potential editable C(s) within the editable window, which was reported from positions 4 to 8 in mammalian cells (Komor et al., 2016). The spacers were assembled into the pnCasPA-BEC plasmid, and the constructed plasmids were transformed into the PAO1 strain. In agreement with our expectation, the Cs at positions 6 and 7 of the *rhlR* spacer, and the C at position 5 of the *rhlB* spacer, were successfully mutated to T with an efficiency of 11/12 for both two genes (Figures 4C, 4D, and 4B). These conversions generated premature stop codons in the genes, resulting in gene inactivation. In addition, this system succeeded in base editing in the *P. aeruginosa* PAK strain with high efficiencies (14/15 for *rhlR* and 17/17 for *rhlB*) (Figures 4E and 4F). After editing, the plasmid could be easily cured by plating the cells on the plate containing sucrose (Figure S4C).

To assess the nonspecific mutator effects of pnCasPA-BEC, we searched spacers similar to the *rhlR* and *rhlB* spacers across the entire genome using sgRNAcas9 software (Xie et al., 2014). We picked the top six similar spacers for each gene (Table S1). Next, we amplified the spacer locus and sent them out for sequencing. The results showed that none of the similar spacer sites were mutated (Table S1).

Next, we applied the pnCasPA-BEC system for base editing in other *Pseudomonas* species, including *P. putida*, *P. fluorescens*, and *P. syringae*. As shown in Figures 5A–5F, all the *cadR* (GenBank: PP_5140)}
and ompR (GenBank: PP_0246) genes in P. putida KT2440, the per (GenBank: GU120326) and aspC (GenBank: FJ485937) genes in P. fluorescens GcM5-1A, as well as the gacA (GenBank: PSPTO_3024) and hrpL (GenBank: PSPTO_1404) genes in P. syringae DC3000 were successfully mutated with high efficiencies, demonstrating the great capacity of the pnCasPA-BEC system for base editing in a variety of Pseudomonas species.

To systematically investigate the editable window of the pnCasPA-BEC system in Pseudomonas species, we designed and assembled eight different spacers containing Cs at positions from 2 to 9 into the pnCasPA-BEC plasmid. Next, we transformed the plasmids into the PAO1 strain and examined the editing efficiencies. The results showed that all the Cs at positions 3 to 8 were mutated to Ts with an efficiency of 100%, whereas the Cs at the position 2 or 9 could not be mutated (Figure S4D). Thereby the possible editable window of the pnCasPA-BEC system was from positions 3 to 8 in the PAO1 strain (Figure S4D). Notably, the adjacent bases of the editable sites greatly affected the editing efficiency. The editing efficiencies of GC and AC were much lower than those of CC and TC (Figures 4C, 4D, S4D, and S4E), in agreement with the results of mammalian cells (Komor et al., 2016).

**DISCUSSION**

We have engineered the CRISPR/Cas9 and the λ-Red recombination systems (pCasPA/pACRISPR) for rapid, precise, and seamless genetic manipulation in P. aeruginosa. We first directly installed the λ-Red system and CRISPR/Cas9 into a single plasmid, but the resulting plasmid failed to improve the editing efficiency (Figures S1E and S1F). From our experiences and others’ work (Jiang et al., 2013; Sun et al., 2018), to achieve a successful editing, the λ-Red recombination proteins probably need to be pre-expressed in bacterial cells before the genome cleavage by CRISPR/Cas9. Thus, when both the CRISPR/Cas9 system and the λ-Red system are present in a single plasmid, the λ-Red system may be expressed too late to repair the double-stranded DNA break generated by the CRISPR/Cas9 system that is expressed at the same time as the λ-Red system in the one-plasmid system. These observations may shed light on the development of CRISPR/Cas9-based genome editing tools in other bacteria.

We systematically investigated the optimal editing conditions of the pCasPA/pACRISPR system. Both circular and linear templates could be used for homology-directed repair (Figures S2B and S2C). Although short linear ssDNA repair template (e.g., 90 bp) could be synthesized easily and used for repair directly,
its repair efficiency was relatively low (Figure S2C). Long circular template (>500 bp) exhibited high repair efficiency (Figure S2B). However, it had to be cloned into the editing plasmid for repair. The pCasPA/pACRISPR system also enabled multiplex gene editing (Figure S3C). However, the complex construction process and low-transformation CFUs may limit its application.

The highly efficient base-editing system pnCasPA-BEC is capable of inactivating genes in a variety of Pseudomonas species. Compared with the recently reported Cpf1-BEC that recognizes AT-rich (TTTV) PAM (Li et al., 2018), the Cas9-BEC that recognizes the NGG PAM is more suitable for base editing in \( P. \) aeruginosa, because the genome of \( P. \) aeruginosa is GC rich. No off-target mutations were detected in the six similar spacers of the \( rhlR \) and \( rhlB \) genes. The editing window of the pnCasPA-BEC system is from positions 3 to 8 in the PAO1 strain, which is slightly different from that of the base editors in the mammalian cells (Komor et al., 2016) and \( S. \) aureus (Gu et al., 2018). Given the ease of use and high efficiency, future engineering of the base-editing system would provide a new way for high-throughput screening in Pseudomonas species.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file. The strains, plasmids, and primers used in this study are listed in Tables S2, S3, and S4, respectively.

SUPPLEMENTAL INFORMATION
Supplemental Information includes Transparent Methods, four figures, and four tables and can be found with this article online at https://doi.org/10.1016/j.isci.2018.07.024.

