

MALVA: genotyping by Mapping-free ALlele detection of known VAriants

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Abstract

The amount of genetic variation discovered and characterized in human populations is huge, and is growing rapidly with the widespread availability of modern sequencing technologies. Such a great deal of variation data, that accounts for human diversity, leads to various challenging computational tasks, including variant calling and genotyping of newly sequenced individuals. The standard pipelines for addressing these problems include read mapping, which is a computationally expensive procedure. A few mapping-free tools were proposed in recent years to speed up the genotyping process. While such tools have highly efficient run-times, they focus on isolated, bi-allelic SNPs, providing limited support for multi-allelic SNPs, indels, and genomic regions with high variant density.

To address these issues, we introduce **MALVA**, a fast and lightweight mapping-free method to genotype an individual directly from a sample of reads. **MALVA** is the first mapping-free tool that is able to genotype multi-allelic SNPs and indels, even in high density genomic regions, and to effectively handle a huge number of variants such as those provided by the 1000 Genome Project. An experimental evaluation on whole-genome data shows that **MALVA** requires one order of magnitude less time to genotype a donor than alignment-based pipelines, providing similar accuracy. Remarkably, on indels, **MALVA** provides even better results than the most widely adopted variant discovery tools.

1 Introduction and Related Work

The discovery and characterization of sequence variations in human populations is crucial in genetic studies. A prime challenge is to efficiently analyze the variations of a freshly-sequenced individual with respect to a reference genome and the available genomic variations data. To reach this goal, the standard pipeline includes aligning sequenced reads with softwares like BWA [19] and Bowtie [17] and then calling the genotypes (*e.g.*, with GATK [22] or bcftools [20]); such an approach, though, can be highly time consuming, thus impractical for clinical applications, where time is often an issue. Typically in diploid organisms variant calling requires SNPs and indel detection and the identification of the pairs of alleles for each position of the studied genome, called genotype.

Recent tools for genotyping and variant calling like GraphTyper [11] and vg [31], which are based on a graph representation of a pan-genome to avoid biases introduced by considering only the information included in a set of genomes [5], are nevertheless heavy in both computational space and time. Moreover, the size of indexes of variation graphs may be subjected to an exponential growth in the number of variants included, and indexes typically require a great deal of computational resources to be updated with newly discovered variants. When the task is to call the genotype in positions where variants have been previously annotated, alignment-free methods come to the aid. Recent mapping-free genotyping tools such as LAVA [30] and VarGeno [34] are word-based methods that, given a list of known SNP loci, call SNPs as either mutant or wild-type up to an order of magnitude faster than the usual alignment-based methods. A major shortcoming of these tools is the large memory requirement, that can easily exceed hundreds of GB of RAM. Their strategy is to create a dictionary for both the reference genome and the SNP list that maps each k -mer to the positions at which it appears, and then to call variants from the reads by evaluating k -mers frequency. FastGT [27] is yet another k -mer-based method to genotype sequencing data: it strongly relies on a pre-compiled database of bi-allelic SNVs and corresponding k -mers, obtained by subjecting the k -mers that overlap known SNVs to several filtering steps. Such filters remove from the database the SNPs for which unique k -mers (*i.e.*, not occurring elsewhere in the reference genome) are not observed, those that are closely located (*i.e.*, that are less than k bases apart), and others: after the filtering steps, only 64% of bi-allelic SNVs survive and are therefore identifiable. These tools implement strategies to represent and analyze SNPs that improve the time performance but, on the other hand, do not allow to model indels and close variants.

Short insertions and deletions of nucleotides (*indels*) are believed to represent around 16% to 25% of human genetic polymorphism [24]. The presence of indels can be associated with a number of human diseases [26, 13, 16]: for instance, cystic fibrosis [26], lung cancer [29], Mendelian disorders [21] and Bloom syndrome [14] are all known to be closely correlated to indels. Indels are particularly challenging to call from NGS data, because mapping is more difficult when the reads overlap with indels [25].

In this paper we introduce MALVA, a rapid, lightweight, alignment-free method to genotype known (*i.e.*, previously characterized) variants, including indels and close SNPs, in a sample of reads. MALVA is a word-based method: each allele of each known variant is assigned a *signature* in the form of a set of k -mers, which allows to efficiently model indels and close variants. The genotypes will be called according to the frequency of such signatures in the input reads. Based on the well-known Bayes' formula, we also design a new rule to genotype multi-allelic variants (*i.e.*, variants such that more than one alternate allele is known): even if such variants are trickier to genotype than bi-allelic ones, we are still able to achieve high precision and recall, as revealed in the real-data experiments we conducted. MALVA directly analyzes a sample leveraging on the information of the variants included in a VCF file, that is the standard format released by the 1000 Genomes Project [32] (1KGP from now on). To the best of our knowledge, MALVA is the first

mapping-free tool able to call indels. Moreover, it proved to be the only such tool capable of handling the huge number of variants included in the latest version of the VCF released by the 1KGP.

The rest of the paper is laid out as follows. In Section 2 we give some definitions and we describe the data structures used by MALVA. In Section 3 we thoroughly detail the methods. In Section 4 we show some implementation details and provide an experimental analysis of MALVA on real data to assess the tool's performance against state-of-the-art tools. Finally, in Section 5, we draw conclusions and sketch possible future directions of our work.

2 Preliminaries

In this section we will describe the fundamental concepts that we will use in this document.

Let Σ be an ordered and finite alphabet of size σ and let $t = c_1, \dots, c_k$, where $c_j \in \Sigma$ for $j = 1, \dots, k$, be an ordered sequence of k characters drawn from Σ , we say that t is a k -mer. When a k -mer originates from a double stranded DNA, it is common to consider it and its reverse-complemented sequence as the same k -mer, and to say that the one that is lexicographically smaller among the two is the *canonical* one. In the following, we will abide by this definition and whenever we refer to a k -mer we implicitly refer to its canonical form. Moreover, to avoid k -mers being equal to their reverse-complement, we will only consider odd values of k .

