# Approaches to estimating inbreeding coefficients in clinical isolates of Plasmodium falciparum from genomic sequence data 

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#### Abstract

A recent genomic characterization of more than 200 Plasmodium falciparum samples isolated from the bloodstreams of clinical patients across three continents further supports the presence of significant strain mixture within infections. Consistent with previous studies, these data suggest that the degree of genetic strain admixture within infections varies significantly both within and across populations. The life cycle of the parasite implies that the mixture of multiple genotypes within an infected individual controls the out-crossing rate across populations, making methods for measuring this process in situ central to understanding the genetic epidemiology of the disease. Peculiar features of the $P$. falciparum genome mean that standard methods for assessing structure within a population - inbreeding coefficients and related $F$-statistics - cannot be used directly. Here we review an initial effort to estimate the degree of mixture within clinical isolates of P. falciparum using these statistics, and provide several generalizations using both frequentist and Bayesian approaches. Using the Bayesian approach, based on the Balding-Nichols model, we provide estimates of inbreeding coefficients for 168 samples from northern Ghana and find significant admixture in more than $70 \%$ of samples, and characterize the model fit using posterior predictive checks. We also compare this approach to a recently introduced mixture model and find that for a significant


minority of samples the F-statistic-based approach provides a significantly better explanation for the data. We show how to extend this model to a multi-level testing framework that can integrate other data types and use it to demonstrate that transmission intensity significantly associates with degree of structure of within-sample mixture in northern Ghana.

## INTRODUCTION

The protozoan parasite Plasmodium falciparum causes most cases of severe malaria and presents one of humanity's most significant public health burdens, killing at least half a million people a year [16, 39]. The parasite's ability to develop resistance to drugs and the rapid spread of that resistance across geographically-separated populations presents a constant threat to international control efforts [23, 27]. While genetic factors play a crucial role in resistance, many aspects of the genetic epidemiology of the parasite remain obscure [33, 35]. The beginnings of a global perspective on the genetic structure of this problem emerged from the analysis of whole-genome sequencing (WGS) data derived from $\sim 200$ parasite genomes collected directly from clinical patients in six countries on three continents [20]. This study provides strong evidence for significant continentalscale structure in the amount of genetic variation present within populations, as well as indicating frequent but variable amounts of within-isolate strain mixture, often referred to as multiplicity of infection (MOI) [8].

Within-isolate $P$. falciparum strain mixture may result from a host individual being infected by a mosquito carrying several distinct strains of the parasite, being infected successively by several mosquitoes carrying single strains, or some combination of the two. The parasites's sexual mating process occurs only in the period immediately after being taken up into the mosquito hindgut as a blood meal implying that mixed infections - either within the host or vector - provide the essential grist for the maintenance of population-level genetic diversity by creating an opportunity for out-crossing [15, 7]. The degree of apparent mixture within an infected individual's bloodstream then would largely depend on the number of infective events and the effective population size of the surrounding $P$. falciparum population [3]. Other effects, such as genetic interactions with the vector, density-dependent selection, non-random mating and host immune response, may also play a role. Consequently, reliable techniques for measuring the structure of mixture within clinical infections will give researchers a quantitative measurement of potential out-crossing - a key determinant of
genetic epidemiological control - and may provide insight into other important biological effects, such as host immune response. This paper presents a collection of statistical methods for capturing the amount of mixture within clinical isolates using WGS data.

Historically, the $P$. falciparum research focuses on MOI, the minimal number of distinct genetic strains identifiable in a clinical sample. Researchers have largely used microsatellite, RFLP, or SNP data to infer the number of strains [8, 1, 3, 18] and have associated these values with morbidity [21, 26], patient age [9, 32], the course of infection in pregnancy [31], the presence of bed nets [10], and a number of other conditions [6]. The focus on the number of strains naturally follows from the limits of earlier genotyping techniques since more subtle features, such as the mixture proportions of strains or recombination events among them, could not be readily identified.

The introduction of WGS data naturally enforces a generalization from MOI into more complex measures of mixture, since, in a strict sense, we find find evidence for some level of MOI in nearly all of our samples. For instance, we present four samples in Figure 1 with within-sample observed allele frequency plotted against the population-level allele frequency for all relevant SNPs. The left plot shows little apparent mixture. Previous technologies limited measurement of the number of genetic types to small regions of the genome, effectively bounding the maximum number of strains that could be identified. This has led to considerations of complexity of infection (COI), where a strain mixture model is used to explain these within sample mixture levels [11, expanding on traditional methods for inferring MOI [17. Below, we show below that this model, while powerful in modeling patterns within some isolates, is not sufficient to explain observed mixture within all samples.