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Pseudomonas putida boosts recombineering efficiency in Aparicio, T., Lorenzo, V.D., and Martínez-García, Jinek, M. (2014). Structural basis of PAM-dependent target DNA recognition by the Cas9 endonuclease. Nature 513, 569–573.

Aparicio, T., Lorenzo, V.D., and Martínez-García, E. (2017). CRISPR/Cas9-based counterselection boosts recombineering efficiency in Pseudomonas putida. Biotechnol. J. 13, 1700161.

Banno, S., Nishida, K, Araroz, T., Mitsunobu, H., and Kondo, A. (2018). Deaminase-mediated multiplex genome editing in Escherichia coli. Nat. Microbiol. 3, 423–429.

Bao, Z., Xiao, H., Liang, J., Zhang, L., Xiang, X., Sun, N., Si, T., and Zhao, H. (2015). Homology-integrated CRISPR-Cas (HI-CRISPR) system for one-step multigene disruption in Saccharomyces cerevisiae. ACS Synth. Biol. 4, 585–594.

Brint, J.M., and Ohman, D.E. (1995). Synthesis of multiple exoproteins in Pseudomonas aeruginosa is under the control of RhlR-RhlI, another set of regulators in strain PAO1 with homology to the autoinducer-responsive LuxR-LuxU family. J. Bacteriol. 177, 7155–7163.

Brosius, J., Effe, M., and Storella, J. (1985). Spacing of the -10 and -35 Regions in the tac promoter. J. Biol. Chem. 260, 3529–3541.

Cao, Q., Wang, Y., Chen, F., Xia, Y., Lou, J., Zhang, X., Yang, N., Sun, X., Zhang, Q., Zhuo, C., et al. (2014). A novel signal transduction pathway that modulates rhl quorum sensing and bacterial virulence in Pseudomonas aeruginosa. PLoS Pathog. 10, e1004340.

Chen, W., Zhang, Y., Ye, W.S., Bae, T., and Ji, Q. (2017). Rapid and efficient genome editing in Staphylococcus aureus by using an engineered CRISPR-Cas9 system. J. Am. Chem. Soc. 139, 3790–3795.

Cong, L., Ran, F.A., Cox, D., Lin, S., Barretto, R., Habib, N., Hsu, P.D., Wu, X., Jiang, W., Marraffini, L.A., et al. (2013). Multiplex genome engineering using CRISPR/Cas systems. Science 339, 819–823.

Datson, K.A., and Wanner, B.L. (2000). One-step inactivation of chromosomal genes in Escherichia coli K-12 using PCR products. Proc. Natl. Acad. Sci. USA 97, 6640–6645.

Ellis, H.M., Yu, D., Ditizio, T., and Court, D.L. (2001). High efficiency mutagenesis, repair, and engineering of chromosomal DNA using single-stranded oligonucleotides. Proc. Natl. Acad. Sci. USA 98, 6742–6746.

Engler, C., Gruetzmacher, R., Kandzia, R., and Marillonnet, S. (2009). Golden gate shuffling: a one-pot DNA shuffling method based on type II restriction enzymes. PLoS One 4, e5553.

Gaudelli, N.M., Komor, A.C., Rees, H.A., Packer, M.S., Badran, A.H., Bryson, D.I., and Liu, D.R. (2017). Programmable base editing of A-T to G-C in genomic DNA without DNA cleavage. Nature 551, 464–471.

Ghanta, K.S., Dakshin, G.A., Mir, A., Krishnamurthy, P.M., Greid, H., Edraki, A., Watts, J.K., Sontheimer, E.J., and Mello, C.C. (2018). 5' modifications improve potency and efficacy of DNA donors for precision genome editing. bioRxiv. https://doi.org/10.1101/354480.

Gibson, D.G., Young, L., Chuang, R.Y., Venter, J.C., Hutchison, C.A., III, and Smith, H.O. (2009). Enzymatic assembly of DNA molecules up to several hundred kilobases. Nat. Methods 6, 343–345.

Gross, H., and Loper, J.E. (2009). Genomics of secondary metabolite production by Pseudomonas spp. Nat. Prod. Rep. 26, 1408–1446.

Gu, T., Zhao, S., Pi, Y., Chen, W., Chen, C., Liu, Q., Li, M., Han, D., and Ji, Q. (2018). Highly efficient base editing in Staphylococcus aureus by an engineered CRISPR RNA-guided cytidine deaminase. Chem. Sci. 9, 3248–3253.

Hauser, A.R., Jain, M., Barmer, M., and McCollery, S.A. (2011). Clinical significance of microbial infection and adaptation in cystic fibrosis. Clin. Microbiol. Rev. 24, 29–70.

Hmelo, L.R., Borle, B.R., Almblad, H., Love, M.E., Randall, T.E., Tseng, B.S., Lin, C., Irie, Y., Storek, K.M., and Yang, J.J. (2015). Precision-engineering the Pseudomonas aeruginosa genome with two-step allelic exchange. Nat. Protoc. 10, 1820–1841.

Hoang, T.T., Karkhoff-Schweizer, R.R., Kutchma, A.J., and Schweizer, H.P. (1998). A broad-host-range Flp-FRT recombination system for site-specific excision of chromosomally-located DNA sequences: application for isolation of unmarked Pseudomonas aeruginosa mutants. Gene 212, 77–86.

Horvath, P., and Barrangou, R. (2010). CRISPR/Cas, the immune system of bacteria and archaea. Science 327, 167–170.

