Pedigree reconstruction from poor quality genotype data

Jinliang Wang

Abstract
Marker genotype data could suffer from a high rate of errors such as false alleles and allelic dropouts (null alleles) in situations such as SNPs from low-coverage next-generation sequencing and microsatellites from noninvasive samples. Use of such data without accounting for mistyping properly could lead to inaccurate or incorrect inferences of family relationships such as parentage and sibship. This study shows that markers with a high error rate are still informative. Simply discarding them could cause a substantial loss of precious information, and is impractical in situations where virtually all markers (e.g. SNPs from low-coverage next-generation sequencing, microsatellites from noninvasive samples) suffer from a similarly high error rate. This study also shows that some previous error models are valid for markers of low error rates, but fail for markers of high error rates. It proposes an improved error model and demonstrates, using simulated and empirical data of a high error rate (say, >0.5), that it leads to more accurate sibship and parentage inferences than previous models. It suggests that, in reality, markers of high error rates should be used rather than discarded in pedigree reconstruction, so long as the error rates can be estimated and used properly in the analyses.

Introduction
Molecular marker data are widely used to reconstruct pedigrees of wild populations in which behaviour data are difficult to obtain (Pemberton 2008). The complete, or most often partial, pedigrees recovered from marker data could then be used to make evolutionary and conservation studies of populations in their natural habitats. Such studies include investigating, for example, the mating system and reproductive success, inbreeding and inbreeding depression, the inheritance of qualitative (e.g. some genetic diseases or abnormalities) and quantitative (e.g. heritability of body weight) traits, and the conservation management of endangered species (e.g. for minimizing loss of genetic diversity). Indeed, molecular pedigrees have contributed tremendously to our understanding and the management of wild species. For example, many socially monogamous bird species are found to have extra-pair paternity (EPP) from genetic marker-based parentage analyses, and the EPP rates range up to 55% across species and between populations within species (Griffith et al. 2002).

Like other types of data, however, molecular marker data are not perfect. In various situations, marker data quality can be rather poor, due to many factors such as genotyping errors (e.g. allelic dropouts), mutations, imperfect markers or genotyping technology (e.g. null or recessive alleles which have no phenotypes) and missing data (Bonin et al. 2004; Pompanon et al. 2005). It has long been recognized that such quality problems have large impacts on genealogical relationship inferences (e.g. Sobel et al. 2002; Douglas et al. 2002), and, if ignored, could cause grossly erroneous inferences. Parentage assignments are particularly vulnerable to genotyping problems, as a parent–offspring pair must share at least one allele identical in state at each locus. A single genotyping error in the pair could cause a genotype mismatch and thus a false parentage exclusion, no matter how many other loci are correctly genotyped and are thus in support of the relationship. Recognizing these, various genotyping error models (e.g. Ott 1993; Ehm et al. 1996; Sobel et al. 2002; Marshall et al. 1998; Wang 2004; Hadfield et al. 2006) have been proposed and integrated into likelihood methods for robust relationship inference. Wang (2004) showed that, with an increasing number of markers suffering from genotyping errors at a low rate, sibship inference quality declines rapidly if the errors are ignored.
but improves rapidly if they are integrated into the likelihood inference framework, respectively.

Previous work focussed on modelling and evaluating the impacts of genotyping errors occurring at a relatively small rate (say, <5%). Such “high” quality data can be obtained from microsatellites genotyped by optimized PCR (polymerase chain reaction) using DNA templates of sufficiently high quality and quantity (e.g. extracted from fresh tissue or blood samples), or from SNPs genotyped by microarrays or from high-coverage next-generation sequencing (NGS) technologies. However, much dirtier data with a high genotyping error rate are now widely generated and used in molecular ecology, evolution, and conservation studies. First, SNPs from low-coverage NGS are now regularly generated for many species. Such data can have an extremely high frequency of false homozygotes produced by random sampling of the two alleles in a heterozygote (Nielsen et al. 2011). A heterozygote (say, cg) would be determined to be a homozygote (cc or gg) with probabilities 75, 54, 36 and 24% when NGS has an average sequencing coverage of 1, 2, 3, and 4 respectively (see below). Second, microsatellites genotyped by PCR from noninvasive samples (e.g. faeces, hair) can also show a high frequency of erroneous genotypes because of multiple causes such as low quantity and quality of DNA, pollution, and the presence of PCR inhibitors (Pompanon et al. 2005). Genotyping errors occurring at a high rate of 37% per locus has been reported (Gagneux et al. 1997) for microsatellites genotyped from noninvasive single shed hair in wild chimpanzees.

