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Digital droplet PCR and IDAA for the detection of CRISPR indel edits in the malaria species Anopheles stephensi

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ABSTRACT
CRISPR/Cas9 technology is a powerful tool for the design of gene-drive systems to control and/or modify mosquito vector populations; however, CRISPR/Cas9-mediated nonhomologous end joining mutations can have an important impact on generating alleles resistant to the drive and thus on drive efficiency. We demonstrate and compare the insertions or deletions (indels) detection capabilities of two techniques in the malaria vector mosquito Anopheles stephensi: Indel Detection by Amplicon Analysis (IDAA™) and Droplet Digital™ PCR (ddPCR™). Both techniques showed accuracy and reproducibility for indel frequencies across mosquito samples containing different ratios of indels of various sizes. Moreover, these techniques have advantages that make them potentially better suited for high-throughput nonhomologous end joining analysis in cage trials and contained field testing of gene-drive mosquitoes.

METHOD SUMMARY
Mosquito DNA was extracted with the Promega Wizard® Genomic DNA Purification Kit protocol and quantified with Qubit® 3.0 following manufacturer protocols. PCR products for IDAA and ddPCR were generated with primers spanning 150–500 bp around the target site. IDAA amplicons were sent directly to COBO Technologies for analysis. ddPCR amplicons were analyzed using the Bio-Rad QX200™ ddPCR system.

KEYWORDS
CRISPR-Cas9 • ddPCR • gene editing • IDAA • mosquitoes • NHEJ quantification

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CRISPR/Cas9 gene-editing technology has transformed the field of genome modification. This system is composed of two fundamental components that interact to form a complex: Cas9 endonuclease and sgRNA, a target-specific RNA that guides Cas9 to the desired genomic DNA target site. Cas9 induces a double strand break at the target site, activating the DNA repair pathways of homology-directed repair (HDR) and nonhomologous end joining (NHEJ). HDR can induce accurate gene repair of one to thousands of base pairs in the presence of a homologous donor molecule, allowing for the correction of point mutations and introduction of exogenous sequences. In contrast, NHEJ produces genetic lesions comprised of random sizes of small insertions or deletions (indels) that alter the target site and can disrupt gene function. The HDR mechanism offers the opportunity to genetically modify large populations of arthropods, among other model organisms, by integrating the Cas9 endonuclease gene, the sgRNA targeting the desired locus and a dominant marker (fluorescent protein). The cassette is autonomous and can replicate to the homologous chromosome through HDR. This process effectively converts a heterozygous organism into a homozygote for the desired synthetic cassette, resulting in a selfish pattern of inheritance [1]. The nature of this type of genetic modification is designated gene drive and has been proposed as a tool for genetically modifying mosquito populations [2,3].

Gene drive in mosquitoes has been proposed as a promising tool for combating malaria and other mosquito-borne diseases, including dengue and zika [4], either by population suppression by spreading a lethal gene in wild-type (WT) mosquito populations to cause population crash or by replacement through the introduction of an anti-pathogen gene into a WT population. Recent progress demonstrated that CRISPR/Cas9 gene-drive-derived systems drive target-specific gene conversion at ≥99.5% efficiency in transgene heterozygotes of the Anopheles stephensi AsMCRkh2 line [5]. Gene drive efficiency depends on the availability of WT or susceptible alleles targeted by the gRNA-directed Cas9 cleavage. When a susceptible chromosome has been mutated by NHEJ, the key nucleotides necessary for gRNA recognition could be mutated or eliminated, thus preventing subsequent HDR-mediated gene conversion in the mosquito germline. An accumulation of NHEJ events has a diminishing effect on the drive, and the mosquito progeny approach Mendelian inheritance of the introduced DNAs due to the generation of drive-resistant loci [5,6]. Methods to detect NHEJ events rely on artificial reporter assays, gel-based systems, Sanger sequencing and deep sequencing [7–9]. None of these methods is suitable for high-throughput screening of NHEJ alleles in samples from multiple, large-cage populations or field trials due to their technical complexity, cost and time or labor required. A resistant Cas9-induced NHEJ allele percentage is considered acceptable when it is lower than the naturally occurring single nucleotide polymorphisms (SNPs) at the target site in the wild population [10]. This percentage can be tolerated while not affecting drive fixation; therefore, NHEJ quantification is an essential parameter during laboratory and field trials. Detecting indels in large populations of mosquitoes over many generations requires a high-throughput method that maximizes efficiency and provides
sensitive, accurate results. To circumvent the difficulties of conventional techniques, we compared two novel techniques, Droplet Digital PCR™ (ddPCR™; Bio-Rad Laboratories, CA, USA) and Indel Detection by Amplicon Analysis (IDAA™; COBO Technologies, Copenhagen, Denmark) for NHEJ quantification in the A. stephensi AsMCRkh2 line carrying a CRISPR/Cas9 gene drive.

