

# A Population Phylogenetic View of Mitochondrial Heteroplasmy

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**ABSTRACT** The mitochondrion has recently emerged as an active player in myriad cellular processes. Additionally, it was recently shown that >200 diseases are known to be linked to variants in mitochondrial DNA or in nuclear genes interacting with mitochondria. This has reinvigorated interest in its biology and population genetics. Mitochondrial heteroplasmy, or genotypic variation of mitochondria within an individual, is now understood to be common in humans and important in human health. However, it is still not possible to make quantitative predictions about the inheritance of heteroplasmy and its proliferation within the body, partly due to the lack of an appropriate model. Here, we present a population-genetic framework for modeling mitochondrial heteroplasmy as a process that occurs on an ontogenetic phylogeny, with genetic drift and mutation changing heteroplasmy frequencies during the various developmental processes represented in the phylogeny. Using this framework, we develop a Bayesian inference method for inferring rates of mitochondrial genetic drift and mutation at different stages of human life. Applying the method to previously published heteroplasmy frequency data, we demonstrate a severe effective germline bottleneck comprised of the cumulative genetic drift occurring between the divergence of germline and somatic cells in the mother, and the separation of germ layers in the offspring. Additionally, we find that the two somatic tissues we analyze here undergo tissue-specific bottlenecks during embryogenesis, less severe than the effective germline bottleneck, and that these somatic tissues experience little additional genetic drift during adulthood. We conclude with a discussion of possible extensions of the ontogenetic phylogeny framework and its possible applications to other ontogenetic processes in addition to mitochondrial heteroplasmy.

**KEYWORDS** somatic evolution; cell lineage; development; phylogeny

**A**S the energy providers of the cell, mitochondria play a vital role in the biology of eukaryotes. Much of the metabolic functionality of the mitochondrion is encoded in the mitochondrial genome, which in humans is ~ 16.5 kb in length and inherited from the mother. While it was long thought that the mitochondria within the human body are genetic clones, it is now recognized that variation of mitochondrial DNA (mtDNA) is common within human cells and tissues. This variation, termed mitochondrial heteroplasmy, is a normal part of healthy human biology (Li *et al.* 2010, 2016;

Rebolledo-Jaramillo *et al.* 2014), but it is also important in human health and disease, being the primary mode of inheritance of mitochondrial disease and playing a role in cancer and aging (reviewed in Wallace and Chalkia 2013; Stewart and Chinnery 2015).

Because of its importance in human health, it is crucial to understand how mitochondrial heteroplasmy is transmitted between generations and becomes distributed within an individual. Heteroplasmy frequencies can change drastically between mother and offspring, owing to a hypothesized bottleneck in the number of segregating units of mitochondrial genomes during early oogenesis (Cree *et al.* 2008). There has been considerable debate about whether the mechanism of this bottleneck involves an actual decrease in the number of mitochondrial genome copies vs. cosegregation of genetically homogeneous groups of mitochondrial DNA (*e.g.*, Jenuth *et al.* 1996; Cao *et al.* 2007; Cree *et al.* 2008; Wai *et al.* 2008; Carling *et al.* 2011). Nevertheless, in order to better

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predict the change in heteroplasmy frequencies between generations, previous studies have sought to infer the size of the oogenic bottleneck, either through direct observation (in mice) of the number of mitochondrial DNA genome copies (Cao *et al.* 2007; Cree *et al.* 2008), or through indirect measurement, making statistical conclusions about the bottleneck size based on observed frequency changes between generations (Millar *et al.* 2008; Hendy *et al.* 2009; Rebolledo-Jaramillo *et al.* 2014; Johnston *et al.* 2015; Li *et al.* 2016). Recently, Johnston *et al.* (2015) have proposed a statistical framework that combines direct observations of mtDNA copy number with genetic variance in order to make inferences about the dynamics of the oogenic bottleneck. In mice, estimates of the physical bottleneck size have ranged from 200 to >1000 (Cao *et al.* 2007; Cree *et al.* 2008; Johnston *et al.* 2015), and in a recent reanalysis of previous data, it was claimed that the minimal bottleneck size may have only small effects on heteroplasmy transmission dynamics, depending on the details of how oogonia proliferate (Johnston *et al.* 2015). In humans, indirect estimates of the effective genetic bottleneck size have ranged from one to 200, depending on the dataset and the statistical methods used to estimate the bottleneck size (Marchington *et al.* 1997; Guo *et al.* 2013).

Surveys of heteroplasmy occurrence in humans have also found that heteroplasmic variants are often more numerous and at greater frequency in older individuals, and that older mothers transmit more heteroplasms to their offspring (Sondheimer *et al.* 2011; Rebolledo-Jaramillo *et al.* 2014; Li *et al.* 2015). It has also been observed that heteroplasmy frequencies vary from one tissue to another within an individual (Rebolledo-Jaramillo *et al.* 2014; Li *et al.* 2015). These observations underscore the fact that heteroplasmy frequencies change not only during oogenesis in the mother, but also during embryogenesis and throughout adult life. Ideally, any indirect statistical inferences made about the bottleneck size or other aspects of heteroplasmy frequency dynamics would account for all sources of heteroplasmy frequency change simultaneously. Such an approach would need to account for the phylogenetic and developmental relationships between sampled tissues in order to make full use of the information contained in the observed heteroplasmy allele frequencies. While a number of studies have employed or developed population-genetic models to study mitochondrial heteroplasmy (*e.g.*, Wonnapijit *et al.* 2008; Hendy *et al.* 2009; Johnston *et al.* 2015; Johnston and Jones 2016), few have considered the phylogenetic relationship between tissues in doing so, often because only a single tissue type is under consideration. Recently, Burgstaller *et al.* (2014) inferred tissue-specific rates of heteroplasmy segregation in artificially heteroplasmic mouse lines, implicitly relating the sampled tissues by a star-like phylogeny. In a study of heteroplasmy frequencies sampled from 11 tissues in unrelated individuals, Li *et al.* (2015) constructed a phylogeny of the tissues but did not combine it with population-genetic analysis.

Here, we describe a model of heteroplasmy dynamics throughout several key stages of human growth and reproduction. Our approach is to model heteroplasmy frequency change as a population-genetic process of genetic drift and mutation that occurs along the branches of an ontogenetic

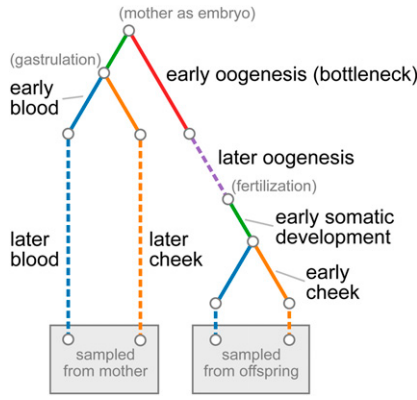
phylogeny, which we define as the tree-like structure relating sampled tissues by their developmental and ontogenetic histories. Our model is similar to typical population-phylogenetic inference models (*e.g.*, Pickrell and Pritchard 2012; Gautier and Vitalis 2013), but it also includes features that are unique to ontogenetic phylogenies. We employ our model in a Bayesian inference procedure that uses Markov chain Monte Carlo (MCMC) to sample from posterior distributions of genetic drift and mutation rate parameters for various developmental processes. After demonstrating the accuracy of our method with simulated data, we apply it to real heteroplasmy frequency data and present new insights into the dynamics of heteroplasmy frequency change in humans.

## Methods

### Ontogenetic phylogenies

We model the mitochondria in tissues sampled from one or more related individuals as a group of populations related by an ontogenetic phylogeny. Along each branch of the ontogenetic phylogeny, heteroplasmy frequencies within some ancestral tissue change due to the action of genetic drift and mutation. We assume that the shape of the ontogenetic phylogeny is given.

