Bioinformatics Advance Access published January 30, 2013

Genetics and population analysis

Bayesian parentage analysis with systematic accountability of genotyping error, missing data, and false matching

Mark R. Christie1*, Jacob A. Tennessen1 and Michael S. Blouin1
1Department of Zoology, Oregon State University, Corvallis, OR, USA

Associate Editor: Dr. Jeffrey Barrett

ABSTRACT

Motivation: The goal of any parentage analysis is to identify as many parent-offspring relationships as possible, while minimizing incorrect assignments. Existing methods can achieve these ends, but require additional information in the form of demographic data, thousands of markers, and/or estimates of genotyping error rates. For many non-model systems, it is simply not practical, cost-effective, or logistically feasible to obtain this information. Here, we develop a Bayesian parentage method that only requires the sampled genotypes in order to account for genotyping error, missing data, and false matches.

Results: Extensive testing with microsatellite and SNP data sets reveals that our Bayesian parentage method reliably controls for the number of false assignments, irrespective of the genotyping error rate. When the number of loci is limiting, our approach maximizes the number of correct assignments by accounting for the frequencies of shared alleles. Comparisons with exclusion and likelihood-based methods on an empirical salmon data set revealed that our Bayesian method had the highest ratio of correct to incorrect assignments.

Availability: Our program SOLOMON is available as an R package from the CRAN website. SOLOMON comes with a fully functional graphical user interface, requiring no user knowledge about the R programming environment. In addition to performing Bayesian parentage analysis, SOLOMON includes Mendelian exclusion and a priori power analysis modules. Further information and user support can be found at https://sites.google.com/site/parentagemethods/.

Contact: christim@science.oregonstate.edu

Supplementary Information: Supplementary data are available at Bioinformatics online.

1 INTRODUCTION

Accurate parentage assignment and pedigree reconstruction are required to make correct inferences for a broad array of study questions (Pemberton, 2008). Parentage methods span a vast gamut of theoretical approaches from fractional to categorical allocation and simple exclusion to sophisticated likelihood-based approaches (Jones and Ardren, 2003; Jones et al., 2010). One area of parentage analysis that has been largely overlooked is a general Bayesian method for categorical allocation. This void is unfortunate as additional sampling or field information can be elegantly incorporated as priors into a Bayesian framework (Hadfield et al., 2006). Furthermore, the information present within the genotypic data itself can be used to calculate a prior analogous to a false discovery rate, which can be useful for the challenges associated with parentage analysis. As an illustrative example, consider a typical kinship data set consisting of 7 microsatellite loci and 750 individuals (Rieseberg et al., 2012). In this data set, a parent and offspring would share at least one allele across all loci following Mendelian inheritance. However, the probability of two unrelated individuals sharing alleles by chance at all loci is not trivial considering that hundreds of thousands of pairwise comparisons are required. Thus, a primary challenge of parentage analysis in natural populations is to correctly identify the true parent-offspring pairs within a data set, while simultaneously excluding any pairs that share alleles by chance.

The challenge of parentage analysis is further exacerbated by missing data and genotyping errors, which can erode the parent-offspring “signal” of sharing at least one allele at all loci (Slater et al., 2000; Vandeputte et al., 2006). Because errors can create an incorrect record of genotypes, true parent-offspring pairs in an empirical data set may not share an allele at all loci despite being the Mendelian expectation. Here, we address the challenges associated with parentage analysis by first calculating the prior probability of a dyad sharing an allele across all numbers of mismatching loci. The calculation of this prior (analogous to a false discovery rate) creates a systematic framework for determining how many loci to let mismatch and does not require any estimates of genotyping error. For each putative pair, we next employ Bayes’ theorem to calculate the posterior probability of a parent-offspring pair being false given the frequencies of shared alleles. Because the probability of sharing common rather than rare alleles is much greater for unrelated pairs, we can compare the frequencies of observed shared alleles to a distribution of alleles shared by unrelated individuals. By combining this information with Bayes’ theorem, we can maximize the identification of true parents and offspring in a data set, while minimizing the number of false assignments. Here, we overhaul the approach of Christie (2010) to (1) account for genotyping error and missing data, (2) reduce the computational time by up to three orders of magnitude as measured in minutes, and (3) allow for one known parent or for known parent-pairs (i.e., known matings), which can substantially increase assignment power. We extensively test this methodology with data drawn from three empirical studies and use an empirical salmon data set to make comparisons to commonly implemented exclusion and likelihood-based methods.