MATERIALS & METHODS

Sample sources

A. stephensi mosquitoes (Indian strain, gift of M. Jacobs-Lorena, Johns Hopkins University) maintained at the University of California, Irvine (UCI) insectary are the source of all insects used in the experiments. The gene-drive line AsMCRkh2 (gene drive) and WT (non-gene drive) mosquitoes were maintained at 27°C with 77% humidity and a 12-h day/night, 30-min dusk/dawn lighting cycle. AsMCRkh2 mosquitoes with indels were recovered from crosses between WT and AsMCRkh2 mosquitoes over 20 generations [11]. The Cas9-targeted sequence, 5′-GATGGTTCCGTTCTACGGGCAGG-3′ (protospacer adjacent motif sequence underlined), is in the gene encoding kynurenine hydroxylase (kh).

DNA extraction & quantification

Genomic DNA extraction was performed using the Wizard® Genomic DNA Purification Kit protocol (Promega, WI, USA) for mouse tails according to the manufacturer’s instructions. Pools of 10 adult mosquitoes were used for DNA extraction. DNA was resuspended in 50 μl of PCR-grade water. DNA extracts were quantified at the UCI Genomics High-Throughput Facility using a Qubit® 3.0 Fluorometer (Thermo Fisher Scientific, MA, USA) following the manufacturer’s instructions. One microliter of DNA extract was analyzed using the Qubit dsDNA HS Assay Kit followed by Qubit 3.0 quantification.

ddPCR drop-off assay

We prepared 25-μl reactions with 12.5 μl Bio-Rad ddPCR 2× Supermix for Probes (No dUTP), 10 μl DNA (0.9 ng/μl), 1.25 μl fluorescein amidite (FAM)/forward (5μM FAM probe, 18-μM forward primer) (Supplementary Table 1) and 1.25 μl hexachlorofluorescein (HEX)/reverse (5μM HEX probe, 18-μM reverse primer) (Supplementary Table 1) in a 96-well PCR plate. Twenty microliters from the PCR reactions were used for droplet generation, each theoretically containing 30,000 haploid genome copies per 20-μl reaction, assuming that one A. stephensi haploid genome is 0.24 pg [12]. Droplets were generated at the UCI Genomics High-Throughput Facility using a Bio-Rad QX200 Droplet Generator following the manufacturer’s instructions; they were then transferred to a Bio-Rad 96-well PCR plate and foil

Table 1. Insertions or deletions quantification in nonhomologous end joining mosquito samples from small-cage trials of the gene drive AsMCRkh2 strain.

