A Markerless Method for Genome Engineering in Zymomonas mobilis ZM4

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Metabolic engineering of the biofuel-producing Zymomonas mobilis is necessary if we are to unlock the metabolic potential present in this non-model microbe. Manipulation of such organisms can be challenging because of the limited genetic tools for iterative genome modification. Here, we have developed an efficient method for generating markerless genomic deletions or additions in Z. mobilis. This is a two-step process that involves homologous recombination of an engineered suicide plasmid bearing Z. mobilis targeting sequences and a subsequent recombination event that leads to loss of the suicide plasmid and a genome modification. A key feature of this strategy is that GFP expressed from the suicide plasmid allows easy identification of cells that have lost the plasmid by using a fluorescence activated cell sorter. Using this method, we demonstrated deletion of the gene encoding lactate dehydrogenase (ldh) and the operon for cellulose synthase (bcsABC). In addition, by modifying the plasmid design, we demonstrated targeted insertion of the crtIBE operon encoding a neurosporene biosynthetic pathway into the Z. mobilis genome without addition of any antibiotic resistance genes. We propose this approach will provide an efficient and flexible platform for improved genetic engineering of Z. mobilis.

Keywords: genome engineering of a non-model bacterium, green fluorescent protein, fluorescence activated cell sorting, recombineering suicide plasmid, biofuels, Zymomonas mobilis

INTRODUCTION

While the pace of research into genetic tools for model organisms has accelerated over the last half century, the development of tools for non-model bacteria has lagged behind. Zymomonas mobilis ZM4, an obligatory fermentative alpha-proteobacterium, is one such organism that lacks convenient tools for genetic engineering. This organism has garnered significant interest as a platform for the biosynthesis of biofuels because of its native ability to efficiently metabolize glucose to ethanol rather than to biomass (He et al., 2014; Yang et al., 2016; Wang et al., 2018). To optimize Z. mobilis’ metabolism and unlock its full potential for industrial fermentation of compounds of interest, methods for quickly and efficiently removing undesirable genes and pathways and adding novel genes and pathways are needed.
Existing genetic tools for *Z. mobilis* are not optimal for creation of strains with multiple engineered loci. A common approach to insertionally inactivate genes relies on host homologous recombination of an antibiotic resistance gene flanked by *Z. mobilis* DNA contained on a suicide plasmid, which, once integrated into the genome, can be screened for events leading to loss of delivery plasmid and gene of interest (Senthilkumar et al., 2004; Strazdina et al., 2018). This approach was enhanced by use of lambda red recombinase to increase the frequency of recombination events (Senthilkumar et al., 2004; Bo et al., 2013). However, the inability to remove the antibiotic resistance cassette quickly limits the number of modifications that can be made to a particular strain especially given the natural resistance of *Z. mobilis* to several antibiotics (Bochner et al., 2010). To overcome this problem, FLP recombinase was used to remove the antibiotic resistance gene to create a markerless gene deletion (Zou et al., 2012). However, introduction and removal of the stable plasmid bearing the FLP recombinase is a time-consuming process. To avoid the problems associated with using antibiotic resistance genes, Xia et al. (2018) made use of a sucrose counter-selection (*sacB*) to accelerate the process of integrated plasmid loss (Xia et al., 2018), but *Z. mobilis* ZM4 is annotated to also have *sacB*, questioning the general utility of this approach. Lastly, a CRISPR/Cas9 system was deployed to remove a native plasmid from *Z. mobilis* (Cao et al., 2017). This technique has not yet been extended to genome modification. Therefore, we sought to develop a flexible and efficient method for genome engineering that would overcome some of the limitations of existing methods.

In this paper, we describe the creation of an efficient, reproducible, and flexible method for making markerless mutations in *Z. mobilis* by homologous recombination. We demonstrate its primary utility by making knockouts of the *ldh* gene (ZMO00256: lactate dehydrogenase) and the *bcsABC* genes (ZMO1083-85; cellulose synthase). We also describe how this technique could be extended to make markerless knock-ins at a desired locus by introducing carotenoid biosynthetic genes, *crtlBE*, from *Rhodobacter sphaeroides* (Takaichi and Maoka, 2015; Su et al., 2018). Herein, we also describe the suicide vector, which forms the basis for this technique. We also describe our unique choice to include the gene encoding GFP in our vector, which, through the use of a transilluminator or fluorescence activated cell sorting (FACS), allows us to track and screen for the presence or absence of the plasmid during intermediate steps in the process. This system should allow the facile and targeted addition or deletion of genes of interest in an iterative manner and thus facilitate the engineering of *Z. mobilis* for desirable traits.