A Bloom filter [1] is a probabilistic space-efficient data structure that represents a set of elements and allows approximate membership queries. The result of such queries may be a false positive but never a false negative. Bloom filters are usually represented as the union of a bitvector of length m and a set of h hash functions $\{H_1, \dots, H_h\}$, each one mapping one element of the universe to one integer in $\{1, \dots, m\}$. Using these data structures, the addition of an element e to the set is performed by setting to 1 the bitvector's cells in positions $\{H_1(e), \dots, H_h(e)\}$, while testing if an element is in the set boils down to checking whether the same positions are all set to 1. Due to collisions of the hash functions, an element can be reported as present in the set even if it is absent. Nevertheless, the false positive rate of a Bloom filter of a set of n elements, with h hash functions and an array of m bits is $(1 - e^{-\frac{hn}{m}})^h$; therefore, to increase the size of the Bloom filter decreases the false positive rate. Due to their simplicity and efficiency, Bloom filters have been applied to multiple problems in bioinformatics, such as representing de Bruijn graphs [2] and counting k -mers in a sample [23].

Let B be a bitvector, the \mathbf{rank}_1 function reports, for each position $i \in \{1, \dots, |B| + 1\}$, the number of 1s from the beginning of B to i (excluded); we refer to such value as $\mathbf{rank}_1(i, B)$. Clearly, $\mathbf{rank}_1(i, B)$ is not defined for $i \leq 0$ and for $i > |B| + 1$, $\mathbf{rank}_1(1, B)$ is 0, and $\mathbf{rank}_1(|B| + 1, B)$ is the number of 1s in B . By using succinct support data structures and by a linear time preprocessing step, it is possible to answer \mathbf{rank}_1 queries in constant time for any position of the bitvector [36].

The difference between the genetic sequence of two unrelated individual of the same species is estimated to be smaller than 0.1% [35]; therefore, it is common to represent the DNA sequence of an individual as a set of differences from a *reference* genome. Indeed, thorough studies [3, 4, 6] of the variations across different individuals encode such information as a VCF (Variant Calling Format) file [8]. In the following, we will call *variant* the information encoded by a data line of a VCF file. Besides the genotype data, we are interested in the information carried by the second, fourth, fifth, and eighth field of a VCF line, namely: (i) field POS that is the position of the variant on the reference, (ii) field REF that is the reference allele starting in position POS, (iii) field ALT that is a list of alternate alleles that in some sample replace the reference allele, and (iv) field INFO that is a list of additional information describing the variant. From the latter list we will get the frequencies of reference and alternate alleles, which are needed to call the genotype of a given

individual. We denote with $\text{POS}(v)$, $\text{REF}(v)$, $\text{ALT}(v)$, $\text{FREQ}(v)$, and $\text{GTD}(v)$ the reference position, reference allele, list of alternate alleles, list of allele frequencies, and genotype data of a variant v , respectively.

The variants we take into account are SNPs (*i.e.*, both REF and all the elements of ALT are single base nucleotides) and indels (REF and at least one element of ALT are not of the same length). Moreover, given an allele a (either reference or alternate) of some variant v , we refer to its sequence of nucleotides as $\text{SEQ}(a)$, *i.e.*, $\text{SEQ}(a)$ is the string that represents a .

Let R be a reference genome and let V be a VCF file that describes all the known variants of R . Since the genotype data provides information on the alleles expressed in each genome, another way of thinking of a VCF file is as an encoding of a set of genomes \mathcal{G} . Each haplotype of the genomes in \mathcal{G} can be reconstructed by modifying R according to the genotype information associated to each variant. For ease of presentation, in the following we use the term genome and haplotype interchangeably, although each genome of a polyploid organism is composed of multiple haplotypes.

Let \mathcal{G} be the set of genomes encoded by a VCF file and let a be an allele of some variant v , we denote by $\mathcal{G}^a \subseteq \mathcal{G}$ the subset of genomes that include a . We say that a variant v is k -isolated if there is no other known variant within a radius of $\lfloor k/2 \rfloor$ from the center of any of its alleles, as formally stated in the following definition.

Definition 1 (k -isolated variant). *A variant v is k -isolated if, for all $a \in \text{ALLELES}(v)$ and $g \in \mathcal{G}^a$, there is no variant $v' \neq v$ with an allele $a' \in \text{ALLELES}(v')$ such that $g \in \mathcal{G}^{a'}$ and either $|\text{BEGIN}_g(a') - \text{CENTER}_g(a)| \leq \lfloor k/2 \rfloor$ or $|\text{CENTER}_g(a) - \text{END}_g(a')| \leq \lfloor k/2 \rfloor$, where $\text{ALLELES}(v) = \text{REF}(v) \cup \text{ALT}(v)$, $\text{BEGIN}_g(a)$ is the position of the first base of a in g , $\text{END}_g(a)$ the position of the last base, and $\text{CENTER}_g(a)$ the position of the $\lceil \frac{|a|}{2} \rceil$ -th base of a in g .*

The procedure we will present in the next section is heavily based on the concept of *signature* of an allele. Intuitively, a signature of the allele a of a variant v is the k -mer centered in a in some genome g in \mathcal{G}^a . Note that, depending on the genomes encoded by the VCF file (specifically, if variants less than k bases apart are known), an allele might have multiple signatures. Moreover, if $\text{SEQ}(a)$ is longer than k bases, the previous definition is not well formed, since there is no k -mer that can be centered in a . In this case, we define the signature of a as the set of its substrings of length k . The following definition formalizes the notion of signature of an allele.

Definition 2 (Signature of an allele). *Let \mathcal{G} be the set of all the genomes encoded by a VCF file V and let k be an odd positive value. Let v be a variant in V , let a be one of the alleles of v , and let $\mathcal{G}^a \subseteq \mathcal{G}$ be the set of the genomes that include a . If $\text{SEQ}(a)$ is longer than k bases, we say that the signature of a is the set of all the substrings of length k of $\text{SEQ}(a)$. If $\text{SEQ}(a)$ is shorter than k bases, we say that $\{x\text{SEQ}(a)y\}$ is the signature of a in a genome g in \mathcal{G}^a if: (i) $x\text{SEQ}(a)y$ is a k -mer, (ii) $|x| = \lfloor \frac{k - |\text{SEQ}(a)|}{2} \rfloor$, (iii) $|y| = \lceil \frac{k - |\text{SEQ}(a)|}{2} \rceil$, (iv) x is a suffix of the sequence that precedes a in g , and (v) y is a prefix of the sequence that follows a in g .*

We will refer to the set of all the possible signatures of an allele a as $\text{SIGN}(a)$ and we say that k is the *length of the signature*. An example of signatures of an allele is shown in Figure 1.