| Number | Sample (cage generation) | ddPCR average indel (%) | IDAA average indel (%) |
|--------|--------------------------|-------------------------|------------------------|
| Indel-1 | A1-G3                    | 100.00                  | 100.00                 |
| Indel-2 | A1-G8                    | 99.97                   | 100.00                 |
| Indel-3 | A1-G14                   | 100.00                  | 100.00                 |
| Indel-4 | A1-G16                   | 100.00                  | 100.00                 |
| Indel-5 | A3-G4                    | 100.00                  | 100.00                 |
| Indel-6 | A3-G7                    | 100.00                  | 100.00                 |
| Indel-7 | A3-G8                    | 100.00                  | 100.00                 |
| Indel-8 | A3-G9                    | 99.97                   | 100.00                 |
| Indel-9 | A3-G10                   | 100.00                  | 100.00                 |
| Indel-10 | B1-G4                   | 100.00                  | 100.00                 |
| Indel-11 | B1-G7                   | 99.97                   | 100.00                 |
| Indel-12 | B1-G9                   | 100.00                  | 100.00                 |
| Indel-13 | B1-G10                  | 99.80                   | 100.00                 |
| Indel-14 | C1-G8                    | 100.00                  | 100.00                 |
| Indel-15 | C1-G11                  | 99.97                   | 100.00                 |

A total of 15 sample pools of ten mosquitoes each were obtained from different cages through several generations [11]. ddPCR and IDAA were used to analyze the same DNA extract of each sample to quantify the total percentage of indel sequences. Analysis was carried out in triplicate (n = 3) with averages shown. The Pearson correlation coefficient is r = 0.77 when comparing similarity trends. Student’s t tests performed for each individual sample yielded no statistical significance (p < 0.05) between the results of both techniques. IDAA and ddPCR are sensitive in detecting multiple types of indels in a pool sample, as there is no significant difference between the results of the two methods and the expected percentage of indel, which is 100% in all samples. Cage numbers refer to those in Pham et al. [11].

ddPCR: Droplet Digital PCR; IDAA: Indel detection by amplicon analysis; Indel: Insertions or deletion.
heat-sealed at 180°C for 5 s. PCR was performed using a Bio-Rad C1000 Touch™ thermal cycler with a 96-deep-well reaction module under the following conditions: 95°C for 10 min, 40 cycles of 94°C for 30 s, 55°C for 1 min and 60°C for 2 min, followed by 98°C for 10 min and a 4°C hold. A 2°C/s ramp rate was used for all steps. Droplets were read using the Bio-Rad QX200 ddPCR system. The data analysis was performed using Bio-Rad QuantaSoft™ Analysis Pro version 1.0.596 in drop-off mode requiring manual cluster designation.

**IDAA assay**

Samples were prepared from 25-μl PCR reactions using 0.5 U of TEMPase (Amplicon, Odense, Denmark) in 1× ammonium buffer with 2.5 mM MgCl₂, 200 μM dNTP, 5% DMSO, 0.25-μM Universal Fam-For, 0.025-μM forward-extension primer and 0.25-μM reverse-extension primer (Supplementary Table 1). PCR conditions included an initial incubation at 95°C for 15 min; 15 cycles of 95°C for 30 s, 72°C for 30 s and 72°C for 30 s, with the annealing temperature decreasing 1°C per cycle beginning from 72°C; and an additional 25 cycles of 95°C for 30 s, 58°C for 30 s and 72°C for 30 s, with 7 min of final extension at 72°C. PCR products were run in 3% agarose gel and analyzed directly. Samples were sent to COBO Technologies for fragment analysis and Profile IT Solutions (New Delhi, India) indel profiling and quantification.

**RESULTS & DISCUSSION**

ddPCR is based on mechanically emulsifying a PCR solution into thousands of nanoliter droplets, effectively monitoring thousands of PCR reactions individually and thereby vastly increasing accuracy and reproducibility. It utilizes two fluorescent probes to discern WT and indel sequences: a HEX probe targets the WT gRNA site, and a FAM probe targets a conserved sequence within the amplicon (Figure 1; Supplementary Table 1). Sequences that are WT will give a fluorescent signal for both probes, and sequences possessing indels at the gRNA cut site display only a FAM signal, with the HEX probe failing to anneal. Each PCR-amplified nanoliter droplet is measured for these fluorescent signals, allowing statistically powerful quantification of indels present in PCR reactions. The alternative technique, IDAA, utilizes triple-primer PCR amplification to fluorescently tag the amplicon that includes the gRNA target site (Figure 1) [13]. Amplicons have their base-pair
length measured by capillary gel electrophoresis; the WT length is used as a standard, and sequence lengths that differ are designated indels. The fluorescent signal allows unbiased quantification of amplicons, and the indel size is capable of being determined, importantly, without dependence on prior knowledge of the nature of the indels induced after Cas9:gRNA targeting.