Our ontogenetic phylogeny model differs in a few important ways from the typical population-phylogenetic likelihood framework. In the typical population-genetic model, each branch is considered to be an independent period of evolutionary history and thus is under control of an independent parameter. In contrast to this, we allow a single parameter to determine the dynamics on multiple parts of the phylogeny, since a single developmental process can act in multiple related individuals, and this developmental process can be assumed to act similarly in each individual. Furthermore, while it is typically assumed that each locus has been transmitted through a single phylogeny and thus has been subject to the same population-genetic forces, we allow the effects of genetic drift and mutation to depend on the age of the sampled individuals. In particular, for certain ontogenetic processes, we model the rate of accumulation of genetic drift and mutation with age. This is motivated by previous observations that heteroplasmic variants segregate and accumulate with time within somatic tissues (Sondheimer *et al.* 2011; Rebolledo-Jaramillo *et al.* 2014; Li *et al.* 2015) and within the germline (Rebolledo-Jaramillo *et al.* 2014; Li *et al.* 2016; Wachsmuth *et al.* 2016). Finally, in the typical population-phylogenetic model, each branch of the phylogeny represents a single period in evolutionary history and is modeled by a single parameter. Because multiple ontogenetic processes of interest can occur along a single branch of an ontogenetic phylogeny, we allow branches on the ontogenetic phylogeny to be broken into multiple distinct ontogenetic processes, controlled by independent parameters. Figure 1 demonstrates these features with an ontogenetic phylogeny representing the relationships between two tissues sampled in both a mother and her offspring.



**Figure 1** Ontogenetic phylogeny for sampled tissues in mother-child duos from Rebolledo-Jaramillo *et al.* (2014). Each color represents a different tissue or developmental process. The leaves of the tree represent the blood and cheek epithelial tissues sampled from the mother and her child. Solid lines show processes modeled by a fixed amount of genetic drift and dashed lines show processes in which genetic drift accumulates linearly with age. The red component, representing early oogenesis, models a single-generation bottleneck with subsequent doubling of the population size back up to a large size. Parenthetical descriptions in gray show the timing of notable developmental events.

Each ontogenetic process in the phylogeny is parameterized by a genetic drift parameter and a mutation rate. The mutation rate is  $\theta = 2N_e\mu$ , where  $N_e$  is the effective size of the relevant cell population and  $\mu$  is the per-replication, per-base mutation rate. Genetic drift can be modeled in one of three ways, namely as a fixed amount of genetic drift, as an explicit bottleneck size, or as a rate of accumulation of genetic drift per year.

### Likelihood calculation

Given ontogenetic tree  $\mathcal{T}$  with  $k$  ontogenetic processes, genetic drift parameters  $\mathbf{b} = \{b_1, \dots, b_k\}$ , and mutation rates  $\boldsymbol{\theta} = \{\theta_1, \dots, \theta_k\}$ , our likelihood is

$$L_{\mathcal{T}}(\mathbf{b}, \boldsymbol{\theta} | \mathcal{D}) = P_{\mathcal{T}}(\mathcal{D} | \mathbf{b}, \boldsymbol{\theta}), \quad (1)$$

where  $\mathcal{D}$  represents the heteroplasmy frequency data. (Below, the  $\mathcal{T}$  subscript is left off for brevity.) Suppose heteroplasmy frequencies were sampled from  $F$  families. Writing  $C_i$  for the number of heteroplasmy sites in family  $i$ ,  $D_{ij}$  for the heteroplasmy frequency data at the  $j$ th heteroplasmy locus (of  $C_i$ ) in family  $i$ , and  $H_{ij}$  for the event that site  $j$  is heteroplasmy in family  $i$ , our likelihood can be written

$$L(\mathbf{b}, \boldsymbol{\theta} | \mathcal{D}) = P(\mathcal{D}; \mathbf{b}, \boldsymbol{\theta}) = \prod_{i=1}^F P(C_i; \mathbf{b}, \boldsymbol{\theta})^{\frac{1}{C_i}} \prod_{j=1}^{C_i} P(D_{ij} | H_{ij}; \mathbf{b}, \boldsymbol{\theta}), \quad (2)$$

where  $P(C_i; \mathbf{b}, \boldsymbol{\theta})$  is the probability of  $C_i$  heteroplasmy variants occurring in family  $i$  and  $P(D_{ij} | H_{ij}; \mathbf{b}, \boldsymbol{\theta})$  is the probability of the observed heteroplasmy data at the  $j$ th heteroplasmy locus in family  $i$ , conditional on heteroplasmy (*i.e.*, polymorphism) in at least one tissue at that locus. We assume that  $C_i$

is Poisson distributed with rate  $G \cdot P(H_{ij}; \mathbf{b}, \boldsymbol{\theta})$ , where  $G$  is the genome size, and  $P(H_{ij}; \mathbf{b}, \boldsymbol{\theta})$  is the probability that site  $j$  is heteroplasmy in family  $i$ . We note that  $P(H_{ij}) = P(H_{ik})$  for each  $j$  and  $k$ ; that is, the probability of heteroplasmy depends on the family (specifically, on the age of individuals in the family) and not on the particular locus.

We penalize the part of the likelihood involving the number of heteroplasmy variants with the parameter  $\alpha$  in order to make inference less sensitive to experimental heteroplasmy detection, which is a nontrivial problem, especially for heteroplasmies segregating at low frequency (Li and Stoneking 2012; Rebolledo-Jaramillo *et al.* 2014). Without such a penalty, the likelihood is too strongly influenced by the number of observed heteroplasmies, a quantity influenced both by false positives—at a rate of up to  $\sim 10\%$  for low-frequency heteroplasmies in Rebolledo-Jaramillo *et al.* (2014)—and by false negatives caused by conservative minimum allele frequency thresholds (1% in Rebolledo-Jaramillo *et al.* 2014). On the other hand, if the number of heteroplasmies is completely absent from the likelihood, such that all information about drift and mutation is taken only from the heteroplasmy frequencies, posterior distributions of mutation rates are sensitive to outlier allele frequencies that do not fit a model of genetic drift and (infrequent) mutation as well. As a compromise, we set the value of this likelihood penalty to  $\alpha = 100$ , which, in effect, artificially reduces the total number of sites considered in this component of the likelihood, such that if, in reality, 500 heteroplasmy sites are observed out of a total of 100,000 sites, the contribution to the likelihood would be the same as if five heteroplasmy sites were observed in a total of 1000 sites.

With our likelihood (2), we implicitly ignore linkage between heteroplasmy sites within a family, even though, in reality, the lack of recombination means that the sites are perfectly linked. We justify this approximation in two ways: first, there are usually few heteroplasmy variants cosegregating in a family [mean 2.6 in Rebolledo-Jaramillo *et al.* (2014), 1.0 in Li *et al.* (2016)], and, second, among heteroplasmy variants cosegregating in a family, most segregate at low frequency, so that changes in the frequency of one heteroplasmy do not greatly affect the frequency of another. Thus, the dynamics at several heteroplasmy sites should closely resemble those of a model in which each site truly segregates independently. This assumption is supported by simulations of non-recombining mitochondrial genomes (see section *Simulation* below). We further assume that heteroplasmy frequencies are independent between families.

A site is determined to be heteroplasmy according to the filtering steps described in Rebolledo-Jaramillo *et al.* (2014), which include filters for mapping quality, base quality, minimum allele frequency (1%), coverage ( $> 1000\times$ ), local sequence complexity, and contamination. Rather than calculate likelihoods based on called allele frequencies, we model binomial sampling error in the number of consensus and alternative reads sampled from a true, unknown allele frequency. Thus  $D_{ij}$  represents the number of consensus and alternative alleles at the

$j$ th heteroplasmic locus in family  $i$ . Conditional on heteroplasmy (*i.e.*, polymorphism), the probability of the observed read counts  $D_{ij}$  at locus  $j$  in family  $i$  is

$$P(D_{ij}|H_i; \mathbf{b}, \boldsymbol{\theta}) = \frac{\sum_{x_{ij}} P(D_{ij}|x_{ij})P(x_{ij}; \mathbf{b}, \boldsymbol{\theta})}{P(H_i; \mathbf{b}, \boldsymbol{\theta})}, \quad (3)$$

where  $x_{ij}$  is the true, unknown, allele frequency at locus  $j$  in family  $i$ . The sum is performed over all possible allele frequencies in the sampled tissues. Both the numerator and the denominator can be calculated using Felsenstein's (1981) pruning algorithm—a dynamic programming algorithm frequently used in likelihood calculations for phylogenetic trees. Details of how we calculated these quantities are given in Appendix A. The pruning algorithm requires calculating allele frequency transition distributions for different genetic drift and mutation parameter values. As described in Appendix B, we achieved this by numerically precomputing allele frequency transition distributions under the discrete-generation Wright-Fisher model, and linearly interpolating between precomputed values. The pruning algorithm also requires a distribution of allele frequencies at the root of the phylogeny, which, in our application (see below), represents the unobservable distribution of heteroplasmy allele frequencies in the mother as an embryo. Following Tataru *et al.* (2015), we use a discretized, symmetric beta distribution with additional, symmetric probability weights at frequencies 0 and 1. The two parameters specifying this distribution are inferred jointly with genetic drift and mutation parameters.