In the following we will leverage on the definition of signature of an allele to detect its presence in an individual without mapping the reads to the reference genome. More precisely, we will analyze whether the k -mers of a given signature are present in the reads and use such information as an *hint* of the presence of the allele. Unlike other approaches [27], Definition 2 admits the presence of the alleles of multiple variants in a single signature, allowing MALVA to manage variants that are not k -isolated. Indeed, the set of signatures of an allele represents all the genomic regions where the allele appears in the genomes encoded by the VCF file.

	Pos	Ref	Alts	Donors																
\mathcal{R}	1	2	3	4	5	6	7	8	9	10	11	12	13	v_1	5	C	AAA	0 1	0 0	1 1
	A	G	A	T	C	C	T	G	C	G	A	A	G	v_2	7	T	G	0 1	0 1	0 1
														v_3	10	G	A,C	0 0	1 2	2 0

Variant	Allele	Signatures
v_2	$a_0 = T$	{ {TCCTGCG}, {TCCTGCA}, {AACTGCC} }
	$a_1 = G$	{ {AACGGCG}, { TCCGGCC } }

Figure 1: Signatures of the alleles of variant v_2 . \mathcal{R} is the reference sequence and the table on the right is a VCF information associated to it, representing 3 variants: an indel (v_1), a bi-allelic SNP (v_2), and a multi-allelic SNP (v_3). The last columns of the VCF file carry the genotype information of 3 individuals. The table at the bottom reports the signatures of each allele of variant v_2 . Note that there are only 5 signatures although 6 haplotypes are encoded by the VCF file since the second haplotype of the first and third individual are the same. We highlighted in red the genotype information associated to the second haplotype of the second genome and the corresponding signature.

3 Methods

In this section we will describe MALVA, the method we designed to genotype a set of known variants directly from a read sample. The general idea of MALVA is to use the frequencies of the signatures of a variant in the sample to call its genotype. The method works under the assumption that given a sample of reads from a genome with standard coverage depth, if an allele is included in the genome then at least one of its signatures must exist as substrings in multiple reads (depending on the coverage depth and the length of the signature). We leverage on this concept to genotype known variants directly from the input reads.

MALVA takes as input a reference genome, a VCF file representing all its known variants, and a read sample; it outputs a VCF file containing the most probable genotype for each variant. The main method is composed of four steps.

In the first step, MALVA computes the set of signatures of length k_s of all the alternate alleles of all the variants in VCF and stores them in the set ALTSIG. In the same step, the signatures of the reference alleles are computed and stored in a second set named REFSIG. For each k_s -mer t of a signature s two weights, one representing the number of occurrences of t in an alternate allele signature and one representing the number of occurrences of t in a reference allele signature, are stored. We will refer to these two values as w_t^A and w_t^R , respectively.

We note that for small values of k_s the probability that the k_s -mers that constitute a signature appear in other regions of the genome is high. Since in the following steps MALVA exploits the signatures' sets of the alleles of each variant to call the genotypes, the presence of conserved regions of the reference genome identical to some signature could lead the tool to erroneously genotype some variants. To get rid of a large amount of wrong calls, in the second step MALVA makes use of the context around the allele to distinguish its signatures from such regions. More precisely, if a k_s -mer of a signature of an alternate allele appears somewhere in the reference genome, MALVA extracts the context of length k_c (with $k_c > k_s$) covering the reference genome region and collects such k_c -mers in a third set (REPCTX).

In the third step, MALVA extracts all the k_c -mers from the sample along with the number of its occurrences. For each k_c -mer t_c that occurs w times in the sample, the k_s -mer t_s that constitutes the center of t_c is extracted. If t_s is found in REFSIG, $w_{t_s}^R$ is increased by w . Moreover, if t_c is

not found in REPCTX and if t_s is in ALTSIG, $w_{t_s}^A$ is increased by w . Otherwise, if t_c is in REPCTX, $w_{t_s}^A$ is not updated since, although its central k_s -mer is identical to some k_s -mer of a signature of an alternate allele of some variant, it is indistinguishable from another region of the genome not covering the variant. We note that when $w_{t_s}^A$ is not updated, our method might miss a variant in the donor and report a false negative, although for large values of k_c this would rarely occur. The rationale behind this choice is to avoid biases due to k_c -mers in conserved regions of the reference genome, preferring not to include an alternate allele in the output whenever ambiguities arise.

Finally, in the fourth step, MALVA uses the weights computed in the previous step to call the genotypes.

In the rest of this Section we will detail each one of the four steps of MALVA.

Signature computation. The first step consists of building the signatures of the alleles of all the variants and adding them either to ALTSIG, if they are the signatures of an alternate allele, or to REFSIG, if they are the signature of the reference allele. If a variant v is k_s -isolated, we build $1 + |\text{ALT}(v)|$ signatures, one for each allele of v . Otherwise, there are some genomes in \mathcal{G} in which there is at least another allele of a variant that lays within a radius of $\lfloor k_s/2 \rfloor$ nucleotides from the center of the allele of v . In practice, this means that we have to look at the genotype data of the variants within such radius: for each allele a of v we reconstruct the k_s bases long portions of the genomes in \mathcal{G}^a that constitute the signatures of a .

As pointed out in Definition 2, if $|\text{SEQ}(a)| \geq k_s$, the signature of a is the set of k_s -mers that appear in $\text{SEQ}(a)$. In this case we extract all such k_s -mers and add them either to REFSIG or ALTSIG. Otherwise, if $|a| < k_s$, we build the k_s bases long substrings of each genome in \mathcal{G}^a centered in a by scanning the VCF file and reconstructing the sequences according to the genotype information it includes. More precisely, let a be an allele of a variant v and let $V = \{v_1, \dots, v_n\}$ be the set of variants such that, for all $1 \leq i \leq n$: (i) $v_i \neq v$, (ii) there exists an allele a_j in $\text{ALLELES}(v_i)$ such that a and a_j are both included in some genome g , and (iii) either $(\text{END}(a_j) < \text{BEGIN}(a)$ and $\text{CENTER}(a) - \lfloor k_s/2 \rfloor \leq \text{END}(a_j)$) or $(\text{END}(a) < \text{BEGIN}(a_j)$ and $\text{CENTER}(a) + \lfloor k_s/2 \rfloor \geq \text{BEGIN}(a_j))$ in g .

