Efficient transposon mutagenesis mediated by an IPTG-controlled conditional suicide plasmid

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Abstract

Background: Transposon mutagenesis is highly valuable for bacterial genetic and genomic studies. The transposons are usually delivered into host cells through conjugation or electroporation of a suicide plasmid. However, many bacterial species cannot be efficiently conjugated or transformed for transposon saturation mutagenesis. For this reason, temperature-sensitive (ts) plasmids have also been developed for transposon mutagenesis, but prolonged incubation at high temperatures to induce ts plasmid loss can be harmful to the hosts and lead to enrichment of mutants with adaptive genetic changes. In addition, the ts phenotype of a plasmid is often strain- or species-specific, as it may become non-ts or suicidal in different bacterial species.

Results: We have engineered several conditional suicide plasmids that have a broad host range and whose loss is IPTG-controlled. One construct, which has the highest stability in the absence of IPTG induction, was then used as a curable vector to deliver hyperactive miniTn5 transposons for insertional mutagenesis. Our analyses show that these new tools can be used for efficient and regulatable transposon mutagenesis in Escherichia coli, Acinetobacter baylyi and Pseudomonas aeruginosa. In P. aeruginosa PAO1, we have used this method to generate a Tn5 insertion library with an estimated diversity of ~ 10^8, which is ~ 2 logs larger than the best transposon insertional library of PAO1 and related Pseudomonas strains previously reported.

Conclusion: We have developed a number of IPTG-controlled conditional suicide plasmids. By exploiting one of them for transposon delivery, a highly efficient and broadly useful mutagenesis system has been developed. As the assay condition is mild, we believe that our methodology will have broad applications in microbiology research.

Keywords: Transposon mutagenesis, IPTG, Conditional suicide plasmid, Escherichia coli, Pseudomonas, Acinetobacter

Background

Transposon mutagenesis is a powerful technique for bacterial genetic and genomic studies. One of the most widely used transposons is derived from Tn5. The Tn5 transposon contains two IS50 elements as inverted terminal repeats (Additional file 1: Figure S1) [1, 2]. Both IS50 and Tn5 can be mobilized by their encoded transposase (Tnp) protein, which recognizes two 19 base pair (bp) sequences at their ends, namely outside end (OE) and inside end (IE), for transposition [2]. OE and IE differ by 7 bp (Additional file 1: Figure S1). As Tn5 insertion is almost completely random, it can insert into any gene in a bacterium. The native Tn5/IS50 is not very active, thus avoiding overt deleterious effect on their hosts, but hyperactive mutants have been engineered as genetic manipulation tools [2, 3]. The most active one contains a mosaic sequence of OE and IE (mosaic end; ME) at the transposon termini and an engineered tnp gene encoding a highly active transposase enzyme (TnpH), which together increase Tn5 transposition by more than 1000-fold.

Transposons for insertional mutagenesis are usually delivered into bacteria through conjugation of a suicide plasmid [4–6]. Insertion mutants are then selected as
the transposons are tagged with an antibiotic-resistance gene. The success of a transposon mutagenesis assay, especially a saturation mutagenesis assay, requires generation of an insertion library with high diversity, which requires efficient plasmid conjugation and transposon transposition. However, conjugation is inefficient in many bacterial species. Occasionally, electroporation has also been used to deliver transposon-containing suicide plasmids for mutagenesis, but low library diversities were often achieved using such approaches [7–9]. To perform efficient transposon mutagenesis in these organisms, temperature-sensitive (ts) plasmids are sometimes used for transposon delivery [10–16]. However, many organisms do not have a ts and easily manipulatable plasmid, and sometimes a ts plasmid in one organism is either non-ts or suicidal in a different organism [10, 14, 17]. In addition, a high temperature is often required to cure the ts plasmids after mutagenesis, which can be inhibitory to cell growth and may result in selection of mutants with adaptive genetic changes [10, 11, 14].

In this study, we have developed an efficient and regulatable transposon mutagenesis tool that exploits an IPTG-controlled conditional suicide plasmid. It contains an RSF1010 replicon, an IncQ-type replication origin that allows plasmid replication in most Gram-negative bacteria, as well as a few Gram-positive bacteria [18]. It is relatively small, so it can be easily modified. To control plasmid replication by IPTG, a second copy of the plasmid-encoded repF repressor gene is cloned downstream of the Escherichia coli tac promoter. For efficient and regulatable transposon mutagenesis, we used miniTn5 (mTn5) transposons and cloned the hyperactive transposase gene downstream of a lac promoter. We show that the resulting constructs can be used for efficient insertional mutagenesis in three different bacterial species. In Pseudomonas aeruginosa PAO1, we show that our system is able to generate a Tn5 insertion library that is almost 2 logs larger than the best library of PAO1 and related Pseudomonas strains previously reported, demonstrating that we have developed a powerful mutagenesis tool that is highly useful for microbiology studies.

