



Bioinformatics

doi.10.1093/bioinformatics/xxxxxx

Advance Access Publication Date: Day Month Year

Original Paper



???Genetics and population analysis???Genome analysis

DEploid: Untangling complexity of infection in

Plasmodium falciparum.

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Associate Editor: XXXXXXX

Received on XXXXX; revised on XXXXX; accepted on XXXXX

Abstract

Motivation: Complexity of infection in the malarial parasite *Plasmodium falciparum* affects key phenotypic traits, including drug resistance and risk of severe disease. Advances in protocols and sequencing technology have made it possible to obtain high-coverage genome-wide sequence data from blood samples taken in the field. However, analysing and interpreting such data is challenging because of the high rate of multiple infections present.

Results: The software package *DEploid* deconvolutes sequences of mixed samples by learning haplotype structure from a reference panel of clonal isolates. It reports the number of strains, the mixing proportions and the haplotypes present in an isolate, allowing researchers to study complexity of infection in malaria with an unprecedented level of detail.

Availability and implementation: The open source implementation *DEploid* is freely available at https://github.com/mcveanlab/dEploid under the conditions of the GPLv3 license. An R version is available at https://github.com/mcveanlab/DEploid-r.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

Malaria remains one of the top global health problems. Transmitted by mosquitoes of the genus *Anopheles*, the majority of malaria related deaths are caused by the *Plasmodium falciparum* parasite (WHO, 2016). Patients are often infected with more than one parasite strain, due to bites from multiple mosquitoes, mosquitoes carrying multiple genetic types or a combination of both. Multiplicity of infection can lead to competitions among co-existing strains and may influence disease development (de Roode et al., 2005), transmission rates (Arnot, 1998) and even the spread of drug resistance (de Roode et al., 2004).

The presence of multiple strains of *P. falciparum* makes fine scale analysis of genetic variation challenging, since genetic differences between the genetic types of this haploid organism will appear as heterozygous loci. Mixed calls also confound methods that exploit haplotype data to detect, among other phenomena, the occurrence of natural selection or recent

demographic events (reference). In light of these difficulties, researchers usually focus on clonal infections or resort to heuristics methods for resolving heterozygous genotypes. The former approach discards valuable information regarding genetic diversity and inbreeding whereas the latter tends to create chimeric haplotypes that are not suitable for analysis, unless mixed calls are very sparse.

Phasing or deconvoluting the strains of a mixed infection is a harder problem than phasing diploid organisms because the levels of mixture within isolates (i.e. the abundance of each genetic type) vary greatly and are unknown. Existing tools for phasing diploid organisms, such as Beagle (Browning and Browning, 2007) and IMPUTE2 (Howie et al., 2009), are not designed to cope with this. Galinsky et al. (2015) and O'Brien et al. (2015) have attempted to address the mixed infection problem by inferring the mixed proportions from allele frequencies within samples, but they do not provide haplotypes and have limited use.

As part of the Pf3k project (Pf3k, 2016), an effort to map the genetic diversity of *P. falciparum* at global scale, we have developed DEploid, a software package for deconvoluting mixed infections. The program

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provides estimates for the number of different genetic types present in the isolate, the proportion or abundance of each strain and their sequences (i.e. haplotypes). To our knowledge, DEploid is the first package able to deconvolute strain haplotypes and provides a unique opportunity for researchers to study inbreeding and complexity of infection, leaving open the possibility to investigate infection history at fine scale.

2 Methods

2.1 Notations

Let's first introduce some notation (see Table 1). Suppose that our data D are the allele read counts of sample j at a given site i, denoted as $r_{j,i}$ and $a_{j,i}$ for reference (REF) and alternative (ALT) alleles respectively. The allele frequencies within sample (WSAF) $p_{j,i}$ and at population level (PLAF) f_i can be calculated by $\frac{a_{j,i}}{a_{j,i}+r_{j,i}}$ and $\frac{\sum_j a_{j,i}}{\sum_j a_{j,i}+\sum_j r_{j,i}}$. Since all data in this section refers to the same sample, we drop the subscript j from now on.

i	Marker index		
j	Sample index		
r	Read count for reference allele		
a	Read count for alternative allele		
f	Population level allele frequency (PLAF)		
n	Number of strains within sample		
l	Sequence length		
\mathbf{w}	Proportions of strains		
\mathbf{h}_i	allelic states of n parasite strains at site i		
$h_{k,i}$	allelic states of k parasite strains k at site i , h_k in short		
p	Observed within sample allele frequency (WSAF)		
q	Unadjusted expected WSAF		
π	Adjusted expected WSAF		
Ξ	Reference panel		
g_p	allelic states of reference strain p		
e	Probability of read error		

Table 1. Notation summary

2.2 Model

We descibe the mixed infection problem by considering the number of strains n, relative abundance of each strain \mathbf{w} and their allelic states \mathbf{h} . Similar to O'Brien et al. (2015), we use a Bayesian approach and define the posterior probabilities of n, \mathbf{w} and \mathbf{h} for some reference panel Ξ and probability of miss read e given data as the following:

$$P(n, \mathbf{w}, \mathbf{h}, \Xi, e|D) \propto L(n, \mathbf{w}, \mathbf{h}, \Xi, e|D) \times P(n, \mathbf{w}, \mathbf{h}, \Xi, e).$$