There are several outstanding questions to be asked and answered about pedigree reconstruction from marker data with high genotyping error rates. First, are current error models valid in handling data with high error rates? Second, what is the highest error rate that current pedigree reconstruction methods can tolerate and thus a marker with such a high error rate is still informative and useful? Cautious researchers usually discard markers with a moderate error rate (say >7%, Hauser et al. 2011) in pedigree analyses. However, these threshold error rates are more or less arbitrary and could be either too low, wasting valuable marker information, or too high, potentially derailing a pedigree analysis.

This study aims to address these questions. I will show that current error models in marker-based pedigree reconstruction can adequately handle markers with a small to moderate genotyping error rate. However, these models fail for markers with a high error rate and lead to erroneous inferences. I will develop an improved error model that is valid and powerful for markers with very high error rates (say, >50%). The old and new models are then checked and compared in inferring full sibship, half sibship and parentage from simulated SNPs of a high dropout rate (mimicking low-coverage NGS) or a high false allele rate, and from simulated microsatellites of a high false allele rate (mimicking PCR of low-quality noninvasive samples). Their performances are also demonstrated by analysing a real dataset. In conclusion, the study advocates the wide application of the new error model, and encourages using rather than discarding markers of poor genotyping quality in marker-based pedigree analysis and other analyses of individual genotypes.

Methods

Throughout the paper, I will use uppercase letters to denote genotypes or true genealogical relationships, and use lowercase letters to denote phenotypes or inferred genealogical relationships.

Error models

Through different mechanisms, mutations, imperfection of markers or genotyping technology (e.g. recessive or null alleles), and genotyping errors could all cause an observed genotype (phenotype) to differ from its underlying true genotype. They have the same effect on data quality and cause essentially the same consequence in inferences. Therefore these problems are not distinguished and are collectively called typing errors hereafter. The essence of an error model is to consider the uncertainty of a phenotype by calculating and using the probabilities of its different genotypes or true genealogical relationships.

Allelic dropouts

Wang (2004) proposed an error model to handle false homozygotes generated by allelic dropouts of microsatellites during PCR. An allelic dropout occurs when PCR fails to amplify one of an individual’s two homologous genes at a locus, leading to a false homozygote phenotype when the underlying genotype is a heterozygote. The model can also handle false homozygotes due to other causes, such as those produced by null alleles (Wang 2018) or by sampling errors in SNP data from low-coverage NGS.

The error model assumes that each of the two homologous genes in a diploid individual drops out during PCR at the same rate $e_1$, and that double dropouts of both genes do not occur (Wang 2004). For a heterozygote genotype $G = A_1A_2$, the model leads to $\text{Pr}(g(G) = 1 − 2e_1, e_1$ and $e_1$ when the observed phenotype is $g = A_1A_2$, $g = A_1A_1$ and $g = A_2A_2$, respectively, where $e_1 = e_1/(1 + e_1)$. For a homozygote genotype $G = A_iA_i$, the model leads to $\text{Pr}(g(G) = 1$ and 0 when $G$ is observed to be $g = A_iA_i$ and any other
phenotypes, respectively. Null alleles can also be handled by this error model and the parameter $e_1$ is equivalent to null allele frequency (Wang 2018).

**False alleles**

In contrast to allelic dropouts that affect heterozygotes only and cause an apparent homozygote excess, the false allele model applies to any genotypes, homozygotes or heterozygotes. It makes two assumptions. First, the two homologous genes at a locus in a diploid individual are independently and equally likely to be incorrectly observed, with rate $e_2$. Second, when an allele is incorrectly observed, it is observed to be any one of the other alleles at an equal probability of $e_2 = e_2/(k-1)$, where $k$ is the observed number of alleles at a focal locus (Wang 2004, 2018). Thus the probabilities that an allele X is observed as such (x) and as one of the other alleles (y, y ≠ x) at the locus are $Prb(x|X) = 1 - e_2$ and $Prb(y|X) = e_2 = e_2/(k-1)$, respectively. These probabilities sum to 1, $Prb(x|X) + (k-1)Prb(y|X) ≡ 1$, as expected.