Results

Construction of IPTG-controlled suicide plasmids

To develop a method for efficient transposon mutagenesis in bacterial species that are difficult to transform and conjugate, we created multiple IPTG-controlled suicide plasmids that have a broad host range (Fig. 1a). The plasmids were derived from pMMB208, which is a conjugatable plasmid containing an RSF1010 oriV (an IncQ-type origin of replication) that can replicate in most Gram-negative bacteria and a few Gram-positive bacteria [18]. Plasmid replication requires three proteins, RepA, MobA/RepB and RepC, which are a helicase, a primase and an oriV-binding protein, respectively. repF encodes a small repressor protein that binds the P4 promoter and controls the repF-repA-repC operon through feedback inhibition [19, 20]. pMMB208 also contains a tac promoter (Pptac), a lacIQ gene and a chloramphenicol resistance marker (CamR). To create a conditional suicide plasmid (pMMB-repF), a second copy of the repF gene was inserted downstream of Ptac. Upon IPTG induction, efficient plasmid loss from transformed E. coli DH10B cells was observed (99.97%; Fig. 1b). As an alternative strategy, we inserted two repA helicase dominant negative mutants, K42A and D139A, downstream of Ptac [21]. Similarly, IPTG was able to induce efficient plasmid loss from the transformed DH10B cells. In fact, plasmid retention rates of the dominant negative mutants (K42A, 4.9 × 10−7; D139A, 1.5 × 10−5) were much lower than that of pMMB-repF (3.5 × 10−4) (Fig. 1b). However, the two repA dominant negative mutants showed significantly lower plasmid stability in the absence of IPTG induction (Fig. 1b), suggesting that plasmid replication is strongly inhibited by leaky expression of the dominant negative mutants, or by spontaneous recombination of the wild-type and the dominant negative repA genes (~ 861 bp direct repeats). Consistent with that, there were ~ 20–30-fold less plasmid isolated from the same amount of cells for the two mutant constructs (Fig. 1c). Therefore, we decided to choose pMMB-repF for further experiments. To kill the cells that still retain the plasmid after IPTG induction, we inserted a sacB counter selection marker into the vector [22], resulting in pMMB-repF/sacB. Indeed, insertion of sacB allows efficient killing of plasmid-containing cells by sucrose (data not shown; also see below).

IPTG-controlled mutagenesis of E. coli by a highly-active mTn5 transposon

A KanR-tagged mTn5 was then inserted in pMMB-repF/sacB for transposon mutagenesis (Fig. 2a) [4]. The mTn5 contains an OE and an IE at the termini. In addition, it contains an uncoupled, lac promoter (Plac)-controlled tnpH gene encoding the hyperactive transposase (TnpH) [3], thus allowing inducible expression of TnpH. E. coli cells transformed with this plasmid, pSNC-mTn5, were cultured in LB media with and without IPTG induction for 24 h. Cells were then analyzed for efficiencies of plasmid loss, sucrose counter selection and transposon insertion (See Methods). The plasmid is stable without IPTG induction, as ~ 91.4% of cells retained the plasmid (CamR) after 24 h culture in the absence of antibiotics (Fig. 2b). In contrast, ~ 2.6 × 10−3 of the cells retained the plasmid post IPTG induction, suggesting that overexpression of the RepF repressor caused efficient plasmid loss. Sucrose counter selection further reduced plasmid-bearing cells (~ 1.6 × 10−6 are CamR; ~ 1600-fold reduction). In
comparison, the percentage of Suc\textsuperscript{R}Kan\textsuperscript{R} cells after IPTG induction was found to be \(\sim 2.3 \times 10^{-4}\) (Tn5-containing), significantly higher than that of Suc\textsuperscript{R}Cam\textsuperscript{R} cells (\(\sim 1.6 \times 10^{-6}\), plasmid-containing), suggesting that Tn5 transposition had occurred efficiently (Suc\textsuperscript{R}Kan\textsuperscript{R}Cam\textsuperscript{S} \(\sim 2.3 \times 10^{-4}\)). Colony restreaking showed that 150/150 Suc\textsuperscript{R}Kan\textsuperscript{R} colonies were Kan\textsuperscript{R}Cam\textsuperscript{S} (Fig. 2c). Colony PCRs, which used two sets of primers (P1 + P2 for detection of Kan\textsuperscript{R}, or mTn5, and P3 + P4 for detection of tac-rep\textsuperscript{F}, or plasmid), confirmed plasmid loss in 10 out of 10 colonies (10/10) (Fig. 2d). Sequence analysis showed that all 13 Suc\textsuperscript{R}Kan\textsuperscript{R} colonies analyzed had different Tn5 insertion sites (Fig. 3a).