### Inference

We take a Bayesian approach to inference. Prior distributions are Log-Uniform ( $5 \times 10^{-4}$ , 3) for genetic drift parameters, measured in generations per  $N_e$  (henceforth “drift units”). For genetic drift parameters specified by a rate of accumulation of drift units per year, the lower (*resp.* upper) limit of the (Uniform) prior distribution limits are divided by the minimum (*resp.* maximum) of the ages by which the rate is multiplied. We did not allow the effects of genetic drift to decrease with age. Prior distributions on bottleneck sizes are Log-Uniform(2500), and, for mutation rate parameters  $\theta = 2N_e\mu$ , the prior distribution is Log-Uniform ( $10^{-8}$ ,  $10^{-1}$ ).

We employ an affine-invariant ensemble MCMC procedure (Goodman and Weare 2010) to sample from posterior distributions, as implemented in the Python package *emcee* (Foreman-Mackey *et al.* 2013). We assess convergence by visual inspection of the posterior traces. Running 500 chains in the ensemble MCMC for 20,000 iterations each, we find good convergence after  $\sim 2500$  iterations, and, thus, discard the first 5000 iterations of each chain as burn-in. With  $\sim 100$  heteroplasmic loci, a run takes 60–80 CPU hours, but, due to the parallel nature of ensemble MCMC, calculations can be efficiently spread across CPUs, so that, on a 20-core compute node, results are obtained in  $\sim 4$  hr. Reported 95% credible intervals (CIs) are intervals of the highest posterior density.

As a way of evaluating the relative support for different ontogenetic models, we estimate Bayes factors (*i.e.*, ratios of posterior evidence integrals) for alternative ontogenetic models of the accumulation of drift within cell lineages. For models  $M_1$  and  $M_2$ , the Bayes factor is

$$BF(M_1, M_2) = \frac{\int p(\theta)L(\theta|\mathcal{D}, M_1)d\theta}{\int p(\theta)L(\theta|\mathcal{D}, M_2)d\theta}, \quad (4)$$

where  $p(\cdot)$  is the prior distribution and  $L(\cdot|\mathcal{D}, M_k)$  is the likelihood under model  $k$ . These posterior evidence integrals are approximated using *emcee*'s (Foreman-Mackey *et al.* 2013) implementation of an approach using thermodynamic integration (see Goggans and Chi 2004).

### Simulation

We performed two sets of simulations to test our inference procedure. The first simulations were performed under the model assumed by our inference procedure. As described above, this model assumes that each locus segregates independently, allele frequency transitions occur according to the Wright-Fisher model of genetic drift and biallelic mutation, and heteroplasmy frequencies in the root of the ontogenetic phylogeny are controlled by the two parameters of a discretized, symmetric beta distribution with extra probability weight at frequencies zero and one. These simulations were performed forward in time using a custom Python script.

The second set of simulations tested how our assumption that loci segregate independently affects inference when the data are simulated from nonrecombining genomes sampled from many different families. These simulations were performed using a custom interface to the simulation package *msprime* (Kelleher *et al.* 2016), which simulates genetic variation under the standard neutral coalescent model with infinite-sites mutation. In these simulations, population sizes and branch lengths are equivalent to those under the forward-time simulations, but at the root of the ontogenetic phylogeny, we assume that ancestral lineages trace their ancestry back in time in a single panmictic population of constant size. Simulations were performed under conditions in which the distribution of the number of heteroplasmic variants per family roughly matched the distribution observed in the data.

### Data

We applied our inference procedure to a publicly available dataset, containing allele frequencies for 98 heteroplasms sampled from 39 mother-offspring duos, originally published by Rebolledo-Jaramillo *et al.* (2014). In this dataset, mitochondria from blood and cheek epithelial cells were sampled from both mother and offspring, resulting in an ontogenetic phylogeny with four leaves, each representing one of the four tissues sampled from a mother-offspring duo. Details of heteroplasmy discovery are described in Rebolledo-Jaramillo *et al.* (2014).

To model the segregation of heteroplasmy frequencies during the ontogeny of the four tissues sampled from each duo, we used the ontogenetic phylogeny shown in Figure 1.

This ontogenetic phylogeny models several life stages. The root of the phylogeny occurs at the divergence of the mother's somatic and germline tissues when she is an embryo. On the branch leading to the somatic tissues in the mother, there is a brief period of early embryonic development before the blood and cheek epithelial cell lineages diverge at gastrulation as members of the ectodermal (cheek epithelial) and mesodermal (blood) germ layers. After diverging at gastrulation, each somatic tissue undergoes independent periods of genetic drift and mutation during later embryogenesis and early growth, and, finally, for each tissue there are independent rates of accumulation of genetic drift and mutation throughout adult life.

On the branch leading to the offspring tissues in the ontogenetic phylogeny in Figure 1, the first stage represented is the period of oogenesis prior to the birth of the mother, when the oogenic bottleneck is thought to occur. This is followed by the oocyte stage, during which we assume the mitochondria accumulate genetic drift and mutation at some rate linearly with the age of the mother before childbirth. At fertilization, this branch undergoes the same period of early somatic development experienced by the mother's somatic tissues prior to gastrulation. Finally, the two somatic tissues of the offspring diverge at gastrulation and go through the same stages of development as the somatic tissues of the mother. For an overview of the events of human development, see, for example, Carlson (2014).

### Effective oogenic bottleneck

Analyzing both simulated and real data, we find that there is limited power to infer the size of the oogenic bottleneck. This is to be expected, given that we also model the subsequent genetic drift of the later stages of oocyte development and in the early developing embryo; each of these three ontogenetic processes occurs along the same branch of the ontogenetic phylogeny of the tissues considered here (Figure 1), which causes their respective contributions of genetic drift to be conflated with one another. We note that the genetic drift parameters of these ontogenetic processes are not truly unidentifiable: power to distinguish genetic drift during the early-oogenesis bottleneck from that of the later maternal germline is provided by the differing effects of genetic drift in mothers of different ages, and power to distinguish the contribution of drift in the early embryo is provided by the fact that this process occurs in both the mother and the offspring. Differences in effective population size (and thus scaled mutation rates) also provide theoretical power to distinguish these parameters, but, nevertheless, we find that these genetic drift parameters tend to become conflated with one another.

As a way of counteracting this conflation, we combine the genetic drift parameters of this branch in the ontogenetic phylogeny into an effective bottleneck size (EBS), summarizing the total genetic drift between mother and offspring. The effective bottleneck is comprised of the oogenic bottleneck *per se*, the accumulation of genetic drift in the oocyte prior to ovulation, and the genetic drift in the embryo between fertilization and gastrulation.