IDAA and ddPCR were tested with a variety of indel mosquito samples to verify their sensitivity toward multiple mutations at the target sites (Figure 1). We examined mosquito samples obtained from a series of small-cage trials of the gene drive AsMCRkh2 strain of the Asian malaria vector mosquito, A. stephensi [5,11]. We analyzed 15 pools of 10 mosquitoes each that were considered to have a NHEJ by phenotype selection (white-eye and DsRed-negative) based on previous data [5]. However, because the kh mutant white-eye phenotype is associated with a recessive mutation, no phenotypic selection was possible until the second generation (G2). Previous work with these NHEJ mosquitoes had shown that Cas9 indel mutations happened at and around the cut site and protospacer adjacent motif sequence, resulting in insertions and deletions of multiple lengths ranging from 1 to 473 bp [11]. With 185 NHEJ individuals analyzed by Sanger sequencing, 50 different types of indels were identified, including three types of 1-bp indels (from 15 individuals, ~8% of 185) and one type of substitution (from 1 individual, ~0.5%) [11]. Two sets of samples were generated: NHEJ samples, which contained only pools of confirmed NHEJ individuals obtained from previous cage experiments with white-eye and DsRed-negative phenotypes to challenge the sensitivity of the two techniques toward different types of NHEJ, and mixed samples, which were generated by using DNA extracted from a mixture of NHEJ mosquito samples with WT mosquitoes at different proportions to quantify the NHEJ proportion in those samples.

Results from both IDAA and ddPCR experiments for the NHEJ samples showed that both techniques were able to detect all mutant sequences in the NHEJ mosquito samples with an indel percentage of 100%. All samples were analyzed by three technical replicates, with the average total percentage of indels shown in Table 1 and Supplementary Table 2. Both IDAA and ddPCR provide a quantitative analysis of total indel percentage, but only IDAA details the indel sizes and their respective proportions in a sample. Based on the IDAA analysis, indels were detected in a range from 1 to 469 bp, including insertions and deletions, thus representing a broad variety of Cas9-induced indels (Table 2). Many samples contained multiple different indels that were quantified for the proportional amount of each indel present in the sample. Sequencing data for some chosen NHEJ individuals are not available here.

| Table 2. Insertions or deletion lengths detected by IDAA. |
| --- |
| Indel source | Frameshift indels (%) | Length (%) |
| | First top indel | Second top indel | Third top indel | Fourth top indel | Fifth top indel |
| A1-G3 | 100 | 2 (52.8) | -2 (47.2) | - | - |
| A1-G8 | 100 | -8 (61.9) | 7 (27.0) | -11 (11.1) | - |
| A1-G14 | 42.8 | -11 (34.5) | -3 (27.4) | -6 (20.1) | 18 (9.7) |
| A1-G16 | 100 | -13 (85.3) | 469 (14.7) | - | - |
| A3-G4 | 51.2 | -48 (48.8) | 11 (22.0) | -13 (17.5) | 8 (7.3) |
| A3-G7 | 100 | 5 (54.0) | 8 (46.0) | - | - |
| A3-G8 | 100 | 1 (62.4) | 11 (37.6) | - | - |
| A3-G9 | 53.9 | -33 (46.1) | 11 (44.7) | 1 (4.9) | -34 (4.3) |
| A3-G10 | 32.1 | -33 (65.5) | 1 (26.2) | -34 (6.0) | - |
| B1-G4 | 14.1 | -6 (85.9) | -7 (13.0) | -29 (1.1) | - |
| B1-G7 | 100 | -4 (51.5) | 1 (27.5) | -14 (11.5) | 8 (9.5) |
| B1-G9 | 100 | -4 (54.7) | 1 (39.9) | -5 (5.5) | - |
| B1-G10 | 100 | -4 (90.7) | -5 (9.3) | - | - |
| C1-G8 | 100 | -10 (51.6) | 2 (20.0) | 2 (19.0) | -14 (9.5) |
| C1-G11 | 100 | -4 (90.4) | -5 (9.6) | - | - |