*To whom correspondence should be addressed.
Table 1. Empirical data sets used to validate the Bayesian parentage method. NL refers to the total number of loci used in the study, NA equals the average number of alleles per locus, and Max equals the frequency of the most common allele in the data set. The retriever data set had a total of 21,115 SNPs of which 200 were randomly selected. References are as follows: beech (Lander et al., 2011), steelhead (Araki et al., 2007), and retriever (Akey et al., 2010).

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Species</th>
<th>Marker</th>
<th>NL</th>
<th>NA</th>
<th>Max</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>European Beech (Fagus sylvatica)</td>
<td>μsat</td>
<td>13</td>
<td>11.08</td>
<td>0.66</td>
</tr>
<tr>
<td></td>
<td>Steelhead Trout (Oncorhynchus mykiss)</td>
<td>μsat</td>
<td>8</td>
<td>34.88</td>
<td>0.19</td>
</tr>
<tr>
<td></td>
<td>Labrador Retriever (Canis lupus familiaris)</td>
<td>SNP</td>
<td>21,115 (200)</td>
<td>2.00</td>
<td>0.98</td>
</tr>
</tbody>
</table>

2 METHODS

We created test data sets of multilocus genotypes with allele frequencies based on the site frequency spectra from three empirical studies. We chose empirical studies featuring three distinct taxonomic groups with two different marker types, SNPs and microsatellites (Table 1). The test data sets were fully characterized such that we knew all true parents and offspring. For drawing comparisons between methods, we used complete genotype data from a summer-run steelhead (Oncorhynchus mykiss) data set (see details below).

2.1 Bayesian parentage method

To identify true parent-offspring pairs, we employed Bayes’ theorem to determine the posterior probability of a putative parent-offspring pair being false given the frequencies of shared alleles. For illustrative purposes, we first consider a scenario with no missing data, genotyping error, or known parents, though we expand upon each of these below. In accordance with Mendelian expectation, each parent-offspring pair will share at least one allele across all loci. If a limited number of loci are employed, then pairs of individuals can share alleles by chance alone. In fact, the rate of false matching increases exponentially with a linear increase in sample size (Christie, 2010). We first calculate a prior equal to the probability of any given putative pair sharing alleles by chance:

\[ \Pr(\phi) = \frac{F_{pairs}}{N_{putative}} \]  

(1)

where \( F_{pairs} \) equals the expected number of false parent-offspring pairs and \( N_{putative} \) equals the total number of putative parent-offspring pairs. Here, we define a “false parent-offspring pair” to be a pair of unrelated individuals that share alleles by chance. A “putative parent-offspring pair” is any pair of individuals that share alleles across all loci and contains all true and false parent-offspring pairs. Thus, if a data set was expected to contain 10 pairs that shared alleles by chance, but was observed to contain 100 pairs, then \( \Pr(\phi) \) would equal 0.1. Estimates for \( \Pr(\phi) \) are constrained to range between 0 and 1. To calculate the expected number of false pairs in a data set, we deviate from the approach presented in Christie (2010) and use simulations rather than allele frequencies. We chose to use simulations because they (1) facilitate the incorporation of genotyping error into a Bayesian framework and (2) substantially expedite the calculation of the posterior probability.

To determine the expected number of false pairs we first calculate allele frequencies across all loci. For each locus separately, we calculate genotype frequencies in accordance with Hardy-Weinberg Equilibrium (HWE) and create a pool of genotypes where the rarest genotype occurs at least 100 times. We next create simulated genotypes by sampling from this pool a number of individuals equal to the number genotyped in the empirical data set (randomly assigning individuals as adults and juveniles). We then make all pair-wise comparisons between adults and juveniles and calculate the number of times each allele is shared. If a shared allele is homozygous in an individual, then that allele is only counted once. If an adult and juvenile are heterozygous for the same alleles, then only the rarer of the two alleles is counted. The number of times that an allele is not shared between an adult and juvenile is also recorded. The user may choose how many simulated data sets (hereafter, “simulations”) per locus that they wish to employ, though we recommend a minimum of 100 simulations for SNPs and 1000 simulations for microsatellites to maximize precision for the posterior probability (Table S1). In the simulations, we examine each locus separately in order to expedite the calculation and reduce the amount of memory allocated by R (R Core Team, 2012).