Jiang, W., Bikard, D., Cox, D., Zhang, F., and Marraffini, L.A. (2013). RNA-guided editing of bacterial genomes using CRISPR-Cas systems. Nat. Biotechnol. 31, 233–239.

Jiang, W., Brueggeman, A.J., Horken, K.M., Pluncik, T.M., and Weeks, D.P. (2014). Successful transient expression of Cas9 and single guide RNA genes in Chlamydomonas reinhardtii. Eukaryot. Cell 13, 1465–1469.

Jiang, Y., Chen, B., Duan, C., Sun, B., Yang, J., and Yang, S. (2015). Multigene editing in the Escherichia coli genome via the CRISPR-Cas9 system. Appl. Environ. Microbiol. 81, 2506–2514.

Jiang, Y., Qian, F., Yang, J., Liu, Y., Dong, F., Xu, C., Sun, B., Chen, B., Xu, X., Li, Y., et al. (2017). CRISPR-Cpf1 assisted genome editing of Corynebacterium glutamicum: Nat. Commun. 8, 15179.

Jinek, M., Chylinski, K., Fonfara, I., Hauer, M., Doudna, J.A., and Charpentier, E. (2012). A programmable dual-RNA-guided DNA endonuclease in adaptive bacterial immunity. Science 337, 816–821.

Komor, A.C., Kim, Y.B., Packer, M.S., Zuris, J.A., and Liu, D.R. (2016). Programmable editing of a target base in genomic DNA without double-stranded DNA cleavage. Nature 533, 420–424.

Li, X., Wang, Y., Liu, Y., Yang, B., Wang, X., Wei, J., Lu, Z., Zhang, Y., Wu, J., Huang, X., et al. (2018). Base editing with a Cpf1-cytidine deaminase fusion. Nat. Biotechnol. 36, 324–327.

Liang, R., and Liu, J. (2010). Scarless and sequential gene modification in Pseudomonas using PCR product flank by short homology regions. BMC Microbiol. 10, 209.

Loeschcke, A., and Thies, S. (2015). Pseudomonas putida-a versatile host for the production of natural products. Appl. Microbiol. Biotechnol. 99, 6197–6214.
Supplemental Information

CRISPR/Cas9-based Genome Editing in

*Pseudomonas aeruginosa* and Cytidine Deaminase-

Mediated Base Editing in *Pseudomonas* Species

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**Transparent Methods**

**Bacterial Strains, Plasmids, Primers and Growth Conditions**
The strains, plasmids, and primers used in this study were listed in tables S2, S3 and S4, respectively. All *E. coli* strains and *Pseudomonas* strains were grown in Luria-Bertani broth (LB). *E. coli* strains and *P. aeruginosa* strains were grown at 37 °C, while *P. putida*, *P. fluorescens*, and *P. syringae* were cultured at 30 °C.

**Plasmid Construction**
The construction of the pCasPA plasmid was performed using the following method. The λ-Red system and the *araB* promoter (P*araB*) were amplified from the pKD46 plasmid, respectively (Datsenko and Wanner, 2000). The gene encoding the Cas9 nuclease was amplified from the pCasSA plasmid (Chen et al., 2017). These three DNA fragments were assembled between the *Kpn*I/*Hind*III sites of the pDN19 plasmid (Li et al., 2013) using Gibson assembly. Next, the *sacB* gene amplified from the pEX18Ap plasmid (Hoang et al., 1998) was inserted into the *Xba*I/*Xho*I sites of the aforementioned plasmid, resulting in the final plasmid pCasPA. The pACRISPR plasmid was constructed by inserting the *trc* promoter along with the sgRNA and the *sacB* gene into the *BamHI*/*Hind*III sites of the pAK1900 vector (Srikumar et al., 1998) via Gibson assembly.

The pnCas9PA-BEC plasmid was constructed using the following procedures. The *rpsL* promoter was amplified from the genomic DNA of the *P. aeruginosa* PAO1 strain. The origin ColE1 and the gene encoding APOBEC1-Cas9D10A were amplified from the pnCasSA-BEC plasmid (Gu et al., 2018). The *sacB* gene was amplified from the pEX18Ap plasmid (Hoang et al., 1998). The origin mSF was amplified from pAK1900 plasmid (Srikumar et al., 1998). The gentamycin-resistance marker was amplified from the pPS858 plasmid (Hoang et al., 1998). The six fragments were assembled into a plasmid via Golden Gate assembly. The *trc* promoter along with the sgRNA was amplified from the pACRISPR plasmid and inserted into the *SalI*/*Xho*I sites of the aforementioned plasmid, resulting in the final pnCasPA-BEC plasmid.