IDAA allows quantification of each indel sequence of different length, giving insight into the indel composition of the DNA extract. IDAA analysis was done in triplicate (n = 3) with the averages of the top five most prevalent indel lengths displayed from each sample source, the same source used for the analysis detailed in Table 1. Total frameshift indel percentages are provided by excluding indel sizes that are divisible by three. Indels found across all samples range from deletions of 48 bp to insertions as large as 469 bp. Identified mutations included insertions (+) and deletions (-) with different lengths as small as 1 bp. The percentage of each indel is shown in parentheses. ddPCR: Droplet Digital PCR, IDAA: Indel detection by amplicon analysis; Indel: Insertions or deletion.
listed in Supplementary Table 3, and these confirmed the sensitivity accuracy of IDAA and ddPCR for different types of indels. Not all indels were identified with Sanger sequencing because of the time and labor costs necessary to extract and sequence all individuals, limited sources of genomic DNA from single mosquito extractions and PCR technical problems. In addition, not all samples were suitable for Sanger sequencing because DNA was extracted from mosquito pools and included a mixture of mutations. This level of complexity reduced the reliability of PCR amplification because not all of the mutations could be amplified with the same efficiency due to variants of different indel frequencies, resulting in nonspecific sequencing errors. Providing sequencing data for each sample via next-generation sequencing would cost, unnecessary and difficult due to the abovementioned reasons regarding the quality of DNA extracts and amplification of different indels in a pooled sample.

Sensitivity and quantification of 1-bp insertions by IDAA can be seen in samples G8A3, G9A3, G10A3, G7B1 and G9B1 (Table 2 & Supplementary Table 3). The same DNA extracts from all samples were used for both techniques, allowing a direct comparison of indel quantification. Because the ddPCR technique designated the same sample extracts at or near 100% indel, it demonstrates that the 1-bp insertions in those samples are being reliably detected (Table 1). This is consistent with prior data supporting the 1-bp indel sensitivity of both IDAA and ddPCR [13,14]. Overall, every indel size discovered by the IDAA method was detected by ddPCR, as it designated all samples as 100% or near 100% indel with no significant differences observed between individual samples and a strong correlation coefficient of 0.73 (Table 1). If ddPCR were insensitive to a certain indel identified in a sample by IDAA, then the total indel percent determined by ddPCR would proportionally reflect an increase in WT percentage. Samples G8A1, G9A3, G7B1, G10B1 and G11C1 were slightly below 100% indel frequency in either technique, and this is likely due to fluorescence anomalies. The nominal WT sequence quantified in these samples (0.03–0.8%) is unreliable because, at its lowest frequency, a true WT allele in a pool of ten indel mosquitoes (20 alleles total) would produce a 5% WT (or 95% indel) proportion, which was not observed.

In order to assess accuracy and replicate a trial scenario in which quantification techniques are employed, 11 pooled samples of NHEJ mosquitoes were made with WT mosquitoes at different ratios of WT:NHEJ mosquitoes (10:0, 9:1, 8:2, 7:3, 6:4, 5:5, 4:6, 3:7, 2:8, 1:9 and 0:10). DNA was used for PCR, and amplicons were subjected to IDAA and ddPCR analysis to determine the indel percentage in each sample. Each ratio was conducted in triplicate (n = 3), and average results were compared between the two techniques. Although deviating from theoretical indel percentages (40% indel for a 6:4 ratio), both techniques demonstrated precision based on producing similar results for each ratio and having a Pearson correlation coefficient of r = 0.99. In addition, values provided by ddPCR and IDAA are also representative of their accuracy; because both deduced the same indel percentage, it is likely close to the actual indel percentage. Student’s t test was performed to compare the measurements at each ratio (*p < 0.05).

dPCR: Droplet Digital PCR; IDAA: Indel detection by amplicon analysis; Indel: Insertions or deletion; NHEJ: Nonhomologous end joining; WT: Wild-type.