We next create a user-defined number of multilocus “genotypes” by using the output of the simulations. Assuming independence across loci, we sample alleles at each locus by the average frequencies that were observed to be shared between two unrelated individuals. Included in the sampling process is a dummy variable that represents the frequency of dyads that did not share an allele. This process simultaneously creates a distribution of frequencies of alleles shared among false parent-offspring pairs, while also creating a distribution of the number of false pairs that share at least one allele at 0, 1, 2, .., L loci, where L equals the total number of genotyped loci. We calculate the expected number of false pairs as:

\[ F_{pairs} = N_{sim} \cdot n_1 \cdot n_2 \]  

(2)

where \( N_{sim} \) equals the frequency of the simulated multilocus genotypes that shared at least one allele at all loci and \( n_1 \) and \( n_2 \) equal the empirical sample sizes of the adults and juveniles. After \( F_{pairs} \) is calculated, the number of observed putative pairs (\( N_{putative} \)) is calculated using Mendelian incompatibility and used to calculate the prior, \( \Pr(\phi) \).

Most, if not all, observed false pairs will share common alleles, since the probability of sharing an allele by chance is approximately proportional to the square of the allele frequency. In contrast, the probability that a true parent-offspring pair will share a particular allele is simply proportional to the allele frequency. Therefore, pairs sharing rare alleles are much more likely to be true parent-offspring pairs. We exploit this principle by employing Bayes’ theorem to calculate the probability of a putative parent-offspring pair being false given the frequencies of shared alleles:

\[ \Pr(\phi | \lambda) = \frac{\Pr(\lambda | \phi) \cdot \Pr(\phi)}{\Pr(\lambda | \phi) \cdot \Pr(\phi) + \Pr(\lambda | \phi^c) \cdot \Pr(\phi^c)} \]  

(3)

where \( \Pr(\phi) \) is calculated as described above and \( \Pr(\phi^c) \) is the complement. \( \Pr(\lambda | \phi) \) equals the probability of sharing the observed alleles given that the putative pair in question is false. We calculate this value for each putative pair using the multilocus “genotypes” where each locus contains a single value representing the frequency of an allele shared by a false pair. To create a distribution of frequencies of shared alleles among false parent-offspring pairs, we multiply these values across all loci (“false-pair products”). We similarly calculate the product of the shared allele frequencies among all putative parent-offspring pairs (“putative-pair products”). To calculate \( \Pr(\lambda | \phi) \) for each putative pair, we count the number of false-pair products that were less than or equal to the observed putative-pair products and divide by the total. Notice that when a putative pair shares the most common alleles across all loci that \( \Pr(\lambda | \phi) = 1 \), and consequently \( \Pr(\phi | \lambda) = \Pr(\phi) \). To calculate \( \Pr(\lambda | \phi^c) \), which is the probability of sharing alleles given that a putative pair is true, we employed the same approach, but use the observed allele frequencies rather than the frequencies at which alleles were shared.

2.2 Genotyping error

Using the simulations, we calculate \( \Pr(\phi) \) for every number of mismatching loci (0, 1, .., L). When \( \Pr(\phi) \) equals unity, the expected number of false pairs equals the total number of putative pairs within the data set. Mathematically speaking, when the prior \( \Pr(\phi) \) equals 1, the posterior, \( \Pr(\phi | \lambda) \), also
Null alleles can be accounted for by loading in adjusted estimates of allele frequencies from programs that specialize with such data types (e.g., MICROCHECKER, van Oosterhout et al. (2006)). To our knowledge, this is the first parentage method that can account for genotyping errors without needing estimates of the genotyping error rate.

2.3 Microsatellites versus SNPs

Using hundreds of thousands to millions of SNPs can allow for the elucidation of first, second and third order relatives (Manichaikul et al., 2010). Nevertheless, for most species it is not yet cost effective to genotype hundreds or thousands of individuals at so many markers. SOLOMON cannot expediently process millions of SNPs, but rather can accommodate large SNP data sets by performing a priori power analyses to determine a minimum number of SNPs for the given sample sizes to capture all true parent-offspring pairs. After a conservative number of SNPs is determined, the appropriate number of loci can be selected.

The precision associated with the posterior probabilities is increased by increasing the number of simulated data sets and genotypes. Because of the greater number of alleles and lower numbers of loci typically found in microsatellite studies, these markers require more simulations than SNPs for comparable levels of precision (see Table S1 for details and guidelines).

