In silico detection of EMS-induced mutations in an Arabis alpina population.



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Abstract

Arabis alpina (Alpine rock-cress weed) is a flowering plant, native to mountainous environments of the northern hemisphere. We analyzed 1,454,931,853 nextgeneration sequencing (NGS) reads from 38 sequenced Arabis alpina mutant individuals which that were mutagenized using the chemical mutagen, ethyl methane sulfphonate (EMS). Using the BWA short reads mapper, BWA, 95% (1,387,167,658) of the NGS reads mapped to Arabis alpina reference genome version 4. Using the SAMtools variant- detection algorithm, SAMtools, we detected a total of 1,457,917 mutations, with an average of 38,366 mutations per sample. Overall, the predicted mutations include 971,252 high-quality single nucleotide polymorphisms (SNPs) and 168,783 high-quality insertions and deletions (INDELs).

Methods

We obtained NGS whole-genome sequencing data for 38 Arabis alpina (Figure 1) EMS-mutagenized individuals from the NCBI Sequenced Reads Archive (SRA) repository. Figure 2 shows our implemented DNA mutation prediction pipeline. Next, after indexing the reference genome (version 4.0) of Arabis alpina by using BWA and SAMtools, we mapped the pre-processed short reads from the mutant genomes to the reference genome by using the BWA short reads aligner. After aligning the reads to the reference genome, SAMtools was used to call the variants which could be single nucleotide polymorphisms (SNPs) or small insertions or deletions (INDELs) from the alignment file. Finally, to predict the effect or impact of the mutations on gene function, we used the snpEff software to annotate the mutations and create the variant call files (VCFs). The annotated VCF file provides valuable information about the variants present in the genomes and their functional impact.



Figure 2. Image of Arabis alpina (Alpine rock-cress weed). Source: https://depositphotos.com/328529872/stock-photo-caucasian-rock-cress-flower-garden.html