Figure 2. Quantification of nonhomologous end joining alleles in AsMCRRk2 mosquito samples by ddPCR and IDAA techniques. DNA was extracted from 15 to 10 mosquito pools. To assess the sensitivity of both techniques, the mosquito pools consisted of a mix of WT and NHEJ mosquitoes at 11 different ratios of WT:NHEJ (10:0, 9:1, 8:2, 7:3, 6:4, 5:5, 4:6, 3:7, 2:8, 1:9 and 0:10). DNA was used for PCR, and amplicons were subjected to IDAA and ddPCR analysis to determine the indel percentage in each sample. Each ratio was conducted in triplicate (n = 3), and average results were compared between the two techniques. Although deviating from theoretical indel percentages (40% indel for a 6:4 ratio), both techniques demonstrated precision based on producing similar results for each ratio and having a Pearson correlation coefficient of r = 0.99. In addition, values provided by ddPCR and IDAA are also representative of their accuracy; because both deduced the same indel percentage, it is likely close to the actual indel percentage. Student’s t test was performed to compare the measurements at each ratio (*p < 0.05).

Table 1: WT:NHEJ mosquito ratios

| WT:NHEJ mosquito ratios | ddPCR | IDAA |
|-------------------------|-------|------|
| 10:0                    | 100   | 100  |
| 9:1                     | 90    | 90   |
| 8:2                     | 80    | 80   |
| 7:3                     | 70    | 70   |
| 6:4                     | 60    | 60   |
| 5:5                     | 50    | 50   |
| 4:6                     | 40    | 40   |
| 3:7                     | 30    | 30   |
| 2:8                     | 20    | 20   |
| 1:9                     | 10    | 10   |
| 0:10                    | 0     | 0    |

Figure 3A. Randomly chosen individuals analyzed with Sanger sequencing confirmed the results obtained by IDAA and ddPCR and showed that the detected indels are accurate (Figure 3B). Some indels were identified by IDAA but were not identified with Sanger sequencing (-16 in G7 and +1 in
**Amplicon size/bp**

- **Relative fluorescence units**

  - **G0**
    - 450 500
    - 15,167.02
    - 14,000
    - 13,000
    - 12,000
    - 11,000
    - 10,000
    - 9000
    - 8000
    - 7000
    - 6000
    - 5000
    - 4000
    - 3000
    - 2000
    - 1000
    - 0