### MATERIALS AND METHODS

Primers, bacterial strains, and plasmids used in this study are listed in Tables 1–3.

### Growth of Bacterial Strains

*Zymomonas mobilis* strains were cultured in rich medium ZRMG (1% yeast extract, 0.2% KH₂PO₄, 2% glucose) or glucose-minimal medium (Martien et al., 2019) at 30°C. *Escherichia coli* strains were grown in Luria-Bertani (LB) medium. Chloramphenicol was used at a final concentration of 120 µg ml⁻¹ for *Z. mobilis* and 20 µg ml⁻¹ for *E. coli* strains. Spectinomycin was used at a final concentration of 100 µg ml⁻¹ for *Z. mobilis* and 50 µg ml⁻¹ for *E. coli* strains.

### Conjugation

*Escherichia coli* donor strain WM6026 (Blodgett et al., 2007) was used to conjugate plasmids into the recipient strain *Z. mobilis* ZM4. WM6026, a *m*-diaminopimelate (DAP) auxotrophic strain, was grown at 30°C aerobically in LB containing DAP (0.1 mM) with appropriate antibiotic. Overnight cultures were then subcultured into the same medium but lacking antibiotic to an OD₆₀₀ of 0.15 and were grown to an OD₆₀₀ of 0.5. *Z. mobilis*

### TABLE 1 | Primer list.

| Primer number | Primer sequence (5’-3’) |
|---------------|------------------------|
| P1            | GTGCCGAGTGATGCTGATG    |
| P2            | CAGTAATCTTGAGGCGGATG   |
| P3            | CAGAAAGCTTAAAGCCCTGG   |
| P4            | CTTAGCCGTATGTTAGCTTCTC |
| P5            | GCTACGCTAAGTGTGACGAGC  |
| P6            | ACTATCCAGGGGTGACAC     |
| P7            | GACCGAGCAACTCTTACAG    |
| P8            | GTCTGCGTTTCTAGCTGCTT   |
| P9            | GCTGCTACGCTTGAATAGTG   |
| P10           | GCGCTTTTCCCTGCTCAT    |
| P11           | GTTATGCAGCGCGTAGCTAGC  |
| P12           | GCTGCACGGCGTATTAGTGTTA |
| P13           | ACCGAGCGCTACTAGTACATT  |
| P14           | TCTACACTGACGCTCTGTTT   |
| P15           | AGCTTACTAAGCTTGATACAT  |
| P16           | GTCTGATATACGCTTGTTT    |
| P17           | AGGCCGAGACGACGACGGCCTT |
| P18           | AGGAGCTGAGTACCTGCTCT   |
| P19           | CTGGCGAGTTACCTGCTG     |
| P20           | AAACCTGTGAAAGCGCAGAC   |
| P21           | TGCGGCTTTCCCACCTTTGGC  |
| P22           | ATGTTATATCCCTGATCAAATACCAAC |
| P23           | TTAATAGAAAGAAGCAGATTTACT|
| P24           | TATTATAAATAAGAAGAATCATG|
| P25           | CCCGAGGTATGCTGTGCTGCTG |
| P26           | TTTTCATGAGGCACAAGATGTTGGTTC |
| P27           | GCAAGGCAATCTCAGCTGATAGG |
| P28           | GTCTGATATCCGTTGCTG     |
| P29           | ATAGCTGATATCCGTTGCTG   |
| P30           | TATAGGCTATATACGCTGCTG  |
| P31           | GCCGAGGCTGTTTCTAGCAGCC |
| P32           | CGCGGCTGCGTTGCTGCTGCTG |
| P33           | CGGTGCTACGCTGCTGCTGCTG |
| P34           | CGTCTGACCTCTTACCTGCTG |
| P35           | CAGGTGCTAGCGCGCTTACAG  |
| P36           | CGAAATACGCGAGCGAGGCC   |
| P37           | ATCTATGCGGTTGACGAGAC   |
| P38           | TCAACACAGGCGTTTACAGC   |
**TABLE 2 | Strain list.**