To combine genetic drift parameterized as a bottleneck with genetic drift parameterized in drift units, we used the approximate relationship  $N_b \approx 2/d$ , where  $d$  is genetic drift in drift units, and  $N_b$  is the bottleneck size. This approximation is justified in Appendix C. Using this relationship, our equation for the EBS has the form

$$N_{be} = \frac{2}{d + \lambda a}, \quad (5)$$

where  $d$  is the summed genetic drift from the oogenic bottleneck *per se* and pregastrulation embryogenesis,  $\lambda$  is the rate of genetic drift accumulation in the oocyte, and  $a$  is the age of the mother at childbirth. Because, in our model, genetic drift accumulates in the oocyte as the mother ages prior to ovulation, the size of the effective bottleneck decreases with age. We summarize this rate of decrease by linearizing (5) between ages 25 and 34, the first and third quartiles of maternal age at childbirth in the dataset from Rebolledo-Jaramillo *et al.* (2014).

### Data availability

Our inference procedure is released under a permissive license in a Python package called *mope*, available at <https://github.com/ammodramus/mope> or from the Python Package Index (PyPI, <http://pypi.python.org/>). As we describe above, our inference procedure requires precomputed transition distributions. These can be generated by the user or downloaded from <https://github.com/ammodramus/mope>. Our simulation scripts are also provided with the inference procedure.

Data from Rebolledo-Jaramillo *et al.* (2014) are available from that paper's supplemental material and from the NCBI Sequence Read Archive ([www.ncbi.nlm.nih.gov/sra](http://www.ncbi.nlm.nih.gov/sra)), accession SRP047378.

## Results

### Application to simulated data

The targets of our inference procedure are genetic drift parameters and population-size-scaled mutation rates for each ontogenetic process in the ontogenetic phylogeny. Genetic drift may be parameterized as a fixed amount of genetic drift (in drift units, *i.e.*, generations/ $N_e$ ), as a rate of accumulation of drift per year, or as a haploid bottleneck size. The scaled mutation rates,  $\theta = 2N_e\mu$  are twice the product of the haploid effective population size  $N_e$  and the per-replication, per-base mutation rate  $\mu$ . Since  $\mu$  can be assumed to be the same in every mitochondrion, the mutation rates can also be interpreted as relative effective population sizes. Two parameters controlling the distribution of allele frequencies at the root of the phylogeny are also inferred.

The inference procedure performed well on data simulated under the model of drift and mutation assumed by the inference procedure. In a simulation of 500 independently segregating sites sampled from two tissues in each of 100 different mothers and their offspring, under parameters producing a total of

110 heteroplasmic variants, the branch lengths and mutation rates were inferred without apparent bias (Figure 2), as were the two root distribution parameters (data not shown). Posterior distributions were generally narrower for genetic drift parameters than for scaled mutation rates, likely corresponding to the fact that there is less information about mutation than genetic drift in the simulated data. Parameters of external branches were inferred more precisely than those of internal branches. Other parameter values produced similar results (Supplemental Material, Figure S1 in File S1).

The procedure also performed well on data generated in simulations that did not assume free recombination between heteroplasmic sites (Figure S2 in File S1). In these simulations, we simulated nonrecombining mitochondrial genomes of 10,000 bp in 30 mother-offspring duos, under parameters resulting in 104 heteroplasmic variants. The  $\sim 3.7$  heteroplasmies per family in these simulations is similar to the  $\sim 2.6$  observed in the data from Rebolledo-Jaramillo *et al.* (2014), supporting our assumption that linkage between heteroplasmic variants within families does not greatly affect inference results. We also tested for robustness against false positive and false negative heteroplasmy detection. Applying our method to simulations with a false negative rate of 0.4 for truly heteroplasmic mutations between frequencies 0.1 and 2%, and a false positive rate of  $3 \times 10^{-5}$  per basepair, so that 14 false negatives and five false positives were produced in a dataset of 101 heteroplasmic loci, the inference procedure was not apparently biased away from the true, simulated parameter values (Figure S3 in File S1).

### Application to real heteroplasmy data

In the application of our method to the heteroplasmy frequency data from Rebolledo-Jaramillo *et al.* (2014) (Figure 3), we find that the posterior distribution of the size of the early oogenesis bottleneck is broad, with a 95% CI spanning from 10.6 to 433.2. As we describe above (see *Effective oogenic bottleneck*), this is unsurprising given that in the assumed ontogenetic phylogeny there are three independent periods of drift and mutation along the branch containing the oogenic bottleneck, namely the early oogenic bottleneck itself, the turnover of mitochondria in the oocyte prior to ovulation, and the period after fertilization but before gastrulation (Figure 1).

To counteract this conflation, we combined the genetic drift into an effective bottleneck. The posterior distribution of the size of this effective bottleneck (*i.e.*, the EBS) was substantially narrower than that of the explicitly modeled bottleneck, with a median of 24.5 (11.6–35.1, 95% CI) for a mother of the mean age in this dataset (Figure 4A). This is in line with the bottleneck size estimate of 32.3 produced by Rebolledo-Jaramillo *et al.* (2014).

In our model, genetic drift accumulates in the oocyte as the mother ages, and thus the size of the effective bottleneck decreases with age of the mother at childbirth. The inferred relationship between age at childbirth and EBS is shown in Figure 4B. At age 18, the median posterior EBS is 26.1 (13.0–36.9, 95% CI), and at

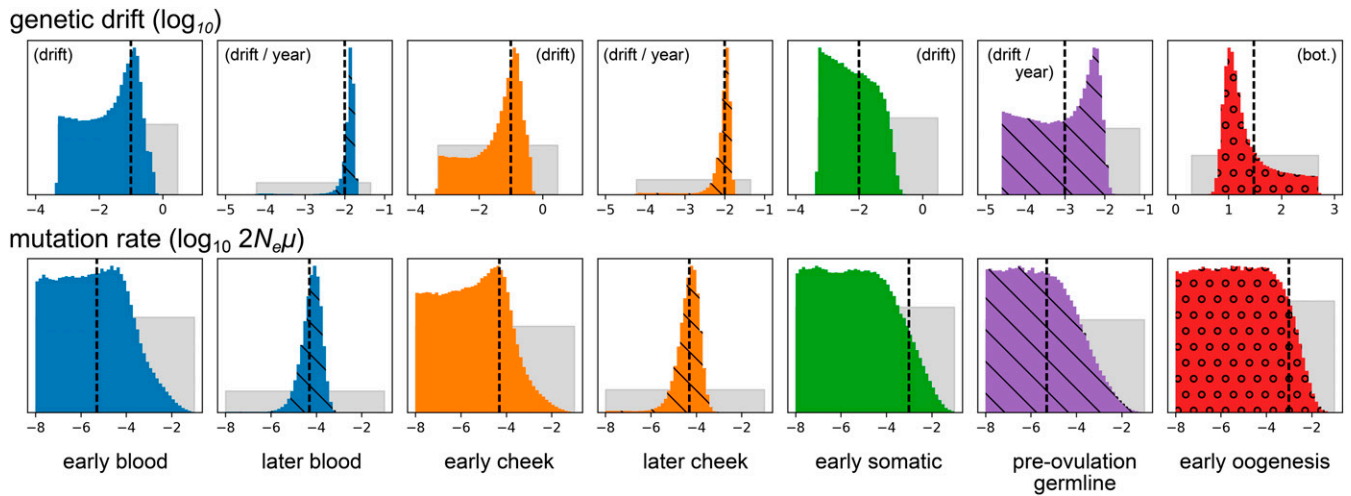
age 40, it is 23.4 (10.5–34.2). The median posterior rate of decrease of the EBS is  $-0.075$  bottleneck units per year, although the central 95% CI for this rate of decrease is broad (0.0–0.34). Given the range of this CI, there is apparently limited information contained in the data about whether or not the EBS decreases with age, or, equivalently, whether genetic drift accumulates meaningfully in the oocyte.

The median posterior rates of genetic drift accumulation in adult somatic tissues were very small, just  $1.0 \times 10^{-3}$  ( $1.0 \times 10^{-3}$ – $1.2 \times 10^{-3}$ , 95% CI) drift units per year for blood, and  $1.0 \times 10^{-3}$  ( $1.0 \times 10^{-3}$ – $1.2 \times 10^{-3}$ ) drift units per year for cheek. These estimates are at the lower limit of what is permissible under our model of genetic drift, which is based upon distributions of allele frequency change in a finite-sized Wright-Fisher model (see Appendix B). On the other hand, the inferred amounts of genetic drift occurring during early development of the somatic tissues was greater: 0.015 (0.0067–0.023, 95% CI) drift units for blood, and 0.0044 ( $5.0 \times 10^{-4}$ –0.011) drift units for cheek, roughly equivalent to bottlenecks of size 136.2 (74.1–247.5) and 457.7 (103.9–2817.1), respectively.