We assume that the proportion of parasite strains \mathbf{w} given n strains, the chosen reference panel Ξ and sequencing error rate are independent from each other, the following expression is then obtained:

$$P(n, \mathbf{w}, \mathbf{h}, \Xi, e|D) \propto L(n, \mathbf{w}, \mathbf{h}, \Xi, e|D) \times P(\mathbf{h}|\mathbf{w}, \Xi, e) \times P(\mathbf{w}|n) \times P(n) \times P(\Xi) \times P(e).$$
(1)

Since we are not making inferences on the reference panel nor error rate, the prior probabilities are fixed as 1s. For convience, we fix the number of strains n and simplify Eqn. (2) as the following:

$$P(\mathbf{w}, \mathbf{h}, \Xi, e|D) \propto L(\mathbf{w}, \mathbf{h}, \Xi, e|D) \times P(\mathbf{h}|\mathbf{w}, \Xi, e) \times P(\mathbf{w}|n).$$
 (2)

2.2.1 Likelihood of data given the WSAF

Let $\mathbf{w} = [w_1, \dots, w_n]$ and $\mathbf{h}_i = [h_{1,i}, \dots, h_{n,i}]$ denote the proportions and alleic state of n parasite strains at site i. We use O'Brien et al. (2015)'s expression for the expected WSAF q_i as:

$$q_i = (\mathbf{w} \cdot \mathbf{h}_i) = \sum_{k=1}^n w_k \cdot h_{k,i}.$$
 (3)

Thus, the likelihood of $L(\mathbf{w},\mathbf{h},\Xi,e|D)$ in Eqn. (2) becomes $L(q_i,\Xi,e|D).$

We adjust the allele frequency q_i by taking into account of read errors e. This implies that the expected allele frequency of 'REF' read as 'ALT' is $(1-q_i)e$, and the expected allele frequency of 'ALT' read as 'REF' is q_ie . Thus, the adjusted WSAF becomes:

$$\pi_i = q_i + (1 - q_i)e - q_i e = q_i + (1 - 2q_i)e. \tag{4}$$

We model over-dispersion in read counts relative to Binomial using a Beta-binomial distribution. Specifically, the read counts of 'ALT' are identically and independently distributed (i.i.d.) with probability v_i , i.e. $a_i \sim Binom(a_i+r_i,v_i)$, and $v_i \sim Beta(\alpha,\beta)$, where $v_i = \alpha/(\alpha+\beta)$. Let $\alpha = c \cdot \pi_i$, $\beta = c \cdot (1-\pi_i)$ and π_i is calculated by Eqn. (4), we can formalise the likelihood of the data using:

$$L(q_i, \Xi, e|D) \propto \frac{\Gamma(a_i + c \cdot \pi_i)\Gamma(r_i + c \cdot (1 - \pi_i))}{\Gamma(c \cdot \pi_i)\Gamma(c \cdot (1 - \pi_i))}.$$
 (5)

2.2.2 Priors

• We use a multinormal prior for the proportions $\mathbf{w}|n$: A multivariate normal variable titre $\mathbf{x}|n=[x_1,\ldots,x_n]$ is proporsed, where x_i is i.i.d. normally distributed from $N(\eta,\sigma^2)$. We then compute w_k as

$$w_k = \frac{\exp(x_k)}{\sum_{k=1}^n \exp(x_k)}.$$

Since x_i s are i.i.d. normally distributed. The prior $P(\mathbf{w}|n) = \prod_{i=1}^{n} \Phi(x_i)$ where $\Phi(x_i)$ is the distribution function of x_i

$$\prod_{k=1} \Phi(x_i), \text{ where } \Phi(x_i) \text{ is the distribution function of } x_i.$$

- For the prior on h|ω, Ξ, we use Li and Stephens (2003)'s hidden Markov model framework as a starting point, with the following modifications made:
- likelihood of data given the expected WSAF rather than the "product of approximate conditionals" (PAC).
- multiple strains with variable proportion rather than two sequences with strict diploidity.
- * simplifying the mutation model with a fixed miss copying operation.

Let ξ_k denote the path in the reference panel Ξ that h_k is copying from; and μ denote the probability of miss copying:

$$\begin{cases} P(\xi_k = h_k) &= 1 - \mu, \\ P(\xi_k \neq h_k) &= \mu, \end{cases}$$

We can therefore express the prior on each h_k as

$$P(h_k|\mathbf{w},\Xi,e) = P(\xi_k = a|\mathbf{w},\Xi,e)(1-\mu) + P(\xi_k = 1 - a|\mathbf{w},\Xi,e)\mu,$$
(6)

where $a\in\{0,1\}$, and 1-a indicates the event that allelic state h_k differs from ξ_k . We can therefore transform the prior on \mathbf{h}_k into the form of a prior on ξ_k , and the prior of the path is simply the transmission probabilities between two loci of a given recombination model. For position i>1, let ρ_i' denote the probability of \mathbf{no} recombination from site i-1 to i, we have $\rho_i'=\exp(-\psi_i)$, where ψ is the recombnation rate. Thus, the probability of recombining from any strain in the panel is $\frac{1-\rho_i'}{|\Xi|}$, where $|\Xi|$ is the size of the panel.