| Strain     | Description | Source or references |
|------------|-------------|----------------------|
| WM6026     | *E. coli lacIq mB3 Δ lacZ4787 hasR14 Δ araBAD567 Δ araBAD658 rph-1 attX::pAE12(ΔoriR6K-cat; Frt5), Δ endA::Frt udrAΔMull::pir attHK::pK1006(ΔoriR6K-cat::Frt5; trfA::Frt) dapA::Frt | Bloedgett et al., 2007 |
| DK505      | *Z. mobilis* Zymomonas mobilis ZM4 ATCC 31281 | GLBRC |
| PK15558    | *Z. mobilis* ldhU::pPK15552 | This work |
| PK15557    | *Z. mobilis* ldhU::pPK15553 | This work |
| PK15569    | *Z. mobilis* ldhU::pPK15564 | This work |
| PK15575    | *Z. mobilis* Δldh::ori tet | This work |
| PK15589    | *Z. mobilis* bcsABC::pPK15538 | This work |
| PK15597    | *Z. mobilis* ΔbcsABC | This work |
| PK15598    | *Z. mobilis* Δldh | This work |

**TABLE 3 | Plasmid list.**

| Plasmid | Description | References or source |
|---------|-------------|----------------------|
| pRL814  | Broad host range plasmid | Ghosh et al., 2019 |
| pACYC184| cat p15A ori tet | Lab collection |
| pPK15252| Gibson assembly of p15A ori and cat (2115 bp of pACYC184 amplified by primers P1 and P2), 500 bp flanking regions of *Z. mobilis ldh (ldhU and ldhDN fragments)* | This work |
| pPK15296| pPK15252 with mob from pSUP202 inserted between cat and ldhDN | This work |
| pPK15303| pPK15296 with RBS element and gfp gene from pRL814 inserted between cat and mob | This work |
| pPK15445| pPK15303 with optimized RBS for gfp | This work |
| pPK15534| pPK15445 with deletion of terminal 100 bp of p15A fragment to remove tet promoter and adjacent ldhU and ldhDN replaced by a Spel site | This work |
| pPK15535| pPK15445 with deletion of terminal 100 bp of p15A fragment to remove tet promoter insertion of Spel site between ldhU and ldhDN | This work |
| pPK15538| pPK15534 with 500 bp regions upstream and downstream of the bcsABC operon inserted at Spel site | This work |
| pPK15564| pPK15535 with crnIEB cassette inserted at Spel site between ldhU and ldhDN | This work |
| pPK15552| pPK15530 with deletion of terminal 100 bp of p15A fragment to remove tet promoter | This work |
| pSUP202  | pMB1 replicon, Ap<sup>R</sup>, Tc<sup>R</sup>, Cm<sup>R</sup>, mob | Simon et al., 1983 |

was incubated in 5.0 ml of ZRMG without shaking at 30°C until late exponential phase was reached. For conjugation, 1.0 ml of cells, adjusted to OD<sub>600</sub> of 0.5, of both recipient and donor was mixed in a 2.0 ml microfuge tube. The cell suspension was briefly centrifuged at 17,300 × g for 30 s, and the supernatant was decanted. The pellet was resuspended in the small amount of remaining medium and placed as a drop on a prewarmed (30–37°C) ZRMG containing 1% tryptone, 0.15 mM DAP agar plate. Plates were incubated overnight at 30°C (12–15 h). Cells were collected from plates by resuspending into 1.0 ml ZRMG liquid media. The conjugation mixture was then vortexed for 10 s, spun at 17,300 × g, and the pellet was resuspended in 1.0 ml ZRMG liquid medium. The suspensions were incubated without shaking at 30°C for 2 h, and 100 µl of the undiluted, 10-fold, and 100-fold diluted cell suspension was plated on antibiotic-containing ZRMG agar plates. Plates were incubated at 30°C.

**Enrichment of Non-fluorescent Cells Using FACS**

A single fluorescent exconjugant colony was grown in liquid ZRMG medium at 30°C without shaking. Twenty microliters of cell suspension from early log phase was diluted in 1 ml sterile 137 mM NaCl, 2.7 mM KCl, 10 mM Na<sub>2</sub>HPO<sub>4</sub>, 1.8 mM KH<sub>2</sub>PO<sub>4</sub>, pH 7.4, and loaded into a Sony MA900 or SH800 FACS equipped with a 70 µm sorting chip according to the manufacturer’s recommendations. Sorting of non-fluorescent events away from the major population of fluorescent cells was done using a 488 nm laser for excitation and 525/50 nm detector to measure emission. Sorted events were collected in 1 ml ZRMG liquid medium, a fraction of which was plated on ZRMG agar plates. Alternatively, one can screen for gfp<sup>-</sup> cells by plating the recombinant cell suspension, grown without selection, on antibiotic-free rich medium plates to obtain ~10,000 colonies and visually screening colonies that have lost fluorescence with a blue light transilluminator.

