SOFTWARE NOTE

InDelGT: An integrated pipeline for extracting indel genotypes for genetic mapping in a hybrid population using next-generation sequencing data

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Abstract
Premise: Although several software packages are available for genotyping insertion/deletion (indel) polymorphisms in genomes using next-generation sequencing data, simultaneously calling indel genotypes across many individuals for use in genetic mapping remains challenging.

Methods and Results: We present an integrated pipeline, InDelGT, for the extraction of indel genotypes from a segregating population such as backcross or F2 lines, or from an F1 cross between outbred species. The InDelGT algorithm is implemented in three steps: generating an indel catalog, calling indel genotypes, and analyzing indel segregation. We demonstrated the use of the pipeline with an example data set from an F1 hybrid population of Populus and successfully constructed the two parental genetic linkage maps.

Conclusions: InDelGT is a practical tool that can quickly genotype a large number of indel markers within a population following Mendelian segregation. The InDelGT pipeline is freely available on GitHub (https://github.com/tongchf/InDelGT).

KEYWORDS
genetic mapping population, genotype calling, insertion/deletion polymorphism (indel), next-generation sequencing (NGS)

Insertion/deletion polymorphisms (indels) are a common and functionally important type of sequence variation (Albers et al., 2011). The insertion of mobile elements such as retrotransposons, slippage in simple sequence replication, or unequal crossovers during meiosis can all lead to indel generation (Britten et al., 2003). Consequently, indels are widely and densely distributed across the genome, with a distribution density second only to single-nucleotide polymorphisms (SNPs); indels are thus much more common than simple sequence repeats (SSRs). Indels can be multiallelic and codominant, potentially providing more genomic information than the typical biallelic SNPs (Lv et al., 2016). Indels in coding regions have a significant impact on the structure and function of the resulting protein, which in turn leads to phenotypic differences in the organism, meaning indels have made a significant contribution to the processes of speciation and phenotypic evolution (Tian et al., 2008). Indels have also become an effective marker system for genetic analysis due to their multiallelic nature, codominant inheritance, and wide genome coverage (Das et al., 2015).

With advances in next-generation sequencing (NGS), a large amount of DNA sequence data can be generated at low cost, facilitating the bioinformatic identification of genetic variants. A number of software programs have been developed for calling indel genotypes from sequence reads, including SOAPindel (Li et al., 2013), Pindel (Ye et al., 2009), VarScan (Koboldt et al., 2009), and GATK (McKenna et al., 2010). Although the analytical processes of these software programs are described in detail in the literature, each process involves multiple steps, with each step requiring the input of multiple parameters and commands. This means the analytical processes of the available...
indel-calling programs are all relatively complicated and difficult to use. Moreover, there is interest in using indels for constructing genetic maps because they are useful in marker-assisted selection and quantitative trait locus mapping (Li et al., 2015; Song et al., 2015); however, no specific tools have been developed for extracting indel genotypes across many individuals in a hybrid population for use in genetic mapping.

Here, we report an integrated pipeline called InDelGT, which can genotype a large number of indel markers in a population following Mendelian segregation to construct genetic linkage maps. InDelGT can be used to perform indel genotyping of traditional backcross (BC) and F2 populations from inbred lines, and we demonstrate its use here in an F1 hybrid population derived by crossing two individuals from two outbred species. InDelGT can complete the entire genotyping process by simply entering a command line, and it can perform multicore or multithreaded parallel computing, which greatly accelerates the indel genotyping process.

**METHODS AND RESULTS**

**Implementation of InDelGT**

InDelGT was implemented with Perl to extract indel genotypes in a hybrid population. It can be run in three steps: (1) generating two parental indel catalogs, (2) calling indel genotypes across a hybrid population, and (3) analyzing indel segregation. A flowchart of the implementation is shown in Figure 1.

**Generating two parental indel catalogs**

In the first step, InDelGT generates parental indel catalogs with the Perl program `parent_genotyping.pl`. First, the paired-end reads of each parent are aligned to a reference sequence using BWA, generating a SAM format file (Li et al., 2009). The SAM files for the two parents are converted to BAM format files using the sorting and indexing process of SAMtools. Subsequently, BCFtools is used to generate BCF files from the BAM files, then to further convert them into VCF files. Finally, the indel genotype of each parent at each site is extracted from its VCF file, ensuring a read depth of at least 3 for an allele and a genotype quality greater than 30. The indel sites from the two parents are merged and saved in a text file (namely ‘parent.cls’) as an indel catalog. With this catalog file, the genotypes of the female and male parents are called and saved in two files, “female_merge.gt” and “male_merge.gt”, respectively.