**Preparation of Electrocompetent Cells**
1 mL overnight culture of *P. aeruginosa* was 1:100 diluted into 100 mL of fresh LB medium and incubated at 37 °C. When the optical density at 600 nm (OD₆₀₀) of the culture reached ~1.0, the cells were harvested by centrifugation at 6000 rpm for 5 min. The supernatant was discarded, and the cells were washed twice with 20 mL of sterile ice-cold 10% v/v glycerol. Finally, the cells were resuspended into 1 mL of 10% v/v glycerol. 50 μL aliquots of the cells were used for the subsequent experiments. Similar procedures as that of the preparation of electrocompetent *P. aeruginosa* cells were utilized to make *P. putida*, *P. fluorescens*, and *P. syringae* electrocompetent cells.
Genome Editing Using the pCasPA/pACRISPR System
The pCasPA plasmid was transformed into *P. aeruginosa* strains using electroporation with the parameters of 2100 V, 100 Ω, 25 μF, 1 mm cuvette (Bio-Rad). The colonies containing pCasPA were selected on a LB agar plate in the presence of 100 μg/mL tetracycline. A colony was picked from the plate and cultured in the LB medium at 37 °C overnight. The next day, 1 mL culture was diluted into 100 mL fresh LB medium and shaken at 37 °C until the OD$_{600}$ of 1.0~1.5. The expression of the Cas9 nuclease and the λ-Red system was induced by the addition of L-arabinose to a final concentration of 2 mg/mL. The culture was further incubated for 2 hours before being prepared as the electrocompetent cells. Next, the pACRISPR plasmid assembled with the spacer and the repair template was electroporated into the electrocompetent cells. After electroporation, the cells were recovered in LB for 1~2 hours at 37 °C and plated onto the LB agar plate containing 100 μg/mL tetracycline and 150 μg/mL carbenicillin. PCR and sequencing were utilized to evaluate the editing efficiency. The editing efficiency was calculated as the number of colonies mutated successfully divided by the number of all the colonies evaluated.

Base Editing Using the pnCasPA-BEC Plasmid
For the base editing in the *P. aeruginosa* strains, a suitable 20-bp spacer sequence before a PAM site was designed and inserted into the *BsaI* sites of the pnCasPA-BEC plasmid using Golden Gate assembly. The constructed plasmid was then transferred into *P. aeruginosa* strains and the colonies were selected on the LB agar plate containing 30 μg/mL gentamycin. The PCR reaction was performed individually using the genomic DNA of colonies as the template, and the PCR products covering the editable site(s) were sent out for sequencing. The base editing of other *Pseudomonas* spp. such as *P. putida*, *P. fluorescens*, and *P. syringae* were performed following a similar procedure, except that these three strains were selected on the LB plate containing 20 μg/mL gentamycin.

Pigment Assay
Different colonies were randomly picked and cultured in 3 mL pyocyanin production broth (PPB) medium (PPB medium: 20 g peptone, 1.4 g MgCl$_2$, 10 g K$_2$SO$_4$, 20 mL glycerol per liter; pH 7.0)(Brint and Ohman, 1995). One colony of the PAO1 wild-type strain was incubated in the same condition as a control. The tubes were shaken at 37 °C for 12~16 h before being photographed.

Data and Software Availability
The plasmids have been deposited in the Addgene (http://www.addgene.org/) with the accession codes 113347 (pCasPA), 113348 (pACRISPR) and 113348 (pnCasPA-BEC).
Detailed protocol for the construction of the pACRISPR plasmid.

(1) Spacer design

Design a suitable 20 nt DNA sequence in the target gene of *P. aeruginosa* via sgRNAcas9 (Xie et al., 2014). Two oligos were designed in the following form:

5’-GTGGNNNNNRRRRRRRRRRRRRRRR-3’

3’-NNNNNNNNRRRRRRRRRRRRRRRRRCAAA-5’

(2) Phosphorylation of the oligos

| Component                                | Volume |
|------------------------------------------|--------|
| 10× T4 DNA ligase buffer (NEB)           | 5 μL   |
| 50 μM Oligo F                           | 2 μL   |
| 50 μM Oligo R                           | 2 μL   |
| T4 polynucleotide kinase (Takara)       | 1 μL   |
| ddH₂O                                    | 40 μL  |
| **Total volume**                         | **50 μL** |

Incubate at 37 °C for 1 hour.

(3) Annealing

2.5 μL 1 M NaCl was added into the phosphorylated product. The phosphorylated oligos were incubated at 95 °C for 3 min and then slowly cooled down to room temperature using a thermocycler. Dilute the annealed oligos 20 folds to the final concentration of 100 nM with ddH₂O.

(4) Golden Gate assembly

Prepare the samples as indicated below:

| Component                                | Volume |
|------------------------------------------|--------|
| 10× T4 DNA ligase buffer (NEB)           | 1 μL   |
| Annealed oligos (100 nM)                | 1 μL   |
| The pACRISPR plasmid (20 nM)            | 1 μL   |
| T4 DNA ligase (NEB)                     | 0.5 μL |
| BsaI-HF (NEB)                           | 0.5 μL |
| ddH₂O                                   | 6 μL   |
| **Total volume**                         | **10 μL** |

Run the reaction protocol in a thermocycler:

| Segment | Temperature | Time  | Cycles |
|---------|-------------|-------|--------|
| 1       | 37 °C       | 3 min | 25     |
|         | 16 °C       | 4 min |        |
| 2       | 80 °C       | 15 min| 1      |
| 3       | 10 °C       | Forever| 1      |

(5) Transformation

The Golden Gate assembly product was transformed into 100 μL *E.coli* DH5α competent cells. The cells were plated onto a LB agar plate containing 50 μg/mL carbenicillin and incubated at 37 °C overnight. The success for the construction of the pACRISPR-NN_spacer plasmid was
verified by PCR and sequencing.

(6) Digest the pACRISPR-NN_spacer plasmid with XbaI and XhoI

| Component                          | Volume |
|------------------------------------|--------|
| 10× CutSmart buffer (NEB)          | 5 μL   |
| 2 μg pACRISPR-NN_spacer plasmid    | xx μL  |
| XbaI (NEB)                         | 2 μL   |
| XhoI (NEB)                         | 2 μL   |
| ddH2O                              | xx μL  |
| Total volume                       | 50 μL  |

Incubate the mixture at 37 °C for 2~3 hours. The digested plasmid was purified by using the SanPrep PCR purification kit (Sangon Biotech, Shanghai, China).