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2.3 Inference

Overall, our method generates Markov chain Monte Carlo (MCMC) samples for the proportions \mathbf{w} and the haplotypes \mathbf{h} given a fixed number of strains. As for the MCMC updates on the prior, we use a Metropolis-Hastings algorithm to sample proportions (\mathbf{w}) given \mathbf{h} ; and use Gib sampler to update \mathbf{h} for a given \mathbf{w} , which are further divided into cases of updating only one haplotype and updating a pair of haplotypes.

2.3.1 Metropolis-Hastings update for proportions

We use a sparse update on $\mathbf{w}|n$, by updating $\mathbf{x}|n$. More specificly, we propose new x_i' s from $x_i' = x_i + \delta x$, where $\delta x \sim N(0, \sigma^2/s)$, and s is a scaling factor. The new proposed proportion is therefore $\frac{\exp(x_k')}{\sum_{k=1}^n \exp(x_k')}$. Since the proposal distribution is symmtrical, we have hasting ratio as 1. A new update is accepted with the probability of:

$$\min\left(1, \frac{P(\mathbf{w}'|n)}{P(\mathbf{w}|n)} \frac{L(\mathbf{w}', \mathbf{h}, \Xi, e|D)}{L(\mathbf{w}, \mathbf{h}, \Xi, e|D)}\right).$$

2.3.2 Gibbs update for single haplotype

We choose haplotype strain s uniformly at random from K strains, considering both cases for updating the state of strain s at position i to 0 and 1, we compute the WSAF and its associated likelihood as follows: Regardless the state of strain s at position i, we firstly remove it from the current WSAF, i.e. subtract $w_s \cdot h_s$ from Eqn. (3), which gives

$$q_{i,-s} = \sum_{k \neq s} w_k \cdot h_k = \text{Eqn.}(3) - w_s \cdot h_s. \tag{7}$$

Therefore, updating strain s of state 0 and 1, so the expected WSAF becomes

$$q_{i,h_s=0} = \text{Eqn.}(7)$$
 (8)

$$q_{i,h_s=1} = \text{Eqn.}(7) + w_s \times 1$$
 (9)

Substitute equations (8) and (9) into Eqn. (5) to compute associated likelihood $L(q_{i,h_s},\Xi,e|D)$, which is expressed as $L(h_s|D)$ in the rest of the paper. Therefore, Eqn. (2) then becomes

$$P(h_s|D) \propto L(h_s|D) \times P(h_s|\mathbf{w}, \Xi, e).$$
 (10)

Instead of computing $P(h_s|D)$ directly, we consider the posterior probabilities of the path p where h_s is copying from:

$$P(g_p|D) \propto L(g_p|D) \times P(g_p|\mathbf{w}, \Xi, e).$$
 (11)

Similar to the prior on $\mathbf{h}|\Xi,e$ (see Eqn.(6)), we can compute the posterior probabilities of h_k by $P(g_p|D)$ via:

$$P(h_s = a|D) = \begin{cases} P(g_p = a|D) \cdot (1 - \mu), & h_s = g_p; \\ P(g_p = 1 - a|D) \cdot \mu, & h_s \neq g_p. \end{cases}$$
(12)

where $a\in\{0,1\}$, and 1-a indicates the event that allelic state h_k differs from g_p . Note that g_p is the same as ξ_k , but indexed differently: g_p is indexed by the reference strain index p from the panel.

In addition to updating the haplotypes from the panel, we take into account of miss copying (see example shown in Fig. 1), which allow the actual genotype differ from the path. Therefore, this method favours mutation rather than recombination for a single-site difference from a reference strain. Our model benefits from combining information from both the reference haplotypes as well as the data. For *de novo* mutations which are not found reference panel, our method will infer mutations based on read count from data. Specific steps are as following:

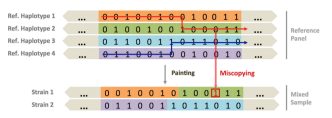


Fig. 1. Illustration of Li and Stephens (2003)'s algorithm. Strain 1 haplotype is made up from reference haplotype segments of 1 and 2; and strain 2 haplotype is made up from reference haplotype segments of 3 and 4. With miss copying, we allow strain states differ from the path: At the third last position of strain 1, the path is copied from reference haplotype 2, with the state of "0".

1. Consider the likelihood as the emission probabilities at site i. Let's use g_p and h_s to denote the allelic state of the copied path and the updated strain respectively.

$$L(g_p = a|D) = L(h_s = a|D) \times P(g_p = h_s) +$$

$$L(h_s = 1 - a|D) \times P(g_p \neq h_s)$$
(13)

where $a \in \{0, 1\}$, and 1 - a indicates the event that h_s takes value that differs from g_p .