**Efficient mutagenesis of Acinetobacter baylyi and P. aeruginosa by a highly-active mTn5 transposon**

Construct pSNC-mTn5 was then tested in two Gram-negative, capsule-bearing bacteria, *A. baylyi* 33,305 and *P. aeruginosa* PAO1 [23, 24]. Comparing to *E. coli* DH10B, transformed *A. baylyi* 33,305 and *P. aeruginosa* PAO1 appeared to lose the plasmid more easily in the absence of IPTG, with \(\sim 56.3\%\) of *A. baylyi* and \(\sim 59.6\%\) of *P. aeruginosa* PAO1 appearing to lose the plasmid after 24 h culture in LB media without antibiotics (Fig. 2e, f). Following IPTG induction, \(\sim 14.7\%\) of *A. baylyi* and \(\sim 3.2\%\) of *P. aeruginosa* retained the plasmid, suggesting that IPTG induced additional plasmid loss from these organisms, although their efficiencies...
were lower than that in DH10B cells. With sucrose counter selection, \(1.6 \times 10^{-7}\) of \(A.\) baylyi remained CamR, indicating that they contained the plasmid (Fig. 2e). Similarly, \(5.4 \times 10^{-7}\) of \(P.\) aeruginosa cells were found to be SucRCamR (Fig. 2f). These results suggest that IPTG and sucrose both contributed in reducing plasmid-bearing cells.

In comparison, the percentages of SucRKanR cells were \(8.4 \times 10^{-6}\) for \(A.\) baylyi and \(3.4 \times 10^{-6}\) for \(P.\) aeruginosa, suggesting that Tn5 transposition occurred in both organisms prior to plasmid loss. Colony restreaking showed that 100/100 Suc\(^R\)Kan\(^R\) colonies are Suc\(^R\)Cam\(^S\), suggesting that efficient plasmid loss had occurred following Tn5 transposition (\(\sim 100\%\) for both; Additional file 2: Figure S2a, c). Loss of plasmids was further confirmed by PCR tests (Additional file 2: Figure S2b, d). As observed in DH10B cells, Tn5 insertion also seemed to be random, as 9/9 \(A.\) baylyi and 13/15 \(P.\) aeruginosa mutants had different Tn5 insertion sites (Fig. 3b, c). The detection of identical
mutant suggests that cell growth ensued following transposon transposition (Fig. 3c), which is common in different transposon mutagenesis assays [5, 6, 25].

**Construction of a Tn5 insertion library of *P. aeruginosa* using the highly-active mTn5 transposon**

To determine whether we can construct a transposon insertion library of *P. aeruginosa* PAO1 with high diversity, ten pSNC-mTn5 transformants of the bacterium were cultured independently and then combined and induced with IPTG to initiate transposon mutagenesis. Following 24 h culture in LB media containing IPTG, ~ 6.4% of cells retained the plasmid (Additional file 3: Figure S3a). The frequencies of Suc\(^r\)Cam\(^r\) and Suc\(^r\)Kan\(^r\) cells in the IPTG-induced culture were found to be ~ 6.5 × 10\(^{-7}\) and ~ 3.5 × 10\(^{-6}\), respectively. Based on the total number of cells cultured and the frequency of SucRKanRCamS cells, the total diversity of the mTn5 insertion library was estimated to be ~ 1.3 × 10\(^7\), which covers the entire gene repertoire (5697) of *P. aeruginosa* PAO1 by ~ 2238 times [24]. To our knowledge, the diversity of this transposon insertion library is bigger than the best transposon insertion library of PAO1 and related strains previously reported (Table 1) [5, 6, 9, 26–32].

An mTn5 with MEs enables generation of a *P. aeruginosa* mutant library with even higher diversity

To determine whether the efficiency of mTn5 transposition can be further improved, we replaced both OE and IE of the mTn5 with MEs (Fig. 4a). The new plasmid, pSNC-mTn5ME, was transformed into DH10B cells. Cell growth (or colony sizes) appeared to be normal, suggesting that basal-level transposition, if any, did not lead to obvious cellular toxicity, which was our initial concern. The behavior of the plasmid and Tn5 transposition efficiency were determined under the same conditions described above. Without IPTG induction, the plasmid remained relatively stable, as ~ 100% of the cells retained the plasmid (Cam\(^r\)). After IPTG induction for 24 h, 1.1 × 10\(^{-3}\) of the cells retained the plasmid, suggesting that RepF overexpression caused efficient plasmid loss. With sucrose counter selection, ~ 2.4 × 10\(^{-5}\) cells remained SucRCamR. Interestingly, the frequency of Suc\(^r\)Kan\(^r\) cells was found to be very high (~ 28.0%), indicating that mTn5ME is much more active than the non-ME version (~ 1200 folds). Restreaking of SucRKanR colonies showed that they were all Kan\(^r\)Cam\(^r\) (100/100) (Additional file 4: Figure S4a), and colony PCRs confirmed plasmid loss (10/10) (Additional file 4: Figure S4b). Sequence analysis of the transposon insertion junctions showed that
all 13 Suc^{R}Kan^{R} colonies analyzed had different Tn5 integration sites (Additional file 4: Figure S4c).