2.4 Validation

We use hypothesis-testing nomenclature to define the null hypothesis as no relationship between a putative parent-offspring pair (i.e., the pair is unrelated). In this framework, a type I error occurs when a putative pair are unrelated, but are falsely identified as a true pair for a given alpha. For example, a type I error would occur if alpha was set to 0.05 and an unrelated adult and juvenile were assigned a Pr(φ) value less than 0.05. Because lower Pr(φ) values represent a reduced probability of sharing alleles by chance, a lower posterior probability represents a reduced probability of committing a type I error. For most methods the type I error should be less than or equal to the chosen alpha, else too many alternative hypotheses will be falsely accepted. A type II error occurs when a true parent-offspring pair are not identified for a given alpha (i.e., Pr(φ | λ) < α for a true parent-offspring relationship). We determined the properties of our method by measuring the type I and type II errors across a range of alpha levels.

To examine the relationship between alpha and type I and II errors, we used the per locus allele frequencies from the empirical studies (Table 1) to construct test data sets. For each of the three empirical studies we created 100 test data sets with 100 adults, 100 juveniles and 50 true parent-offspring pairs. The adult and juvenile genotypes were created in accordance with Hardy-Weinberg Equilibrium (HWE). The parents and offspring were created by randomly selecting 50 adults and 50 juveniles and, for each pair, randomly copying one allele from the adult to the juvenile at each locus. For each of the 100 test data sets, the posterior probabilities were calculated and type I and type II errors were identified. Precision of the posterior probability was calculated by measuring the range of posterior probabilities across identical pairs from 100 replicate runs of a single test data set from each of three study species (Table S1). We also created test data sets with varied numbers of unrelated individuals and offspring per parent (Tables S2 and S3).

We examined the effects of genotyping error by introducing errors into the test data sets. We defined the genotyping error rate as the proportion of all alleles that were called incorrectly (Bonin et al., 2004; Pompanon et al., 2005). To add error to the test data sets, we randomly sampled a single allele position from the multilocus data set. We treated the data set as a matrix with m rows and n columns and randomly selected allele a_{mn}. We next replaced allele a_{mn} with a randomly selected allele from the same locus. This process was repeated until the desired genotyping error rate was obtained. Because alleles were randomly selected, an allele chosen to contain an error could be replaced with the same allele. We chose genotyping error rates of 0, 0.005, 0.01 and 0.03 because they encompass the average documented error rates for SNPs and microsatellites (Pompanon et al., 2005; Saunders et al., 2007).
2.5 Number of known parents

The approach presented above is general in that no information about the sample of adults is required. We expanded the above approach to two specific parentage applications. First, we expanded the method to situations where one parent is known and it is possible to genotype the parent and their offspring. For example, many young mammals remain closely associated with their mothers. After genotyping both the mother and their offspring, it is possible to exclude the maternal alleles from the offspring. This reduces the number of alleles to search for in putative fathers and can greatly increase the power for assignment (Christie et al., 2011; Jamieson and Taylor, 1997). Second, we expanded the approach to include known parent-pairings, where it is known which males mated with which females. For example, captive-breeding and livestock programs often specifically cross certain males to females and keep detailed records of such pairings. Knowing which females and males are paired can substantially increase assignment power because it (1) reduces the number of pair-wise comparisons and (2) each allele in the offspring must match one allele in each parent. To allow researchers to take advantage of the increased power and reduce type I error from such study designs, we appropriately modified the simulation and posterior probability calculation algorithms. We tested these modified approaches with 100 test data sets created from the European beech study because it had the lowest power of the three data sets (and thus the most to gain from additional information). For validation purposes we set the genotyping error rate to 1% and created 100 mothers and 100 fathers, each of which produced a single offspring.

2.6 Siblings and other relatives

Although full-siblings differ from parents and offspring in the way that alleles are shared by descent (Blouin 2003), they can share alleles across large numbers of loci, particularly when including alleles that are shared by chance. This is only a concern if full siblings can occur in both the sampled adults and juveniles (e.g., species with lengthy and overlapping generations), and if they occur at high frequency. To account for full-siblings, we additionally calculate a modified Bayesian prior that includes alleles that are both identical-by-state and identical-by-descent. This modification results in a more conservative test that prevents full-siblings from being assigned as parent-offspring pairs. We tested both the modified and unmodified approach on data sets as described above, but where we introduced pairs of full siblings as 5, 15, 25, and 50 percent of the sampled individuals. Additionally, we tested whether more distant kinship pairs (e.g., aunts/uncles to nieces/nephews, half-siblings) would be falsely identified as parent-offspring pairs.