The posterior distributions of scaled mutation rates were broad, and, thus, limited information about the relative population sizes of different developmental and adult tissues is contained in the heteroplasmy frequency data. This is unsurprising given that the problem is similar to attempting to infer population size history from  $\sim 100$  single-nucleotide polymorphisms (SNPs). A high scaled mutation rate ( $2N_e\mu > 10^{-4}$ ) is (relatively) most supported in oogenesis, reflecting the observation of possibly *de novo* mutations in the dataset. However, the 95% CI of each developmental process spans several orders of magnitude (at least  $10^{-8} < 2N_e\mu < 10^{-5}$ ), so firm conclusions cannot be drawn.

We assessed the fit of our model to the real heteroplasmy data by simulating data under the maximum *a posteriori* (MAP) parameter values and comparing to the real data. Comparing the marginal distribution of allele frequencies in the sampled tissues (*i.e.*, the marginal site-frequency spectrum) from the actual data to the MAP simulation data, we find that the marginal distribution of allele frequencies is similar between the two datasets (Figure 5A), as is the distribution of absolute differences between each pair of sampled tissues (Figure 5B).

In order to use Bayes factors (4) to compare the support for different ontogenetic phylogenies, we calculated the posterior evidence integral for the ontogenetic phylogeny in Figure 1, as well as for two additional ontogenetic phylogenies differing in their assumptions about how genetic drift accumulates in somatic tissues (Figure S4 in File S1). The first additional model (termed “fixed,” Figure S4A in File S1), assumes that all genetic drift and mutation particular to each somatic tissue occurs early during development, and that there is no additional drift accumulating later in life. The second, (“linear,” Figure S4B in File S1), assumes that genetic drift and mutation accumulate linearly with age in somatic tissues. Our original model (Figure 1) we term “both,” since it assumes that



**Figure 2** Posterior distributions of genetic drift and mutation parameters inferred from data simulated under the model assumed by the inference procedure. The top and bottom rows depict genetic drift and mutation rate parameters, respectively. Gray distributions depict prior distributions, and colored distributions depict posterior distributions. Colors match the colors of the ontogenetic processes in Figure 1. Distributions hashed with diagonal lines correspond to processes with drift parameterized by rates of accumulation of genetic drift with age. (That is, they correspond to the dashed lines in Figure 1.) The circles in the red posterior distributions indicate that this process is modeled by an explicit bottleneck. All parameters are  $\log_{10}$ -transformed, and the distributions correspond to these transformed variables. Vertical dashed lines show the simulated parameter values. Not shown are the two parameters controlling the allele frequency distribution at the root of the phylogeny, which were inferred with comparable accuracy.

genetic drift both occurs in a fixed quantity during early development and accumulates later in life.

We find that the “fixed” model is more supported than the “both” or “linear” models, with the approximate log-evidence values of the “fixed,” “both,” and “linear” models being  $-1704 \pm 3$ ,  $-1764 \pm 4$ , and  $-1816 \pm 3$ , respectively. In the “both” model, in which there is both a period of genetic drift and mutation in the somatic tissues during early development, the inferred rates of drift accumulation are at the minimum allowed by the inference method ( $\sim 10^{-3}$  drift units per year). This, together with the fact that the best-supported model does not include the accumulation of genetic drift in adult somatic tissues, suggests that there is very little additional genetic drift occurring after birth in the two somatic tissues considered here.

## Discussion

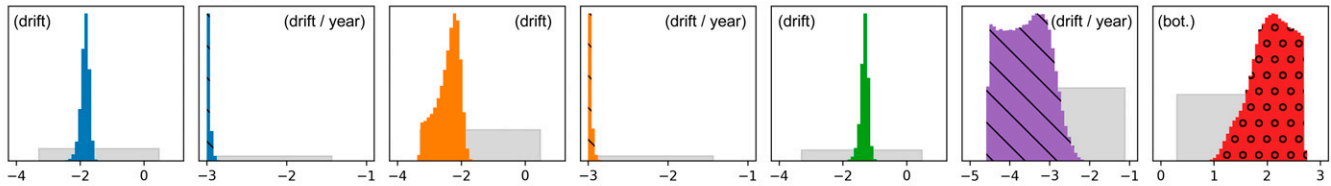
Because we modeled genetic drift during multiple ontogenetic processes between embryogenesis in the mother and the sampling of tissues in the child, our estimate of the size of the oogenic bottleneck *per se* was imprecise, with a broad 95% CI (10.6–433.1). This is concordant with a recent analysis of the time-evolution of heteroplasmy variance in mouse oocytes, which concluded that the actual minimal bottleneck size is difficult to determine, and may have only limited impact on overall heteroplasmy dynamics during oogenesis (Johnston *et al.* 2015). However, our estimates of the EBS (median 24.5, 95% CI: 11.6–35.1) are similar to other recent estimates of the oogenic bottleneck size, including an estimate of 32.3 in a previous analysis of the data used in this study (Rebolledo-Jaramillo *et al.* 2014), and a previous estimate of nine in Li *et al.* (2016).

Our inference framework allows for the size of the effective oogenic bottleneck to decrease with the age of the mother as genetic drift accumulates in the oocyte. We found a broad posterior distribution of the rate by which the EBS decreases in the oocyte (roughly 0.00–0.34 bottleneck units per year, 95% CI), demonstrating that, with the 39 mother-child pairs and 98 heteroplasmic variants in the dataset we analyzed (Rebolledo-Jaramillo *et al.* 2014), there is insufficient information obtained by our model to determine whether genetic drift accumulates with age in the oocyte. In the future, sampling more individuals and tissues, and with larger pedigrees, it may be possible to provide stronger statistical evidence for or against genetic drift occurring in the oocyte; this will potentially be informative on the question of how mitophagy and mitochondrial turnover are involved in oocyte aging—a topic of interest in the study of human fertility (see Zhang *et al.* 2017).

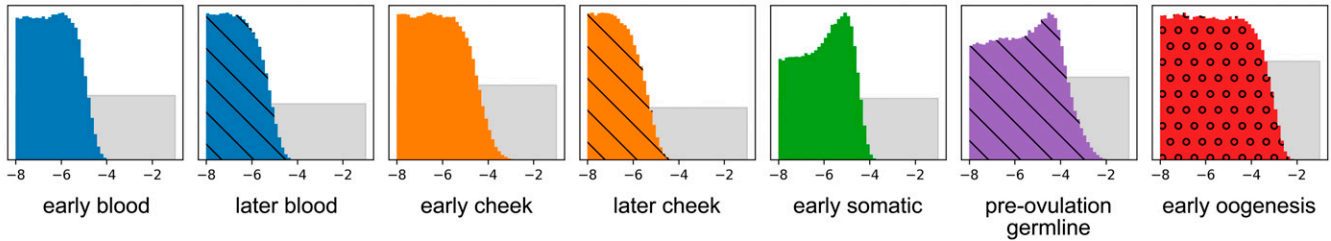
In addition to the effective bottleneck between mother and offspring, we also quantified genetic drift occurring during the embryonic development of the blood and cheek epithelial lineages. We found that the embryonic genetic drift of heteroplasmy frequencies specific to these tissues was less than the effective between-generation bottleneck but still appreciable, with median posterior estimates of the EBSs being 136.2 (74.1–247.5, 95% CI) and 457.7 (103.9–2817.1) for blood and cheek epithelial cells, respectively.