Given a , we use the genotype information stored in the VCF file to retrieve the haplotypes in which it is included, *i.e.*, a subset of the haplotypes in \mathcal{G}^a , and build the set V . Using V we gather all the alleles that precede and succeed a in the selected haplotypes and we use them, together with the reference sequence, to reconstruct *on the fly* the k_s -mer that covers a , by interposing reference substrings and allele sequences. Doing so, we don't need to reconstruct the whole haplotypes but we only analyze and reconstruct the required k_s -mers when needed.

Once all the k_s -mers have been constructed, they are added to REFSIG if a is the reference allele, to ALTSIG if it is an alternate allele.

Detection of repeated signatures. This step is aimed to detect and store in set REPCTX all the k_c -mers of the reference sequence whose central k_s -mer is included in some signature of some alternate allele, $k_c > k_s$. REPCTX will be used in a further step to discard alternate alleles that might be erroneously reported as expressed by MALVA only because they cannot be told apart from other identical regions of the reference sequence. To compute REPCTX, we extract all the k_c -mers of the reference sequence and test whether their central k_s -mer is in ALTSIG. If so, we add the k_c -mer to REPCTX to report that the k_s -mer is indistinguishable from some k_s -mer that is included in the signature of an alternate allele. The set REPCTX is then used in the next step as illustrated below. An example comprising the first two steps is shown in Section B of the Appendix.

Alleles' signatures weights computation. In the third step, MALVA computes how many times the k_s -mers of each signature appear in the dataset. First, MALVA extracts all the k_c -mers of the read sample and tests their existence in REPCTX to check whether their central k_s -mer cannot be told apart from some repetition in the reference genome. Then, given a k_c -mer t_c that occurs w times in the read sample, the k_s -mer t_s that constitutes its center is extracted. If t_s is found in REFSIG, *i.e.*, t_s is the signature of the reference allele of some variant, the weight $w_{t_s}^R$ is increased by w . Moreover, if t_c is not found in REPCTX and t_s is in ALTSIG, *i.e.*, k_s -mer t_s is uniquely associated to an alternate allele of some variant, the weight $w_{t_s}^A$ is increased by w . Conversely, if t_c is in REPCTX, $w_{t_s}^A$ is not updated. The last scenario happens when t_s is identical to the signature of an alternate allele of some variant (indeed, t_s is in ALTSIG), but even the enlarged context t_c (and consequently t_s) appears somewhere else in the reference genome.

Genotype calling. In the last step, MALVA uses the allele frequencies stored in the INFO field of the VCF file and the weights of the signatures computed in the previous step to call the genotype of each variant. To this aim, we extend the approaches proposed in the literature for bi-allelic variants [30, 34] to multi-allelic variants.

Let v be a variant with $n - 1$ alternate alleles. The number of possible distinct genotypes is $\binom{n}{2} + n = \frac{n(n+1)}{2}$, that is one *homozygous reference* genotype, $\binom{n}{2}$ *heterozygous* genotypes, and $n - 1$ *homozygous alternate* genotypes. We will refer to the homozygous reference genotype as $G_{0,0}$, to the heterozygous genotypes as $G_{i,j}$ with $0 \leq i < j \leq n - 1$, and to the homozygous alternate genotypes as $G_{i,i}$ with $1 \leq i \leq n - 1$. Following well-established techniques [30, 22, 18], we compute the likelihood of each genotype $G_{i,j}$ by means of the Bayes' theorem. Given the observed coverage C , we compute the posterior probability of each genotype as:

$$P(G_{i,j}|C) = \frac{P(G_{i,j})P(C|G_{i,j})}{P(C)}$$

that, by the law of total probability, can be expressed as:

$$P(G_{i,j}|C) = \frac{P(G_{i,j})P(C|G_{i,j})}{\sum_{p=0}^{n-1} \sum_{q=p}^{n-1} P(G_{p,q})P(C|G_{p,q})}$$

To calculate this probability, we compute the *a priori* probabilities of each genotype $G_{i,j}$ ($P(G_{i,j})$) and the *conditional probability* of the observed coverage given the considered genotype ($P(C|G_{i,j})$). The Hardy-Weinberg equilibrium equation ensures that for each variant v , $(\sum_{i=0}^{n-1} f_i)^2 = 1$, where $f_i = \text{FREQ}(v)[i]$, *i.e.*, the frequency of the i -th allele of v . We recall that $\text{FREQ}(v)$ is stored in the INFO field of the VCF file. The *a priori* probability of each genotype $G_{i,j}$ is therefore computed as follows:

$$P(G_{i,j}) = \begin{cases} f_i^2 & \text{if } i = j \\ 2f_i f_j & \text{otherwise} \end{cases}$$

To compute the conditional probability $P(C|G_{i,j})$, it is first necessary to compute the *coverages* of the alleles of the variant. Without loss of generality, let a_0 be the first allele of the variant, *i.e.*, a_0 is the reference allele with index 0. We recall that $\text{SIGN}(a_0)$ is the set of signatures of allele a_0 and that each signature is a set of one or more k -mers. We also recall that, in the previous step, for each k -mer t that belongs to some signature we computed two weights, namely w_t^R and w_t^A . Given a signature $s \in \text{SIGN}(a_0)$, we define its weight as the mean of the weights associated to the k -mers it contains, *i.e.*, $\frac{\sum_{t \in s} w_t^R}{|s|}$ where $|s|$ denotes the number of k -mers contained in signature s . Since

the same allele may exhibit more signatures, we define the coverage c_0 of allele a_0 as the maximum value among the weights of its signatures, *i.e.*, $\max\{\frac{\sum_{t \in s} w_t^R}{|s|} : s \in \text{SIGN}(a_0)\}$. This formula can be easily modified to compute the coverage of an alternate allele (c_i for $i \geq 1$) by switching w_t^R with w_t^A . The coverage c_i of an allele a_i of a variant is thus computed as follows:

$$c_i = \begin{cases} \max\{\frac{\sum_{t \in s} w_t^R}{|s|} : s \in \text{SIGN}(a_0)\} & \text{if } i = 0 \\ \max\{\frac{\sum_{t \in s} w_t^A}{|s|} : s \in \text{SIGN}(a_i)\} & \text{otherwise} \end{cases}$$