**PCR Amplification of DNA**

For PCR amplification of DNA fragments to be used for cloning or sequence verification, the high fidelity Q5 polymerase from New England Biolabs was used. For screening or confirming genome modifications, GO<sup>Taq</sup> Flexi DNA polymerase (Promega) was used. For colony PCR, a single colony was picked and resuspended in 50.0 µl of distilled water. 2.0 µl of the suspension was used as a template for PCR amplification. Cell lysis was achieved by heating the PCR reaction mixture at 98°C for 30 s in a thermocycler before amplification steps. Annealing temperature and elongation time were followed as directed by the manufacturer’s specifications. Standard techniques were used for analysis or purification of DNA fragments or other molecular biology methods.

**Construction of Suicide Plasmid (pPK15534)**

A conjugatable suicide plasmid (pPK15534) was constructed in several steps by PCR amplification of indicated DNA fragments followed by Gibson assembly using NEBuilder<sup>®</sup> HiFi DNA Assembly Cloning Kit. DNA fragments containing the p15A origin of replication (ori) and a chloramphenicol resistance gene (cat) were amplified from pACYC184 (Chang and Cohen, 1978), and 500 bp regions upstream (ldh<sub>U</sub>) and downstream (ldh<sub>DN</sub>) of the ldh coding region were amplified from the
Z. mobilis genome, using primer pairs P1-P6, P2-P3, and P4-P5, respectively, to generate pPK15252. A plasmid mobilization element, mob, was amplified from pSUP202 (Simon et al., 1983) using primers P8-P9 and cloned at the junction of \textit{ldh\textsubscript{DN}} and \textit{cat} in pPK15252, amplified by primers P7-P6 to generate pPK15296. A DNA fragment encoding the superfolder fluorescent protein, GFP, driven by the T7A1 promoter and a ribosome binding site was amplified from pRL814 (Ghosh et al., 2019) using primers P11-P12 and added to pPK15296, amplified by primers P6-P10 to generate pPK15303. The ribosome binding site was subsequently optimized using Salis' algorithm (Salis et al., 2009; Borujeni et al., 2014) to improve GFP fluorescence in \textit{Z. mobilis} by PCR amplification of pPK15303 using primers P13 and P14, which contain optimized RBS sequence in overhang regions to generate pPK15445. An orphan \textit{tet} promoter was also removed and \textit{SpeI} cloning site by Gibson assembly of two fragments of pPK15445 generated by a pair of primers P19-P18 and P17-P20, to generate pPK15535. Finally, to create a vector for cloning any target DNA (pPK15534), the orphan \textit{tet} promoter, \textit{ldh\textsubscript{UP}} and \textit{ldh\textsubscript{DN}}, were removed and replaced by a \textit{SpeI} cloning site by Gibson assembly of a fragment of pPK15445 generated by a pair of primers P15-P16.

### Measurement of the Carotenoid Neurosporene

To measure the production of neurosporene, we followed an established protocol (Wraith et al., 1978). Briefly, cells were grown in ZRMG media to an OD\textsubscript{600} of 0.5, the cell pellet from 2 ml culture was resuspended in 100 \mu l water, mixed with 1 ml of 7:2 mixture of acetone:methanol solvent, and spun for 3 min at 21,000 \times g. The absorbance between 300 and 550 nm of the resulting supernatant was recorded on a Perkin Elmer Lambda 25 spectrophotometer.

### RESULTS

#### Overview

To create a more flexible system for deleting or adding genes iteratively to \textit{Z. mobilis}, we rationalized that a recombination-based method was needed that did not require permanent introduction of antibiotic resistance genes into the genome (i.e., markerless modifications) but still could be rapidly screened for desired events. Creation of a conjugatable suicide vector containing GFP (pPK15534) allowed a more convenient method for tracking the loss of the recombinating suicide plasmid from strains, resulting in markerless gene deletions or insertions. Here, we describe the development and function of this system with three examples.