At the same time, we inferred that there is little accumulation of genetic drift in adult somatic tissues. This may seem to contradict previous observations that heteroplasmy becomes more numerous with age (*e.g.*, Rebolledo-Jaramillo *et al.* 2014; Li *et al.* 2016). If the effective population size of the somatic stem cells supporting mitotic somatic tissues

## genetic drift ( $\log_{10}$ )



## mutation rate ( $\log_{10} 2N_e\mu$ )



**Figure 3** Inference results for real heteroplasmy frequency data. The top row shows results for genetic drift parameters, and the bottom row shows posterior distributions for scaled mutation rates. Distributions hashed with diagonal lines correspond to processes with drift parameterized by rates of accumulation of genetic drift with age. (That is, they correspond to the dashed lines in Figure 1.) The circles in the red posterior distributions indicate that this process is modeled by an explicit bottleneck. All parameters are  $\log_{10}$ -transformed, and the depicted distributions correspond to these transformed variables. Distributions are not drawn to a common vertical axis.

is larger than the effective population size during embryogenesis or the maternal germ line, an accumulation of genetic drift with age would produce additional *de novo* somatic heteroplasms. On the other hand, if effective population sizes of somatic stem cells are smaller than effective population sizes during early development, a longer period of genetic drift in adulthood would result in fewer heteroplasmic loci, as genetic variation is lost due to ongoing genetic drift in a smaller population. Here, the posterior distributions of population-scaled mutation rates are too broad to permit anything to be concluded about the relative sizes of relevant stem cell populations.

There are several ways our inference procedure could be extended. Our model assumes selective neutrality, but it is possible, or even likely, that neutral population-genetic models do not completely describe the dynamics of heteroplasmy frequency change. Studies of heteroplasmy occurrence in humans have found a relative lack of nonsynonymous heteroplasmic mutations (Rebolledo-Jaramillo *et al.* 2014; Ye *et al.* 2014), or an excess of nonsynonymous mutations at low vs. high frequencies (Li *et al.* 2016), suggesting purifying selection. However, evidence for biased transmission of the major heteroplasmic allele over the minor allele has been inconsistent, with one recent study finding no systematic difference in heteroplasmy allele frequency between other offspring (Li *et al.* 2016), while the original publication of the data analyzed here did find transmission to be biased toward the major allele at nonsynonymous sites (Rebolledo-Jaramillo *et al.* 2014). Other studies have also found evidence for positive selection acting on heteroplasmies in somatic tissues, observing repeated occurrence of tissue-specific and allele-specific heteroplasmies in many unrelated individuals (Samuels *et al.* 2013; Li *et al.* 2015). Studies in mice have also indicated that hetero-

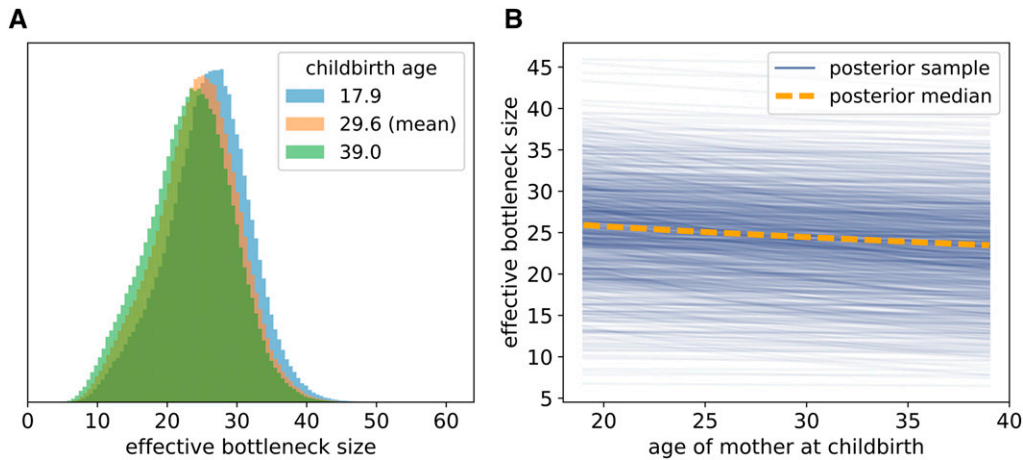
plasmies may be under natural selection in many instances (*e.g.*, Fan *et al.* 2008; Stewart *et al.* 2008; Sharpley *et al.* 2012; Burgstaller *et al.* 2014).

It is possible that the systematic biases in model fit represented in Figure 5 are caused by unaccounted-for natural selection. For example, compared to the observed distribution of heteroplasmy frequencies, the MAP model parameters produce an overabundance of intermediate-to-high-frequency heteroplasmies in blood tissues (Figure 5). Hypothetically, this could be caused by purifying selection against harmful heteroplasmic mutations in blood, which could skew the distribution of heteroplasmy frequencies toward zero. If selection tends to act on only a single heteroplasmic variant at a given time (*i.e.*, if clonal interference between different heteroplasmic alleles is rare), the method we present here could potentially be adapted to make inferences about natural selection in place of mutation. We leave this for future work.

We note that, in a recent study finding repeated convergent heteroplasmy in specific tissues in humans, and, thus, evidence of positive selection on heteroplasmy (Li *et al.* 2015), the subjects under consideration were deceased, and, thus, older than those considered by Rebolledo-Jaramillo *et al.* (2014); if selection on mitochondrial heteroplasmy intensifies with age, this may explain the lack of such repeated convergence in Rebolledo-Jaramillo *et al.* (2014).

We chose to model heteroplasmy allele frequency dynamics with the Wright-Fisher population model from population genetics. This model is well-studied and, thus, facilitates interpretation, and it is general in the sense that many different population-genetic models of reproduction closely resemble the Wright-Fisher model when population sizes are at least moderately large (Ewens 2004). The Wright-Fisher model





**Figure 4** Posterior samples of the EBS for mothers of different ages. (A) Posterior distribution of the effective, between-generation, bottleneck size for younger, older, and median-aged mothers. (B) Relationship between mother's age at childbirth and the EBS. The orange dashed line shows how the median EBS varies with age at childbirth. The solid blue lines show posterior samples from the relationship between EBS and age at childbirth, with each having the form of (C.4), where the genetic drift parameters in this equation are jointly sampled from the posterior distribution. A total of  $n = 1000$  lines sampled

from the posterior are plotted. We note that each line necessarily decreases with mother birth age due to our assumption that genetic drift accumulates at some rate in the oocyte [see (C.4)]; what varies from one line to another is the rate at which the EBS decreases due to this accumulation of genetic drift.

does not include mechanistic details of mtDNA dynamics such as the hypothesized segregation of mtDNA copies in genetically homogeneous nucleoids (e.g., Cao *et al.* 2007; Khrapko 2008), or mitochondrial fission, fusion, degradation, and duplication. The coarse effects of many of these mechanistic details are likely to be captured by the Wright-Fisher model through appeals to the concept of an effective population size, just as the details of reproduction of many classical models of reproduction from population genetics can often be reduced to a change in the effective population size of the Wright-Fisher model (Ewens 2004; Wakeley 2009). Here, if we were to include these effects in our model, there would likely be very little power to infer their properties, as sample sizes are small ( $n < 100$  heteroplasmies). In larger, and differently structured, datasets, there may be greater power to infer mechanistic details of mitochondrial proliferation.