(7) Gibson assembly

Select ~500bp DNA sequence of the upstream and downstream of the target gene, respectively. The two DNA fragments were PCR-amplified from the genomic DNA of the *p. aeruginosa* strains with a 20~40 bp overlap of the digested pACRISPR-NN_spacer plasmid. The overlap could be created via PCR with primers that contain a 5’ end that is identical to an adjacent segment and a 3’ end that anneals to the target sequence.

5’ primer of the upstream is in this form:

5’- tttagatctgtccatacccatggTCTAGANNNNNNNNNNNNNNNNNNNNNNNNN -3’

3’ primer of the downstream is in this form:

5’-tctgaatggcgggagtatgaaaagtCTCGAGNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNN -3’

3’ primer of the upstream and 5’ primer of the downstream should have 30~40 bp overlap between each other.

The two DNA fragments were assembled into the digested pACRISPR-NN_spacer plasmid using Gibson assembly.

| Component                          | Volume |
|------------------------------------|--------|
| NEBuilder HiFi DNA Assembly Master Mix (NEB) | 10 μL   |
| 20 fmol XbaI/XhoI digested pACRISPR-NN_spacer plasmid | xx μL  |
| 20 fmol upstream of the target gene | xx μL  |
| 20 fmol downstream of the target gene | xx μL  |
| ddH2O                              | xx μL  |
| Total volume                       | 20 μL  |

The reaction solution was incubated at 50 °C for 1 hour.

For the gene insertion, external 20 fmol DNA fragment of the trc promoter or rpsL promoter was added into the reaction solution.

The Gibson assembly product was transformed into 100 μL *E.coli* DH5α competent cells. The cells were plated onto a LB agar plate containing 50 μg/mL carbenicillin. The successful construction of the plasmid pACRISPR-NN was verified by PCR and sequencing.

The construction of pnCasPA-BEC_spacer followed the same protocol of steps 1~4. The plasmid was selected on the LB agar plate containing 15 μg/mL gentamycin.
Figure S1. The single plasmid system pCasPAGm and pCasPAGm-Red deleted genes in *P. aeruginosa* with different efficiencies, related to Figure 1. (A) Map of the one-plasmid genome editing plasmid pCasPAGm. *trc* promoter, the sgRNA expression promoter; *rpsL* promoter, the Cas9 protein expression promoter; *Bsa*I sites, Golden Gate assembly of spacers; *Xba*I and *Xho*I sites, Gibson assembly of repair arms; mSF, A broad-host-range origin amplified from pAK1900 vector; ColE1, a replication origin for *E. coli*; Gm, the gentamycin-resistance marker in *E. coli* and *P. aeruginosa*; sacB, the counter-selectable marker for fast curing of the plasmid after editing. (B) The pCasPAGm system enabled the deletion of *wspF* gene in the PAO1 strain with the efficiency of 9/11. (C) The pCasPAGm system failed to delete *nalD* gene in the PAO1 strain. (D) The pCasPAGm system failed to delete *rhlR* gene in the PAO1 strain. (E) Map of the pCasPAGm-Red plasmid. The plasmid was constructed by the insertion of the λ-Red system into the pCasPAGm plasmid. (F) The pCasPAGm-Red system failed to delete the *nalD* gene in the PAO1 strain.
Figure S2. The pCasPA/pACRISPR system-mediated genome editing in the *P. aeruginosa* PAO1, related to Figure 3. (A) 1 μg pACRISPR (empty plasmid), pACRISPR-*rhlR*_spacer (assembled with a 20-nt *rhlR* gene spacer), pACRISPR-*rhlR*_repair (assembled with the *rhlR* gene repair arms), and pACRISPR-*rhlR* (assembled with both spacer and repair arms) plasmid was electroporated into the PAO1 strain containing pCasPA after L-arabinose induction, respectively. The pACRISPR-*rhlR* plasmid was also transformed into the pCasPA-containing PAO1 strain without arabinose induction as a control. (B) The
editing efficiency varies when different lengths of circular repair templates (assembled into the pACRISPR plasmid) were utilized for *nalD* gene deletion in the PAO1 strain. (C) The editing efficiency varies when different lengths of linear repair templates were utilized for *nalD* gene deletion in the PAO1 strain.
Figure S3. The pCasPA/pACRISPR system enabled highly efficient Genome editing in the P. aeruginosa PAO1 and PAK strains, and can be easily cured after editing, related to Figure 3. (A) pCasPA/pACRISPR-mediated deletion of the rsaL, algR and rhlB genes in the PAO1 strain with the efficiencies of 12/12, 5/12 and 8/12, respectively. The “ck” lane was the PCR product from the wild-type strain as a control. (B) The pCasPA/pACRISPR system successfully deleted 3kb DNA fragment in the rhlR gene locus with the efficiency of 12/12, while it failed in the deletion of 5kb and 10kb DNA fragments in the same locus. (C) The pCasPA/pACRISPR system enabled
multiplexed gene deletion. The *rhlR* and *lasR* genes were deleted simultaneously in a single transformation with the efficiency of 10/12. (D) The transformation CFUs of multiplexed gene deletion. Only 10~20 colonies were observed on the LB plate after electroporation with 1 μg of the pACRISPR-*rhlR-lasR* plasmid. (E) The sgRNA locus of the four escaped transformants from the PAO1ΔnalD plate could not be amplified. The “ck” lane was the PCR product from the mutant that was successfully edited as a control. (F) pCasPA/pACRISPR-mediated deletion of the *rsaL, algR* and *lasR* genes in the PAK strain with the efficiencies of 10/12, 8/12 and 6/12, respectively. (G) Much fewer colonies grew on the plate in the presence of sucrose (right) than that grew on the plate without sucrose (left). (H) The colonies picked from the plate containing sucrose failed to grow on the plates containing 100 μg/mL tetracycline (middle) or 150 μg/mL carbenicillin (right). Colonies only grew on the plate without antibiotics, indicating both the pCasPA and the pACRISPR plasmids were successfully cured.
Figure S4. The C→T base editing plasmid pnCasPA-BEC in *P. aeruginosa*, related to Figure 4. (A) Map of the C→T base editing plasmid pnCasPA-BEC. APOBEC1-nCas9, a fusion protein composed of a deaminase APOBEC1 at the N terminus and a Cas9 nuclease Cas9D10A at the C terminus; trc promoter, a strong promoter to drive the expression of the sgRNA; rpsL promoter, the APOBEC1-nCas9 protein expression promoter; BsaI sites, Golden Gate assembly of spacers; mSF, A broad-host-range origin from pAK1900 vector; ColE1, a replication origin for *E. coli*; Gm, the gentamycin-resistance marker in *E. coli* and *P. aeruginosa*; sacB, the counter-selectable marker for fast curing of the plasmid after editing. (B) The *rhlR* gene of the PAO1 strain was effectively inactivated by the pnCasPA-BEC system. The mutation efficiency of *rhlR* W108 to stop codon was 11/12 confirmed by pigment assay. (C) The pnCasPA-BEC plasmid could be easily cured after editing. Six individual colonies picked from the plate containing sucrose were streaked onto the LB agar plates in the presence (right) or absence (left) of gentamycin. Colonies only grew on the plate without gentamycin, indicating the pnCasPA-BEC plasmid was successfully cured. (D) Examination of the editable window of the pnCasPA-BEC system. A representative sequencing chromatogram for the target locus was shown. The C(s) in the spacer were colored red, and the editable C(s) were indicated with red triangles. (E) The kind of adjacent bases of editable sites affected the editing efficiency greatly.
Table S1. The sequence and position of the spacer of *rhlR* and *rhlB* genes and their top six similar spacer sites, related to Figure 4.