2.7 Comparison with existing methods

We next analyzed empirical data by examining paternity assignments for four run-years of summer-run steelhead collected from the Hood River, Oregon. This is a new dataset that has not been previously analyzed. Tissue samples from all returning anadromous steelhead were collected as the fish were passed over the Powerdale dam en route to their spawning grounds. The dam was a complete barrier to migrating fish. All 1702 summer-run steelhead were genotyped at the same 8 polymorphic loci used in the winter-run steelhead examples above (Araki et al., 2007). This data set presents a rigorous test for two reasons. First, not all candidate fathers were sampled because resident steelhead (i.e., rainbow trout) that remained above the dam could also have sired offspring (Christie et al., 2011). Second, any given offspring may have aunts and uncles competing for parental assignments (Olsen et al., 2001).

Direct and equitable comparisons between parentage methods can be challenging because each method represents different theoretical approaches. Furthermore, each method often makes different assumptions and requires different input information. We first used Mendelian incompatibility (exclusion) to assign offspring to putative fathers. We allowed one locus to mismatch to account for genotyping error. We next used the most frequently used parentage program, CERVUS 3.03 (Kalinowski et al., 2007; Marshall et al., 1998), to perform the same assignments. CERVUS employs a simulation procedure to determine the significance of likelihood scores for candidate parent-offspring pairs. This program requires the estimates of three parameters: (1) the number of candidate parents, (2) the proportion of candidate parents sampled and (3) the genotyping error rate. Because we did not have estimates of these parameters (they require substantial observational data), we set the number of candidate parents to the number of adults sampled in our data set and chose a small
Fig. 3. Relationship between the number of used SNPs and the percentage of true parent-offspring pairs that were correctly identified in the retriever data sets. Genotyping error rates were varied from 0 to 3%, and all parent-offspring pairs were correctly identified with 250 SNPs. Notice that small amounts of error do not substantially affect the assignment rate with intermediate numbers of loci.

and large proportion of candidate parents sampled (0.1 and 0.9, respectively). We set our genotyping error rate to 1%, which is the default setting, and included assignments with 95% or higher confidence. Lastly, we used SOLOMON to analyze the same sets of samples, using an alpha of 0.05.

To verify our assignments with these three methods, we genotyped all individuals at 5 additional microsatellite loci (see SI for details). To determine which pairs were definitively true, we performed exclusion at all 13 loci and allowed for one locus to mismatch. For matches at both 12 and 13 loci, the average expected number of false pairs was less than one. For all three methods we measured the total number of assignments and the total number of correct assignments as determined by comparison to the pairs identified with the additional loci.

3 RESULTS
3.1 Validation

For all three empirical studies used to generate test data sets, the type I error rate was always equal to or less than the desired alpha (Fig. 2). The beech data sets had the highest type II error rate (lowest power) of the three studies. The steelhead data sets had a lower type II error rate, despite having 5 fewer loci than the beech study. Thus, in these two cases, increased marker polymorphism resulted in greater power for parentage analysis than did additional loci. Lastly, the retriever study with 200 SNPs had the lowest type II error rate (highest power), further confirming that SNPs can be useful markers for parentage analysis (Anderson and Garza, 2006).

The inherent tradeoffs between type I and II errors revealed that there is a marked decrease in type II error (increase in power) by changing the alpha threshold from 0.001 to 0.01. Further increases in alpha from 0.01 to 0.1 yielded marginal increases in power for the steelhead and retriever data sets, but provided consistent increases in power for the beech data set. In general, a good tradeoff between type I and II errors can be obtained by setting alpha at 0.05, but this value should ultimately be decided by weighing the relative risks of committing type I and II errors for a particular study (Sokal and Rohlf, 1994). Not surprisingly, the likelihood of committing type I errors increases with low-power data sets that have high values for the prior. As such, we recommend reporting both the prior and posterior probabilities.

Table 2. Comparison of Exclusion, CERVUS, and SOLOMON on a summer-run steelhead data set. ‘Adults/Juvs’ represents the sample sizes of adults and their putative offspring, respectively. ‘Assigned’ refers to the total number of assignments. ‘Correct’ refers to the number of assignments that were correct after genotyping all putative pairs at 5 additional loci. For CERVUS, we estimated the proportion of candidate parents sampled to be 0.1 or 0.9, though we did not possess demographic estimates of this parameter (results for 0.9 are presented in parentheses).