### Demonstration of an Efficient Conjugation System for Introducing DNA Into \textit{Z. mobilis}

To implement a gene-modification system our first step was to establish a reproducible and efficient conjugation method for introducing plasmids into \textit{Z. mobilis} from \textit{E. coli}. We took advantage of an existing \textit{E. coli} donor strain that possesses a chromosomally encoded copy of the RP4 conjugation machinery able to transfer plasmids containing a mobilization element (\textit{mob}) and is auxotrophic for DAP (Blodgett et al., 2007). When this \textit{E. coli} donor strain containing the broad host range plasmid pRL814 was mated with \textit{Z. mobilis} for an optimum period of 12–14 h at 30°C on a solid medium (Figure 1), we found a reproducible transfer frequency of 0.1% per input recipient cells; conjugations performed in liquid medium were generally unsuccessful, which was expected since conjugation by RP4 (IncP-type) machinery is enhanced on solid medium (Bradley et al., 1980). Importantly, this conjugation efficiency was robust enough to support the use of a suicide-plasmid homologous-recombination-based integration method; exconjugants should appear at a 100–1000-fold less frequency than that obtained with a stable plasmid to account for the additional recombination event.

#### Designing a Mobilizable Suicide Plasmid pPK15534 for \textit{Z. mobilis}

We designed the suicide plasmid to easily track genomic recombination events (Figure 2). The origin of replication, \textit{oriC}, was selected from pACYC184, which was known not to replicate in \textit{Z. mobilis}. The RP4 \textit{mob} element from pSUP202 was chosen to enable conjugation from an \textit{E. coli} donor strain (Simon et al., 1983). A single \textit{SpeI} restriction site was introduced to facilitate cloning of target DNA to direct recombination of the plasmid into the \textit{Z. mobilis} genome. Inclusion of \textit{cat} on the plasmid backbone allowed selection of chloramphenicol resistance to identify single crossover recombination events in \textit{Z. mobilis}. The superfolder \textit{gfp} (Pedelacq et al., 2006), also on the plasmid...
FIGURE 2 | Scheme for deleting genes in Z. mobilis. Deletion of a gene in Z. mobilis is accomplished by cloning 500 bp of DNA from upstream (UP) of and downstream (DN) of the gene to be deleted into the suicide plasmid pPK15534. Next, the plasmid is introduced into Z. mobilis by conjugation and single crossover homologous recombination events are selected for with chloramphenicol. Shown here is a crossover that occurred at the UP location, but recombination at the DN location is equally possible. Lastly, growing cells without selection allow detection of a second recombination event that results in plasmid loss by screening for the loss of fluorescence from GFP. Wild-type and deletion mutant genotypes were distinguished by PCR amplification.
backbone, allowed direct tracking of the plasmid and sorting by FACS. The latter enabled rapid enrichment of non-fluorescent cells that had undergone additional recombination events and lost the plasmid.

**pPK15534 Can Be Used as a Vehicle to Efficiently Delete Genes in Z. mobilis**

As a proof of principle, we tested whether we could efficiently delete an average size gene, choosing the 1.7 kb \textit{ldh} gene encoding lactate dehydrogenase (\textit{ldh}, ZMO0256), since its function should be dispensable in \textit{Z. mobilis} (Zou et al., 2012; Strazdina et al., 2018; Jones-Burrage et al., 2019). Our suicide plasmid (pPK15534) was engineered to contain two 500 bp sequences, immediately flanking but not including the target gene (pPK15535) (Figure 2). These sequences should direct homologous recombination of the plasmid into the genome, crossing over either upstream or downstream of the target gene within the genome. Following conjugation of the plasmid into \textit{Z. mobilis}, we isolated \textit{Cm} \textsuperscript{R} exconjugants at a frequency of $10^{-5}$, a reasonable frequency for a single recombination event following conjugation. We confirmed the integration location of

![Diagram of gene deletion workflow](image)