F-statistics, measurements of the departure of allelic heterozygosity observed within a population from those expected at Hardy-Weinberg equilibrium (HWE), make for a natural approach to quantifying the structure within mixed samples [37, 24]. HWE specifies the distribution of alleles assuming panmixia, a population exhibiting perfectly random mating with an absence of mutation, migration, drift, selection or other effects [40]. F-statistics calibrate the empirical allele distribution within a subpopulation against those expected under HWE, ranging from a value of one (no mixture) to zero (perfect mixture). In the context of comparing the parasites' genetic diversity within a single infected individual relative to the local geographic population (and absent any geographic structuring of the population, i.e. the Wahlund effect), these statistics effectively become inbreeding
coefficients. For a sample, $i$, we refer to this value as $f_{i}$.
These methods have proven to be an effective and extremely popular means for investigating species' population structure from both allelic and genomic data [37, 30, 38]. However, standard methods assume specific ploidy structures incommensurate with WGS data from P. falciparum and so cannot be used directly. The critical difference is that, within a human host, P. falciparum exists only in the haploid stage of its life-cycle [15]. Since short read WGS data cannot yet capture full haplotypes, individual reads cannot be uniquely identified with their strain of origin. Without being able to associate reads to individual $P$. falciparum strains, we cannot see any 'out-of-the-box' use of standard $F$-statistics approaches with this new data.
[20] provides an initial estimator for inbreeding coefificents using WGS based on the slope of a modified regression line between the expected heterozygosity assuming population-level HWE and the observed heterozygosity within a sample. [3] explores the connection between this estimator and standard MOI approaches by comparing estimates from WGS with MOI values inferred by genotyping the msp-1 and msp-2 genes, showing strong correlation between these values in their sample sets. They note that the correlation is strongest at high $f_{i}$ and low MOI values, where samples are close to being unmixed, with weaker correlation among more mixed samples, suggesting increasing divergence between these models. This estimator, while providing an effective first effort, does not clearly connect to the larger statistical tradition around $F$-statistics. This paper seeks to clarify this estimator by placing it more firmly within this larger discussion.

This paper proceeds as follows. First, we provide an overview of data collection and layout our notation. We present the initial estimator employed by [20] for estimating $f_{i}$ and provide two additional frequentist estimators and detail their connection to classical $F$ statistics. We then proceed to describe a Bayesian approach for these statistics that builds on the Baldings-Nichols model together with an inference scheme and framework for hierarchical modeling. We use this construction to show that observed transmission intensity, a measure of the amount of infective mosquito activity in the surrounding environment, significantly associates with changes in mixture among 168 northern Ghanaian samples. We then show that, in comparison with a COIL-like approach [11], the Bayesian $F$-statistic is a more powerful explanatory model for a substantial fraction of samples. We conclude with a discussion of the strengths and limitations of our approaches, and possible future directions for modeling within-sample mixture using WGS.

## 1 Model and Data

### 1.1 Data and preparation

The WGS data come from Illumina HiSeq sequencing applied to P. falciparum extracted from 235 clinical blood samples collected from infected patients from the Kassena-Nankana district (KND) region of northern Ghana. This publication uses data from the MalariaGEN Plasmodium falciparum Community Project on www.malariagen.net/projects/parasite/pf Collection occurred over approximately 2 years, from June 2009 to June 2011. The full sequencing protocol and collection regime are described in [20, 2]. After quality control measures, sequencing was performed on 235 samples, and, following a documented protocol using comparison against world-wide variation, 198, 181 single-nucleotide polymorphisms (SNPs) were called within each sample 20]. Each call provides the number of reference and non-reference read counts observed at each variant position within the genome, mapped to the 3D7 reference [12]. Positions that exhibited no variation within the KND samples, any level of missingness (no read counts observed), or minor allele frequency less than 0.05 were excluded. Samples that possessed more than 4000 SNPs called with fewer than 10 read counts were also excluded, following an observed inflection point. These cleaning measures left 1470 SNPs in 168 samples. We observe little apparent population structure among the samples, evidenced either principal components analysis or a neighbor-joining tree of the pairwise difference among samples, as in Supplementary Figure S1.

### 1.2 Notation

We label the samples $i=1, \cdots, N$ and the SNPs by $j=1, \cdots, M$, with $N=194$ and $M=1,470$ if all samples and all SNPs are considered. In some contexts below $M$ may be the number of SNPs within a chromosome, which should be clear by context. At SNP $j$ within sample $i$, we observe $r_{i j}$ reads that agree with the reference, and $n_{i j}$ reads that are different from the reference. We write $p_{i j}$ for the allele frequency for reference allele for SNP $j$ in sample $i$ and estimate it via the maximumlikelihood estimator (MLE) for proportions: $\hat{p}_{i j}=\frac{r_{i j}}{r_{i j}+n_{i j}}$. Similarly, we write $p_{j}$ as population-level reference allele frequency for each SNP and estimate according to the across-sample MLE:

$$
\widehat{p}_{j}=\sum_{i=1}^{N} n_{i j} / \sum_{i=1}^{N}\left(r_{i j}+n_{i j}\right) .
$$

To slightly streamline our notation, we relabel the inbreeding coefficient, $F_{i s}$, for each sample $i$ as $f_{i}$. We provide Table 1 as a reference to the reader for notation.