<table>
<thead>
<tr>
<th>Runyear</th>
<th>Adults/Juvs</th>
<th>Method</th>
<th>Assigned</th>
<th>Correct</th>
</tr>
</thead>
<tbody>
<tr>
<td>2001</td>
<td>201/227</td>
<td>Exclusion</td>
<td>79</td>
<td>38</td>
</tr>
<tr>
<td>2001</td>
<td>201/227</td>
<td>CERVUS</td>
<td>35 (98)</td>
<td>23 (37)</td>
</tr>
<tr>
<td>2001</td>
<td>201/227</td>
<td>SOLOMON</td>
<td>29</td>
<td>27</td>
</tr>
<tr>
<td>2002</td>
<td>343/285</td>
<td>Exclusion</td>
<td>141</td>
<td>90</td>
</tr>
<tr>
<td>2002</td>
<td>343/285</td>
<td>CERVUS</td>
<td>47 (151)</td>
<td>39 (78)</td>
</tr>
<tr>
<td>2002</td>
<td>343/285</td>
<td>SOLOMON</td>
<td>63</td>
<td>61</td>
</tr>
<tr>
<td>2003</td>
<td>144/216</td>
<td>Exclusion</td>
<td>73</td>
<td>49</td>
</tr>
<tr>
<td>2003</td>
<td>144/216</td>
<td>CERVUS</td>
<td>44 (83)</td>
<td>34 (49)</td>
</tr>
<tr>
<td>2003</td>
<td>144/216</td>
<td>SOLOMON</td>
<td>28</td>
<td>28</td>
</tr>
<tr>
<td>2004</td>
<td>90/196</td>
<td>Exclusion</td>
<td>56</td>
<td>36</td>
</tr>
<tr>
<td>2004</td>
<td>90/196</td>
<td>CERVUS</td>
<td>32 (65)</td>
<td>27 (35)</td>
</tr>
<tr>
<td>2004</td>
<td>90/196</td>
<td>SOLOMON</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>All years</td>
<td>778/924</td>
<td>Exclusion</td>
<td>349</td>
<td>213</td>
</tr>
<tr>
<td>All years</td>
<td>778/924</td>
<td>CERVUS</td>
<td>158 (397)</td>
<td>123 (199)</td>
</tr>
<tr>
<td>All years</td>
<td>778/924</td>
<td>SOLOMON</td>
<td>140</td>
<td>136</td>
</tr>
</tbody>
</table>
In all three data sets, genotyping error increased the number of type II errors. Because the retriever data set could allow for the greatest number of mismatching loci (Fig. 1), this data set was the least affected by genotyping error. In general, genotyping error rates of 0.005 or 0.01 did not drastically increase the type II error rate. A genotyping error rate of 3%, however, did result in substantial increases in type II error for all three data sets. We further examined the tradeoff between genotyping error rates and power in the retriever data set. All data sets, regardless of the genotyping error rate, identified all true parent-offspring pairs with 250 loci (Fig. 3). As expected, the number of loci required to identify all true parent-offspring pairs increased with an increase in the genotyping error rate.

Additional samples of a single known-parent or information about putative parent-pairings greatly reduced the type I and II error rates (Fig. 4). Both the type I and type II errors were highest when no known parents were sampled. Having a known sample of one of the parents or knowing the parent-pairs reduced the type II error by nearly 60% for the beech study. Thus, when possible, we recommend collecting this additional data in order to maximize power for parentage analysis.

In general, errors of simulated full siblings that were split between adult and juvenile files did not get assigned in large numbers until they represented more than 25% of the individuals in a data set (Table S4). Adjusting the prior for alleles that were identical-by-state as well as those that were identical-by-descent resulted in fewer sibling pairs with a posterior probability less than 0.05 (Table S5). Accounting for alleles that are identical-by-descent comes at the cost of assigning true parents, however, as it can be difficult to distinguish between full-siblings and parent-offspring pairs with genotyping errors with limited numbers of loci. As such, we recommend using the modified sibling approach only when large numbers of siblings are expected to be sampled. Other levels of relationship, that share fewer alleles than full-sibs (e.g., aunts/uncles to nieces/nephews) were not falsely identified using the unmodified approach.