2. For the prior on $g_p|\mathbf{w},\Xi,e$, it is simply the Li and Stephens (2003) HMM transition probabilities from position i-1 to i: the probabilites ρ_i' and $\frac{1-\rho_i'}{|\Xi|}$ reflect no recombination and recombination events respectively. Therefore, we have

$$P(g_p|\mathbf{w},\Xi,e) = \rho_i' \cdot P_{i-1}(g_p|D) + \frac{1-\rho_i'}{|\Xi|} \cdot \sum_{x \in \Xi} P_{i-1}(g_x|D).$$
(14)

Combine equations (13) and (14), the posterior probability of path (reference strain) p at position i is then obtained (Eqn. (11)).

3. Given the posterior probabilities of path (reference strain) *p* at position *i*, we sample the path backwards to generate a MCMC sample for the path that strain *s* copied from. We start at the end position, first propose if a recombination events have had happened with the probabilities proportional to

$$\begin{cases} \rho_i' \cdot P_i(g_p | D) & \text{no recombination,} \\ (1 - \rho_i') \cdot \sum_{x \in \Xi} P_{i-1}(g_x | D) & \text{recombination,} \end{cases}$$

if it recombines, sample path backwards according $P_i(g_p|D)$.

4. Lastly, we generate a MCMC sample of the haplotype from the MCMC sample of the copied path using the probabilities expressed in Eqn.(12).

2.3.3 Gibbs update for a pair of haplotypes

In order to improve the MCMC mixing, we consider pairs of haplotypes and update both strain simultaneously. Suppose random sampling two strains to update, namely, s_1 and s_2 . Similarly to Eqn. (7), we first remove their states from the WSAF:

$$\begin{aligned} q_{i,-s_1,-s_2} &= \sum_{k \neq s_1,s_2} w_k \cdot h_k \\ &= \text{Eqn.} (3) - w_{s_1} \cdot h_{s_1} - w_{s_2} \cdot h_{s_2} \end{aligned} \tag{15}$$



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Considering all four possible combination of genotypes, we have

$$q_{i,h_{s_1}=0,h_{s_2}=0} = \text{Eqn.} (15)$$
 (16)

$$q_{i,h_{s_1}=0,h_{s_2}=0} = \text{Eqn.} (15) + w_{s_1} \times 1$$
 (17)

$$q_{i,h_{s_1}=0,h_{s_2}=1} = \text{Eqn.} (15) + \cdot w_{s_2} \times 1 \tag{18} \label{eq:18}$$

$$q_{i,h_{s_1}=0,h_{s_2}=1} = \text{Eqn.}(15) + w_{s_1} \times 1 + w_{s_2} \times 1$$
 (19)

Substitute expressions. (16) to (19), into Eqn. (5), we then obtain their associated likelihood $L(q_{i,h_{s_1},h_{s_2}},\Xi,e|D)$, which is denoted as $L(h_{s_1},h_{s_2}|D)$ in the rest of the paper.

Thus, similar to Eqn. (??), we obtain the following posterior probability, and sample the state (genotype) of strains s_1 and s_2 simultaneously at site i:

$$P(h_{s_1}, h_{s_2}|D) \propto L(h_{s_1}, h_{s_2}|D) \times P(h_{s_1}, h_{s_2}|\mathbf{w}, \Xi, e).$$
 (20)

Similarly to the previous section, instead of working directly with posterior probability $P(h_{s_1},h_{s_2}|D)$, we consider the posterior probabilities of the path p_1 and p_2 where s_1 and s_2 are copying from respectively:

$$P(g_{p_1},g_{p_2}|D) \propto L(h_{p_1},h_{p_2}|D) \times P(g_{p_1},g_{p_2}|\mathbf{w},\Xi,e). \tag{21} \label{eq:21}$$

Then we can compute the posterior probabilities of the pair h_{s_1} and h_{s_2} from

$$P\left(\begin{array}{c} h_{s_{1}} = a, \\ h_{s_{2}} = b \end{array} \middle| D\right) = \begin{cases} P\left(\begin{array}{c} g_{p_{1}} = a, \\ g_{p_{2}} = b \end{array} \middle| D\right) \cdot (1 - \mu) \cdot (1 - \mu), \\ P\left(\begin{array}{c} (g_{p_{1}} = a, \\ g_{p_{2}} = 1 - b \end{array} \middle| D\right) \cdot (1 - \mu) \cdot \mu, \\ P\left(\begin{array}{c} (g_{p_{1}} = 1 - a, \\ g_{p_{2}} = b \end{array} \middle| D\right) \cdot \mu \cdot (1 - \mu), \\ P\left(\begin{array}{c} (g_{p_{1}} = 1 - a, \\ g_{p_{2}} = 1 - b \end{array} \middle| D\right) \cdot \mu \cdot \mu, \end{cases}$$

where $a,b\in\{0,1\}$, and 1-a indicates the event that allelic state h_{s_1} differs from g_{p_1} , and 1-b indicates the event that allelic state h_{s_2} differs from g_{p_2} .