**Calling indel genotypes across a hybrid population**

In the second step, the Perl program `progeny_genotyping.pl` is used to call the indel genotypes of all progeny with the catalog file generated above. The calling procedures are similar to the generation of the parental indel catalog. After generating the BCF files for each individual, the VCF files are generated according to the indel catalog by using BCFtools. Next, indel genotyping is performed according to the VCF or BCF files. A heterozygous indel genotype can be easily determined for a sample from the records in its VCF file; however, when calling a homozygous genotype at an indel site, we need to know the site start and end positions in the parental indel catalog. Only when every position from start to end is homozygous can we consider the indel genotype to be homozygous. Finally, the indel genotypes at all indel sites are saved for each sample. The file contains the indel site positions, allele nucleotides, and genotypes.

**Analysis of indel segregation**

In the last step, InDelGT uses the Perl program `segregation_ratio.pl` to identify the segregation types of indels, to count the number of individuals with each genotype, and to calculate a P value to test whether each indel follows Mendel’s law of segregation. The indel sites that deviate from the Mendelian segregation ratio (P < 0.01) and have more than a certain percentage (e.g., 10%) of samples missing genotypes in the progeny are filtered out. The result files are saved in text files that vary for the different populations. For a BC or F2 population, InDelGT will only produce one result file because there is only one type of segregation in the population. By contrast, for a hybrid population between two outbred species, up to seven result files will be produced, corresponding to the seven segregation types: ab × aa, aa × ab, ab × cc, ac × cc, ac × bc, ab × ac, and ab × cd, where the first two letters of each type indicate the genotype of the female parent and the last two are the genotype of the male parent.

**Use of InDelGT**

InDelGT can run on Linux systems. To run InDelGT, users must download and install the necessary software packages, including BWA (Li and Durbin, 2009), SAMtools (Li et al., 2009), and BCFtools (Li et al., 2009). Moreover, a parameter setting file must be created, namely ‘parameters.ini’, which consists of three parts: folders, parameters, and data files (Figure 2). The first part, ‘folders’, gives the paths to software packages and data files, including BWA, SAMtools, BCFtools, and InDelGT itself, as well as the reference sequence file and FASTQ files. The second part, ‘parameters’, includes the minimum score of indel genotypes, the minimum score of the mapping quality of the reads, the percentage of missing genotypes allowed, the minimum P value allowed for testing the segregation ratio at an indel site, and the number of threads used for parallel computing. The third part, ‘data files’, shows the FASTA file of a reference sequence and the first and second read files of
FIGURE 1  Schematic diagram of the InDelGT workflow. Three steps are required to run InDelGT: (1) generating a parental indel catalog, (2) calling indel genotypes across a hybrid population, and (3) analyzing indel segregation. The blue rounded rectangles represent the internal program written in Perl. The ordinary yellow rectangles indicate the external software packages used in InDelGT, including BWA, SAMtools, and BCFtools. The green cut-corner rectangles represent text files containing the intermediate and final results.
the two parents and all progeny in FASTQ format. After installing the external software packages and preparing the parameter file, users can run InDelGT with a simple command line 'perl InDelGT.pl -o' followed by an output. InDelGT is freely available at https://github.com/tongchf/InDelGT. For users to quickly grasp the use of InDelGT, we have also made test data available online including a reference sequence file and the FASTQ files of two parents and 20 progeny.