| Gene | Site | Sequence | Position     | Mutation efficiency |
|------|------|----------|--------------|---------------------|
| rhlB | WT   | cgttcaggeagcgcggcgcgtgg | 3891315-3891293 | 11/12               |
|      | M1   | cCataTgggGgceggcgcgtgg | 841017-841039  | 0/4                 |
|      | M2   | cgtcCgCgcagccTgcgctgg | 3984809-3984831| 0/4                 |
|      | M3   | cgtcagggTCgceggAcAgctgg| 1565062-1565040| 0/4                 |
|      | M4   | cgtcaCgceagcegCcACctgg | 4873175-4873197| 0/4                 |
|      | M5   | cgtcCgCcagccTgcgGCgg  | 5005635-5005613| 0/4                 |
|      | M6   | TgatcaggTcGgceggTgcegg| 5534902-5534924| 0/4                 |
| rhlR | WT   | cgttcagcagcatcgcgttgg  | 3890323-3890345| 11/12               |
|      | M1   | cgttccagaCcatcAgcGCAGg | 743768-743790  | 0/4                 |
|      | M2   | cgtCtcGCgcatcgcgcgtgg | 1905399-1905421| 0/4                 |
|      | M3   | cgtGCcTGagcaCcegctgcgg | 5090604-5090626| 0/4                 |
|      | M4   | cgtCGTagTgcatcgcgcAcctgg| 591063-591085  | 0/4                 |
|      | M5   | cgttcGagaTeCtcGgcttg  | 2514335-2514357| 0/4                 |
|      | M6   | cgttcTGgCgcaGceggcGctgg| 2737948-2737926| 0/4                 |

The mismatched bases are labeled with the thicker capital letter.
Table S2. Bacterial strains used in this study. Related to Fig. 1, Fig. 2, Fig. 3, Fig.4, Fig. 5, and Transparent Methods.