Specificly, we carry out the following steps:

1. Compute the likelihood of the allelic state pairs of the path given data:

$$L\left(\begin{array}{c}g_{p_{1}}=a,\\g_{p_{2}}=b\end{array}\middle|D\right) = L\left(\begin{array}{c}h_{s_{1}}=a,\\h_{s_{2}}=b\end{array}\middle|D\right) \times P\left(\begin{array}{c}g_{p_{1}}=h_{s_{1}},\\g_{p_{2}}=h_{s_{2}}\end{array}\right) + \\ L\left(\begin{array}{c}h_{s_{1}}=a,\\h_{s_{2}}=1-b}\middle|D\right) \times P\left(\begin{array}{c}g_{p_{1}}=h_{s_{1}},\\g_{p_{2}}\neq h_{s_{2}}\end{array}\right) + \\ L\left(\begin{array}{c}h_{s_{1}}=1-a,\\h_{s_{2}}=b\end{array}\middle|D\right) \times P\left(\begin{array}{c}g_{p_{1}}\neq h_{s_{1}},\\g_{p_{2}}=h_{s_{2}}\end{array}\right) + \\ L\left(\begin{array}{c}h_{s_{1}}=1-a,\\h_{s_{2}}=1-b\end{array}\middle|D\right) \times P\left(\begin{array}{c}g_{p_{1}}\neq h_{s_{1}},\\g_{p_{2}}\neq h_{s_{2}}\end{array}\right)$$

$$(23)$$

where

$$\begin{split} &P(g_{p_1} = h_{s_1}, g_{p_2} = h_{s_2}) = (1 - \mu) \cdot (1 - \mu), \\ &P(g_{p_1} \neq h_{s_1}, g_{p_2} = h_{s_2}) = \mu \cdot (1 - \mu), \\ &P(g_{p_1} = h_{s_1}, g_{p_2} \neq h_{s_2}) = \mu \cdot (1 - \mu), \\ &P(g_{p_1} \neq h_{s_1}, g_{p_2} \neq h_{s_2}) = \mu \cdot \mu, \end{split}$$

and $a,b\in\{0,1\}$, and 1-a indicates the event that allelic state h_{s_1} differs from g_{p_1} , and 1-b indicates the event that allelic state h_{s_2} differs from g_{p_2} .

2. For the prior of the allelic state pairs of the path, we consider all possible pair of the copying strain, we take into account of the possibility of one strain recombines and the other does not with the probability of $\rho_i' \cdot \frac{1-\rho_i'}{|\Xi|}$; both recombines, with the probability of $\rho_i' \cdot \frac{\rho_i'}{|\Xi|}$, assuming that recombination events of two copying strains are independent from each other. Thus, we have the prior of the path as

$$P(g_{p_{1}}, g_{p_{2}} | \mathbf{w}, \Xi, e) = P_{i-1}(g_{p_{1}}, g_{p_{2}} | D) \cdot \rho'_{i} \cdot \rho'_{i} + \sum_{x \in \Xi} P_{i-1}(g_{p_{1}}, g_{x} | D) \cdot \rho'_{i} \cdot \frac{1 - \rho'_{i}}{|\Xi|} + \sum_{y \in \Xi} P_{i-1}(g_{y}, g_{p_{2}} | D) \cdot \rho'_{i} \cdot \frac{1 - \rho'_{i}}{|\Xi|} + \sum_{x,y \in \Xi \cdot \Xi} P_{i-1}(g_{x}, g_{y} | D) \cdot \frac{1 - \rho'_{i}}{|\Xi|} \cdot \frac{1 - \rho'_{i}}{|\Xi|}.$$

$$(24)$$

Combine equations (23) and (24), Eqn. (21) is then obtained.

3. Given the posterior probabilities of path pair p1, p2 at position i, we sample the path backwards to generate a MCMC sample for the path pair that strain s1 and s2 copied from. We start at the end position, first propose if a recombination events have had happened with the probabilities proportional to

$$\begin{cases} \sum_{x \in \Xi} P_{i-1}(g_{p_1}, g_x \big| D) \cdot \frac{1 - \rho_i'}{|\Xi|} \cdot \rho_i', & p_1 \text{ recombines,} \\ P_{i-1}(g_{p_1}, g_{p_2} \big| D) \cdot \rho_i' \cdot \rho_i', & \text{no recombination} \\ \sum_{y \in \Xi} P_{i-1}(g_y, g_{p_2} \big| D) \cdot \frac{1 - \rho_i'}{|\Xi|} \cdot \rho_i', & p_2 \text{ recombines,} \\ \sum_{x,y \in \Xi \cdot \Xi} P_{i-1}(g_x, g_y \big| D) \cdot \frac{1 - \rho_i'}{|\Xi|} \cdot \frac{1 - \rho_i'}{|\Xi|}, & \text{both recombine.} \end{cases}$$

If both strains recombine, sample the path, according to $P_i(g_{p_1},g_{p_2}|D)$. If one of the strains recombine, sample the path according to the marginal probability of $P_i(g_p|D)$.