### 1.3 A previous estimator for $f_{i}$, and two alternatives

In [20], the authors provide an initial approach to estimating $f_{i}$. We refer to this estimator as $f_{i}^{(m)}$ to contrast it with subsequent estimators. This method relies on minor allele frequencies rather than reference allele frequency, which we mark with a tilde so that $p_{j}$ becomes $\widetilde{p}_{j}$. The two quantities are naturally related so that $\widetilde{p}_{j}=p_{j}$ if $p_{j}<0.5$ and $\widetilde{p}_{j}=1-p_{j}$ otherwise. $\widetilde{p}_{i j}$ and $p_{i j}$ are related in the same fashion and we continue to use hats to denote estimates. The estimator $f_{i}^{(m)}$ proceeds sample by sample, so we will consider a generic sample $i$. The estimator first partitions alleles into 11 equally-spaced bins based on their minor allele frequency: $(0,0.05), \cdots,(0.45,0.50)$. Within each bin, $b$, the averaged expected heterozygosity assuming population-level HWE is calculated by

$$
H_{e}(b)=\frac{1}{M_{b}} \sum_{k \in b}^{M_{b}} 2 \cdot \hat{\tilde{p}}_{k} \cdot\left(1-\hat{\tilde{p}}_{k}\right),
$$

where $M_{b}$ is the number of SNPs within bin $b$. The averaged observed heterozygosity within each bin and each sample is calculated by

$$
H_{o}(b, i)=\frac{1}{M_{b}} \sum_{k \in b}^{M_{b}} 2 \cdot \hat{\tilde{p}}_{i k} \cdot\left(1-\hat{\tilde{p}}_{i k}\right) .
$$

Finally, $\hat{f}_{i}^{(m)}$ is calculated as $1-\beta$ where $\beta$ is the slope found by regressing the $H_{o}(b, i)$ values against $H_{b}^{e}$ values centered within their respective allele frequency bins and constrained to pass through the origin.

The binning procedure stabilizes the estimator against influence by the low frequency alleles that dominate the samples. Consequently, this has the result of biasing the estimates towards high frequency alleles. We can remove this effect by discarding the binning procedure in favor of directly regressing observed heterozygoity for each SNP against the expected value, still constrained to pass through the origin. This provides a closed for expression for a new estimator, $f_{i}^{(r)}$, as

$$
\hat{f}_{i}^{(r)}=1-\frac{\sum_{j=1}^{M} \hat{\tilde{p}}_{j} \cdot\left(1-\hat{\tilde{p}}_{j}\right) \cdot \hat{\tilde{p}}_{i j} \cdot\left(1-\hat{\tilde{p}}_{i j}\right)}{\sum_{j=1}^{M} \hat{\tilde{p}}_{j}^{2} \cdot\left(1-\hat{\tilde{p}}_{j}\right)^{2}} .
$$

We can also create a similar estimator but one more transparently derived from the ideas underpinning traditional $F$-statistics in the following way. For a single SNP $j$, suppose $f_{i}$ to be the fraction of the population-level heterozygosity equal to the difference between the population-level heterozygosity, $H_{j}^{p}$ and the sample-level heterozygosity, $H_{j}^{i}$ that is,

$$
f_{i} \cdot H_{j}^{p}=H_{j}^{p}-H_{j}^{i} .
$$

Dividing through by $H_{j}^{p}$ gives an estimate for $f_{i}$ for the SNP $j$. Averaging across all SNPs, and taking the ratio of expectations to be the expectation of the ratios, gives the estimator

$$
\hat{f}_{i}^{(d)}=1-\frac{\sum_{j=1}^{M} \tilde{p}_{i j}\left(1-\tilde{p}_{i j}\right)}{\sum_{j=1}^{M} \tilde{p}_{j}\left(1-\tilde{p}_{j}\right)} .
$$

For each of these estimators, a corresponding variance calculation is possible. For $f_{i}^{(i)}$ and $f_{i}^{(r)}$ these can be made by re-coursing to known properties of regression lines. For $f_{i}^{(d)}$, a delta approximation can be used. However, we instead employ a more convenient bootstrap approach to capture the variance in the estimates for confidence intervals. For the Bayesian estimates presented below, we can establish credible intervals based on the inferred posterior distribution.

Figures 1 and Table 2 compare the $f_{i}$ estimates made by the initial, regressed and direct estimators. In Figure 1, we present the estimates for four samples, together with the SNP data and the binned values from the initial estimator. For the direct estimator, we construct the line shown by connecting the origin with the $(x, y)$ point of the denominator and numerator of Equation 1.3. The other two estimators' lines come naturally from their regression procedure. The slope of each line is $1-f_{i}$ for that estimator. As shown in Table 2, the correlation of the three estimators is greater than 0.98. In particular, the direct and regressed estimates differ by at most $1 \%$ across all samples. The initial estimator produces values that are almost invariably slightly lower than the two other estimates, by as much as $15 \%$ of the higher value for highly mixed samples.