3.2 Empirical data
Across all four run-years of our summer-run steelhead data set, we found that using simple exclusion for 7 of 8 loci (i.e., allowing one locus to mismatch) resulted in a high type I error rate. Using exclusion, a total of 349 offspring were assigned to a father, of which 213 were later confirmed to be true assignments with genotyping at the 5 additional loci (Table 2). Thus, exclusion produced a total of 136 false assignments yielding a type I error rate of 0.39. CERVUS had type I error rates of 0.22 and 0.49 when we set the estimates of the proportion of candidate parents sampled to 0.1 and 0.9, respectively. In contrast, SOLOMON had a type I error rate of 0.029 for an alpha set to 0.05. Consistent with the results from the test data sets (see Figs. 2.4), varying the alpha in this empirical data set resulted in an observed type I error less than or equal to alpha in all 4 years (Table S6). It is worth noting that in some years CERVUS had a higher number of false assignments than exclusion because the program sometimes allowed for up to two loci to mismatch.

Previous studies have shown that the performance of CERVUS is robust and we suspect that the possible presence of aunts and uncles among the candidate parents coupled with an unknown percentage of sampled parents provided challenging conditions. In general, SOLOMON performed favorably by minimizing the number of false assignments while maximizing the number of correct assignments (Table 2).

4 DISCUSSION
Accurate parentage assignments are necessary in order to appropriately address a wide range of research questions (Jones and Ardren, 2003; Pemberton, 2008). Here, we provide a Bayesian method that can account for genotyping error, missing data, and false matches without requiring estimates of any non-genetic parameters (i.e., all analyses simply use the provided genotypic data). These methods can be applied to a vast array of data sets ranging from samples of large, wild, populations with unknown numbers of sampled parents to carefully controlled crosses with detailed pedigree records. To our knowledge, this is the first parentage program that does not require direct estimates of genotyping error. This solution represents a significant advance because choosing the appropriate method for estimating genotyping error rates can be ambiguous and is further obfuscated by the different types of genotyping errors that can occur (Pompanon et al., 2005). Furthermore, the estimation of error rates typically involves the genotyping of additional (or duplicate) samples, which is costly from both a time and monetary standpoint. Because this method was designed with a null hypothesis of no relationship, it may not be ideally suited for data sets with large numbers of related individuals. Future improvements could include specifying different null hypotheses of relationship and evaluating them in a likelihood-based framework.

Our analyses revealed that, for a given data set, the Bayesian approach approximately minimizes false assignments while maximizing the number of correct assignments. The number of true parent-offspring relationships correctly identified depends upon the sample sizes, the number of loci, the allele frequencies, and the genotyping error rate. For a given marker set, larger sample sizes rapidly increase the number of pairs that share alleles by chance (Christie 2010) and increases in genotyping error can diminish power (Fig. 2, Fig. 3). Furthermore, the number and frequency distribution of alleles at each locus contribute to the rate of false matching. Uniform allele frequencies result in the greatest power for parentage analysis, but are rarely observed in genetic markers. On the other hand, SNPs with a minor allele frequency less than 1% will contribute little information to the elucidation of parent-offspring pairs. Given the multitude of factors that contribute to false matching and reduced power, we suggest that researchers conduct a priori power analyses before designing a study that involves parentage analysis. Such power analyses can dictate precisely how many loci would be required for given sample sizes.

We provide a module for a priori power analysis as part of our program SOLOMON, which is available as a freely distributable R package (R Development Core Team, 2012). SOLOMON is run with a graphical user interface (GUI) written with the TL/TCK package provided by R. SOLOMON performs the described Bayesian parentage analysis for data sets with no known parents, one known parent, or known parent-pairs. Using an Intel core i7™ processor with eight gigabytes of RAM, the average run-time was 11 minutes for the beech data sets, 8 minutes for the steelhead data set, and 13 minutes for the retriever data set (with larger sample sizes resulting in increased run times). Furthermore, the program performs exclusion for the three types of parentage analysis, and the exclusion interfaces allow for user-defined numbers of loci to mismatch. In summary, the Bayesian approach implemented in SOLOMON can be applied to a wide variety of data sets resulting in robust parentage assignment.
ACKNOWLEDGEMENTS

We acknowledge Zaid Abdo, Chris Sullivan, and the Center for Genome Research and Biocomputing at Oregon State University for helpful contributions. We also thank the reviewers for comments that greatly benefited this manuscript. Funding: This research was supported by a grant to M.S. Blouin from the Bonneville Power Administration.

REFERENCES