  - **G1**
    - 450 500
    - 30,000
    - 20,000
    - 10,000
    - 0

  - **G7**
    - 450 500
    - 30,000
    - 20,000
    - 10,000
    - 0

  - **G9**
    - 450 500
    - 30,000
    - 20,000
    - 10,000
    - 0

**B**

- **WT**
  - GCATGACGCAGTTCCGTTTCATGAGAAGGTGACGTAGC
  - +4 GCATGACGCAGTTCCGTTTCATGAGAAGGTAGC
  - +1 GCATGACGCAGTTCCGTTTCATGAGAAGGTGACGTAGC
  - +8 GCATGACGCAGTTCCGTTTCATGAGAAGGTGACGTAGC
Figure 3. Tracing and quantification of insertions or deletions in AsMCRkh2 mosquito samples over generations (see facing page). G0 and G1 mosquito samples were chosen randomly, and samples from G7 through G10 were all individuals carrying NHEJ alleles selected by phenotype (white-eyed/ DsRed-negative). (A) G0: ‘Founder’ individuals show the baseline for the IDAA profile. Both males and females present a WT sequence at the target site shown by a yellow peak (because all female G0s are WT, whereas even though G0 males were a combination of transgenic and WT males only WT [and NHEJ alleles, if there were any] were amplified by PCR). G1: First-generation offspring display expected low frequency of indels in sample pools of WT and low-frequency NHEJ individuals. Red-dotted line zoom-in inserts display the rarely occurring NHEJ indel events in the population (<0.8%), and the black and gray triangles indicate spectra peaks of indels. Two types of indels, -11 and -4, were identified in G1. IDAA and ddPCR allowed the analysis of a large number of samples from G0 and G1, which was required to determine NHEJ allele-generated frequency. G7–G10: White-eyed phenotype mosquitoes with homozygous NHEJ alleles were selected from different generations. Although phenotypically similar, the variable peak heights indicate that G7 individuals represent a heterogeneous population, with different types of indels, whereas G9 (near equal peak heights) and G10 (single peak) represent a homogeneous population with only one indel (-4) selectively carried to subsequent generations. WT alleles are distinguished by yellow peaks when present in the spectra; when absent, yellow triangles above the spectra panels are used to reference the WT location. Frameshift-causing indels are indicated with peaks color-coded in blue. (B) Sanger sequencing in mosquitoes from G7 (11 individuals), G9 (5 individuals) and G10 (16 individuals) show results comparable with the IDAA findings. Three types of indels (-4, +1, +8) were identified in G7 mosquitoes, whereas only one type of indel (-4) was present in G9 and G10 (Supplementary Table 3).

ddPCR: Droplet Digital PCR; IDAA: Indel detection by amplicon analysis; Indel: Inserts or deletion; NHEJ: Nonhomologous end joining; WT: Wild-type.
samples with indels are rare, ddPCR or IDAA can be coupled with sequencing to acquire sequence data while maintaining high-throughput efficiency.

IDAA and ddPCR showed sensitive and reproducible detection of NHEJ events in mosquito samples from cage experiments. Both techniques offer a more efficient analysis of indel quantification in a cost- and time-saving manner, and they can be used for efficient analysis of gene-drive mosquito populations for quantifying NHEJ. Thus, they possess the qualifications to determine factors that will influence gene drive in cage trials or field releases.

FUTURE PERSPECTIVE

As CRISPR-Cas9-based gene drives are being widely developed for applications in vector control, ecology conservation and pest management, cage trials and field trials will likely become regulatory checkpoints for deploying gene drives in living organisms into the field. A high-throughput yet cost-effective method to determine NHEJ alleles compared with HDR for drive efficiency is necessary for the study of gene-drive behaviors in big population samples format.

SUPPLEMENTARY DATA

To view the supplementary data that accompany this paper please visit the journal website at www.future-science.com/doi/suppl/10.2144/btn-2019-0103

AUTHOR CONTRIBUTIONS

R Carballar-Lejarazu: Study conception and design, data analysis and interpretation, manuscript drafting and revision, approval for publishing and agreement to be accountable for the study and manuscript. A Kelsey: Study design and investigation, data collection, analysis and interpretation; manuscript drafting and revision; approval for publishing; and agreement to be accountable for the study manuscript. TB Pham: Study design and investigation; data collection, analysis and interpretation; manuscript drafting and revision; approval for publishing; and agreement to be accountable for the study and manuscript. EP Bennett: IDAA indel profiling and quantification, and manuscript revision. AA James: Data interpretation and manuscript revision.

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FINANCIAL & COMPETING INTERESTS DISCLOSURE

This study was funded by Bill and Melinda Gates Foundation (OPP1160739) and the University of California, Irvine Malaria Initiative. This technical support was not sponsored by Cobo Technologies or Bio-Rad Laboratories. EP Bennett declares that a patent application covering the IDAA method is pending, and he acts as a scientific advisor for Cobo Technologies. The authors have no other relevant affiliations or financial involvement with any organization or entity with a financial interest in or financial conflict with the subject matter or materials discussed in the manuscript apart from those disclosed.

No writing assistance was utilized in the production of this manuscript.

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