We then determined whether construct pSNC-mTn5ME would also be more active in A. baylyi 33,305 and in P. aeruginosa PAO1. For A. baylyi, the efficiencies of plasmid loss were higher for pSNC-mTn5ME than for pSNC-mTn5, both in the absence and presence of IPTG induction (Figs. 2e and 4c). Sucrose counter selection was highly effective for both constructs (Figs. 2e and 4c). As in E. coli, mTn5ME was found to be much more active than the non-ME version (~140 fold higher; Suc^{R}Kan^{R} cells: ~1.2 × 10^{-5} for mTn5ME vs. ~8.4 × 10^{-6} for mTn5). Similarly, colony restreaking of Suc^{R}Kan^{R} cells showed that 100/100 colonies are Kan^{R}Cam^{S} (Additional file 5: Figure S5a), and plasmid loss was further verified by PCR (Additional file 5: Figure S5b). 9/14 colonies were found to have different Tn5 insertion sites (Additional file 5: Figure S5c).

For PAO1, efficiencies of plasmid loss (~IPTG) were found to be similar for both pSNC-mTn5 and pSNC-mTn5ME (Figs. 2f and 4d), and sucrose counter selection was also effective for pSNC-mTn5ME (Fig. 4d). As in E. coli and in A. baylyi, mTn5ME was found to be more active than mTn5 in PAO1 (~11 fold higher; Suc^{R}Kan^{R} cells: ~3.9 × 10^{-5} for mTn5ME vs. ~3.4 × 10^{-6} for mTn5) (Figs. 2f and 4d). In the colony restreaking assay, 100/100 Suc^{R}Kan^{R} colonies were found to be Kan^{R}Cam^{S} (Additional file 6: Figure S6a), and PCR assays further confirmed plasmid loss (10/10) (Additional file 6: Figure S6b). Sequence determination showed that 7/10 colonies tested had different Tn5 insertion sites (Additional file 6: Figure S6c).

We then determined whether pSNC-mTn5ME would be a better construct than pSNC-mTn5 for transposon saturation mutagenesis in P. aeruginosa. Ten transfectants were randomly picked for Tn5 insertion library construction using the protocol described above. About 0.87% of cells retained the plasmid after IPTG induction, and the frequency of Suc^{R}Cam^{R} cells was found to be 6.8 × 10^{-8}. In comparison, the frequency of Suc^{R}Kan^{R} cells was found to be ~9.5 × 10^{-5}, suggesting that efficient mTn5ME transposition has occurred. Based on the total amount of cells cultured and the mTn5ME transposition efficiency (~9.5 × 10^{-5}), the diversity of the mTn5ME insertion library was estimated to be 1.02 × 10^{8}, which is ~3 logs larger than the best PAO1 transposon insertion library previously reported and ~2 logs larger than a Tn5 insertion library of P. aeruginosa MPAO1, a derivative of PAO1 with ~0.2% genetic variation [6, 33]. This new library is by far the biggest transposon insertion library of PAO1 and related species ever reported (Table 1). The size of our new library is enough to cover the entire gene repertoire of PAO1 by ~18,000 times. Colony restreaking (100) and PCR tests (10) confirmed plasmid loss in all the Suc RKanR clones analyzed (Fig. 4f, g), and 34/46 clones tested had different Tn5 insertion sites (Fig. 4h). Thus, the independent clones in the library (library diversity) are estimated to be ~7.5 × 10^{7}.

**Discussion**

We have developed a new transposon mutagenesis system that is efficient, regulatable, easy-to-use, and broadly useful. We believe it will be especially useful for functional genomics studies of Gram-negative bacteria that are difficult to transform and conjugate, such as certain capsule-containing bacteria, obligate anaerobes, and possibly obligate intracellular pathogens as well. The advantage of this method relies on the following features: (i) A broadly-functional plasmid replicon; (ii) Replication of the plasmid is regulated by IPTG; (iii) The inclusion of the sacB gene for counter selection; (iv) A highly-active/hyperactive transposon; (v) Regulatable expression of the

