The need for high-quality oocyte mitochondria at extreme ploidy dictates germline development

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ABSTRACT

Selection against severe mitochondrial mutations is facilitated by germline processes, lowering the risk of genetic diseases. How selection works is disputed: experimental data are conflicting and previous modelling work has not clarified the issues. Here we develop computational and evolutionary models that compare the outcome of selection at the level of individuals, cells and mitochondria. Using realistic de novo mutation rates and germline development parameters, the evolutionary model accurately predicts the observed prevalence of mitochondrial mutations and diseases in human populations. We show that biogenesis of high-quality mitochondria at extreme ploidy in mature oocytes can only be achieved under realistic parameters through selective pooling of mitochondria into the Balbiani body. The principal mechanisms debated in the literature, bottlenecks and follicular atresia, fail to predict these clinical data, because neither process effectively eliminates mitochondrial mutations under realistic conditions. Our findings explain the major features of female germline architecture, notably the longstanding paradox of over-proliferation of primordial germ cells followed by massive germ cell loss. The near-universality of these processes across animal taxa makes sense in light of the need to maintain mitochondrial quality at extreme ploidy in mature oocytes, in the absence of sex and recombination.

Keywords: Balbiani body, bottleneck, germline, mitochondria, mitochondrial mutation, mtDNA, oogenesis
INTRODUCTION

In mammals, mitochondrial gene sequences diverge at 10-30 times the mean rate of nuclear genes [1, 2]. This difference is typically ascribed to a faster underlying mutation rate and limited scope for purifying selection on mitochondrial genes, given negligible recombination and high ploidy [3]. At face value, weak selection against mitochondrial mutations might seem to be consistent with the high prevalence of mitochondrial mutations (~1 in 200 people) [4] and diseases (~1 in 5000 births) [5] in the general population. But it is not consistent with the strong signal of purifying selection [6], evidence of adaptive change [7] and codon bias [8] in mitochondrial genes, nor with the low transmission rate of severe mitochondrial mutations between generations [9-11]. Despite the high rate of sequence divergence, female germline processes apparently facilitate selection against mitochondrial mutations, but the mechanisms are disputed and poorly understood [12].

Here we develop computational and evolutionary models that compare three hypotheses of germline mitochondrial inheritance and selection: (i) mitochondrial bottlenecks and selection at the individual level; (ii) follicular atresia, which imposes selection on primordial germ cells during development; and (iii) selective transfer of mitochondria into the Balbiani body. Our approach incorporates plausible modes of selection against mitochondrial mutations, given the high (but not constant) ploidy of mitochondrial DNA (mtDNA) through all stages of germline development. We also incorporate important factors neglected in earlier work, in particular the input of de novo mitochondrial mutations and their segregation over multiple rounds of germ-cell division. This provides a realistic model of mutation, segregation and selection allowing the three hypotheses to be tested against the observed levels of mitochondrial mutation and disease in human populations [4, 5].
The idea that mitochondrial mutations are winnowed through a tight germline bottleneck is pervasive in the literature and has long been held to explain sharp changes in mutation load between generations [13-15]. The exact size of the bottleneck is unclear, with estimates from human studies spanning two orders of magnitude, between 10 and 1000 [14, 16-18]. Bottlenecks generate variance in mutation loads among the resulting germ cells, and tighter bottlenecks produce greater variance, offering scope for selection against mitochondrial mutations at the level of the individual [19-21]. The problem with this line of thinking is that it ignores two other forces. First, gametes are produced through multiple rounds of cell division, leading to repeated rounds of mitochondrial segregation, which in itself generates considerable variance [22]. Second, bottlenecks induce greater input of de novo mutations as more rounds of replication are required to regenerate the extreme ploidy of mitochondrial DNA in mature oocytes. By applying realistic segregation dynamics and mutational input, we evaluate the impact of these forces on the value of bottleneck size on individual fitness.