Despite these differences, these estimators provide strongly consistent portraits of the $f_{i}$ values for the samples in our data. However, they all possess two less-than-desirable properties: they rely on a separate estimate of the allele frequency; and cannot be easily incorporated into a more involved analysis for use in hypothesis testing. Researchers will presumably seek to use estimates of
$f_{i}$ as a means of testing clinical or epidemiological differences between subpopulations. A preferable approach would simultaneously allele frequency across all SNPs and the inbreeding coefficient for each sample, as well as permitting extension to more complex modeling contexts. We submit that our Bayesian models below satisfy these requirements.

### 1.4 Bayesian model framework

We present two models, with the second as a multi-level extension of the first. In the first, we estimate inbreeding coefficients comparable to the above estimators but employing the BaldingsNichols model [4]. In the second model, we show how we can exploit the more flexible Bayesian approach to estimate these values inside of a nested structure that allows us to test how different transmission regimes affect inbreeding coefficients. In both cases, we make several simplifying assumptions. We treat SNPs as being unlinked (i.e. no linkage disequilibrium) and assume that individual parasites within a sample represent a random sample of the surrounding population. We also assume that read counts are sampled identically, independently, and represent an unbiased sample of variation at each position. We will discuss the evidence for and against these assumptions and possibilities for modeling extensions in the discussion.

### 1.4.1 Likelihood and priors

Our approach adapts the Balding-Nichols model of allele frequency within inbred subpopulations to the specific context of $P$. falciparum WGS data [5. In P. falciparum the relevant subpopulation is the collection of parasites within a clinical sample. For sample $i$ and SNP $j$, we assume that each read count arises as an identical and independent Bernoulli process with the probability of a reference read given by the unobserved reference allele frequency $p_{i j}$. Conditional upon an inbreeding coefficient $f_{i}$ and a population-level allele frequency $p_{j}$, the Balding-Nichols model gives the allele frequency $p_{i j}$ as a Beta distribution:

$$
p_{i j} \sim \mathcal{B}\left(\frac{1-f_{i}}{f_{i}} p_{j}, \frac{1-f_{i}}{f_{i}}\left(1-p_{j}\right)\right) .
$$

Since the read counts are assumed to be i.i.d, $p_{i j}$ is drawn from a Beta, and the probability of the data is binomial, we use the conjugacy of these distributions to eliminate the dependence on
the unknown $p_{i j}$ and give a Beta-binomial distribution for the likelihood:

$$
\begin{equation*}
\mathbb{P}\left(r_{i j}, d_{i j} \mid p_{j}, f_{i}\right)=\binom{r_{i j}+n_{i j}}{n_{i j}} \frac{\mathrm{~B}\left(n_{i j}+\frac{1-f_{i}}{f_{i}} p_{j}, r_{i j}+\frac{1-f_{i}}{f_{i}}\left(1-p_{j}\right)\right)}{\mathrm{B}\left(\frac{1-f_{i}}{f_{i}} p_{j}, \frac{1-f_{i}}{f_{i}}\left(1-p_{j}\right)\right)}, \tag{1}
\end{equation*}
$$

where $\mathrm{B}(\cdot, \cdot)$ is the beta function. Since we assume independence by site and by sample, the complete likelihood of the data, $\mathcal{D}$, conditional upon the inbreeding coefficients for all samples, $\mathbf{f}=\left(f_{1}, \cdots, f_{N}\right)$ and the allele frequency for all SNPs $\mathbf{p}=\left(p_{1}, \cdots, p_{M}\right)$ becomes

$$
\mathbb{P}(\mathcal{D} \mid \mathbf{f}, \mathbf{p})=\prod_{i=1}^{N} \prod_{j=1}^{M} \mathbb{P}\left(r_{i j}, r_{i j} \mid f_{i}, p_{j}\right)
$$

In this first model, where we seek to estimate only the inbreeding coefficients for a set of samples, prior specification is straight-forward. The only prior information we have about the $f_{i}$ values suggests that high levels of inbreeding are common but not obligatory in west African populations, and we quantitatively interpret this as a uniform prior on each $f_{i}$. We place a uniform prior on each allele frequency, although we have eliminated rare variants as part of data cleaning described in Section 2.1.

### 1.4.2 A hierarchical extension

For nearly all samples we possess additional metadata on the assessed transmission intensity (TI) in the KND area at the time of P. falciparum sample collection. Field researchers categorize TI as low, medium or high based on the perceived probability of infection from observed mosquito counts, temperature, precipitation, and number of malaria cases entering area clinics. We write $c_{i}$ for the TI of sample $i$, with $c_{i} \in\{0,1,2,3\}$, with 0 representing no record and 1,2 , and 3 denoting low, medium and high transmission, respectively. The collection of all $c_{i}$ 's we write as $\mathbf{c}$.