| Table 1 Comparison of transposon insertion libraries of P. aeruginosa strains |
|-----------------------------|-----------------------------|-----------------------------|
| Pseudomonas strains         | Total mutants               | Transposon                  | Method         | References |
| P. aeruginosa PAO1          | ~ 75,000,000                | mTn5ME                     | IPTG-curable   | This study |
| P. aeruginosa PAO1          | ~ 10,000,000                | mTn5                       | IPTG-curable   | This study |
| P. aeruginosa PAO1          | ~ 80,000                    | Himar1                     | Conjugation    | [26]        |
| P. aeruginosa MPAO1         | ~ 18,500                    | Tn5                        | Conjugation    | [27]        |
| P. aeruginosa PAO1          | ~ 217,725^c                 | Tn5                        | Conjugation    | [6]         |
| P. aeruginosa PAO1          | ~ 100,000                   | Tn5                        | Conjugation    | [28]        |
| P. aeruginosa MPAO1         | ~ 45,409                    | Tn5                        | Conjugation    | [5]         |
| P. aeruginosa PAO1SR        | ~ 1,000,000                 | Himar1                     | Conjugation    | [29]        |
| P. aeruginosa PA14          | ~ 300,000^d                 | Himar1                     | Conjugation    | [30]        |
| P. aeruginosa PA14          | ~ 38,976                    | Himar1                     | Conjugation    | [31]        |
| P. aeruginosa QR1           | ~ 15,000                    | Himar1                     | Conjugation    | [32]        |

^a A PAO1 derivative with ~0.2% genetic changes  
^b A streptomycin resistant derivative of PAO1 with unknown changes  
^c Combination of 16 experiments  
^d Combination of 100 independent conjugations
Fig. 4 (See legend on next page.)
hyperactive transposase gene; (vi) mTn5 and mTn5ME transposons insert almost completely randomly in different bacteria (Additional file 7: Figure S7) [34, 35]. In addition, the relatively small sizes of the RSF1010-based plasmids also facilitate their transformation and conjugation. Similar to transposon mutagenesis using ts plasmids, our system does not depend on efficient plasmid transformation and conjugation, and requires as few as one transformant or conjugated cell for transposon saturation mutagenesis. Using this new tool, we have generated a Tn5 transposon insertion library of P. aeruginosa PAO1 with a diversity of ~ 10^8, which is ~ 2 logs larger than the liverring gene targeting systems, e.g., TargeTrons, λ such as for allelic exchange or as curable vectors for de-we developed should have many other applications, the IPTG-controlled conditional suicide plasmids that addition to their utilities in transposon mutagenesis, platforms that have been used for transposon mutagen-

**Conclusion**

In this work, we have developed a number of IPTG-controlled conditional suicide plasmids that contain the broad-host-range RSF1010 origin. Using one of the constructs to deliver a hyperactive mTn5 transposon, we showed that this system can be used for efficient mutagenesis of different bacterial species. As the assay condition is mild and the host range of the RSF1010 plasmid is extremely wide, we believe that our methodology will have broad applications in microbiology research.

**Methods**

**Bacterial strains and growth conditions**

E. coli DH10B was purchased from Invitrogen. A. baylyi (ATCC 33305) and P. aeruginosa PAO1 (ATCC BAA-47) were purchased from ATCC. Unless stated otherwise, all the strains were grown at 37 °C in Luria Broth (LB) liquid media with agitation at 200 rpm or on LB plates with 1.5% agar. For sacB counter selection, we used LBNS plates (LB no salt: 1% Tryptone, 0.5% yeast extract and 1.5% agar) supplemented with 10% sucrose (Fisher Scientific). Appropriate antibiotics and concentrations were used to select for bacterial cells that are antibiotic resistant. E. coli DH10B: chloramphenicol (Cam; Gold Biotechnology), 25 μg/ml; kanamycin (Kan; Fisher Scientific), 50 μg/ml A. baylyi: Cam, 10 μg/ml; Kan, 10 μg/ml. P. aeruginosa PAO1: Cam, 250 μg/ml; Kan, 500 μg/ml.

**Plasmid construction**

To construct plasmid pMMB-repF, we PCR-amplified the repF gene from pMMB208 [18]. The PCR fragment was digested with HindIII and PstI, and inserted at the corresponding sites of pMMB208, downstream of the tac promoter (Ptac). To construct plasmid pMMB-repF/sacB, the sacB gene and its promoter were PCR amplified from plasmid pRE112 [22] and inserted between the uniqueSac1 and KpnI sites of pMMB-repF.

To construct plasmids pMMB-repAK42A and pMMB-repAD139A, repA genes containing K42A and D139A mutations were generated in two-step PCRs from plasmid