| Strains | Description | Reference |
|---------|-------------|-----------|
| **E. coli** | | |
| DH5α | F^- Φ80lacZΔM15 Δ(lacZYA-argF) U169 recA1 endA1 hsdR17 (rK-, mK+) phoA supE44 λ- thi-1 gyrA96 relA1 | Lab stock |
| **P. aeruginosa strain** | | |
| PAO1 | Wild-type | Lab stock |
| PAO1ΔrhlR | PAO1 rhlR gene deleted | This study |
| PAO1ΔnalD | PAO1 nalD gene deleted | This study |
| PAO1ΔarsaL | PAO1 rsaL gene deleted | This study |
| PAO1ΔalgR | PAO1 algR gene deleted | This study |
| PAO1ΔrhlB | PAO1 rhlB gene deleted | This study |
| PAO1ΔlasR | PAO1 lasR gene deleted | This study |
| PAO1ΔlasR-rhlR | PAO1 lasR and rhlR genes both deleted | This study |
| PAO1ΔrhlR3k | PAO1 3kb DNA fragment deleted in the rhlR gene locus | This study |
| PAO1-trc | PAO1 trc promoter insertion upstream of the rhlA gene | This study |
| PAO1-epsL | PAO1 epsL promoter insertion upstream of the rhlA gene | This study |
| PAO1rhlR W108 to stop | PAO1 rhlR W108 mutation to stop codon | This study |
| PAO1rhlB Q249 to stop | PAO1 rhlB Q249 mutation to stop codon | This study |
| PAK | Wild-type | (Li et al., 2013) |
| PAKΔarsaL | PAK rsaL gene deleted | This study |
| PAKΔalgR | PAK algR gene deleted | This study |
| PAKΔlasR | PAK lasR gene deleted | This study |
| PAKrhlR W108 to stop | PAK rhlR W108 mutation to stop codon | This study |
| PAKrhlB Q249 to stop | PAK rhlB Q249 mutation to stop codon | This study |
| **Other Pseudomonas spp.** | | |
| P. putida KT2440 | Wild type | Lab stock |
| KT2440cadR Q92 to stop | P. putida KT2440 cadR Q92 mutation to stop codon | This study |
| KT2440ompR Q129 to stop | P. putida KT2440 ompR Q129 mutation to stop codon | This study |
| P. fluorescens GcM5-1A | Wild type | (Feng et al., 2015) |
| GcM5-1Aper Q374 to stop | P. fluorescens GcM5-1A per Q374 mutation to stop codon | This study |
| GcM5-1AspC Q164 to stop | P. fluorescens GcM5-1A aspC Q164 mutation to stop codon | This study |
| P. syringae DC3000 | Wild type | Lab stock |
| DC3000gacA Q113 to stop | P. syringae DC3000 gacA Q113 mutation to stop codon | This study |
| DC3000hrpL Q29 to stop | P. syringae DC3000 hrpL Q29 mutation to stop codon | This study |
Table S3. Plasmids used in this study. Related to Fig. 1, Fig. 2, Fig. 3, Fig. 4, Fig. 5, and Transparent Methods.

| Plasmids         | Description                                                                 | Reference                                |
|------------------|-----------------------------------------------------------------------------|------------------------------------------|
| pAK1900          | Ap’, broad-host-range cloning vector                                         | (Srikumar et al., 1998)                  |
| pDN19            | Tc’, *E. coli-P. aeruginosa* shuttle cloning vector                          | (Li et al., 2013)                        |
| pEX18Ap          | Ap’, oriT+ sacB+, gene replacement vector with MCS from pUC18               | (Hoang et al., 1998)                     |
| pPS858           | Ap’, Gm’; the vector containing gentamycin-resistance marker                | (Hoang et al., 1998)                     |
| pCasSA           | Km’, Cm’; the *S. aureus* genome editing vector                              | (Chen et al., 2017)                      |
| pACRISPR         | Ap’, a sgRNA expression plasmid for targeting a specific sequence           | This study                               |
| pACRISPR-rhlRSpacer      | pACRISPR derivative with the spacer of the *rhlR* gene                     | This study                               |
| pACRISPR-rhlRRepair          | pACRISPR derivative with the repair arms of the *rhlR* gene                 | This study                               |
| pACRISPR-rhlR       | pACRISPR derivative for *rhlR* deletion                                     | This study                               |
| pACRISPR-nalDSpacer      | pACRISPR derivative with *nalD* spacer                                      | This study                               |
| pACRISPR-nalD-45bp     | pACRISPR derivative for *nalD* deletion with 45bp+45bp repair               | This study                               |
| pACRISPR-nalD-100bp    | pACRISPR derivative for *nalD* deletion with 100bp+100bp repair             | This study                               |
| pACRISPR-nalD-200bp    | pACRISPR derivative for *nalD* deletion with 200bp+200bp repair             | This study                               |
| pACRISPR-nalD-500bp    | pACRISPR derivative for *nalD* deletion with 500bp+500bp repair             | This study                               |
| pACRISPR-nalD-1kb      | pACRISPR derivative for *nalD* deletion with 1kb+1kb repair                 | This study                               |
| pACRISPR-rsal        | pACRISPR derivative for *rsal* deletion                                      | This study                               |
| pACRISPR-algR        | pACRISPR derivative for *algR* deletion                                      | This study                               |
| pACRISPR-rhlB        | pACRISPR derivative for *rhlB* deletion                                      | This study                               |
| pACRISPR-lasR        | pACRISPR derivative for *lasR* deletion                                      | This study                               |
| pACRISPR-rhlR3k      | pACRISPR derivative for deletion of 3kb fragment in *rhlR* locus            | This study                               |
| pACRISPR-rhlR5k      | pACRISPR derivative for deletion of 5kb fragment in *rhlR* locus            | This study                               |
| pACRISPR-rhlR10k     | pACRISPR derivative for deletion of 10kb fragment in *rhlR* locus           | This study                               |
| pACRISPR-lasR-rhlR    | pACRISPR derivative for deletion of *lasR* and *rhlR* genes                 | This study                               |
| pACRISPR-trc         | pACRISPR derivative for *trc* promoter insertion                            | This study                               |
| pACRISPR-rpsL        | pACRISPR derivative for *rpsL* promoter insertion                            | This study                               |
| pnCasPA-SEC         | Gm’, *Pseudomonas* spp. genome base-editing vector                          | This study                               |
| pnCasPA-SEC-rhlRsp   | pnCasPA-SEC derivative with *rhlR* spacer, W108 mutation to stop codon       | This study                               |
| pnCasPA-SEC-rhlBsp   | pnCasPA-SEC derivative with *rhlB* spacer, Q249 mutation to stop codon       | This study                               |
| pnCasPA-SEC-C2sp     | pnCasPA-SEC derivative containing spacer with C at position 2               | This study                               |
| pnCasPA-SEC-C3sp     | pnCasPA-SEC derivative containing spacer with C at position 3               | This study                               |
| pnCasPA-SEC-C4sp     | pnCasPA-SEC derivative containing spacer with C at position 4               | This study                               |
| pnCasPA-SEC-C5sp     | pnCasPA-SEC derivative containing spacer with C at position 5               | This study                               |
| pnCasPA-SEC-C6sp     | pnCasPA-SEC derivative containing spacer with C at position 6               | This study                               |
| pnCasPA-BEC-C7sp         | pnCasPA-BEC derivative containing spacer with C at position 7 | This study |
|--------------------------|---------------------------------------------------------------|------------|
| pnCasPA-BEC-C8sp         | pnCasPA-BEC derivative containing spacer with C at position 8 | This study |
| pnCasPA-BEC-C9sp         | pnCasPA-BEC derivative containing spacer with C at position 9 | This study |
| pnCasPA-BEC-GC4          | pnCasPA-BEC derivative containing spacer with GC at position 4 | This study |
| pnCasPA-BEC-GC5AC8       | pnCasPA-BEC derivative containing spacer with GC at position 5 and AC at position 8 | This study |
| pnCasPA-BEC-GC5          | pnCasPA-BEC derivative containing spacer with GC at position 5 | This study |
| pnCasPA-BEC-AC5          | pnCasPA-BEC derivative containing spacer with AC at position 5 | This study |
| pnCasPA-BEC-cadRsp       | pnCasPA-BEC derivative with cadR spacer, Q92 mutation to stop codon | This study |
| pnCasPA-BEC-ompRsp       | pnCasPA-BEC derivative with ompR spacer, Q129 mutation to stop codon | This study |
| pnCasPA-BEC-persp        | pnCasPA-BEC derivative with per spacer, Q374 mutation to stop codon | This study |
| pnCasPA-BEC-aspCsp       | pnCasPA-BEC derivative with ompR spacer, Q164 mutation to stop codon | This study |
| pnCasPA-BEC-gacAsp       | pnCasPA-BEC derivative with gacA spacer, Q113 mutation to stop codon | This study |
| pnCasPA-BEC-hrpLsp       | pnCasPA-BEC derivative with hrpL spacer, Q29 mutation to stop codon | This study |