We extend the previous model to also model the relationship between the distribution of inbreeding coefficients and TI by constructing a model of the inbreeding coefficients in terms of the c. Conditional upon $c_{i}$, we assume that each $f_{i}$ is drawn independently from a Beta distribution with parameters $\alpha_{c_{i}}$ and $\beta_{c_{i}}$. There are consequently four $\alpha$ values and four $\beta$ values and we label the vectors $\boldsymbol{\alpha}$ and $\boldsymbol{\beta}$. We then decompose the posterior distribution of the unobserved parameters conditional upon the read count data and TI values by noting that

$$
\begin{align*}
\mathbb{P}(\mathbf{f}, \mathbf{p}, \boldsymbol{\alpha}, \boldsymbol{\beta} \mid \mathcal{D}, \mathbf{c}) \propto & \mathbb{P}(\mathcal{D}, \mathbf{c} \mid \mathbf{f}, \mathbf{p}, \boldsymbol{\alpha}, \boldsymbol{\beta}) \cdot \mathbb{P}(\mathbf{f}, \mathbf{p}, \boldsymbol{\alpha}, \boldsymbol{\beta}) \\
& \mathbb{P}(\mathcal{D} \mid \mathbf{c}, \mathbf{f}, \mathbf{p}, \boldsymbol{\alpha}, \boldsymbol{\beta}) \cdot \mathbb{P}(\mathbf{c}, \mathbf{f}, \mathbf{p}, \boldsymbol{\alpha}, \boldsymbol{\beta}) . \tag{2}
\end{align*}
$$

As the introdcution of the additional structure does not affect the probability of the read count data, we retain the same likelihood as in Equation 3.

The dependency of $\mathbf{f}$ on $\mathbf{c}, \boldsymbol{\alpha}$, and $\boldsymbol{\beta}$ we specified above. Together with the assumption that the remaining parameters are otherwise independent of each other these facts decompose Equation 2 to

$$
\mathbb{P}(\mathbf{f}, \mathbf{p}, \boldsymbol{\alpha}, \boldsymbol{\beta} \mid \mathcal{D}, \mathbf{c}) \propto \mathbb{P}(\mathcal{D} \mid \mathbf{f}, \mathbf{p}) \cdot \mathbb{P}(\mathbf{f} \mid \mathbf{c}, \boldsymbol{\alpha}, \boldsymbol{\beta}) \cdot \mathbb{P}(\mathbf{p}) \cdot \mathbb{P}(\mathbf{c}) \cdot \mathbb{P}(\boldsymbol{\alpha}) \cdot \mathbb{P}(\boldsymbol{\beta}) .
$$

It remains to specify the four prior terms on the right-hand side of the equation. We assume that the prior distribution should be the same as in the previous model. We take the observations of $\mathbf{c}$ to have probability one since they are the researchers' own assessment technique. For $\boldsymbol{\alpha}$ and $\boldsymbol{\beta}$, we assume that they are drawn independently from an exponential distribution with mean one, except $\alpha_{0}$ and $\beta_{0}$, corresponding to the unrecorded category. For those parameters we fix the values to one, ensuring a uniform prior on the corresponding $f_{i}$ 's.

### 1.4.3 Inference

We use a Metropolis-Hastings Markov chain (MCMC) approach to inference. The MetropolisHastings ratio gives the probability that a proposed parameter update $x^{\prime}$ will be accepted from a current state $x$ with probability $\alpha$ such that

$$
\alpha=\min \left(\frac{\mathbb{P}\left(x^{\prime}\right)}{\mathbb{P}(x)} \cdot \frac{\mathbb{P}\left(x^{\prime} \rightarrow x\right)}{\mathbb{P}\left(x \rightarrow x^{\prime}\right)}, 1\right)=\min \left(\alpha_{1} \cdot \alpha_{2}, 1\right) .
$$

The first ratio is that the posterior probabilities of $x$ and $x^{\prime}$, and we write this as $\alpha_{1}$. The second ratio gives probability of choosing the current state from the proposed state over the reverse move and we write this $\alpha_{2}$. Since $\alpha_{1}$ constitutes assessment of the likelihood and the prior functions that can be calculated directly from the likelihood and priors above, we subsequently only consider $\alpha_{2}$. We denote proposed parameters with an apostrophe.
$\mathbf{f}$ - We randomly select $i$ and propose $f_{i}$ from $\mathcal{B}\left(\alpha_{c_{i}}, \beta_{c_{i}}\right)$, leading to $\alpha_{2}=\frac{\mathcal{B}\left(f_{i} \mid \alpha_{c_{i}}, \beta_{c_{i}}\right)}{\mathcal{B}\left(f_{i}^{\prime} \mid \alpha_{c_{i}}, \beta_{c_{i}}\right)}$.
$\mathbf{p}$ - We randomly select $j$ and then draw the proposed parameter $p_{j}$ from the uniform prior, leading to $\alpha_{2}=1$.
$\boldsymbol{\alpha}, \boldsymbol{\beta}$ - For both of these parameters, we randomly select individual components and propose new values directly from the prior distribution, leading to $\alpha_{2}=\frac{\exp (-x)}{\exp \left(-x^{\prime}\right)}$ where $x$ and $x^{\prime}$ are the current and proposed state of the relevant component.