 Lastly, we generate a MCMC sample of the haplotypes from the MCMC sample of the copied paths using the probabilities expressed in Eqn.(22).

2.4 Implementation details

- **Fixed number of strains**. We assume there are more strains than actually present, starting the MCMC chain with a fixed n. As the proportion values drop, we "zero-out" "noisy" strains. For example, suppose that n=5 and the inferred proportions $\mathbf{w}=[0.848, 6.36e-05, 6.81e-05, 0.152, 3.31e-05]$, we drop the strains with proportions less than 0.01, and keep the first and the forth strains.
- Parameters used. In practice, we set the parameters c=100 (Eqn. (5)); $\eta=0$, $\sigma^2=3$ and s=40 (sections 2.2.2 and 2.3.1).
- Update with linkage disequilibrium information. We assume a uniform recombination map, genetic distances between loci i and i+1 are computed by $G_i = D_i/Morgan$ where D_i denotes the physical distance between loci i and i+1 in nucleotide, Morgan is the average morgan distance, which we use 1500000. Suppose the recombination rate ψ_i is given by $\psi_i = N_e G_i$, with $N_e = 10$ being the effective population size. Note that we scale the probabilities with the number of haplotypes in the reference panel.
- Update without LD When assume there is infinite recombination events happening between two adjacent loci, the LD structure hardly









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play any role in the inference. In which case, we can assume independence between site i and i+1. In particular, we use the PLAF as the prior of $P(h_s|\mathbf{w},\Xi,e)$ in Eqn. (10). Further more, assume independence between s_1 and s_2 in Eqn. (20), we compute $P(h_{s_1},h_{s_2}|\mathbf{w},\Xi,e)$ as a product of $P(h_{s_1}|\mathbf{w},\Xi,e)$ and $P(h_{s_2}|\mathbf{w},\Xi,e)$, where $P(h_{s_*}|\mathbf{w},\Xi,e) = PLAF$.

- Model selection Since the final iteration of the MCMC is taken as a point estimate to infer the haplotypes and proportion, the deconvolution process is repeated with different random seeds. We use the lowest deviance information criterion (DIC) to select the best fit model (best chain for the same inferred n). The DIC is calculated from the generated MCMC simulation, and penalised by the average deviance. More specifically, We compute the deviance by $D_{\mathbf{w},\mathbf{h}} = -2\log(L(\mathbf{w},\mathbf{h}|D))$ and $DIC = 2\bar{D} D_{\mathbf{w},\mathbf{h}}$, where \bar{D} is the average deviance of the MCMC chain.
- Pf3k Deconvolution. When deconvolute the Pf3k(Pf3k, 2016) project
 data, we assume clonal sample haplotypes capture the diversities of
 haplotype structures given all samples. In particular, we deconvolute
 clonal samples without LD, then use the deconvoluted clonal
 haplotypes as the reference strains for start, and deconvolute the rest
 mixed samples.

3 Validation and Performance

As a validation set we used a set of *in vitro* mixtures created by Wendler (2015) to simulate mixed infection, which is an ideal validation data set in our use. In this data set, DNA was extracted from four laboratory parasite lines: 3D7, Dd2, HB3 and 7G8, and mixed with different proportions of mixed infection (see Table 2 in brackets), and submitted to the MalariaGEN pipeline (MalariaGEN, 2008) for Illumina sequencing and genotyping pipeline (Manske et al., 2012).

Note that this data set only contains two clonal samples. Due to its limited size, they are not ideal for constructing a reference panel. Moreover, the *P. falciparum* genetic crosses project (Miles et al., 2016) finds that due to sequencing error or applying different variant calling methods, genotype calls vary at the same position given the same strain of *P. falciparum*. Thus we apply inference methods to mutiple samples that contains the same parasite strains, to infer the genotypes of a reference strain.

Infer haplotypes for Dd2 strain. Since 3D7 is the reference strain, we can assume that strain Dd2 is the only source of 'ALT' reads in samples from PG0389-C to PG0394-C. Assume markers are independent from each other, let y be the read count for 'ALT' allele and x be the total read count weighted by the Dd2 mixing proportion (see Table 2 in brackets), we use regression model $y=\beta_0+\beta_1x$ to infer the Dd2 genotype: 1 if β_1 is significant with p-values below 0.001; 0 otherwise.

Infer haplotypes for HB3 and 7G8. Similarly, for sample from PG0398-C to PG0415-C, we let variables x_1 , x_2 be the coverages weighted by the mixing proportions of HB3 and 7G8 respectively; use regression model $y = \beta_0 + \beta_1 x_1 + \beta_2 x_2$ to infer the genotypes of HB3 and 7G8: HB3 is 1 if β_2 is significant with p-values below 0.001; 0 otherwise; similarly for 7G8.