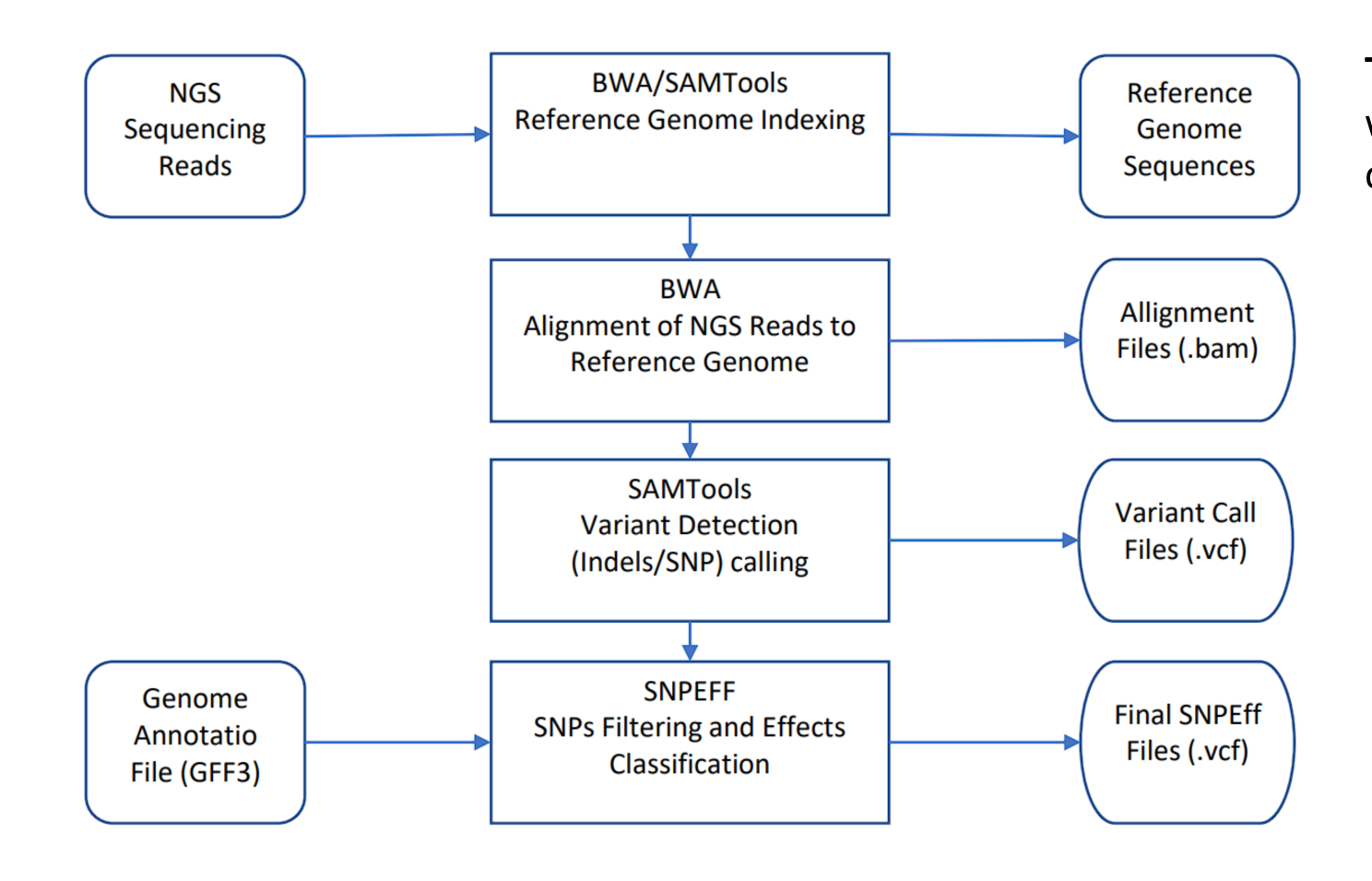


Figure 1. Pseudocode describing the Arabis alpina annotation algorithm.

Results

We obtained 1,454,931,853 NGS which reads of 1,387,167,658 mapped to the Arabis alpina reference genome, with an average of 38,287,680 reads per mutant genome (Table 1). Overall, we predicted 1,457,917 EMSinduced mutations including 222,578 INDELs and 1,235,339 SNPs (Table 2). Annotation of mutations revealed 17,830 high impact mutations and 50,153,551 moderate impact mutations including an average of 469 high impact mutations per mutant (Table 3).

Table 1. Number of reads after using bwa software.

Total Number of Reads All Mapped Reads Paired Mapped Reads **Average Number of Reads**

References:

- Li, Heng, and Richard Durbin. "Fast and accurate short read alignment with Burrows–Wheeler transform." bioinformatics 25.14 (2009): 1754-1760.
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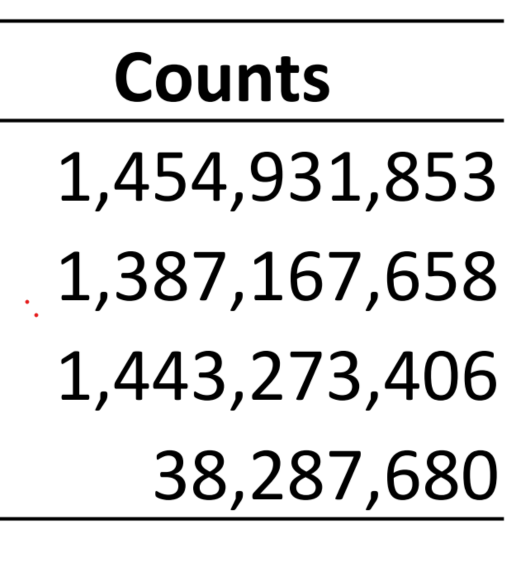




Table 2. The nearly 1,4 million variants of mutation were detected, more than 1,1 million of them are high quality.

Variants	•
All Variants	1,4
INDELS	
SNPs	1,2
Q20 Variants	
Q20 INDELs	, , ,
Q20 SNPs	Ç

Table 3. From total of 38 Arabis alpina individuals we detected 17,830 high impact mutations with an average of 469 and 50,153,551 moderate impact *mutations with an average of 1,319,830.*

SNP Impact	Total	Mean
High Impact	17,830	469
Stop Gained	232	6
Stop Lost	3,120	82
Splice Site Donor	2,688	71
Splice Site Acceptor	452	12
Moderate Impact	50,153,551	1,319,830
Missense	162,199	4,268
Low Impact	26,637	701
Silent	26,637	701
Start Gained	19,889	523
Modifier	165,276	4,349