By extending the approach adopted in [30], we consider each $P(C|G_{i,j})$ to be multinomially distributed. Given a homozygous genotype $G_{i,i}$, we assume to observe the i -th allele, which is the correct one, with probability $1 - \varepsilon$ (where ε is the expected error rate) whereas the other $n - 1$ alleles (the erroneous ones) with probability $\frac{\varepsilon}{n-1}$ each. Hence, we compute the conditional probability of an homozygous genotype as:

$$P(C|G_{i,i}) = \binom{c_i + C_E}{c_i} (1 - \varepsilon)^{c_i} \left(\frac{\varepsilon}{n-1}\right)^{C_E}$$

where C_E is the total sum of the coverages of the erroneous alleles, *i.e.*, $C_E = \sum_{j \in \{0, \dots, n-1\} \setminus \{i\}} c_j$. For what concerns heterozygous genotypes, we assume to observe the correct alleles, *i.e.*, the i -th and the j -th allele, with equal probability $\frac{1-\varepsilon}{2}$ whereas the other $n - 2$ erroneous alleles with probability $\frac{\varepsilon}{n-2}$ each. We compute the conditional probability of an heterozygous genotype as follows:

$$P(C|G_{i,j}) = \binom{c_i + c_j + C_E}{c_i + c_j} \binom{c_i + c_j}{c_i} \left(\frac{1-\varepsilon}{2}\right)^{c_i} \left(\frac{1-\varepsilon}{2}\right)^{c_j} \left(\frac{\varepsilon}{n-2}\right)^{C_E}$$

where, again, C_E is the sum of the coverages of the erroneous alleles, *i.e.*, $C_E = \sum_{p \in \{0, \dots, n-1\} \setminus \{i,j\}} c_p$.

Finally, after computing the posterior probability of each genotype, MALVA outputs the genotype with the highest likelihood.

4 Implementation and Experiments

In this section we will describe the implementation details of MALVA and we will provide an experimental analysis on real data. All the analyses were performed on a 64 bit Linux (Kernel 4.4.0) system equipped with four 8-core Intel[®] Xeon 2.30GHz processors and 256GB of RAM.

Implementation details. MALVA is implemented in C++ and it is freely available at <https://github.com/AlgoLab/malva>. Bloom filters were implemented as the union of a bitvector and a single hash function H. Although it is not conventional, in most cases to use a single hash function has similar results as using multiple ones, as noticed by other authors [33, 34]. To check this claim, while developing the tool we tested whether using multiple hash functions would improve the results by extending the Bloom filters to count-min sketches [7]. As expected, the deterioration of the performance far outweighed the gain in precision and recall (that was less than 0.1%). Moreover, to use a single hash function allows us to store w_t^R and w_t^A efficiently for each k -mer t . Indeed, note that once all the signatures of all the alternate alleles have been added to ALTSIG, the latter is only used to check whether some k_s -mer is part of a signature, *i.e.*, it becomes static. By representing ALTSIG as a Bloom filter B_{ALTSIG} we can create an integer array CNTS of size $\text{rank}_1(|B_{\text{ALTSIG}}| + 1, B_{\text{ALTSIG}})$ to store the weights of each k -mer compactly and, if a k -mer t of a signature s is in ALTSIG (*i.e.*, if $B_{\text{ALTSIG}}[H(s)] = 1$) we can access its weight by accessing $\text{CNTS}[\text{rank}_1(H(s), B_{\text{ALTSIG}})]$. In a

Dataset	Tool	P_{SNP}	R_{SNP}	P_{INDEL}	R_{INDEL}	Time (hh:mm)	RAM (GB)
HalfGenome	MALVA	92.9%	90.7%	85.3%	80.9%	03:19	30
	VarGeno	97.5%	88.1%	39.5%	0.1%	02:31	52
	bcftools	91.2%	94.8%	44.9%	55.4%	24:35	6
	GATK	91.7%	95.1%	53.2%	79.9%	34:43	32
FullGenome	MALVA	91.6%	89.7%	84.4%	80.1%	07:31	37
	VarGeno	-	-	-	-	-	-
	bcftools	91.6%	94.4%	44.7%	54.6%	54:23	9
	GATK	92.0%	94.7%	53.2%	79.2%	73:36	33

Table 1: Accuracy and efficiency results on the `HalfGenome` and `FullGenome` datasets. For each dataset, we reported the values of Precision (P) and Recall (R) obtained by the considered tools on both SNPs and indels. The efficiency results are shown in terms of wall clock time and peak memory usage. `VarGeno` could not complete the analysis of the `FullGenome` dataset, thus we did not report its results on this dataset.

nutshell, after adding all the alternate alleles to B_{ALTSIG} , we *freeze* it, build a rank data structure over it, compute the number of ones, and create the `CNTS` array of the correct size. Similarly, we implemented `REPCTX` as a Bloom filter B_{REPCTX} using a single hash function. Conversely, `REFSIG` was implemented as a simple hash table, because the number of elements it stores is usually smaller than the number of elements stored in `ALTSIG`. The bitvectors, the rank data structure, and the `CNTS` array were implemented using the `sdsl-lite` library [12]. We pose an upper limit of 255 to the value of each cell of the `CNTS` array, so as to store each counter using only 8 bits.

Finally, instead of scanning all the k_c -mers in the read sample, we used `KMC3` [10] to efficiently extract them and counting their occurrences. Therefore, in step 3 `MALVA` parses the output of `KMC3` and updates the counts for each k_s -mer accordingly.

Experiments. We performed an experimental analysis on real data to evaluate the real feasibility of our method, comparing `MALVA` to one mapping-free method and to two different alignment-based pipelines. Among the mapping-free methods proposed in literature we chose `VarGeno`, as it is an improved version of `LAVA` that provides better efficiency and accuracy [34]. For completeness, we included in our evaluation the two most widely used alignment-based pipelines, denoted by `bcftools` and `GATK`, respectively. The former consists of an alignment step performed with `BWA-MEM` [19] followed by a variant discovering step performed using `bcftools` [18]. The latter consists of an alignment step performed with `BWA-MEM` and a variant discovering step performed with `GATK` [22], as recommended by the *GATK Best Practices* [9].