3.1 Accuracy

3.1.1 Proportions and number of strains

To validate our method we applied our program DEploid to 27 lab-mixed *in vitro* samples. As described in section 2.3.1, we start by assuming at most three strains present in the mixtures; and discard strains with less than 1% relative abundance. Our method successfully recovers the proportions

3D7	Dd2	HB3	7G8
88.5 (90)	11.5 (10)	0	0
79.8 (80)	20.2 (20)	0	0
66.1 (67)	33.9 (33)	0	0
31.2 (33)	68.8 (67	0	0
18.4 (20)	81.6 (80)	0	0
9.1 (10)	90.1 (90)	0	0
0	33.6 (33.3)	35 (33.3)	31.3 (33.3)
0	25.9 (25)	26.1 (25)	48 (50)
0	14.7 (14.3)	15.3 (14.3)	69.9 (71.4)
0	0	45.1+54.9 (100)	0
0	0	56.7+40.9 (99)	2.4(1)
0	0	39.5+57.5 (95)	3 (5)
0	0	33.3+56.7 (90)	10 (10)
0	0	85.2 (85)	14.8 (15)
0	0	80.1 (80)	19.3 (20)
0	0	75.4 (75)	24.6 (25)
0	0	70.6 (70)	29.4 (30)
0	0	61 (60)	39 (40)
0	0	50.5 (50)	49.5 (50)
0	0	40.1 (40)	59.2 (60)
0	0	30.1 (30)	69.1 (70)
0	0	25.9 (25)	73,4 (75)
0	0	21.4 (20)	78.5 (80)
0	0	15.2 (15)	84.8 (85)
0	0	3.8 (5)	96.2 (95)
0	0	0(1)	29.9+70.1 (99)
0	0	0	30.0+70.0 (100)
	88.5 (90) 79.8 (80) 66.1 (67) 31.2 (33) 18.4 (20) 9.1 (10) 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	88.5 (90) 11.5 (10) 79.8 (80) 20.2 (20) 66.1 (67) 33.9 (33) 31.2 (33) 68.8 (67 18.4 (20) 81.6 (80) 9.1 (10) 90.1 (90) 0 33.6 (33.3) 0 25.9 (25) 0 14.7 (14.3) 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	88.5 (90) 11.5 (10) 0 79.8 (80) 20.2 (20) 0 66.1 (67) 33.9 (33) 0 31.2 (33) 68.8 (67) 0 18.4 (20) 81.6 (80) 0 9.1 (10) 90.1 (90) 0 0 33.6 (33.3) 35 (33.3) 0 25.9 (25) 26.1 (25) 0 14.7 (14.3) 15.3 (14.3) 0 0 45.1+54.9 (100) 0 0 56.7+40.9 (99) 0 0 39.5+57.5 (95) 0 0 33.3+56.7 (90) 0 0 85.2 (85) 0 0 85.2 (85) 0 0 85.4 (75) 0 0 75.4 (75) 0 0 75.4 (75) 0 0 70.6 (70) 0 0 70.6 (70) 0 0 40.1 (40) 0 0 30.1 (30) 0 0 25.9 (25)

Table 2. Inferred percentages (true in brackets) of the mixed samples. Our model overfits clonal sample PG0398-C as a mixture of two strains each with the abundance proportions of 45.1% and 54.9%. We then find that both inferred haplotypes are referring to the same strain HB3. Hence we use the "+" sign to denote the overfitting events, and both haplotypes are inferring the same strain.

with haplotypes of the input (see Table 2). The deviation between our proportion estimates and the truth is at most 2%.

Our model overfits the noisy lab-mixed sample with additional strains. Note that in Table 2, we infer six of the HB3 and 7G8 mixtures as mixing of three, two of which haplotypes have subtle difference with the same parasite line, but overall vastly different from the last strain. The subtle variation is caused by few heterozygous sites with high coverage resulting high leverage in our model (supplemental material Figure S2.3(a)). The source of the noisy markers are possibly from sequencing or variant calling process, which are not recalabriated by our program.

We experimented deconvoluting the 27 lab-mixed samples with the following reference panels:

- panel I: five Asian and five African clonal strains from the Pf3k(Pf3k, 2016) data base: PD0498-C, PD0500-C, PD0660-C, PH0047-Cx, PH0064-C, PT0002-CW, PT0007-CW, PT0008-CW, PT0014-CW, PT0018-CW.
- panel II: panel I with the addition of HB3;
- panel III: panel II with the addition of 7G8;
- panel IV: panel III with the addition of Dd2;
- panel V: 3D7, HB3, 7G8 and Dd2 strains.
- panel VI: panel I with the addition of six (three each) clonal strains from Asia and Africa: PH0193-C, PH0283-C, PH0305, PT0060-C, PT0146-C and PT0158-C.