**FIGURE 3** | Deletion of \textit{ldh} gene. (A) Workflow for enrichment of non-fluorescent cells and screening for the modified genome. (B) Plot from FACS of \textit{Z. mobilis} strain PK15557 that has suicide plasmid (pPK15535) recombined into the genome. The X-axis represents fluorescence intensity and Y-axis represents the total number of sorted events. The sorting gate for collecting the non-fluorescent cells (red peak) is indicated by the bar. (C) Top plate shows the fluorescence image of colonies from FACS sorted non-fluorescent cells plated on non-selective media. Bottom plate shows the same colonies viewed with visible light. (D) The location of primers used to screen non-fluorescent candidates for wild-type (wt) and deletion genotypes by PCR is indicated by arrows. (E,F) Agarose gel electrophoresis of products from PCR amplification of 10 non-fluorescent-candidate colonies. Selected size markers are indicated on the left (MR). (E) Amplification using primers P34 and P35 yielded a 1.25-kb DNA fragment for the \textit{ldh} deletion whereas the wt \textit{ldh} DNA fragment was 2.97 kb. Strains that still possess the plasmid integrated into the genome (Rec) are expected to have a band of 7.8 kb. (F) Amplification using P34 and P35 yielded a DNA fragment of 0.8 kb, confirming those strains that still possess the \textit{ldh} gene, while strains in which \textit{ldh} has been deleted (six total) lacked the equivalent amplified fragment. Labels are the same as in panel E. Similar results were obtained when this experiment was biologically replicated three times.
FIGURE 4 | Deletion of bcsABC operon. Non-fluorescent deletion candidates for bcsABC were isolated using the same workflow as outlined for ldh and screened for deletions or wild-type (wt) alleles by PCR amplification. (A) The location of primers used to distinguish between the wt and the deletion genotype is indicated by arrows. (B,C) Agarose gel electrophoresis of products from PCR amplification of five non-fluorescent candidate colonies. (B) Amplification using primers P22 and P23 yielded a DNA fragment of 1.4 kb for the deletion of bcsABC (three total), while wt cells are expected to produce an amplicon of ~10 kb, which was poorly amplified under our reaction conditions. (C) To verify the genotype of wt strains, a second PCR experiment using primers P21 and P22 was included. In this case, wt strains are expected to yield an amplicon of 0.7 kb while the deletion strain should not produce a band. Similar results were obtained from six biological replicates of this workflow.

To test the phenotype of our mutant, we compared flocculation of the wt (left) and deletion strain (right) after growth for 15 h in glucose minimal media under aerobic conditions. Flocs are indicated by the arrow. (E) Growth curve of wt and mutant strains grown in glucose minimal media under anaerobic conditions.

one representative recombinant (PK15557) by PCR amplification across ldh using a pair of primers annealing at a region further upstream and downstream of the plasmid ldh targeting sequences (Figure 3D). The recombinant strain (shown in Figure 3E) also showed minor bands for both the genotype of wild type and the deletion of ldh indicating that these variants were present in the population at a low level.

To identify the rare cells in the population that underwent an additional crossover to lose the integrated plasmid, we took advantage of the fact that these cells will also be non-fluorescent because of the loss of gfp. We grew the recombinant strain for seven cell doublings without chloramphenicol selection and then enriched for the desired population of gfp− cells by using a fluorescence-activated cell sorter (FACS). Cells were sorted to obtain 10,000 events that were binned into the gate indicative of non-fluorescent cells (Figures 3A,B). We typically observed that viability of the sorted cells was consistently 20% or less but only those sorted into the gfp− bin. Possibly, in the nested gate that we use for gfp− cells, unwanted dead cells or debris of size similar to the Z. mobilis cells also appear. Nevertheless, we routinely obtain hundreds of colonies after plating the non-fluorescent cells. Visualization of the plates with an imaging system showed that >90% of the colonies were non-fluorescent (Figure 3C).

To distinguish crossover events that regenerate wild type ldh from the ldh deletion, we used PCR amplification (Figures 3D–F). Of 10 non-fluorescent colonies screened by PCR, 6 isolates had a DNA fragment of the size expected for the ldh deletion strain (Figures 3D–F), whereas four were wild type. We showed that the deletion endpoints were as predicted from our plasmid design by Sanger sequencing of the ldh gene region (Supplementary Figure S1). Thus, because the FACS sorting step takes just a few minutes, this strategy for enriching for non-fluorescent cells and then screening non-fluorescent colonies by PCR amplification allowed rapid identification of strains containing the deletion of interest with no other markers.

**Deletion of bcsABC Operon**

Next, we tested whether we could efficiently delete a larger size operon, choosing the 8.3 kb bcsABC operon, encoding functions for cellulose synthesis, since its function is known to be dispensable in Z. mobilis (Deutschbauer et al., 2014; Xia et al., 2018; Jones-Burrage et al., 2019). In addition, elimination of Z. mobilis bcsABC should minimize flocculation in liquid medium, which hinders accurate measurements of optical density (OD600) (Figure 4). Our suicide plasmid was engineered to contain two 500 bp sequences, immediately

Continued
flanking the bcsABC operon (pPK15538). The plasmid was introduced by conjugation, and recombinants were confirmed by PCR. Following culturing without selection, we sorted cells to obtain 10,000 non-fluorescent events and plated for single colonies. Of five non-fluorescent colony candidates screened by PCR, three were confirmed to have the desired deletion of the bcsABC operon (Figure 4). We showed that the deletion endpoints were as predicted from our plasmid design by Sanger sequencing of the bcs gene region (Supplementary Figure S1). This example shows that deletion of a longer sequence does not affect the efficiency of obtaining deletions. To determine if the strain lost the flocculating phenotype, we grew the wild type and the deletion strain in minimal medium without shaking. As expected, wild type produced flocs while the deletion strain did not produce flocs (Figure 4D). Further, growth curves with the deletion strain showed a smooth curve unlike wild type (Figure 4E).