`MALVA` was run setting k_s equal to 47, k_c equal to 53, ϵ equal to 0.1%, Bloom filters size equal to `8GB`, and considering the a priori frequencies of the alleles of the EUR population, since the individual under analysis is part of it.

We tested the tools using the Illumina WGS dataset of the well-studied NA12878 individual provided by the Genome In A Bottle (GIAB) consortium [38]. We chose this individual because the variant calls provided are highly reliable and can be effectively used to assess the precision and the recall of the considered methods. We downloaded the alignments of its 30x downsampled version and used `SAMtools` [20] to extract the corresponding `FASTQ` file, obtaining 696,168,435 150bp-long reads. As reference genome and set of known variants, we used the GRCh37 primary assembly and the `VCF` files provided by Phase 3 of the 1KGP [32]. These `VCF` files contain a total of 84,739,838

variants, the phased genotype information of 2504 individuals, and the a priori frequency of each allele of each variant of 5 populations. We note that **VarGeno** requires a different formatting of the fields describing the a priori frequencies of the alleles than the ones in the VCF file provided by the 1KGP. Thus, we formatted the input files as required before running **VarGeno**.

VarGeno could not complete the analysis of this dataset, from now on denoted by **FullGenome**, on our server. To test whether **VarGeno** crashed due to excessive memory usage, we tried to run it on the same instance on a cluster with 1TB of RAM, but nevertheless it could not complete the analysis, crashing after 20 minutes. In order to include **VarGeno** in our evaluation, we chose 12 chromosomes to create a smaller dataset, denoted by **HalfGenome**, that thus contains some half of the variants and the reads of the **FullGenome** dataset.

Each tool was evaluated in terms of variant calling accuracy and efficiency (wall time and memory usage). We note that some steps of the previously cited tools can use multiple threads to improve the time performance (namely, **KMC3** for **MALVA**, **BWA-MEM** for **bcftools** and **GATK**, and the variant discovery steps of **GATK**). Whenever we had this choice, we provided 4 threads to each tool. We used **hap.py** [15], the tool developed for the evaluation of variant callers in the recent *PrecisionFDA Truth Challenge*¹, and the `/usr/bin/time` system tool to gather the required data.

Table 1 shows the results obtained by the considered tools on both the **FullGenome** and the **HalfGenome** datasets. We point out that **hap.py** computes precision and recall considering only non-reference VCF records (*i.e.*, non 0/0 calls).

As expected, **MALVA** and **VarGeno** are one order of magnitude faster than **bcftools** and **GATK**. Indeed, **MALVA** and **VarGeno** required 3.5 and 2.5 hours to analyze the **HalfGenome** dataset, respectively, while **bcftools** and **GATK** required 24.5 and 34.5 hours. We note that half of the time required by **bcftools** and one third of the time required by **GATK** was spent running **BWA-MEM**, that completed its task in 12.5 hours (using 4 threads). The same trend applies to the analysis of the **FullGenome** dataset, on which each tool required roughly twice the time required on the **HalfGenome** dataset. A qualitative representation of each tool's running times is shown in Figure 2.

For what concerns the memory usage, **bcftools** proved to be the least memory intensive approach, requiring less than 10GB of RAM on both datasets to map the reads and less than 1GB of RAM to call the variants. **MALVA** and **GATK** showed similar memory requirements, with **GATK** showing almost no difference between the two analyses and **MALVA** increasing the memory consumption by only 23% for the bigger dataset. **VarGeno** required slightly less than twice the amount of memory required by **MALVA** on the **HalfGenome** dataset.

Precision and recall of all the tools varied little over the two datasets, proving that the number of variants and reads only slightly affects their accuracy. As expected, **bcftools** and **GATK** achieved the best recall for non-homozygous reference SNPs due to the mapping step, that provides a more precise coverage of the alleles and allows to better discern repeated regions of the reference genome. **VarGeno** achieved the lowest recall obtaining 2% less recall than **MALVA**, that in turn called correctly 90.7% of the SNPs. On the other hand, **MALVA**, **bcftools**, and **GATK** achieved comparable precision on SNPs, whereas **VarGeno** obtained the highest one. Overall, on non-homozygous reference SNPs, **VarGeno** seems to be the most conservative tool among those tested, as it prefers not to call SNPs when there is any uncertainty. On the contrary, **MALVA**, in avoiding the loss of any potentially interesting information, deliberately prefers to detect any potential alternate allele in the donor, at the cost of a slight loss in precision.

Remarkably, on indels **MALVA** obtained significantly better recall than **bcftools** and better precision than any other tool. As expected, since the method of **VarGeno** is not designed to manage indels, it was only able to genotype a negligible percentage of them. **bcftools** showed a

¹<https://precision.fda.gov/challenges/truth>

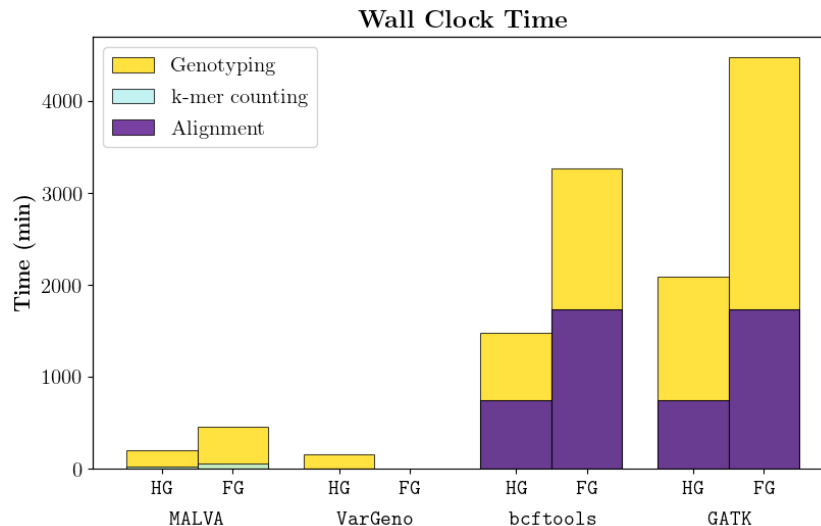


Figure 2: Times required by each tool to analyze both datasets, partitioned by steps performed. For ease of presentation, we denoted the **FullGenome** dataset as FG whereas the **HalfGenome** dataset as HG. Note that we did not include **VarGeno** running time on the **FullGenome** dataset since it crashed after 20 minutes.

very low precision and recall on indels, whereas **GATK** achieved a recall similar to **MALVA** but a low precision. The low precision achieved by the alignment-based tools is mainly due to the difficulties in aligning reads that overlap with indels.