Genotyping of indel markers in poplar: A case study using InDelGT

To demonstrate the function of InDelGT, we performed a real data analysis using two species in the genus Populus L. (poplar). Poplar has a small genome and high heterozygosity, making this an appropriate taxon to use for the evaluation of the computational performance of InDelGT. In previous studies, we performed restriction site–associated
DNA sequencing (RAD-seq) of the two parents and 117 progeny in an F1 hybrid population derived from a cross between *P. deltoides* W. Bartram ex Marshall and *P. simonii* Carrière (Mousavi et al., 2016; Tong et al., 2016). The raw data are available from the National Center for Biotechnology Information (NCBI) Sequence Read Archive (SRA) database (http://www.ncbi.nlm.nih.gov/Traces/sra) using the accession numbers listed in Appendix S1. After performing quality control on the data using the NGS QC toolkit (Patel and Jain, 2012), these RAD-seq data were used to call indel genotypes with InDelGT. We selected the *Populus trichocarpa* v4.0 (Tuskan et al., 2006) reference sequence to enable the calling of indel genotypes across all individuals, including the two parents. We set the genotype quality to be greater than 30, the coverage depth for each allele in a heterozygote as not less than 3, and the allele coverage depth in a homozygote as not less than 5. Additionally, we filtered out those indel sites for which more than 10% of the genotypes were missing data and with *P* values less than 0.01 in the data sets. According to the segregation types, the final genotype data were classified and saved in different text files. Consequently, a total of 50,532 indel sites were identified between the two parents, of which 2805 followed different Mendelian segregation ratios (*P* > 0.01) in the F1 hybrid population. The indel genotype data included the segregation types *ab × aa*, *aa × ab*, *ab × ab*, and *ab × ac*, with totals of 1539, 1262, three, and one occurrence, respectively (Table 1).

To validate the indel genotype data used for genetic mapping, we constructed two parental genetic linkage maps using the two indel data sets from the segregation types *ab × aa* and *aa × ab*. The results showed that the indels with the segregation type *ab × aa* were assigned into 19 linkage groups based on the maternal *P. deltoides* genome under a logarithm of odds (LOD) threshold ranging from six to 12 (Appendix S2), which was consistent with the karyotype of *Populus* (2n = 38) (Eckenwalder, 1996). In the same way, the indels with the segregation type *aa × ab* were divided into 20 linkage groups based on the paternal *P. simonii* genome under a LOD threshold ranging from six to 11 (Appendix S3). The maternal genetic linkage map contained 1125 indels spanning 3146.05 cM, with an average genetic distance between adjacent indels of 2.84 cM. The paternal genetic linkage map contained 1024 indels with a genetic distance of 3760.90 cM, and an average genetic distance of 3.75 cM.

| Female genotype | Male genotype | Expected segregation ratio | Number of indels |
|-----------------|---------------|----------------------------|------------------|
| ab              | aa            | 1:1                        | 1539             |
| aa              | ab            | 1:1                        | 1262             |
| ab              | ab            | 1:2:1                      | 3                |
| ab              | ac            | 1:1:1:1                    | 1                |
| Total           |               |                            | 2805             |

Table 1: Mendelian segregation patterns and the numbers of indel markers identified.

Table 2: The primer sequences of the eight indel markers developed and used in this study.
Experimental validation of indel genotypes

We randomly selected eight indels to verify the accuracy of our indel genotyping for 12 progeny using polyacrylamide gel electrophoresis (PAGE) technology (Table 2). First, the Oligo 7 program (https://www.oligo.net) was used to design specific primers based on the sequences flanking the indel sites. Genomic DNA was extracted from mature leaves of each sample using a Plant Genomic DNA Kit (Tiangen Biotech, Beijing, China), and the DNA fragments were amplified using PCR. The PCR was performed in a 10-μL reaction volume containing 0.05 U of Taq DNA polymerase, 1 μL of 10× PCR buffer (Takara Bio, Kusatsu, Japan), 1 μL genomic DNA (30 ng/μL), 0.3 μL MgCl2 (25 mM), 0.5 μL dNTPs (10 mM), 0.4 μL of each primer (10 μM), and 6.35 μL of double-distilled water. The PCR program began with denaturation at a temperature of 94°C for 3 min; followed by 30 cycles of 94°C for 30 s, 58°C for 30 s, and 72°C for 30 s; with a final extension at 72°C for 7 min. The PCR products were separated using electrophoresis on 8% denatured polyacrylamide gels run at 180 V for 1.5 h, and then detected using silver staining. Finally, we observed that

FIGURE 3 The PCR profiles of eight indels in 12 individuals. M represents the indel marker, and red numbers indicate that the electrophoretic bands of the individual are not consistent with its genotype called from InDelGT.
all but five genotypes were amplified, of which 86 (95%) genotypes were consistent with the results of the InDelGT genotyping (Figure 3). This experimental result showed that InDelGT has a high level of accuracy in determining indel genotypes.