We examined the autocorrelation of the log-posterior and find that the lag was minimal (Supplementary Figure S2). As a secondary check, we ran chains both for all of the chromosomes individually, as well as all together. Since we treat SNPs as independent, the performance of the model should be unaffected if indeed the model performs similarly across chromosomes. We find that across all chromosomes performance is nearly identical, with greater than $95 \%$ correlation among estimates.

### 1.5 Implementation

All code was implemented in the R computational environment [28]. The set of scripts implementing each of the estimators, the MCMC algorithm, and visualizations, together with subsets of the data sets are available at github.com/jacobian1980/pfmix. This repository includes a manual and workflow for completing analyses using these approaches. All materials are released under a Creative Commons License.

## 2 Results

### 2.1 Comparison with frequentist estimates

In Table 2 we compare between the Bayesian model estimates and each of the frequentist estimators for each of the 168 samples, observing strongly consistent estimates for all samples. We note that the least consistency among estimates on highly mixed samples, although still strongly similar. The Bayesian estimates are noticeably more distinct from the frequentist estimates than they are from each other. In general, the Bayesian estimates are less extreme than their frequentist comparators, likely due to mitigated influence from low-frequency variants. We also observe consistent estimates of variation among the two estimation procedures. For the Bayesian estimate, variation was determined using the maximum a posteriori (MAP) parameters and the properties of the Beta-binomial distribution. For the frequentist estimators, we employed 250 bootstrap samples on the set of SNPs.

### 2.2 Temporal changes in Northern Ghana

We describe above additional metadata collected at the time of the samples, the perceived transmission intensity (PTI). This categorical data gives a measure of the frequency of malaria transmission
and incorporates the rate of malaria cases presenting at the clinic, amount of standing water, irrigation status, and other factors. Given the role that transmission intensity is believed to play in the process of out-crossing, researchers may naturally hypothesize that the $f_{i}$ value of samples is partially determined by PTI. Similar effects have been reported in a variety of investigations [14, 29].

We plot the $95 \%$ credible interval of $1-f_{i}$ values for each sample across the two year period of collection with the PTI coded by the color (Figure 22). We find evidence for significant mixture $\left(f_{i}>0.95\right)$ in more than $70 \%$ of samples $(119 / 168)$. We plot $1-f_{i}$ to show highly mixed sample as having high $1-f_{i}$ values. The plot suggests that the PTI at samples collection corresponds to $f_{i}$, with high PTI yielding low values, low PTI giving high values, and moderate PTI somewhere in between. In a frequentist framework, this hypothesis could be tested either by pairwise comparison of means or by ANOVA. Grouping samples by PTI, we plot the distribution of MAP $f_{i}$ values in Figure 3(upper right), noting that there appears some difference in distribution across the groups. However, pairwise comparison of the mean or variance between groups indicates no difference among the groups, even at a liberal signficance level of 0.1.

In Figure 2, we see the distribution of $\alpha$ and $\beta$ values for each PTI suggests the posterior distribution differs between categories, although in a more complex way than a simple shift in mean. Our hierarchical model allows us to test this hypothesis in a different fashion, using Bayes factors, a measure of the support provided in the data for comparable models [19. We can consider the hypothesis as a comparison between two models that we label $M_{0}$ and $M_{1}$. Under $M_{0}$, the $\alpha$ and $\beta$ values are equal for all categories of PTI, i.e. $\alpha_{1}=\alpha_{2}=\alpha_{3}$ and $\beta_{1}=\beta_{2}=\beta_{3}$. This describes the situation where the distribution for $f_{i}$ is constant across PTI categories. Under $M_{1}$, the $\alpha$ and $\beta$ values are unconstrained, and the $f_{i}$ distribution may vary by PTI class. Notice that $M_{0}$ is nested within $M_{1}$. Because of this, we may use the Savage-Dickey ratio to calculate the Bayes factor. Using the methods set forth in [36], we calculate the Bayes factor using a standard kernel density estimator, as in [34]. The Bayes factor is 11.52 , indicating $M_{1}$ provides a a moderately preferable explanation for the data relative to $M_{0}$.