Overall, **MALVA** proved to be an accurate and efficient alternative to mapping-based pipelines for variant calling, achieving good results both on SNPs and indels. The experimental evaluation shows the usefulness of the formalization of signature of an allele, of the extension to multi-allelic SNPs and indels, and of the ability to manage variants in dense genomic regions. A more in-depth comparison of **MALVA** and **VarGeno** is provided in Section A of the Appendix.

5 Conclusions and Future Directions

In this article, we presented **MALVA**, the first efficient mapping-free genotyping tool that is able to handle multi-allelic variants and indels. We compared **MALVA** with **VarGeno**, the state-of-the-art mapping-free genotyping tool, showing that our method is less memory intensive, achieves better recall, handles dozens of millions of variants effectively, and provides correct genotypes even for indels. We also compared our tool with two variant discovery pipelines, namely **GATK** and **bcftools**, showing that **MALVA** is an order of magnitude faster while achieving better accuracy on indels and similar accuracy on SNPs.

MALVA proved to be able to efficiently manage a huge amount of variants like those provided by the 1000 Genome Project (80 millions of variants) and to handle multi-allelic variants and indels. These fundamental features allow our method to exploit the whole information in input, without filtering out any data that might be crucial in successive analyses. Most notably, **MALVA**'s ability to genotype indels allows to apply mapping-free techniques to many clinical contexts, including screens for genetic predispositions for disease linked to the presence of indels [28, 37].

Future steps will be devoted to improving the efficiency of **MALVA** by exploiting the parallel architecture of modern machines, and to extending the method to genotype trios.

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Real GT	Called GT				
	HomoRef	HetRef	HomoAlt	HetAlt	Uncalled
HetAlt	57	93	163	991	0
HomoAlt	25842	23850	697174	117	0
HetRef	90030	1047430	31956	139	0
HomoRef	37806840	69737	2442	17	0

(a) MALVA

Real GT	Called GT				
	HomoRef	HetRef	HomoAlt	HetAlt	Uncalled
HetAlt	0	0	0	0	1304
HomoAlt	2238	7811	632822	0	104112
HetRef	78451	950511	12049	0	128544
HomoRef	36087283	20507	947	0	1770299

(b) VarGeno

Figure 3: Comparison between real genotype (provided by the 1000 Genomes Project) and genotype called by each tool. HomoRef stands for Homozygous Reference, HetRef stands for Heterozygous Reference, HomoAlt stands for Homozygous Alternate, HetAlt stands for Heterozygous Alternate, and Uncalled means that the given variant was not called by the tool.

A Comparison of MALVA and VarGeno output

To assess whether the tools under analysis produce some systematic error, we considered the **HalfGenome** dataset and we performed a more in-depth analysis of the SNPs in the **VCF** output by **MALVA** and **VarGeno**. For each tool, Figure 3 reports the number of correct genotypes output, grouping them in *homozygous reference* (i.e., 0|0), *heterozygous reference* (i.e., 0|1, 0|2, and so on.), *homozygous alternate* (i.e., 1|1, 2|2, and so on), and *heterozygous alternate* (i.e., 1|2, 1|3, 2|3, and so on). As stated in Section 4, we recall that the precision and recall output by **hap.py** do not consider homozygous reference genotypes, thus the analysis we present in this section allows us to better understand the behavior of the tools. Since **VarGeno** is not able to manage indels, we decided not to include them in this analysis.

Consistently with the precision and recall results of **hap.py**, **MALVA** detects between 5% and 10% more correct variants than **VarGeno** in all classes, at the cost of producing more erroneous calls. We note that overall **VarGeno** filters out 2,004,259 of the 39,796,878 SNPs in the truth.

Both tool show similar pattern in erroneous calls. More precisely erroneously genotyped homozygous reference variants were mostly genotyped as heterozygous reference and, vice-versa, erroneously genotyped heterozygous reference variants were mostly genotyped as homozygous reference. On the other hand, erroneous homozygous alternate variants in the donor were mostly genotyped as heterozygous reference by **VarGeno** whereas **MALVA** evenly distributed the errors between homozygous reference and heterozygous reference calls. Finally, erroneous heterozygous alternate variants in the donor were mostly genotyped as homozygous alternate variants by **MALVA**, meaning that the method proposed in this paper was able to detect the fact that the allele was not the reference allele but it called one of the two alternate alleles of the donor erroneously. Figure 3 shows a comparison between real genotype (provided by the 1000 Genomes Project) and genotype called by **MALVA** and **VarGeno** on SNPs. Figure 4 shows the same data row-normalized.

Overall, the errors produced by both tools were “partial” errors in the sense that they rarely mis-call both alleles of the donor.

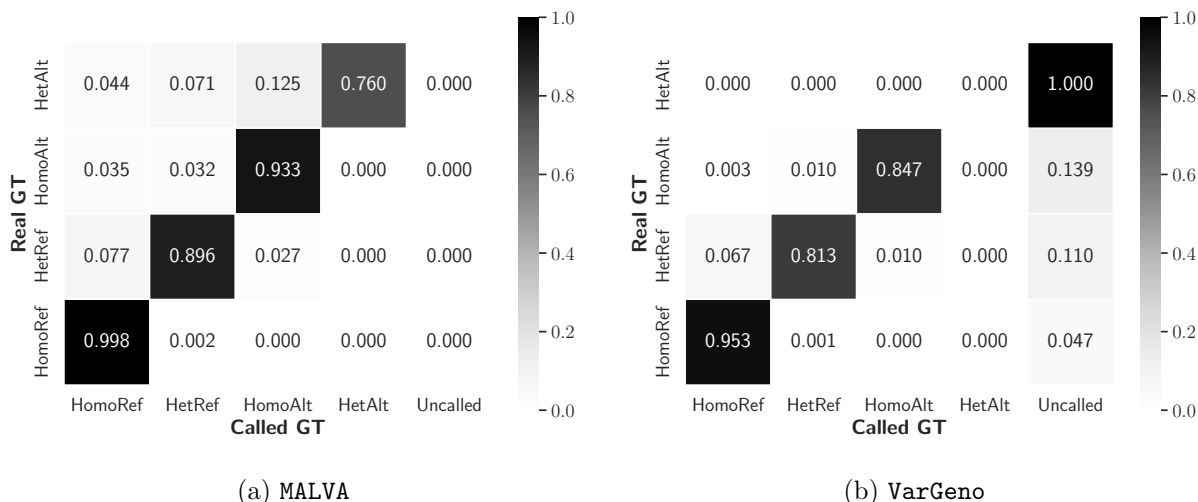


Figure 4: Comparison between real genotype (provided by the 1000 Genomes Project) and genotype called by each tool, normalized by rows.