**Insertion of crtIEB Genes Into the Genome**

For strain engineering, a critical tool is to be able to equip bacteria with novel pathways to produce products of interest. Thus, we also tested whether we could use our suicide plasmid to deliver heterologous genes and recombine them into the genome. An advantage of this method is that these added genes would be stably maintained on the genome and would require no added antibiotics for selection. We demonstrate this by replacing the ldh gene with the heterologous crtIEB operon (CRT) which produces an easily detected orange-pigmented carotenoid, neurosporene (Figure 5A) from *R. sphaeroides* (Su et al., 2018).

In this case, we placed the crtIEB genes from *R. sphaeroides* between the ldh targeting sequences on plasmid pPK15564 (Figure 5B). We also optimized RBSs (Salis et al., 2009; Borujeni et al., 2014) upstream of each gene. The plasmid was introduced by conjugation, and recombinants were confirmed by PCR. Following culturing without selection, we sorted cells to obtain 10,000 non-fluorescent events and plated for single colonies. Visualization of the plates indicated that some of the non-fluorescent colonies were also distinctly orange (Figure 5C). PCR screening of two non-fluorescent pigmented colonies confirmed the insertion of the operon at the ldh gene site (Figures 5D,E). We showed that the insertion junctions were as predicted from our plasmid design by Sanger sequencing (Supplementary Figure S1). Spectral analysis of acetone–methanol extracts of these strains showed an absorbance spectrum characteristic of neurosporene (Figure 5F; Takaichi and Maoka, 2015).

**Optimization of Expression of Genes in Z. mobilis**

We initially encountered insufficient expression of gfp in Z. mobilis for differentiating fluorescent from non-fluorescent colonies or cells. The ribosome binding site for gfp, originally obtained from a stable vector pRL814 was sufficient to produce fluorescent colonies from a plasmid but not when integrated

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**FIGURE 5** | Insertion of a functional CRT cassette. (A) A heterologous pathway for production of the carotenoid neurosporene by introduction of the crtIEB genes from *Rhodobacter sphaeroides*. Dotted arrows indicate the reactions mediated by the *R. sphaeroides* genes products. (B) Schematic of suicide plasmid used to deliver crtIEB to *Z. mobilis* ldh. UP and DN are the same 500 bp sequences used to delete the ldh gene. (C) Non-fluorescent candidates containing crtIEB replacing *Z. mobilis* ldh were isolated using the same workflow as outlined for ldh and screened for insertions first by the presence of orange colored colonies. (D) The location of primers used to distinguish between the wt and the insertion genotype is indicated by arrows. (E) Agarose gel electrophoresis of products from PCR amplification of orange colored colonies. Amplification using primers P34 and P35 yielded a DNA fragment of 5.4 kb for the insertion of crtIEB, whereas wt cells are expected to produce an amplicon of 2.9 kb. Selected size markers are indicated on the left (Mx). (F) Absorption spectrum of acetone:methanol extracts from orange colored colonies containing crtIEB compared to wt. Peak maxima characteristic of neurosporene are marked. We examined more than five isolates for insertion of the CRT cassette in independent experiments, and measured neurosporene from three different isolates. We conducted two independent experiments both resulting in the successful insertion of the CRT cassette at the ldh locus.
into the genome (Z. mobilis PK15556, Figure 6B). To improve the robustness of the GFP fluorescence screening, the ribosome binding site of gfp was modified guided by the results from the RBS calculator for maximizing predicted RBS strength (Salis et al., 2009; Borujeni et al., 2014). The difference between the original sequences and the optimized ribosome-binding site that we used to optimize expression of gfp for our suicide plasmid is shown in Figure 6A. Similar optimization of the crtIEB genes using the RBS calculator was required to detect pigmented colonies for neurosporene production (data not shown).

**DISCUSSION**

This work describes the development of a new flexible method for genome engineering in Z. mobilis, and how this method can be used for both knocking out genes and introducing novel sequences. We demonstrated the general utility of our genome engineering method by creating two targeted gene deletion strains and by adding a functional heterologous operon to a defined position in the Z. mobilis genome. The suicide plasmid and general approach should also be useful to other non-model organisms that benefit from the simple broad host range conjugation delivery system and recombination-based strategy.