The above model proves to be powerful in accounting for mistypings in marker data for sibship and parentage inferences (Wang 2004; Wang and Santure 2009; Hadfield et al. 2006), and in inferring the mistyping rate $e_2$ at a locus given a reconstructed pedigree (Wang 2018). However, this is true only when $e_2$ is not very high (say, <0.3) and $k$ is not small. Otherwise, the model could lead to, counterintuitively, a decreasing uncertainty of a phenotype with an increasing value of $e_2$, and thus suboptimal or incorrect results. To understand this, consider the case of $k = 2$. According to the above error model, the probabilities that an allele X is observed to be itself, x, and the alternative allele, y, are $Prb(x|X) = 1 - e_2$ and $Prb(y|X) = e_2$, respectively. With an increasing value of $e_2 > 0.5$, $Prb(x|X)$ becomes increasingly smaller than $Prb(y|X)$. In the extreme case of $e_2 = 1$, $Prb(x|X) = 0$ and $Prb(y|X) = 1$. This problem as exemplified above yields two counterintuitive results. First, an observation becomes increasingly less uncertain when $e_2$ gets closer to 1. Second, an observed allele is inferred to be more likely to be any one of the other alleles than the observed one, and this probability increases to 1 when $e_2$ increases to 1. Although this problem diminishes with an increasing $k$, it does not disappear completely even when $k$ is large. In the literature, quite a few error models with subtle differences were proposed (e.g. Sancristobal and Chevalet 1997; Sieberts et al. 2001; Sobel et al. 2002), but all suffer from this type of problems.

An improved model for false alleles is to change the second assumption to that, when an allele mutates or is mistyped, it is observed to be any allele, including itself, at an equal probability of $e_2 = e_2/k$. In other words, an observed allele is independent of the underlying true allele when a mistyping occurs. Thus the probabilities that an allele X is observed as itself, x, and as one of the other alleles, y (y ≠ x), at the locus are $Prb(x|X) = 1 - e_2 + e_2$ and $Prb(y|X) = e_2$, respectively, where $e_2 = e_2/k$. These probabilities sum to 1, $Prb(x|X) + (k-1)Prb(y|X) ≡ 1$, as expected. This error model ensures that the probability an allele is correctly observed is always not smaller than that it is incorrectly observed to an alternative allele. With $e_2$ increasing towards the limiting value of 1, the uncertainty of an observation always increases, and $Prb(x|X)$ always decreases but is never smaller than $Prb(y|X)$. For the case of $k = 2$, $Prb(x|X) = 1 - e_2/2$ and $Prb(y|X) = e_2/2$. The minimum value (0.5) of $Prb(x|X)$ is equal to the maximum value (0.5) of $Prb(y|X)$, which occurs at the highest possible value of $e_2 = 1$. This error model was used in evaluating marker information content (Wang 2006), and some variants were also used in detecting mistypings in pedigree genotyping data (e.g. Ehm et al. 1996).

**Error-penetrance function**

Given the two error (dropouts and false alleles) models and error rates ($e_1$, $e_2$), it is straightforward to derive the probability that a genotype is observed to be any phenotype. These error-penetrance functions were shown in Wang (2004) for the above dropout model and the old false allele model. They are good approximations when the rates of both types of errors are small, but become increasingly inappropriate and could cause a loss of relationship inference accuracy when error rates are substantial (as shown below).

For the improved false allele model, these error-penetrance functions are listed in Table 1, derived by assuming that, when both false alleles and dropouts occur to

| Genotype | Phenotype | Penetration function ($Prb(g|G)$) |
|----------|-----------|----------------------------------|
| $G_{st}$  | $g_{st}$  | $(1 - e_2 + e_2^2)/(1 - e_1)$    |
| $G_{st}$  | $g_{st}$  | $(e_1 + (1 - e_1)e_2)(1 - k + e_1 + e_2)/(1 - e_1)$ |
| $G_{st}$  | $g_{su}$  | $e_2(e_2 - e_1e_2 + 2e_1)/(1 - e_1)$  |
| $G_{st}$  | $g_{uv}$  | $2e_2^2(1 - e_1)/(1 + e_1)$    |
| $G_{st}$  | $g_{su}$  | $(1 - e_2 + 2e_2)e_2(1 - e_1)/(1 + e_1)$ |
| $G_{st}$  | $g_{ss}$  | $(1 - e_2 + 2e_2)e_2(1 - e_1)/(1 + e_1)$ |
| $G_{st}$  | $g_{uu}$  | $2e_2(1 - e_2)e_1 + e_2^2$    |
| $G_{st}$  | $g_{uu}$  | $2e_2^2(1 - e_1)/(1 + e_1)$    |

The third column gives the probability of a phenotype (second column, g) given a genotype (first column, G), $Prb(g|G)$. A locus has k codominant alleles. Allele indexes s, t, u, v indicate different alleles. Allelic dropout rate and false allele rate are $e_1$ and $e_2$, respectively, and $e_1 = e_1/(1 + e_1), e_2 = e_2/k$. 

**Table 1 Error-penetrance functions**
the same genotype, they occur in that order. These functions are exact regardless of the values of $\varepsilon_1$ and $\varepsilon_2$, and regardless of the number of alleles, $k$, at a locus. For a locus with $k$ codominant alleles, there are $k(k+1)/2$ possible genotypes and the same number of phenotypes. The probabilities of these phenotypes for any given genotype always sum to 1, which can be confirmed by using the functions listed in Table 1.