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**Fig. 4** Generation of a P. aeruginosa insertion library with pSNC-mTn5ME. a Diagram of pSNC-mTn5ME, a derivative of pSNC-mTn5 that has MEs instead of OE and IE at the termini of mTn5. b Plasmid and transposon retention frequencies in E. coli DH10B. Results were average of three independent experiments, and bars represent mean ± SD (*p < 0.0001 and **p = 0.0004 by unpaired t-test). Colony restreaking and PCR assays are shown in Additional file 4: Figure S4. c Plasmid and transposon retention frequencies in A. baylyi. Results were average of three independent experiments, and bars represent mean ± SD (*p = 0.0029 and **p = 0.0006 by unpaired t-test). Colony restreaking and PCR assays are shown in Additional file 5: Figure S5. d Plasmid and transposon retention frequencies in P. aeruginosa PAO1. Results were average of three independent experiments, and bars represent mean ± SD (*p < 0.0001 and **p = 0.0065 by unpaired t-test). Colony restreaking and PCR assays are shown in Additional file 6: Figure S6. e Plasmid and transposon retention frequencies in the P. aeruginosa PAO1 mutant library generated with pSNC-mTn5ME. f Colony restreaking. 100/100 Suc^+Kan^+ colonies of the mTn5ME library of P. aeruginosa were found to be Kan^+Cam^+. 50 are shown here. g Colony PCR of ten restreaked clones in (f) with the indicated primers. All were mTn5-positive and plasmid-negative. h Transposon insertion sites of 46 mutant clones from the mTn5ME insertion library of P. aeruginosa. Identical clones are shown only once, with their duplication numbers indicated in parenthesis.

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**Fig. 4**: Generation of a P. aeruginosa insertion library with pSNC-mTn5ME. **a** Diagram of pSNC-mTn5ME, a derivative of pSNC-mTn5 that has MEs instead of OE and IE at the termini of mTn5. **b** Plasmid and transposon retention frequencies in E. coli DH10B. Results were average of three independent experiments, and bars represent mean ± SD (*p < 0.0001 and **p = 0.0004 by unpaired t-test). Colony restreaking and PCR assays are shown in Additional file 4: Figure S4. **c** Plasmid and transposon retention frequencies in A. baylyi. Results were average of three independent experiments, and bars represent mean ± SD (*p = 0.0029 and **p = 0.0006 by unpaired t-test). Colony restreaking and PCR assays are shown in Additional file 5: Figure S5. **d** Plasmid and transposon retention frequencies in P. aeruginosa PAO1. Results were average of three independent experiments, and bars represent mean ± SD (*p < 0.0001 and **p = 0.0065 by unpaired t-test). Colony restreaking and PCR assays are shown in Additional file 6: Figure S6. **e** Plasmid and transposon retention frequencies in the P. aeruginosa PAO1 mutant library generated with pSNC-mTn5ME. **f** Colony restreaking. 100/100 Suc^+Kan^+ colonies of the mTn5ME library of P. aeruginosa were found to be Kan^+Cam^+. 50 are shown here. **g** Colony PCR of ten restreaked clones in (f) with the indicated primers. All were mTn5-positive and plasmid-negative. **h** Transposon insertion sites of 46 mutant clones from the mTn5ME insertion library of P. aeruginosa. Identical clones are shown only once, with their duplication numbers indicated in parenthesis.
Plasmid pSNC-mTn5 was constructed in multiple steps. First, plasmid pUT-mTn5Km/lacEZ was constructed from plasmid pUT-mTn5Km [4]. It contains a lac promoter-driven hyperactive transposase gene (tnpH) that has E54K, M56A and L372P mutations [3]. In addition, inside the mTn5 transposon, the inverted repeats flanking the kanamycin resistance marker (KanR) were deleted [4]. The entire KanR-mTn5 transposon and Plac-tnpH, was then PCR amplified and cloned at the XbaI site of pMMB-repF/sacB, resulting in plasmid pSNC-mTn5. It has an OE and an IE at the termini of the mTn5. Plasmid pSNC-mTn5ME was derived from pSNC-mTn5 by replacing both OE and IE with MEs.

Characterization of IPTG-induced plasmid loss and transposon mutagenesis

To test IPTG-induced plasmid loss of pMMB-repF, pMMB-repA4242A and pMMB-repAD1739A, single colonies of E. coli DH10B cells transformed with the plasmids were inoculated into 5 ml LB + Cam media and cultured at 37 °C for ~14 h (h). After measuring OD600, 1 ml of each culture was pelleted by centrifugation and washed with 500 μl of fresh LB to remove antibiotics. Cells were then resuspended in 1 ml LB. An aliquot was added to 5 ml LB (final OD600 = 0.001) with and without 1 mM IPTG and cultured at 37 °C for 24 h. 1 ml of the IPTG-induced samples was then pelleted, washed with 500 μl LB, and resuspended in 1 ml LB. Serial dilutions of the samples (±IPTG) were plated on LB and LB + Cam plates to evaluate plasmid loss. Plasmid retention frequencies were calculated as ratios of cfu (colony forming units) on LB + Cam plates and those on LB plates.

To perform transposon mutagenesis in E. coli DH10B, single colonies of pSNC-mTn5 and pSNC-mTn5ME transformants were cultured in 5 ml LB + Cam media overnight at 37 °C. Cells were then pelleted and washed as above to remove antibiotics, and an aliquot was inoculated to 5 ml LB (final OD600 = 0.001) in a 14 ml culture tube and grown at 37 °C for 24 h with and without 1 mM IPTG induction. A 1 ml aliquot of the IPTG-induced samples was then pelleted, washed with 500 μl LBNS, and resuspended in 1 ml LBNS. Serial dilutions of the samples (±IPTG) were plated on appropriate plates to evaluate plasmid loss and mTn5 (or mTn5ME) transposition as in the assays for E. coli and P. aeruginosa.