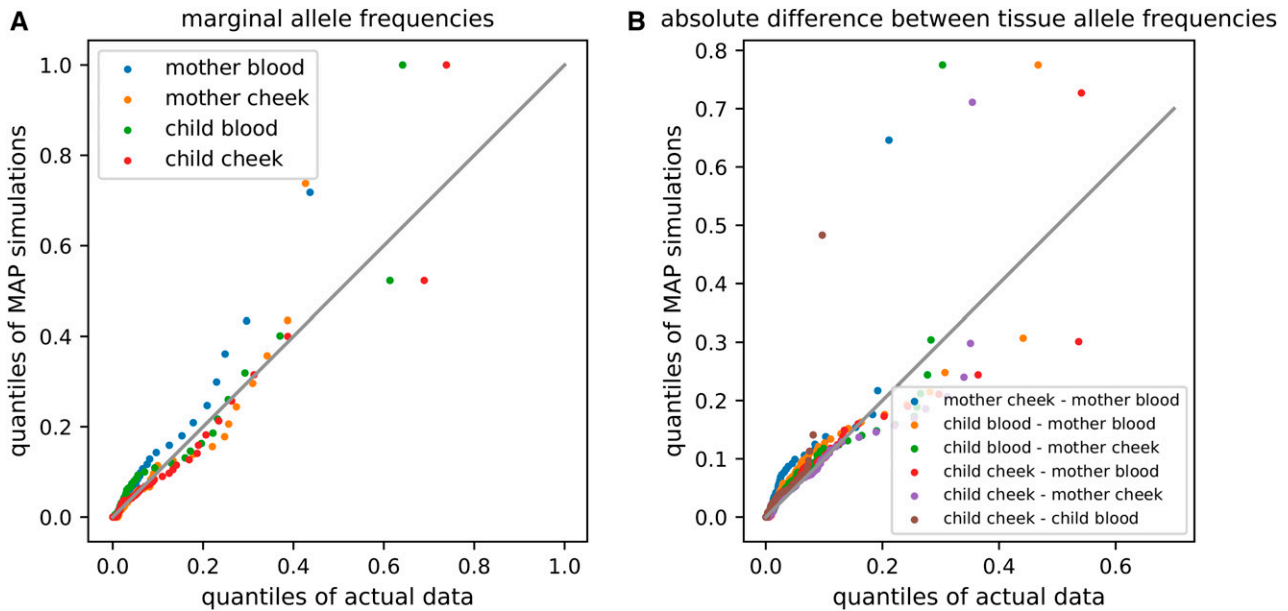
In a study of mitochondrial heteroplasmy transmission between the mothers and children of two-parent-child trios from the Netherlands, Li *et al.* (2016) found support for a variable bottleneck size, where the size of the bottleneck for a particular heteroplasmic locus is randomly sampled from a distribution. The model we present here also allows for variable bottleneck sizes, but it assumes a particular relationship between the effective oogenic bottleneck size and the age of the mother. As discussed above, our inference is inconclusive about whether or not the bottleneck size is variable with age. A variable bottleneck size, independent of mother's age, could also be implemented in our inference framework by integrating over the distribution of bottleneck sizes during the calculation of allele frequency transition distributions. In this case, like Li *et al.* (2016), we would be inferring the parameters of the bottleneck size distribution rather than a single bottleneck size. We leave this as an opportunity for future investigation.

Johnston *et al.* (2015) have recently used a detailed, mechanistic model of mitochondrial duplication, degradation,

and partitioning to study mitochondrial dynamics during oogenesis. The authors applied their model to data on the time evolution of heteroplasmy frequency variance and mtDNA copy number variation during oogenesis in mice, finding that the size of the oogenic bottleneck is just one contributor to the final variance in heteroplasmy frequencies after oogenesis is complete, and that their analysis is inconclusive about the fine details of segregation in nucleoids (except that nucleoids are not very large and genetically homogeneous). This work is broadly in agreement with the present study and is complementary in that it analyzes just one phase of ontogeny (namely, oogenesis) and makes use of time series observations of heteroplasmy frequencies in mice rather than heteroplasmy frequencies in multiple somatic tissues in adult humans.

However, it is still possible that the dynamics of heteroplasmy frequency change do not meet the basic assumptions of any population-genetic model. Any population-genetic model of heteroplasmy would assume that the germ cells or somatic stem cells giving rise to heteroplasmic variation would compete with one another for reproduction, or at least be chosen randomly for transmission or reproduction. If instead, for example, there exists a cellular mechanism of quality control, such that nonheteroplasmic eggs are given priority in ovulation and tend to be ovulated before heteroplasmic eggs, the number of transmitted heteroplasmies would increase with mother's age, but the dynamics would not be completely described by any population-genetic model that assumes random mating (with or without natural selection) and competition among egg cells for offspring. Other such mechanisms of heteroplasmy propagation could be imagined. Even if standard population-genetic models cannot adequately describe heteroplasmy frequency change, modeling heteroplasmy frequency changes on an ontogenetic phylogeny would still be a valid approach.

We assume that the shape of the ontogenetic phylogeny relating the sampled tissues is known. For the dataset from Rebolledo-Jaramillo *et al.* (2014), this is an appropriate



**Figure 5** Quantile-quantile comparison of real heteroplasmy data from Rebolledo-Jaramillo *et al.* (2014) and data simulated under maximum *a posteriori* parameter estimates inferred from this data. (A) compares marginal distributions of allele frequencies in each tissue, and (B) compares distributions of absolute differences in allele frequency between tissues. Each dot represents a sequential percentile of the distributions being compared. Following Rebolledo-Jaramillo *et al.* (2014), alleles were polarized such that the minor allele in the mother (averaged across her two tissues) was denoted as the focal allele.

assumption, since the two somatic tissues in the mother must be most closely related to one another, just as the two somatic tissues of the offspring must be most closely related to one another. For other datasets, differing in the number or identity of the sampled tissues, there may be less of an *a priori* expectation for the shape of the ontogenetic phylogeny. While there is a general understanding of the major divisions of tissues during development, the embryonic origins and lineage of somatic germ cell populations are not straightforward and still being established (*e.g.*, Boisset and Robin 2012; Fuentealba *et al.* 2015; Romagnani *et al.* 2015). The current model could easily be extended to ontogenetic phylogenies for families with two or more offspring. For families with more than two offspring, the genealogy of the oogonia eventually giving rise to the offspring would be unknown. This part of the phylogeny could be inferred jointly with other parameters, or, depending on the inferred rate of genetic drift in the female germ lineage (here  $1.6 \times 10^{-3}$  drift units per year), it could be assumed that no genetic drift occurs between the birth of the youngest and oldest children.

The topology of the ontogenetic phylogeny could also be made more complicated by admixture, which is not included in our inference framework. Admixture could result from biological processes, such as contributions to a mitotic tissue from distinct, isolated adult stem cell niches, or from physical sampling of an organ containing multiple tissues derived from distinct developmental lineages. Conceptually, our ontogenetic phylogeny approach could be extended to work with admixture graphs (Patterson *et al.* 2012; Pickrell and Pritchard 2012) by adapting the pruning algorithm for calcu-

lating likelihoods to the dependence structure introduced by admixture. However, given the small size of current heteroplasmy frequency datasets compared to large whole-genome SNP datasets, detecting admixture with *f*-statistics (Patterson *et al.* 2012; Peter 2016), or a more typical population phylogeny inference procedure (*e.g.*, Treemix, Pickrell and Pritchard 2012), would likely be more suitable.

The inference framework we present here should be applicable in future studies of heteroplasmy dynamics in humans and other organisms. Our software *mope* is flexible with respect to the pedigree of the sampled individuals, and, thus, is suitable for studies of heteroplasmy both across several generations and within unrelated individuals. Flexibility is also given with respect to the number of tissues sampled—even studies of just a single tissue may benefit from modeling multiple ontogenetic processes (*e.g.*, Li *et al.* 2016). Our fully Bayesian inference method provides a natural way of quantifying uncertainty, which is important in studies of heteroplasmy as the number of polymorphic loci is often small compared to other genomic studies. Finally, *mope* allows the user to choose the ontogenetic processes to place in the ontogenetic phylogeny; in the current version allele frequency changes for each such ontogenetic process occur according to the neutral Wright-Fisher model, but processes governed by other dynamics (*e.g.*, selection, mutation) could be implemented by modifying the freely available source code.

The ontogenetic phylogeny framework may also be useful in areas other than the study of mitochondrial heteroplasmy. In particular, in the study of the dynamics of cancer evolution,

heterogeneous progression in samples of many tumors may necessitate modeling per-day rates of genetic drift and mutation (or natural selection) rather than fixed amounts common to all tumors. Our inference procedure could also be used in the typical population phylogenetic setting to infer the divergence history of a group of populations, but this application is limited by the relatively small number of loci [ $< O(1000)$ ] that our method can accept due to the computational costs of likelihood evaluations with the pruning algorithm. A maximum-likelihood implementation of our model, requiring fewer likelihood evaluations, may be applicable to genome-scale SNP data, possibly comparing to Kim Tree (Gautier and Vitalis 2013) and SpikeyTree (Tataru *et al.* 2015).