### Simulated pedigrees

There are so many possible two-generation pedigrees in reality that even a simulation study can only consider a few, hopefully representative, ones. In general, half sibship (HS) is much more difficult to infer than full sibship (FS) and parent–offspring relationship (PO), because genotypes of HS have more uncertainties in IBD (identical by descent) sharing and HS is difficult to distinguish from unrelated (UR). The focus of this study is to investigate whether the current and the improved error models can handle high mistyping rates, and whether markers of extremely high error rates are still informative for relationship inference and thus should be used rather than discarded or not. Therefore, I choose to analyse two pedigrees with modest difficulty to infer.

The first pedigree contains eight sets of full sibships, with set $i$ ($i = 1$–8) having $2^{3-i}$ identical sibships and each sibship having $2^{i-1}$ offspring coming from the same pair of parents. The sampled pedigree has thus a total number of 1024 offspring distributed in 255 full sibships, many being very small (e.g. 128 sibships with each having just 1 offspring and 64 sibships with each having only 2 offspring) and a few being very large (e.g. the largest sibship has 128 offspring). Many singletons and small sibships mean a high potential of type I errors (i.e. non-sibs being inferred to be sibs), and very large sibships mean also a high chance of type II errors (i.e. sib family being split and sibs being inferred to be non-sibs).

The simulated pedigree has in total 255 sibships, each being produced by two singly mated and unrelated parents. Among the 255 true fathers, 15 are selected at random to include in a candidate father sample, which also contains 85 unrelated males. Thus the probability that a true father is sampled is about 0.12, and this probability is used in paternity analysis. Among the 448 true mothers, 15 are selected at random to include in a candidate mother sample, which also contains 85 unrelated females. Thus the probability that a true mother is sampled is about 0.03, and this probability is used in paternity analysis.

### Simulation of SNPs from NGS

I simulated a number of $L$ variable loci equally spaced in a genomic region of $d$ Morgans in genetic map length. Each locus was assumed to have two alleles, with allele frequencies drawn at random from a uniform distribution in the range (0, 1). Given the simulated pedigree, the genotype of each individual at each locus was simulated following Mendelian inheritance laws. Linkage and recombination was simulated by assuming no crossover interference, using Haldane’s mapping function $r = \frac{1}{4}(1 - e^{-2d})$, where $r$ is the recombination rate. For each simulated individual at each locus, the number of reads, $n$, was drawn from a Poisson distribution with the mean equal to the specified sequencing coverage, $c$. To simulate sequencing errors, the allele of each read was changed to any of the two alleles (i.e. including itself) at an equal rate $\varepsilon_2/2$, where $\varepsilon_2$ is the false allele rate. When $n = 0$, the genotype was determined as missing. When the $n (>0)$ reads at a locus of an individual display one or two allele types, the genotype is determined as a homozygote or heterozygote of the observed alleles, respectively.

In the simulations, I assumed $L = 100, 1000$ and 10,000, $d = 1, 5$, and $10M$, $\varepsilon_2 = 0.5, 1$, and $2\%$, and $c = 1, 2, 3, 4, 5$ and 10. In analysing the data by Colony (Wang 2004; Jones and Wang 2010), the simulated value of $\varepsilon_2$ was used as the false allele rate at a locus, and the dropout rate at a locus is calculated as

$$e_1 = \frac{\sum_{n=1}^{\infty} 0.5^{n-1} e^n \exp(-c)/n!}{1 - \exp(-c)} = \frac{2}{1 + \exp(c/2)}. \tag{1}$$

When $c = 1, 2, 3, 4, 5$ and 10, Eq. (1) gives a dropout rate of $e_1 = 0.75, 0.54, 0.36, 0.24, 0.15$ and 0.01 approximately.

### Simulation of SSRs from noninvasive sampling

I simulated a number of $L$ unlinked microsatellites, each having $k$ alleles in a uniform frequency distribution. Due to factors such as the presence of inhibitors, pollutions, and the low quality and quantity of DNA extracted from non-invasive samples (such as faeces), a high dropout rate ($e_1$) and a high other error rate ($\varepsilon_2$) are possible (Gagneux et al. 1997; Pompanon et al. 2005). Because dropouts were
considered in NGS data, I focused on false alleles by keeping a constant and low dropout rate of $e_2 = 0.01$ and by considering a wide range of $e_2$ values (0.01-0.64) for each locus. The values of $k$ used in simulations were 10 and 2, for microsatellites and SNPs, respectively.