To verify plasmid loss in cells with potential transposition events, 100–150 SucRKanR colonies in each assay were then restreaked on LB + Kan and LB + Cam plates. In addition, presence of the transposon and the plasmid was determined by colony PCR, in which 25 μl reaction containing 25 mM TAPS-HCl (pH 9.3), 50 mM KCl, 2 mM MgCl2, 1 mM β-mercaptoethanol, 1x GC enhancer, 0.2 mM dNTPs, 0.1 μl of Q5 polymerase (2 u/μl NEB), 1 μl of resuspended cells, and 150 ng each of the primers (final concentration ~ 0.5 μM; see figure legends and Additional file 9: Table S2 form oligos used). PCRs were performed using the following condition: 1x (94 °C, 2 min); 25x (94 °C, 30 s; 50 °C, 30 s; 72 °C, 1 min); 1x (72 °C, 10 min); 1x (4 °C, hold).

Determination of transposon insertion sites

Transposon insertion sites in bacterial chromosomes were determined by arbitrarily primed PCR, in which transposon junctions were amplified in two steps [5, 39]. Bacterial cells were resuspended in 10–20 μl of deionized water and 1 μl was used directly as the PCR template. In the first PCR step, the reaction was performed using a specific primer annealing to the transposon region (Tn5Km1) and a semi-degenerate primer (BDC1) that anneals to many sites on the bacterial chromosome. In the second step, aliquots of the first-round PCR products were amplified using a primer annealing to the transposon region (Tn5Km2), slightly closer to the
insertion junction, and a non-degenerate primer (BCD2) that anneals to the constant region of the BDC1-derived sequence. PCRs were carried out under the conditions described above. PCR products from Step 2 were resolved in a 2% agarose gel and major products were gel-purified for sequencing to determine Tn5 insertion sites.

**Construction of transposon insertion libraries of *P. aeruginosa* PAO1**

To construct an mTn5 (or mTn5ME) insertion library of *P. aeruginosa* PAO1, plasmid pSNC-mTn5 (or pSNC-mTn5ME) was first electroporated into the bacterial cells. Ten transformants were cultured independently in 5 ml LB + Cam + Kan media at 37 °C for ~14 h. Equal amount of each sample (equivalent to 0.5 OD600 x 1 ml) was then combined, pelleted, washed with 500 µl LB, and the pellet was resuspended in 1 ml LB. An aliquot of the mixture was then inoculated into 500 ml LB supplemented with 10 mM IPTG in a baffled flask (final OD600 = 0.01) and shaken vigorously (300 rpm) at 37 °C for 24 h to perform transposon mutagenesis. The cells were then pelleted by centrifugation and washed with 250 ml LBNS medium. The pellet was resuspended in 50 ml LBNS medium and serial dilutions were plated on LB, LB + Cam, LBNS+ 10% sucrose, LBNS+ 10% sucrose+Cam, and LBNS+ 10% sucrose+Kan plates to determine plasmid loss, mTn5 (mTn5ME) transposition, and total library diversity. Arbitrary PCR and DNA sequencing were then performed to determine Tn5 insertion sites.

**Determination of mTn5 and mTn5ME target site preferences in *P. aeruginosa* PAO1**

To determine if mTn5 and mTn5ME have any target site preferences in *P. aeruginosa* PAO1, we generated sequence logos of their insertion sites in the bacterium using the WebLogo server (https://weblogo.berkeley.edu/logo.cgi). In total, 40 mTn5 insertion sites and 41 mTn5ME insertion sites were used for the analysis.

### Additional files

**Additional file 1:** Figure S1. Tn5 transposons. (a) Full-length Tn5. The full-length Tn5 contains two inverted IS50 elements at its ends. Only one of them encodes an active Tnp and an Ini (Inhibitor of Tnp). KanR, kanamycin-resistance gene; SgR, streptomycin-resistance gene; and B1R, bleomycin-resistance gene. (b) mTn5s. Top, an mTn5s with an OE and an IE at the termini. Bottom, an mTn5s with MEs at the ends. (c) Comparison of OE, IE and ME with their polymorphisms highlighted in red. (TIF 4125 kb)

**Additional file 2:** Figure S2. Confirmation of mTn5 transposition events in *A. baylyi* and in *P. aeruginosa*. (a) Colony restreaking assay of *A. baylyi*. 100 Suc+KanR colonies of *A. baylyi* were restreaked on LB + Kan and LB + Cam plates, and all were found to be KanRCamR. 30 were shown here. (b) Colony PCR of 10 restreaked *A. baylyi* clones with the indicated primers. All were mTn5-positive and plasmid-negative. (c) Colony restreaking assay of *P. aeruginosa*. 100 Suc+KanR colonies of *P. aeruginosa* were restreaked on LB + Kan and LB + Cam plates, and all were found to be KanRCamR. 30 are shown here. (d) Colony PCR of ten restreaked *P. aeruginosa* clones with primers indicated in the diagram. All were mTn5-positive and plasmid-negative. (TIF 6384 kb)