Follicular atresia is another force widely considered to be critical in maintaining oocyte quality [23-25]. In many animal species, including humans, the number of germ cells declines dramatically in the foetus between mid-gestation (~20 weeks in humans) when there are 7-8 million oocytes, to late gestation, when at least two thirds of these are lost, leaving a reserve of 1-2 million at birth [26]. Oocyte loss continues throughout the life of an individual, eventually leading to the depletion of the ovarian pool and loss of reproductive function at menopause [27-29]. This attrition has historically been ascribed to cell death during oocyte maturation [30, 31], but more recent findings implicate the apoptotic loss of
'nurse cells' during the genesis of primary oocytes [32]. In either case, differential oocyte loss offers scope for between-cell selection. However, the basis for between-cell selection has long been questioned, on the grounds that it seems unlikely that 70–80% of oocytes have low fitness as a result of mitochondrial mutations [33]. We therefore test whether selection against oocytes with higher loads of mitochondrial mutations during follicular atresia is capable of giving rise to the distribution of mutations observed in humans.

A more recent interpretation of germ-cell loss links it to the formation of the Balbiani body [34], a prominent feature of female germlines across invertebrates and vertebrates [32, 35, 36], including humans [34]. In the mouse and other mammals, proliferating germ cells form clusters of 5-8 cells that establish cytoplasmic bridges, through which around half the mitochondria from each nurse cell are streamed into the Balbiani body of the primary oocyte [34]. Cytoplasmic transfer is an active cytoskeletal process that depends in part on the membrane potential of discrete mitochondria [37, 38], offering scope for purifying selection through the preferential exclusion of dysfunctional mitochondria. The remaining nurse cells, now denuded of half their mitochondria, undergo apoptosis [34]. We consider the consequence of different strengths of selection at the level of mitochondrial function in the production of the Balbiani body.

To systematically distinguish between the predictions of these three different hypotheses, under a range of reasonable parameter values, we use a computational model to evaluate the patterns of mutation load generated over a single generation in each case. We then use an evolutionary model to compare the predictions of our computational model with clinical data on the prevalence of mutations and disease from human studies. Our results show that
germline bottlenecks and follicular atresia cannot alone explain the observed prevalence of mitochondrial mutations or disease. Only the selective pooling of high-quality mitochondria into the Balbiani body is likely to account for the clinical data. This process also pleasingly clarifies the longstanding paradox of germ-cell over-proliferation followed by massive loss that is almost universally conserved in the female germline of animal taxa.

RESULTS

Computational model including realistic de novo mutational input

The computational model follows the distribution of mitochondrial mutations across a single generation, using model parameters derived from human data [39] (Figure 1). The zygote is assumed to have ~500,000 copies of mitochondrial DNA (exact number $2^{19}$), which are randomly partitioned to the daughter cells at each cell division. We assume independent segregation of mitochondria with one mtDNA per mitochondrion, and do not consider complications that arise from the packaging of multiple mtDNA copies per mitochondrion [14].

Mitochondrial replication is not active during early embryo development [40], so the mean mitochondrial number per cell approximately halves with each division (Figure 1B). After 12 cell divisions a random group of 32 cells form the primordial germ cells (PGC) [41], with a mean of 128 mitochondria per PGC. Mitochondrial replication resumes at this point [39, 40]. Each mitochondrion (and mtDNA) doubles prior to random partitioning at each cell division. With probability $\mu$, one of the daughter mitochondria acquires a new deleterious mutation through a copying error. We consider $\mu$ in the range $10^{-9}$ to $10^{-7}$ per base pair per cell division, consistent with the range of estimates for the female germline, and assume no
back mutations (see Methods). Point mutations during replication are the dominant form of mutation in mtDNA, so we do not consider damage from other sources such as oxidative damage [42]. Mitotic proliferation of PGCs gives rise to ~8 million oogonia, which are reduced to ~1 million primary oocytes during late gestation (Figure 1) [39, 40]. Proliferation is followed by a quiescent phase during which the mitochondria in primary oocytes are not actively replicated. Mutations accumulate far more slowly during this phase, which persists over decades in humans [40, 43]. For simplicity, we assume no mutational input during this period (not marked in Figure 1). At puberty, the primary oocytes mature through clonal amplification of mitochondria back to the extreme ploidy in mature oocytes (~500,000 copies; Figure 1B) [44]. The same copying error mutation rate $\mu$ is applied during this process.