**Simulation of SNPs with high and variable false allele rates**

I simulated data of a number ($L = 100, 500$) of unlinked SNPs with high false allele rates, $e_2$, to compare the accuracies of the old and new false allele models for full-sib, half-sib and parentage inferences. I also simulated unlinked SNPs with variable $e_2$ across loci to investigate the effects of the variation in $e_2$ on relationship inference. In both cases, the allelic dropout rate at each locus was fixed at 0.01. For variable $e_2$, I assumed that $e_2$ followed Beta($\alpha$, $\beta$) distribution, and a value drawn from the distribution was taken as $e_2$ for a locus. The mean and variance of a variable in Beta($\alpha$, $\beta$) distribution are $\mu = \alpha / (\alpha + \beta)$ and $\sigma^2 = \alpha \beta / (\alpha + \beta + 1)$ respectively. For a given mean $\bar{e}_2$ and variance $\sigma^2$ of $e_2$ in Beta($\alpha$, $\beta$), therefore, the parameter values $\alpha$ and $\beta$ can be derived as

$$\alpha = \bar{e}_2^2 (1 - \bar{e}_2) / \sigma^2 - \bar{e}_2^2 \quad \text{and} \quad \beta = (\bar{e}_2 - 1) (\bar{e}_2^2 + \sigma^2 + \bar{e}_2) / \sigma^2,$$

respectively. I used $\bar{e}_2 = 0.2$ and $\sigma^2 = \{0.001, 0.002, 0.004, 0.008, 0.016, 0.032, 0.064\}$ in the simulations.

**An ant dataset**

This dataset was used in studying the mating frequency of an ant species, *Leptothorax acervorum* (Hammond et al. 2001). A total number of 377 ant workers (diploid) were sampled from 10 colonies, with each of 6 colonies contributing 45 workers and the remaining 4 colonies contributing 47, 44, 9, and 7 workers to the sample. Each sampled colony was known to be headed by a single (diploid) queen mated with a single (haploid) male; thus the sampled workers from the same colony and different colonies were full siblings and non-siblings respectively. Each sampled worker was genotyped at up to six microsatellite loci, which had a number of observed alleles varying between 3 and 22.

To investigate the effects of genotyping errors and the ability of different error models in handling the errors, the worker genotype data were modified by introducing each type (allelic dropouts or false alleles) of errors separately and at varying rates. For each type and rate ($e = 0.05, 0.1, 0.15, 0.2, 0.3, 0.4, 0.5, 0.6, 0.7$) of errors, a number of 100 replicate datasets were generated from the original dataset. To simulate dropouts at rate $e$, the genotype at each locus of each individual was examined. If it had a homozygote or missing genotype, then no changes were made. If it had a heterozygous genotype, then a uniformly distributed random number $R$ was generated. If $R \leq e$, a dropout occurred and the genotype was reset to a homozygote of one allele taken at random from the two alleles of the heterozygote. Otherwise, the genotype was not changed. Similarly, the false allele model was applied to each allele in a genotype at each locus of each worker at a rate $e$ (details below). Accuracy of reconstructed pedigree was assessed by comparing the known and estimated pedigrees.

**Data simulation and analyses**

For each parameter combination, a number of 100 replicate datasets were simulated and analysed by the pedigree reconstruction program Colony (Jones and Wang 2010) using either the old or/and the new error models. Note false allele rate is defined and modelled differently in the old and new models. In simulations, false alleles were generated following the new false allele model, and the simulated false allele rate, $e_2$, was used in analysing the data by the new error model. The same data were analysed also by the old error model, with false allele rate $e_2$ being reduced to $e_2' = (1 - 1/k)e_2$ for a locus with $k$ alleles. In presentations of the analysis results, $e_2$ (rather than $e_2'$) was used, bearing in mind that the actual value used by the old error model was $e_2' = (1 - 1/k)e_2$.

**Assessments of inference quality**

The quality of reconstructed pedigrees was assessed by evaluating the frequency that a pairwise relationship was correctly inferred. Among the 1024 offspring in a sample from simulated pedigree one, there are two types of pairwise relationships, full siblings (FS) and non-siblings (NS). The accuracy measures are thus $Prb(fs|FS)$ and $Prb(ns|NS)$, the frequencies that a random FS and non-sib (NS) pair are correctly inferred as such (i.e. fs and ns). Between the offspring and candidate parents, there are two types of pairwise relationships, PO and unrelated (UN). I use the frequencies that a dyadic relationship (PO or UN, in a total of $1024 \times 200 = 204,800$ pairs) is correctly inferred, $Prb(po|PO)$ and $Prb(un|UN)$, to assess parentage assignment accuracy. For simulated pedigree two, there are two types of relationships among offspring, half siblings (HS) and non-siblings (NS), and two types of relationships between offspring and candidate parents, PO and unrelated (UN). Accuracy is thus measured by $Prb(hs|HS)$, $Prb(ns|NS)$, $Prb(polPO)$, and $Prb(unUN)$.