### 2.3 Comparison with COI model

A recently introduced mixture model attempts to model within-sample allele frequency variation, in keeping with the MOI tradition within Pf genetics [11, 17]. To contrast this model with the one
presented here, we use the BIC to compare model fit between the two for each of the 168 clinical samples. Unfortunately, the implementation of this model was not designed for WGS data, so we rely on a reduced version of the model in [25] that amounts to an equivalent model, detailed in the Appendix. For each sample,

### 2.4 Posterior predictive assessment

While $F$ statistics provide a convenient way to summarize the degree of heterogeneity in a clinical sample, researchers may also be interested in the degree that the model captures the complexities of the biological mixture process. We examine this discrepancy using posterior predictive checks (PPC) [22, 13]. PPCs measure the discrepancy between predictive data and the observed data by some discrepancy measure, for which here we take as a $\chi^{2}$ statistic. For each point in the posterior, we generate a realization of data from that model. By sampling from the posterior and generating data for each sample, we create a predictive data distribution, $y^{\mathrm{pred}}$. We then use the $\chi^{2}$ statistic to generate a $p$-value comparing the observed data. For each SNP, we also plot the distribution of predicted SNP data versus and the observed value, across the allele frequency (see Figure 4 for examples).

The PPCs indicate that the model performs best for nearly unmixed samples and a subset of highly mixed samples, where fit appears strong. For a majority of samples, the fit is reasonable for a large section of SNPs but poor for a noticeable subset of variants. The PPCs also indicate that a zero-inflation in the data is not fully accounted for within the model. Taken across all samples, this suggest that the $F$-statistic model is insufficient to fully capture the within-sample heterogeneity. However, the model provides strong fit, better even than the COI model above, for a certain subset of samples and SNPs, indicating that a similar admixture process likely contributes to observed data patterns.

## 3 Discussion

This work presents a number of related approaches to inferring inbreeding coefficients, and connects them to an extensive body of research on multiplicity of infection (MOI) in $P$. falciparum suggesting the importance of MOI in characterizing the epidemiology of malaria. We provide the attendant code and workflows in an open-source platform for other researchers to implement these methods.

In developing the model, we make a number of assumptions about the underlying structure of the read count data and the biological mixing process that may affect our inference. For the read count data, we assume that read counts are unbiased and the SNPs are unlinked. While short read data can be biased in several ways, previous research indicates that mixture proportions calculated by ratios of read counts is largely unbiased [20]. However, P. falciparum exhibits significant linkage disequilibrium on scales significantly larger than the average distance between neigboring SNPs in our data. While we do not expect this violation to bias our estimates as this absence of independence likely occurs roughly evenly across the genome.

A more troublesome assumption is embedded in the underlying structure of the $F$-statistic. An $F$-statistic measures the departure of the observed number of heterozygotes relative to those expected under Hardy-Weinberg equilibrium. In the context of mixed $P$. falciparum infections, the equilibrium assumptions - random mating, no selection, large population size, genetic isolation - are likely each violated at some level. For example, the mixture within a sample may be the result of a small number of founding individuals or be strongly selected by the human immune system. Without a more general approach to understanding the mixing process, we cannot anticipate the robustness of our estimates to this sort of misspecification. While looking at the SNP plots (e.g. Figure 1) indicate that the $f_{i}$ values inferred do correspond qualitatively to their apparent degree of mixture, the PPC analysis suggests that the model does not always capture the data's full complexity.

We suspect that as the genomic data enables more elaborate statistical models for mixed infections to develop that these considerations will become increasingly important to the biological community. In our presentation of PPCs, we only discuss model fit in a statistical fashion although there may be biological implications as well. Genes or regions of the genome that are either more or less mixed relative to the levels observed in the remainder of the genome could indicate either positive or negative selective pressure from the host immune system, intraspecific competition, or other processes. Examining the PPCs produced we find no strong indications of these effects. This may be because there is no signal to be discovered, or because the underlying model is too simple to allow these distinctions to emerge or the signature is not apparent without more involved statistical approaches. The COIL approach and other recent work [11, 25] indicate that strain-based mixtures may be a complementary line of inquiry. However, the substantial minority of samples for which the inbreeding model did provide the most powerful explanation, strongly suggests that
considerations of inbreeding or similar processes will have to be included in the next generation of statistical models.

## Appendix: COI inference

Following the approach in [11], we implemented a finite mixture model for the data. As the method described there does not easily accomodate the amount of data in our samples ( $\sim 1,500$ SNPs), we relied on a reduced version of the model presented in [25. As in [11], the model presumes that a finite number of strains $K$ give rise to $2^{K}$ 'bands' of the within-sample allele frequency owing different combinations of the present strains. Following the presentation in [25], for SNP $j$ within sample $i$ showing read counts $\left(r_{i j}, n_{i j}\right)$, the within-sample allele frequency within band $r$ is given by

$$
q_{i j r}=\sum_{k=1}^{K} w_{k} \cdot \mathbf{1}_{\left\{s_{k} \in r\right\}},
$$

where $w_{k}$ is the sample proportion for strain $s_{k}$ and $\mathbf{1}$. is an indicator function. Supposing that read counts are i.i.d. conditional upon their band of origin, this leads to Beta-binomial likelihood given $r$,