Currently, deletion of genes in Z. mobilis relies on recombineering strategies to replace a gene typically with an antibiotic resistance marker cassette. Removal of this marker to allow subsequent genome modification has been demonstrated by a recombination reaction using FLP recombinase (Zou et al., 2012) when the marker is flanked with FRT sequences. However, this approach is time consuming due to the additional steps needed to introduce and remove the stable plasmid encoding FLP recombinase. In addition, FRT sequences remain in the genome providing sites for unwanted recombination, which could negatively impact iterative strain design. Counter selection with sacB on sucrose media (Xia et al., 2018) is feasible but relies on the desired phenotype or the native Z. mobilis sacB from not negatively impacting the sucrose selection. Our approach of screening for the loss of GFP fluorescence should not negatively impact the desired phenotype as far as we are aware.

We also showed that the frequency of a single crossover event in Z. mobilis averages 0.3%, which indicates that growth of a recombinant strain for just a few cell doublings generates sufficient cell mass to screen for a second recombination event. FACS facilitated the easy isolation of these rare events from the total population by sorting. Overall, this approach allows creation of strains containing a markerless addition or removal of a gene in two recombination events ad infinitum. This approach will be extremely useful in introducing genes for new pathways into Z. mobilis, which can be added at any position in the genome. By simply altering the Z. mobilis sequences flanking the genes of interest on the suicide plasmid, genes can be added to replace native ones, can be incorporated into native operons, or as stand alone operons, demonstrating the versatility of this approach. Our method is also not limited to FACS availability, since one can screen for non-fluorescent colonies by plating the recombinant cell suspension grown without selection and visually identify colonies that have lost the fluorescence using an imager.

**CONCLUSION**

In summary, our method has several highly desirable attributes; namely, it is markerless, it can be greatly accelerated with FACS, and it leverages the simple and efficient process of conjugation. Critical to this technique’s success, especially for use with FACS, was our optimization of the expression of gfp by RBS optimization. This enabled the visualization of fluorescent colonies and sorting of cells by FACS from a single, chromosomally inserted copy of gfp. We expect our method to be of value to genetic studies in the field, and our introduction of the use of FACS to enrich for desired phenotypes to be of exceptional value to Z. mobilis studies in the future. Finally, this method provides a simple approach for metabolic engineering where iterative strategies for genome engineering are necessary to build strains with desirable traits through sequence addition or deletion for producing molecules of interest.
DATA AVAILABILITY STATEMENT

All datasets generated for this study are included in the manuscript/Supplementary Files.

AUTHOR CONTRIBUTIONS

PL and PK conceived the overall study. PL, PK, RL, and IG designed the experiments. PL, FW, and YL carried out the experiments. PK and PL drafted the manuscript and all authors agreed on the final version of the manuscript.

FUNDING

This work was supported by the Great Lakes Bioenergy Research Center and the United States Department of Energy, Office of Science, Office of Biological and Environmental Research under Award Numbers DE-SC00018409 and DE-FC02-07ER64494.

ACKNOWLEDGMENTS

We thank Dr. Timothy Donohue and Rachelle Lemke for providing R. sphaeroides 2.4.1 genomic DNA as a source of crtIEB genes; Dr. William W. Metcalf for strain WM6026; Dr. Yaoping Zhang for plasmid pSUP202; and Dr. Vatsan Raman for use of his FACS. We also thank Dr. Tomas Gverdzys, Dr. Isabel Askensay, Dr. Erin Mettett, Dr. Yaoping Zhang, and Dr. Donna Bates for proofreading and valuable suggestions on this manuscript. We are also thankful to Laura C. Vanderploeg for helping in image capturing and making figures.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2019.02216/full#supplementary-material

FIGURE S1 | Sequence of the modified genome of Z. mobilis strains. To verify the structure of the insertion or deletion strains, we sequenced across the junction regions of genomic DNA. (A) For the idh deletion strain, genomic DNA was amplified using primer P34 and P35 and sequenced using primer P34. (B) For deletion of tcsABC, genomic DNA was amplified using primer P22 and P23 and sequenced using primer P22. (C,D) For insertion of the CRT cassette at the idh locus, we amplified upstream of the CRT cassette using primer P34 and P35 and sequenced using primer P37 and P35 (D) and sequenced using primer P35. Colored arrows indicate junctions of deletion or insertion of heterologous sequence. Top line represents expected sequence of the DNA fragment and the bottom line represents the sequencing data.
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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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