To measure the overall accuracy of the inferred relationship among offspring, I calculate $Prb(OO) = Prb(fsFS) \times Fre(FS) + Prb(nsNS) \times Fre(NS)$, where Fre(FS) and Fre(NS) are the frequencies of FS pairs and NS pairs, respectively, in the offspring sample. Therefore, $Prb(OO)$ is the frequency that the relationship of a pair of individuals drawn at random from the
offspring sample is correctly inferred. For pedigree two, Prb (OO) is calculated similarly, replacing Prb(fslFS) and Fre(FS) by Prb(hsHS) and Fre(HS), respectively. Similarly, the overall accuracy of the inferred relationship between offspring and candidate parent samples is calculated by Prb(PO) = Prb(polPO) + Prb(unlUN) × Fre(UN).

Accuracy is assessed similarly for the real dataset, using Prb(fslFS) and Prb(nslNS), as the sibship among sampled offspring is known.

Results

Comparison of false allele models

When false allele rate (ɛ2) is low (i.e. ɛ2 < 0.1), the new and old error models yield almost identical results, as is expected. This is true for both sibship and parentage assignments, no matter the markers are microsatellites (k = 10) or SNPs (k = 2) and whether the number of markers used in the inference is high (L = 40 and 80 for microsatellites, L = 100 and 400 for SNPs, Fig. 1) or low (e.g. L = 10 for microsatellites or L = 50 for SNPs, data not shown). With an increasing ɛ2 above 0.2, however, the new error model gives increasingly more accurate inferences than the old model (Fig. 1). The contrast between models is especially startling when both ɛ2 and L are large.

Figure 1 also demonstrates that markers of extremely high error rates (ɛ2) are still informative about pedigrees when the proper new error model is applied. At the high false allele rate of ɛ2 = 0.64 for example, the accuracy of sibship and parentage can still reach 0.9 when L = 40 microsatellites are used by the new error model. However, this is not true under the old error model. This means that markers of extremely high error rates are still informative for relatedness under the new error model, but are uninformative under the old error model.

Similar results were obtained in using SNPs of various false allele rates for reconstructing half-sib families (Fig. 2). Again the new and old error models are indistinguishable when ɛ2 < 0.2. With increasing values of ɛ2 > 0.2, the new model becomes more accurate in inferring non-sibs (NS) among offspring and in identifying unrelated (UR) pairs of offspring and candidate parents, while the old model becomes more accurate in inferring half-sibs (HS) among offspring and in inferring parentage (PO). However, because NS and UR relationships are almost always much more frequent than HS and PO relationships in a dataset, the overall accuracy measured by the frequency that the relationship of a random pair of individuals is correctly inferred is higher for the new model than the old model. Basically, the old model results in over-assignments of HS and PO relationships, and infers many false HS and PO relationships.

SNP data of different dropout rates

The accuracy of both sibship and parentage assignments increases with an increasing average coverage, c (Fig. 3). While Prb(fslFS) quickly reaches a plateau (at 0.94) at a low coverage of c = 2, Prb(polPO) increases with c without abating even when c = 10. The low accuracy for both sibship and parentage inferences when c is low is because marker information is rather limited, and is not because the error model is inadequate. With c = 1, the dropout rate, 0.75, is so high that only 25% of heterozygotes are observed as such. At this high error rate, 100 SNPs simply do not have sufficient information for pedigree reconstruction. Indeed, 1000 SNPs distributed evenly on a chromosome segment of 5 Morgans yield almost perfect reconstructed pedigrees, with Prb(fslFS) and Prb(polPO) being 0.999 and 0.936 when c = 1, and 1.0 and 1.0 when c = 2.

Figure 3 also shows that parentage inference is more affected by false homozygotes (due to dropouts or sampling errors in NGS) than sibship inference. With 100 SNPs, sibship inference accuracy becomes almost plateaued when c reaches 2, but parentage inference accuracy is still increasing rapidly even when c reaches 10. Similarly, parentage inference is also more sensitive to false allele rates than sibship inference (Fig. 1).

Number of SNPs with high false allele rates

The new false allele model gives increasingly more accurate inferences of both full sibship and parentage with an increasing number of SNPs of different high false allele rates (Fig. 4). This again proves that markers, even at a high false allele rate of 0.4, are still informative for pedigree reconstruction and more markers lead to more accurate inferences.

Variation of false allele rates among loci

The inferences of FS and PO relationships are affected differently by the variation in ɛ2 among loci (Fig. 5). With an increasing variance of ɛ2 in a beta distribution, Prb(fslFS) is almost constant while Prb(polPO) increases. At a high variance (say, 0.06), many loci have either high ɛ2 values (say, >0.8) or low ɛ2 values (say, <0.05). At a low variance (say, 0.002), all loci have ɛ2 values closely centred around the mean of ɛ2 = 0.2. It seems that markers of low false allele rates are more important in accurately inferring parentage than sibship.