$$
\begin{equation*}
\mathbb{P}\left(n_{i j}, r_{i j} \mid r, q_{i j r}, \nu\right)=\binom{n_{i j}+r_{i j}}{n_{i j}} \cdot \frac{\mathrm{~B}\left(n_{i j}+q_{i j r} \cdot \nu, r_{i j}+\left(1-q_{i j r}\right) \cdot \nu\right)}{\mathrm{B}\left(q_{i j r} \cdot \nu,\left(1-q_{i j r}\right) \cdot \nu\right)}, \tag{3}
\end{equation*}
$$

where B is the beta function and $\nu$ is an inverse variance parameter. Assuming no population structure within the local population, we can then write the probability of a SNP being in band $r$ as binomial random variable with $C_{r}$ being the number of non-reference allele states present in band $r$, that is, $\mathbb{P}\left(\operatorname{SNP} j \in\right.$ band $\left.r \mid p_{j}\right)=p_{j}^{C_{r}} \cdot\left(1-p_{j}\right)^{2^{K}-C_{r}}$. By summing over all bans, we get a likelihood independent of $r$,

$$
\mathbb{P}\left(r_{i j}, n_{i j} \mid q_{i j}, p_{j}, \nu, K\right)=\sum_{r=1}^{2^{K}} p_{j}^{C_{r}} \cdot\left(1-p_{j}\right)^{2^{K}-C_{r}} \cdot \mathbb{P}\left(n_{i j}, r_{i j} \mid r, q_{i j r}, \nu\right) .
$$

Assuming independence across SNPs yields a product over $j$ as the full data likelihood. Inference is performed in a Bayesian fashion using standard MCMC approaches, detailed in [25].

## Competing interests

The authors declare that they have no competing interests.

## Author's contributions

JO'B designed and implemented the study and wrote the manuscript. RL provided visualization of the data and model results. LA-E collected the data, contributed to the study design, and edited the manuscript.

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## Tables

| Parameter | Description |
| :--- | :--- |
| $j=1, \cdots, M$ | index over number of SNPs, $M$ |
| $i=1, \cdots, N$ | index over number of samples, $N$ |
| $d_{i j}=\left(r_{i j}, n_{i j}\right)$ | Read count data in sample $i$ at variant $j$ for reference and non-reference counts. |
| $p_{j}$ | population-level non-reference allele frequency for SNP j |
| $\widehat{p}_{j}$ | estimate of non-reference allele frequency for SNP j |
| $\tilde{p}_{j}$ | minor-allele frequency for SNP j |
| $\widehat{\tilde{p}}_{j}$ | estimate of minor allele frequency for SNP j |
| $p_{i j}$ | within-sample non-reference allele frequency for SNP j in sample i |
| $f_{i}$ | Inbreeding coefficient |
| $H_{o}(b, i)$ | Observed heterozygosity for sample $i$ in bin $b$. |
| $H_{e}(b)$ | Expected heterozygosity for bin $b$. |
| $\widehat{f_{i}^{*}}$ | Estimator of $f_{i}$ by method $*$. |
| $\mathbf{f}, \mathbf{p}$ | Vector of $f_{i}$ and $p_{j}$ 's. |
| $\alpha_{i}, \beta_{i}$ | Parameters of beta distribution by tranmission intensity group $i$. |
| $\mathbf{c}$ | Vector of parameters for PTI. |

Table 1: Notation for parameters used throughout the manuscript. Note that additional parameters in the Appendix are not included.

| Initial | Regressed | Direct | Bayesian |
| :---: | :---: | :---: | :---: |
| 1.000 | 0.999 | 0.996 | 0.930 |
| - | 1.000 | 0.998 | 0.930 |
| - | - | 1.000 | 0.929 |
| - | - | - | 1.000 |

Table 2: Correlation coefficient among the four inbreeding estimators across 168 samples.

## Figures



Figure 1: Raw SNP data for four representative samples with initial, regressed, and direct estimates of $f_{i}$ overlaid. Grey dots represent individual SNPs with $x$-axis showing expected heterozygosity under HWE and $y$-axis showing observed heterozygosity.


Figure 2: $95 \%$ credible intervals for $1-f$ over the study interval, colored by tranmission intensity.
Unmixed samples correspond tot $1-f<0.05$ (grey dashed line).


Figure 3: Comparisons of the inferred $f$ values under mild, moderate and high transmission intensity. (Upper left) Frequency of binned $f$ values by tranmission intensity. (Upper right) Boxplot of $f$ values by transmission intensity. (Bottom) $90 \%$ credible interval of posterior density by transmission intensity.


Figure 4: Posterior predictive distributions for two representative samples: $f=0.45$ above; $f=0.54$ below.

## Supplementary Figures



Figure S1: Observed population structure by principal components (upper left, upper right, lower left panels) and neighbor-joining tree.


Figure S2: Log-likelihood for thinned MCMC chain (left) and autocorrelation for same chain (right).