Ap', ampicillin resistant; Tc', tetracycline resistant; Gm', gentamycin resistant; Km', kanamycin resistant; Cm', chloramphenicol resistant.
REFERENCES

Brint, J.M., and Ohman, D.E. (1995). Synthesis of multiple exoproducts in *Pseudomonas aeruginosa* is under the control of RhlR-RhlII, another set of regulators in strain PA01 with homology to the autoinducer-responsive LuxR-LuxI family. J. Bacteriol. 177, 7155-7163.

Chen, W., Zhang, Y., Yeo, W.S., Bae, T., and Ji, Q. (2017). Rapid and efficient genome editing in *Staphylococcus aureus* by using an engineered CRISPR/Cas9 system. J. Am. Chem. Soc. 139, 3790-3795.

Datsenko, K.A., and Wanner, B.L. (2000). One-step inactivation of chromosomal genes in *Escherichia coli* K-12 using PCR products. Proc. Natl. Acad. Sci. U. S. A. 97, 6640-6645.

Feng, K., Li, R., Chen, Y., Zhao, B., and Yin, T. (2015). Sequencing and analysis of the *Pseudomonas fluorescens* GcM5-1A genome: a pathogen living in the surface coat of *Bursaphelenchus xylophilus*. PLoS One 10, e0141515.

Gu, T., Zhao, S., Pi, Y., Chen, W., Chen, C., Liu, Q., Li, M., Han, D., and Ji, Q. (2018). Highly efficient base editing in *Staphylococcus aureus* using an engineered CRISPR RNA-guided cytidine deaminase. Chem. Sci. 9, 3248-3253.

Hoang, T.T., Karkhoff-Schweizer, R.R., Kutchma, A.J., and Schweizer, H.P. (1998). A broad-host-range Flp-FRT recombination system for site-specific excision of chromosomally-located DNA sequences: application for isolation of unmarked *Pseudomonas aeruginosa* mutants. Gene 212, 77-86.

Li, K., Xu, C., Jin, Y., Sun, Z., Liu, C., Shi, J., Chen, G., Chen, R., Jin, S., and Wu, W. (2013). SuhB is a regulator of multiple virulence genes and essential for pathogenesis of *Pseudomonas aeruginosa*. Mbio 4, e00419.

Srikumar, R., Kon, T., Gotoh, N., and Poole, K. (1998). Expression of *Pseudomonas aeruginosa* multidrug efflux pumps MexA-MexB-OprM and MexC-MexD-OprJ in a multidrug-sensitive *Escherichia coli* strain. Antimicrob. Agents Chemother. 42, 65-71.

Xie, S., Shen, B., Zhang, C., Huang, X., and Zhang, Y. (2014). sgRNAcas9: a software package for designing CRISPR sgRNA and evaluating potential off-target cleavage sites. PLoS One 9, e100448.