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## Appendix A

### Likelihood Calculation

Briefly, the pruning algorithm calculates, for each node  $n$  in the phylogeny and each frequency  $f_j$  at node  $n$ , the probability  $P(\mathcal{D}_{(n)} | x_{(n)} = f_j)$ , where  $\mathcal{D}_{(n)}$  is the data at all the leaves collectively having  $n$  as their most recent common ancestor, and  $x_{(n)}$  is the heteroplasmy allele frequency at node  $n$ . The algorithm proceeds up the tree, from the leaves to the root, using the fact that

$$P(\mathcal{D}_{(n)} | x_{(n)} = f_j) = \prod_c \sum_k P(x_{(c)} = f_k | x_{(n)} = f_j) P(\mathcal{D}_{(c)} | x_{(c)} = f_k). \quad (\text{A.1})$$

child of  $n$

The probability  $P(x_{(c)} = f_k | x_{(n)} = f_j)$  is the probability of transitioning from allele frequency  $f_j$  in node ( $n$ ) to  $f_k$  in node ( $c$ ), a child of ( $n$ ). This probability is calculated using the discrete-generation Wright-Fisher model, as explained in Appendix B.

Here, and below, the current genetic drift parameters  $\mathbf{b}$  and mutation rates  $\boldsymbol{\theta}$  are implied. We model the probability of the data at leaf (*i.e.*, sampled tissue) node  $l$  as the binomial likelihood

$$P(\mathcal{D}_{(l)} | x_{(l)} = f_j) = \binom{C_l}{h_l} f_j^{h_l} (1-f_j)^{C_l-h_l}, \quad (\text{A.2})$$

where  $C_l$  and  $h_l$  are, respectively, the total coverage and number of alternative alleles in that tissue.

Given each  $P(\mathcal{D}_{(r)} | x_{(r)} = f_j)$  for root node  $r$ , the overall likelihood is

$$P(\mathcal{D}_{(r)}) = \sum_j P(x_{(r)} = f_j) P(\mathcal{D}_{(r)} | x_{(r)} = f_j). \quad (\text{A.3})$$

The probabilities  $P(x_{(r)} = f_j)$  are given by the heteroplasmic allele frequency distribution at the root, a discretized symmetric beta distribution with additional weight at frequencies 0 and 1, the parameters of which are inferred jointly with the genetic drift and mutation parameters.

The probability of heteroplasmic polymorphism [cf. denominator of Equation (3)] can be calculated as

$$P(H_i; \mathbf{b}, \boldsymbol{\theta}) = 1 - P(\mathcal{D} | \text{all leaves } 0) - P(\mathcal{D} | \text{all leaves } 1), \quad (\text{A.4})$$

with the second two terms giving the probability of the read count data in all the sampled tissues given that allele frequencies are all 0 or 1, respectively.

## Appendix B

### Calculating Allele Frequency Transition Distributions

The pruning algorithm requires distributions of allele frequency transitions along a branch. Our approach to calculating allele frequency transition probabilities is simple and intuitive: we precalculate transition distributions under the discrete-generation Wright-Fisher model using numerical matrix multiplication on a grid of generations and mutation rates. To obtain a transition distribution that was not precomputed, we linearly interpolate between precomputed distributions. Using a haploid population size of  $N = 2000$  in our Wright-Fisher model calculations, we obtain a satisfactory approximation to numerically exact Wright-Fisher transition probabilities by precomputing distributions at just 207 different generations, ranging from one to 20,000, and 44 mutation rates, with  $\theta = 2N_e\mu$  ranging from zero to  $7.5 \times 10^{-2}$ . For ontogenetic processes modeled by a single-generation bottleneck with subsequent expansion, we precompute allele-frequency transition distributions for 48 bottleneck sizes ranging from two to 500, linearly interpolating between bottleneck sizes for distributions that are not precomputed.

Rather than use each  $(2001 \times 2001)$  transition matrix in its entirety, we combine discrete allele frequencies into 121 bins, with bins unevenly distributed between zero and one such that low and high frequencies are more represented than intermediate frequencies. We bin allele frequencies according to the following scheme: Let  $P = \{P_{ij}\}$  be a  $(2001 \times 2001)$  allele frequency transition matrix for a Wright-Fisher model with  $N = 2000$ , with  $P_{ij}$  being the probability of transitioning from frequency  $i$  to  $j$ . Let  $Q = \{Q_{k,l}\}$  be a  $(121 \times 121)$  binned transition matrix. If  $(a_1, \dots, a_m)$  are frequencies in bin  $k$ , and  $(b_1, \dots, b_n)$  are frequencies associated in bin  $l$ , then

$$Q_{k,l} = \begin{cases} \sum_{x=1}^n P_{(m+1)/2, b_x} & m \text{ odd} \\ \sum_{x=1}^n \left( \frac{1}{2} P_{m/2, b_x} + \frac{1}{2} P_{m/2+1, b_x} \right) & m \text{ even.} \end{cases}$$

## Appendix C

### Calculation of the EBS

We define the effective bottleneck between mother and offspring as the combined genetic drift occurring during the early oogenic bottleneck, the turnover of mitochondria in the maternal germline prior to ovulation, and the first few cell divisions after fertilization but before gastrulation. We combined the effects of genetic drift during these processes by (1) translating all drift parameters into units of generations per effective population size ( $g/N_e$ , “drift units”), (2) summing the drift in these units, and (3) translating this summed drift back into units of an instantaneous bottleneck. Since we assumed that bottlenecks occurred for just a single generation followed by doubling back up to a large population size (here,  $N = 2000$ ), we determined that the relationship between drift  $d_g$  measured in drift units and  $N_b$ , an instantaneous bottleneck size, is close to

$$d_g = \sum_{i=0}^n \frac{1}{N_b 2^i}, \quad (\text{C.1})$$

where  $n = \log_2(N/N_b)$  is the number of generations it takes for the population size to double back up to the original population size.

For  $N_b \ll N$ , this sum is well approximated by the integral

$$d_g \approx \int_{-\frac{1}{2}}^{\log_2(N/N_b)} \frac{dt}{2^t N_b} = \frac{N\sqrt{2} - N_b}{NN_b \ln 2} \approx \frac{\sqrt{2}}{N_b \ln 2} \approx \frac{2}{N_b}. \quad (\text{C.2})$$

The lower limit of integration follows from an interpretation of (C.1) as a midpoint Riemann sum, improving accuracy. Thus, we also have

$$N_b \approx \frac{2}{d_g}. \quad (\text{C.3})$$

For a mother of age  $a$ , the EBS is thus

$$N_{be} = \frac{2}{\frac{2}{N_b} + a\lambda_g + d_s}, \quad (\text{C.4})$$

where  $N_b$  is the early oogenesis bottleneck size,  $\lambda_g$  is the rate at which genetic drift accumulates in the maternal germline, and  $d_s$  is the amount of genetic drift occurring after fertilization but before gastrulation.

We confirmed (C.2) and (C.3) by finding, for different bottleneck sizes  $N_b$ , the amount of drift  $d_g$  that minimized the total variation distance between the allele frequency transition distributions specified by  $d_g$  and  $N_b$ :

$$\hat{d}_g(N_b) = \operatorname{argmin}_{d_g} \frac{1}{2} \sum_i |p_{d_g}(i) - q_{N_b}(i)|. \quad (\text{C.5})$$

Here  $p_{d_g}$  is the probability transition distribution for drift parameterized by  $d_g$  drift units, and  $q_{N_b}$  is the probability transition distribution for drift parameterized by bottleneck size  $N_b$ . Minimizing (C.5) for different values of  $N_b$  shows that our approximation (C.2) closely follows the numerical translation minimizing the total variation distance (Figure S5 in File S1).