**DISCUSSION**

In this paper, we report an integrated pipeline, InDelGT, for the extraction of indel genotypes in a hybrid population using next-generation DNA sequencing data. InDelGT can be used to implement the complex process of genotyping indels across multiple individuals in a population, such as traditional BC and F2 populations, as well as F1 hybrid populations between outbred species. The actual operation is very convenient, and the user can complete the entire calculation process with only one command line. In addition, InDelGT can not only extract indel genotype data but also perform Mendelian segregation ratio tests on the identified indel sites. To demonstrate the effectiveness of InDelGT, we performed indel genotyping on 117 progeny and two parents of an F1 hybrid population derived from a cross between *P. deltoides* and *P. simonii*. The results showed that these indel markers could be divided into five segregation types. When the genetic linkage maps of the two parents were constructed using the two most abundant indel segregation types, these segregating markers were divided into 19 linkage groups or 20 linkage groups within a large LOD threshold range, which largely matched the karyotype of *Populus*.

The coverage depth of reads is an important factor affecting the accuracy of indel genotyping. A higher read coverage depth can reduce the error rate of detecting indels. Compared with small indels (1–5 bp), large indels (>5 bp) will complicate or distort the correct mapping of reads, so detecting large indels requires a higher coverage depth (Fang et al., 2014). The determination of heterozygotes requires a higher coverage depth than that of homozygotes (McKernan et al., 2009). Ajay et al. (2011) suggested that when a read alignment rate of 85% is reached, the average sequencing coverage depth needs to be more than 40×. Fang et al. (2014) believed that 60× is a more ideal average coverage depth for detecting indels to obtain more accurate indel genotypes. In the current study, the average genome coverage depth of reads for each progeny was approximately 20× (Appendix S1). Because the paired-end reads were from the restriction sites of EcoRI, they covered at most 20% of the poplar genome (Tong et al., 2016); therefore, the actual coverage depth may have reached as high as 100×, which is sufficient to satisfy the requirements of indel genotyping. On the other hand, it is worth emphasizing that InDelGT is specifically designed to extract indel genotypes across a mapping population. Unlike other software packages (Koboldt et al., 2009; Ye et al., 2009; McKenna et al., 2010; Li et al., 2013), the genotype data generated from InDelGT are filtered to follow the Mendelian segregation ratio, which could further improve the data accuracy overall. Our experimental validation demonstrated that InDelGT can generate high-quality indel genotype data for genetic linkage mapping.

**CONCLUSIONS**

InDelGT is a practical tool that can quickly genotype a large number of indel markers largely following Mendelian segregation in a hybrid population. While the example here focuses on an F1 hybrid population from crosses between outbred species, InDelGT can also be used with traditional BC and F2 populations. InDelGT is a useful tool for various studies in genetics, genomics, and marker-assisted breeding, especially in forest trees.

**AUTHOR CONTRIBUTIONS**

C.T. and Z.P. conceived the research; Z.P., Z.L., J.Z., and S.B. participated in field surveys; Z.P., Z.L., and J.Z. analyzed the data. Z.P. wrote the initial manuscript, and all authors contributed to revising and editing the text. All authors approved the final version of the manuscript.

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**DATA AVAILABILITY STATEMENT**

The InDelGT pipeline is available for download on GitHub (https://github.com/tongchf/InDelGT), along with test data including a reference sequence file and the FASTQ files of two parents and 20 progeny.

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**Supporting Information**

Additional supporting information can be found online in the Supporting Information section at the end of this article.

**Appendix S1.** The RAD-seq data information for the two parents and 117 progeny in the hybrid population of *Populus deltoides* and *P. simonii*.

**Appendix S2.** The genetic map of linkage groups 1–19 for the maternal *Populus deltoides*. The length and name of each linkage group are presented above and below, respectively. Each indel is named by its position on the reference genome.

**Appendix S3.** The genetic map of linkage groups 1–20 for the paternal *Populus simonii*. The length and name of each linkage group are presented above and below, respectively. Each indel is named by its position on the reference genome.

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