Ant data

Allelic dropouts have little effect on the accuracy of reconstructed sibship (Fig. 6). This is true even when dropout rate is extremely high (ɛ1 = 0.7). However, sibship
inference is affected substantially by false alleles, even when they occur at a moderate rate (say, $\varepsilon_2 = 0.2$).

The results were shown for the modified false allele model, but similar results were obtained from the old model. This is because the two models differ substantially in accuracy only when false allele rate is high and the number of loci is large. This dataset of only six microsatellites with many missing genotypes has simply very little information for pedigree reconstruction when false allele rate is high.

**Discussion**

This study shows that how to properly model genotyping errors becomes important in marker-based pedigree reconstruction when such errors occur at a high rate, and when the number of markers is not small. Some error models in the literature (e.g. Sancristobal and Chevalet 1997; Sieberts et al. 2001; Sobel et al. 2002; Wang 2004) work well when error rate is low, but become increasingly inappropriate and inadequate with an increasing error rate. The fundamental problem of these models is that the uncertainty of a phenotype can counterintuitively decrease when error rate approaches its maximal value of 1. The problem becomes more acute with a decreasing polymorphism (i.e. number of alleles or genotypes) at a locus, and with an increasing number of marker loci. It also shows that a modified error model reduces the problem and leads to more accurate sibship and parentage assignments even when error rates are very high.

Integrating proper error models, the likelihood method implemented in Colony (Wang 2004; Wang and Santure 2009) can yield highly accurate sibship and parentage assignments even when markers of the same high error rates are used, as shown in Figs. 1–4. This is especially true for allelic dropouts, and almost perfect relationship inferences could be obtained with either a moderate number of SNPs from very low-coverage sequencing (i.e. high dropout rate,
variable false allele rate ($\varepsilon_2$, $x$-axis), were used for reconstructing the simulated half-sib pedigree. Each locus was assumed to have alleles in a uniform frequency distribution.

At a high false allele rate (say $\varepsilon_2 = 0.64$), many markers are needed to obtain reasonably good parentage and sibship assignments (Figs. 1, 4). When just a few markers are used, a moderate rate of false alleles can still cause a substantial loss of inference accuracy (Fig. 6). This is because allelic dropouts do not change allele frequencies, but false alleles could homogenize allele frequencies. At a high value of $\varepsilon_2$ at a locus with $k$ alleles, the observed allele frequencies calculated from the sampled genotypes (assuming no genotyping errors) would converge to $1/k$, irrespective of the true underlying allele frequencies. This biased allele frequency estimate calculated and used by Colony would mislead to suboptimal reconstructed pedigrees. Although it

---

**Fig. 2** A comparison of the new and old error models for half sibship and parentage inference accuracy as a function of false allele rate. A number of 500 SNPs, each with a fixed dropout rate of 0.01 and a

**Fig. 3** Sibship and parentage inference accuracy ($\Prb(\text{fs}|FS)$ and $\Prb(\text{po}|PO)$, $y$-axis) as a function of average coverage ($c$, $x$-axis). A number of 100 or 1000 SNPs distributed evenly on a chromosome segment of 5 Morgans in genetic map length were used in reconstructing the simulated full-sib pedigree, with each SNP having two alleles of frequencies drawn from a uniform distribution. The sequencing error rate for each read was 0.01. $\Prb(\text{ns}|NS)$ and $\Prb(\text{un}|UN)$ are always close to 1 and are thus not plotted for clarity.

---

**Fig. 4** Sibship ($\Prb(\text{fs}|FS)$, lower panel) and parentage ($\Prb(\text{po|PO})$, upper panel) inference accuracy as a function of the number of SNPs ($L$, $x$-axis). A variable number of unlinked SNPs, each with two alleles in a uniform frequency distribution, were used in reconstructing the simulated full-sib pedigree. The dropout error rate was fixed at $\varepsilon_2 = 0.01$ while the false allele rates simulated and used were $\varepsilon_2 = 0.1, 0.2, 0.3$, and 0.4 for each locus. $\Prb(\text{ns}|NS)$ and $\Prb(\text{un|UN})$ are always close to 1 and are thus not plotted for clarity.
could iteratively refine allele frequencies by using the reconstructed pedigrees, genotype data, and genotyping error rate data, this ability is rather limited because it relies on sufficient information to reconstruct the pedigree with few errors. When $\varepsilon_2$ is high and the number of markers is low, the reconstructed pedigree is always far from the truth and thus the refined allele frequencies are also far from the true values. As a result, all inferences (including pedigree, allele frequencies, etc.) are grossly erroneous.