B Example k -mers weight computation

In this section we present an example of computation of the weights associated with the signatures' k_s -mers. Figure 5 shows an example composed of three variants and two reads. In this example the values of k_s and k_c are set to 7 and 11, respectively. Subfigure (a) shows the 26-bases long reference sequence. Subfigure (b) reports on the left two bi-allelic variants (v_1 and v_2) and one multi-allelic variant (v_3), and on the right the signatures of each allele of v_2 . Subfigure (c) shows the elements of **ALTSIG** and **REFSIG** related to v_2 . We note that the second signature in **ALTSIG** is composed of a single k_s -mer (t_s , equal to **TCCGGCG**) that appears in the reference genome, starting from position 17. Thus, the k_c -mer starting in position 15 and ending in position 25 (t_c , equal to **GATCCGGCGAA**) is added to **REPCTX**. Subfigure (d) shows two 11-bases long reads including t_s , extracted from position 3 and 15 of the donor. Clearly, only r_1 should contribute to the detection of the alternate allele of v_2 in the donor, since r_2 was sequenced from another position of the genome (*i.e.*, $w_{t_s}^A$ should be equal to 1 in this case). To this aim, **REPCTX** comes to an aid; indeed, when analyzing r_1 the k_c -mer covering t_s is extracted (*i.e.*, the whole read) and its inclusion in **REPCTX** is tested. Since **TATCCGGCGTA** is not in **REPCTX** and t_s is in **ALTSIG**, $w_{t_s}^A$ is increased by one. On the other hand, since **GATCCGGCGAA** is in **REPCTX**, the occurrence of t_s in r_2 is not considered in $w_{t_s}^A$, thus avoiding to erroneously overestimate the frequency of allele a_1 of v_2 .

We note that on one hand this approach allows us to avoid overestimating the frequencies of some alternate allele but, on the other hand, it produces two major side effects. The first one is that some allele might be underestimated by **MALVA**; indeed, if the k_c -mer covering an alternate allele in a donor is equal to a k_c -mer in the genome it will not be detected. The second side effect is that **MALVA** might overestimate the frequency of some allele due to identical signature. Indeed, suppose that the signature of some alternate allele a_i of another variant $v_j \neq v_2$ is equal to the signature of alternate allele a_1 of variant v_2 . It is obvious that the weights of the k_s -mers of the two signatures will be identical and that the occurrences of both the alleles will concur towards their final value, overestimating it.

Although the two side effects pose some limit to the method proposed in this paper, they arise rarely and we think they are a fair price to pay to avoid biases introduced by the reference genome.

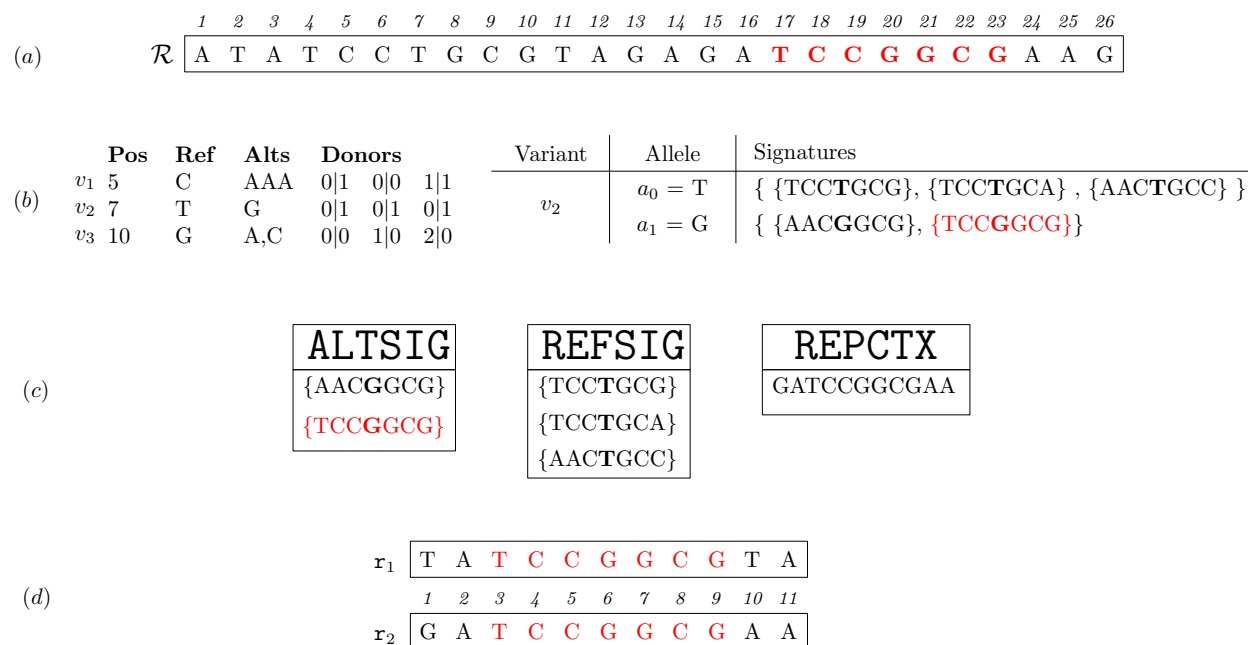


Figure 5: Example with 3 variants and two reads. Subfigure (a) shows a reference genome of 26 bases, Subfigure (b) reports 3 variants and the signatures of each allele of variant v_2 , Subfigure (c) reports the subsets of ALTSIG, REFSIG, and REPCTX including the elements related to v_2 , and Subfigure (d) presents two reads of length 11.