**Additional file 3:** Figure S3. A transposon insertion library of *P. aeruginosa* PAO1 generated with pSNC-mTn5. (a) Plasmid and transposon retention frequencies of the mTn5 insertion library of *P. aeruginosa* PAO1. (b) Colony restreaking assay. 100 random Suc+KanR colonies were restreaked on LB + Kan and LB + Cam plates. 100/100 were found to be KanRCamR and 30 are shown here. (c) Colony PCR of ten restreaked clones in (b). All were found to be mTn5-positive and plasmid-negative. (d) mTn5 insertion sites of 37 mutant clones from the transposon insertion library of *P. aeruginosa*. Identical clones are only shown once, and their numbers are indicated in parenthesis. (TIF 1363 kb)

**Additional file 4:** Figure S4. Confirmation of mTn5ME transposition events in *E. coli* DH10B. (a) Colony restreaking. 100 random Suc+KanR colonies of *E. coli* were restreaked on LB + Kan and LB + Cam plates. 100/100 were found to be KanRCamR and 50 restreaked colonies are shown here. (b) Colony PCR of ten restreaked clones in (a). All were found to be Tn5-positive and plasmid-negative. (c) Sequence analysis shows that 8/14 *A. baylyi* clones had different Tn5 insertion sites. (TIF 7051 kb)

**Additional file 5:** Figure S5. Confirmation of mTn5ME transposition events in *A. baylyi* 33,305. (a) Colony restreaking. 100 random Suc+KanR colonies of *A. baylyi* were restreaked on LB + Kan and LB + Cam plates. 100/100 were found to be KanRCamR and 50 restreaked colonies are shown here. (b) Colony PCR of ten restreaked clones in (a). All were found to be Tn5-positive and plasmid-negative. (c) Sequence analysis shows that 9/14 *A. baylyi* clones had different Tn5 insertion sites. (TIF 6938 kb)

**Additional file 6:** Figure S6. Confirmation of mTn5ME transposition events in *P. aeruginosa* PAO1. (a) Colony restreaking. 100 random Suc+KanR colonies of *P. aeruginosa* PAO1 were restreaked on LB + Kan and LB + Cam plates. 100/100 were found to be KanRCamR and 50 restreaked colonies are shown here. (b) Colony PCR of ten restreaked clones in (a). All were found to be Tn5-positive and plasmid-negative. (c) Sequence analysis shows that 7/10 *P. aeruginosa* clones had different Tn5 insertion sites. (TIF 6280 kb)

**Additional file 7:** Figure S7. Target site preferences of mTn5 and mTn5ME in *P. aeruginosa*. (a) Sequence logo of mTn5 insertion sites generated with WebLogo. 40 target sequences were analyzed. The 9 bp duplicated sequences adjacent to the OE are shown. There is a slight preference for certain nucleotides at several positions. (b) Sequence logo of mTn5ME insertion sites. 41 target sequences were analyzed. The 9 bp duplicated sequences adjacent to an ME are shown. It appears that mTn5ME has less nucleotide preference at the duplicated target sequence than mTn5 in *Pseudomonas*. (TIF 3834 kb)

**Additional file 8:** Table S1. Comparison of conditional suicide vector-based transposon mutagenesis strategies used in Gram-negative bacteria (TIF 7782 kb)

**Additional file 9:** Table S2. List of primers used. (TIF 2042 kb)

### Abbreviations

bp: Base pair; Cam: Chloramphenicol; IE: Inside end; IPTG: Isopropyl-β-D-1-thiogalactopyranoside; Kan: Kanamycin; LB: Luria Broth; LBNS: LB no salt; ME: Mosaic end; OD: Optical density; OE: Outside end; rt: Temperature-sensitive

### Acknowledgements

We thank Drs. Mark McIntosh, David Pintel and Donald H. Burke for helpful discussions.

### Funding

This work was supported by the University of Missouri startup fund to H.G. The funding body had no role in the design of the study and collection, analysis, and interpretation of data and in writing the manuscript.

### Availability of data and materials

The dataset supporting the conclusion of this article are available from the corresponding author on reasonable request.
Authors’ contributions

SSN, JH and HG conceived the study and designed the experiments; SSN, JH, SYZ, IZ, LBG, AS, CVS and FR performed the experiments; SSN and HG analyzed and wrote the manuscript. All authors have read and approved the manuscript.

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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Received: 12 July 2018 Accepted: 16 October 2018
Published online: 24 October 2018

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