We consider three different forms of selection on mitochondria: selection at the level of the organism, cells, or mitochondria. We apply selection at the level of the organism on the zygotic mutation load. Selection at the level of cells or mitochondria is applied during culling at late gestation when primary oocytes are produced. Each of these processes can be captured by modifications of the computational model, allowing easy comparison between them. The model extends earlier modelling work that considered segregational variation of a fixed burden of existing mutations [13, 19-21] but neglected the input of new mutations during PGC proliferation and oocyte maturation, as well as the loss of germ cells during late gestation. The analysis here shows the importance of considering these additional processes governing the population of mitochondria in germline development.
Germline bottleneck increases variance but introduces more de novo mitochondrial mutations

The effect of a bottleneck was assessed in the model by allowing $b$ extra rounds of cell division without replication during early embryonic development (e.g., two extra rounds shown in Figure 2A). Each additional cell division leads to an average reduction of $(0.5)^b$ mitochondria in PGCs compared to the base model. We then held mitochondrial numbers at this lower value through the period of PGC proliferation, consistent with some views of the bottleneck [45]. Tighter bottlenecks at this early developmental stage generate greater segregational variance in mutation load between cells (Figure 2B). This increase in variance persists and is enhanced through PGC proliferation to the production of primary oocytes and ultimately in mature oocytes (Figure 2B). The bottleneck not only creates a wider spread of mutation number per cell, but also the possibility that cells can be mutation free even when initiated from a zygote that contains significant numbers of mutations (Figure 2B). Bottlenecks in themselves do not change the mean mutation load, as they occur before the start of mitochondrial replication (i.e. at PGC specification; Fig. 2B) [40]. But oocyte maturation requires the expansion of mitochondrial number back to half a million. Cells starting with lower numbers must therefore undergo more rounds of replication, and hence will accumulate more de novo mutations. So, the mean mitochondrial mutation load in mature oocytes increases with tighter bottleneck size, albeit this effect is small with low mutation rates ($\mu = 10^{-8}$; Fig. 2B). Nonetheless, the tension between variance and mean determines the overall selective consequence of the bottleneck.

The advantage that the bottleneck brings depends on how selection acts against the mutation load carried by an individual. Based on the observed dependence of mitochondrial
diseases on mutation load [46-48], in which more serious phenotypes typically manifest only at high mutant loads of >60% [46-48], it is thought that individual fitness is defined by a concave fitness function, indicative of negative epistasis (Figure 2C). This assumes that each additional mitochondrial mutation causes a greater reduction in fitness beyond that expected from independent effects. In other words, low mutation loads have a relatively trivial fitness effect, whereas higher mutation loads produce a steeper decline in fitness.

The change in mutation load (Δm) over a single generation after individual selection was measured against 5 mean bottleneck sizes (B = 128, 64, 32, 16, 8), for three initial mutation loads (m₀) and three mutation rates (μ). The bottleneck shows an ambiguous relationship with fitness, dependent on the inherited mutation load (m₀). For the estimated mutation rate (μ = 10⁻⁶) there is always an increase in mutation load in individuals who inherit low or medium mutation loads (m₀ = 0.001, 0.01; Figure 2D). This increase in load becomes more deleterious with a tighter bottleneck (Figure 2D). The bottleneck only confers a benefit among individuals who inherit a high mutation load (m₀ = 0.1; Figure 2D), where the advantage of greater variance outweighs the increase in de novo mutation load. If the mutation rate is lower (μ = 10⁻⁹), bottlenecks have little effect except when severe, where they again cause an increase in mutation number in individuals with low or medium mutation loads (m₀ = 0.001, 0.01; Fig. S1A). In individuals with high mutation load (m₀ = 0.1) only tighter bottlenecks (B = 16, 8) are beneficial (Fig. S1A). If the mutation rate is higher (μ = 10⁻⁷) the pattern shifts slightly. At low or medium mutation loads (m₀ = 0.001, 0.01) all bottleneck sizes (B = 128, 64, 32, 16, 8) accumulate many more de novo mutations (Fig. S1B). Only individuals with high inherited mutation loads (m₀ = 0.1) benefit from a bottleneck, and even then only when the bottleneck is at its tightest (B = 8) (Fig.
S1B). In sum: even though bottlenecks generate greater variance, they impose the need for additional rounds of replication during oocyte maturation, resulting in greater de novo mutational input. This makes tight bottlenecks advantageous only for individuals who inherit high mutation loads, but not for the great majority of the population, where the prevalence of mitochondrial mutations is generally between 0.001 and 0.01 [4, 14].