An implication for practical marker-based pedigree analysis is that markers with high error rates should still be used rather than discarded, if the error rates can be estimated. As shown in this study, markers with an allelic dropout rate as high as 0.7 or a false allele rate as high as 0.5 are still informative about genealogy and contribute to the reconstruction of pedigrees. This is great news for situations where high-quality genotype data are out of reach (e.g. microsatellite genotyping from poor noninvasive samples), and where cost and other considerations override data quality (e.g. SNPs from low-coverage NGS). This is also good news for the situation where most loci can be genotyped accurately but a few suffer from a high rate of errors. For example, some microsatellites could have a high null allele frequency or a high dropout rate (due to the use of primers developed for closely related species, for example), even when high-quality DNA extracted from fresh tissue or blood samples is used in PCR. In such situations, genotype data of the problematic loci should still be used in pedigree reconstruction; discarding the data is unnecessary and could cause a substantial loss of precious information.

The new false allele model is implemented in the current version (2.0.6.5) of Colony software downloadable from https://www.zsl.org/science/software/colony.

**Compliance with ethical standards**

**Conflict of interest** JW declares that he has no conflict of interest.

**Publisher’s note:** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

**Open Access** This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.
References

Bonin A, Bellemain E, Bronken Eidesen P, Pompanon F, Brochmann C, Taberlet P (2004) How to track and assess genotyping errors in population genetics studies. Mol Ecol 13:3261–3273

Douglas JA, Skol AD, Boehnke M (2002) Probability of detection of genotyping errors and mutations as inheritance inconsistencies in nuclear-family data. Am J Human Genet 70:487–495

Ehm MG, Kimmel M, Cottingham Jr RW (1996) Error detection for genetic data, using likelihood methods. Am J Human Genet 58:225–234

Gagneux P, Woodruff DS, Boesch C (1997) Microsatellite scoring errors associated with noninvasive genotyping based on nuclear DNA amplified from shed hair. Mol Ecol 6:861–868

Griffith SC, Owens IPF, Thuman KA (2002) Extra pair paternity in birds: a review of interspecific variation and adaptive function. Mol Ecol 11:2195–2212

Hadfield JD, Richardson DS, Burke T (2006) Towards unbiased parentage assignment: combining genetic, behavioural and spatial data in a Bayesian framework. Mol Ecol 15:3715–3730

Hammond RL, Bourke AFG, Bruford MW (2001) Mating frequency and mating system of the polygynous ant, Leptothorax acervorum. Mol Ecol 10:2719–2728

Hauser L, Baird M, Hilborn RAY, Seeb LW, Seeb JE (2011) An empirical comparison of SNPs and microsatellites for parentage and kinship assignment in a wild sockeye salmon (Oncorhynchus nerka) population. Mol Ecol Resour 11:150–161

Jones O, Wang J (2010) COLONY: a program for parentage and sibship inference from multilocus genotype data. Mol Ecol Resour 10:551–555

Marshall TC, Slate J, Kruuk LEB, Pemberton JM (1998) Statistical confidence for likelihood-based paternity inference in natural populations. Mol Ecol 7:639–655

Nielsen R, Paul JS, Albrechtsen A, Song YS (2011) Genotype and SNP calling from next-generation sequencing data. Nat Rev Genet 12:443–451

Ott J (1993) Detecting marker inconsistencies in human gene mapping. Human Hered 43:25–30

Pemberton JM (2008) Wild pedigrees: the way forward Proc Roy Soc Lond B: Biol Sci 275:613–621

Pompanon F, Bonin A, Bellemain E, Taberlet P (2005) Genotyping errors: causes, consequences and solutions. Nat Rev Genet 6:847–846

Sancristobal M, Chevalet C (1997) Error tolerant parent identification from a finite set of individuals. Genet Res 70:53–62

Sieberts S, Wijsman EM, Thompson EA (2001) Relationship inference from trios of individuals in the presence of typing error. Am J Human Genet 70:170–180

Sobel E, Papp JC, Lange K (2002) Detection and integration of genotyping errors in statistical genetics. Am J Human Genet 70:496–508

Wang J (2004) Sibship reconstruction from genetic data with typing errors. Genetics 166:1963–1979

Wang J (2006) Informativeness of genetic markers for pairwise relationship and relatedness inference. Theor Popul Biol 70:300–321

Wang J (2018) Estimating genotyping errors from genotype and reconstructed pedigree data. Methods Ecol Evol 9:109–120

Wang J, Santure AW (2009) Parentage and sibship inference from multilocus genotype data under polygamy. Genetics 181:1579–1594