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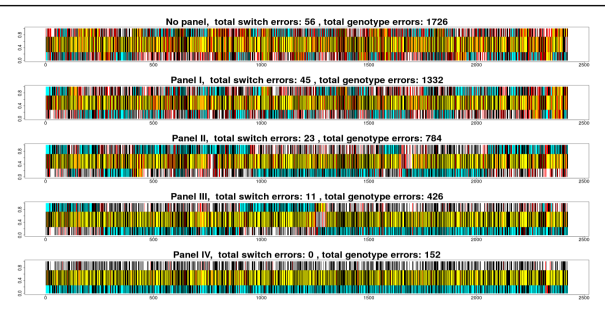


Fig. 2. Haplotypes comparison of sample PG0396-C chromosome 14 deconvolution without any reference strain (top) versus with using reference panels I to IV (from the second to the bottom). Black bars indicate alternative alleles; red bars mark wrongly inferred positions. The yellow, cyan and white background label the haplotype segments from strains 7G8, HB3 and Dd2 respectively. The switch errors are obtained by counting the changes of a strain mapped to reference strains; the genotype errors are the discordance between the strain and the mapped reference segments.

In all cases we estimated the number and proportion of strains accurately, for example Figure 2 y-axises show the proportions of strains Dd2/7G8/HB3 as approximately $\frac{1}{4}/\frac{1}{2}/\frac{1}{4}$.

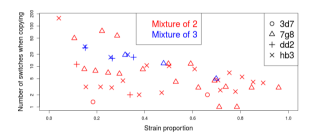
3.1.2 Haplotypes

Our accuracy assessment for inferred haplotypes take into account of both switch errors and genotype discordance, which reflect to the recombination and miss copying events in the method section. Intuitively, one may suggest to use the Li and Stephens model to compute the viterbi path or posterior probabilities to assess the how different our inferred haplotypes differ from the truth. However, we find that such methods overestimate switch errors at short segments of sequence due to variable reference panel quality.

Moreover, we want to assess the switch errors by taking into account of switches from the other haplotypes. Note that in Figure 2, the top and bottom strains have similar proportions, which are difficult to phase without a perfect reference panel. One may flip parts of the two strains to resolve the errors. Unfortunately, our Li and Stephens implementation focuses on a single strain (two strains at the most), the viterbi path or posterior probabilities fail to align the switches of all strains. Therefore, we have taken a heuristic approach by dividing the inferred haplotypes into segments with length of 50, then mapped onto reference strains of panel V according to the indices. The switch errors are obtained by counting the changes of a strain mapped to reference strains; the genotype errors are the discordance between the strain and the mapped reference segments.

From our assessment of haplotype inference, we conclude:

- The inference of relative proportions do not seem to be affected by the use of a reference panel or its quality (Figure 2).
- The accuracy of the haplotype is dependent on having an appropriate reference panel (Figure 2).
- The strain proportion affects the haplotype inference (see Fig 3). Our method infer strains with proportions over 20% accurately, but struggles with minor strains due to insufficient data, in particularly at sites when the minor strain should be alternative allele, and the dominant strain should be the reference allele (see Figure 3).



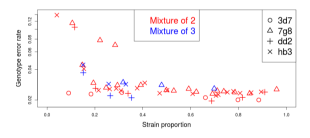


Fig. 3. Error rates vs. strain proportions. We use reference panel V to deconvolute all 27 samples. Each marker represent a deconvoluted haplotype with 18,570 sites.

3.2 Run-time

The complexity of our program is $\mathcal{O}(lm^2)$ (see Fig 4), where l and m are the number of reference strains and sites respectively. In practice, we divide Pf3k samples into several geographical regions and perform deconvolution, with ten most different local clonal strains as reference panel. The run time of deconvoluting a field sample range between 1 and 6 hours, depending on the number variants in a sample: For example, it takes $5\frac{1}{2}$ hours to process sample QG0182-C over 372,884 sites.





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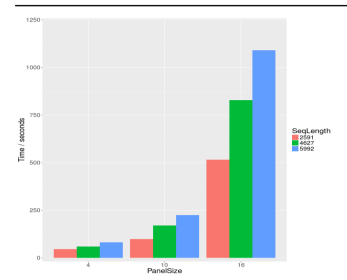


Fig. 4. As a demonstration, we deconvolute chromosome 14, chromosomes 13 and 14, chromosomes 12 to 14 of sample PG0412-C with reference panels I, V and VI. The runtime is almost linear respect to the number of sites; and shows quadratic trend against the number of reference strains.

4 Discussion

The program DEploid and its analysis pipeline is originally developed for *P. falciparum* studies. With some specific minor parameter changes, DEploid can be used for deconvolute *Plasmodium vivax* sequence data (Pearson et al., 2016). The framework is suitable for deconvoluting mixed genomes with unknown proportions. It can thus be extended to a wider range of applications, such as deconvoluting cancer tumour cell genomes or Ebola virus genomes.

Acknowledgements

We thank valuable insights and suggestions from Roberto Amato, John O'Brien, Richard Pearson, and Jason Wendler for providing the data of artificial samples. We thank Zam Iqbal for naming the program DEploid.

Funding

This project is funded by the Wellcome Trust grant [100956/Z/13/Z].

Conflict of Interest: none declared.

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