**Follicular atresia cannot be explained by realistic selection against cells with high mitochondrial mutation loads**

In the analysis of bottlenecks above, the culling of ~8 million oogonia to 1 million primary oocytes at the end of PGC proliferation was assumed to be a random process (Figure 2A). This loss has a minimal effect on the mean and variance in frequency of mitochondrial mutations in germ cells, given the large numbers involved (and no effect at all when averaged over a population). However, the loss of ~80% of oocytes via follicular atresia during late gestation has long been puzzling and could arguably reflect selection against cells with higher mutation loads.

To analyse follicular atresia, cell-level selection was applied to oogonia at the end of PGC proliferation (Figure 3A). PGs vary in mutation frequency due to both the random segregation of mutants during the multiple cell divisions of proliferation and the chance input of new mutations during mtDNA replication. In principle, we assume that between-cell selection is governed by a negative epistatic fitness function (Figure 3B) similar to that thought to apply at the individual level, and vary selection from linear ($\xi = 1$), weak ($\xi = 2$) to strong epistasis ($\xi = 5$). Positive epistasis ($\xi < 1$), whereby a single point mutation produces a steep loss of fitness, but additional mutations have less impact (i.e. mutations
are less deleterious in combination), seems biologically improbable, so we do not consider it here.

The effect of cell selection during follicular atresia was calculated as the change in mutation frequency for individuals carrying different mutation loads ($m_0$) over a single generation, given standard values for de novo mutations ($\mu = 10^{-8}$) and bottleneck size ($B = 128$). Under strong negative epistasis ($\xi = 5$), only the few cells with very high mutation loads (generated by segregation) are eliminated. Cell-level selection does not reduce mutation load, even for individuals with a high initial frequency of mutations ($m_0 = 0.1$; Figure 3C). Cell-level selection is more effective with weak epistasis ($\xi = 2$) or linear selection ($\xi = 1$) as this makes cells with lower mutation loads more visible to selection, and has a greater benefit in individuals carrying higher initial mutation loads (Figure 3C). However, in individuals who inherit low or medium mutation load ($m_0 = 0.001, 0.01$) cell selection offers a minimal constraint against mutation input. The only case in which cell selection produces a benefit is with high mutation load ($m_0 = 0.1$) under linear selection ($\xi = 1$) (Figure 3C). This pattern holds for a lower mutation rate ($\mu = 10^{-9}$; Figure S2A), while there is no benefit at all at a higher mutation rate ($\mu = 10^{-7}$; Figure S2B).

**The Balbiani body pools high-quality mitochondria and restricts de novo mutation input**

An alternative interpretation of atresia lies in the formation of the Balbiani Body, a nearly universal feature of female germlines in animals [49]. We model the developmental process giving rise to the Balbiani body by assuming that cysts of 8 oogonia form at the end of PGC proliferation (Figure 4A) as is typical in mammalian development [32, 50]. Cells within a cyst are derived from a common ancestor (i.e. via 3 consecutive cell divisions). At the 8-cell
stage, intercellular bridges form between the oogonia. These allow cytoplasmic transfer of a proportion of mitochondria ($f$) from each cell to join the Balbiani body of the single cell destined to become the primary oocyte (Figure 4A). The mitochondria that undergo cytoplasmic transfer are sampled at random (without replacement), with different weights for wildtype ($p_{wt}$) and mutant ($p_{mut}$) mitochondria, until $f$ have moved to the Balbiani body. The oogonia that donate their cytoplasm to the primary oocyte are now defined as nurse cells, and undergo programmed cell death – atresia (Figure 4A).

The model shows that two benefits accrue from cytoplasmic transfer. The first benefit of mitochondrial transfer into the Balbiani body is that pooling increases the number of mitochondria in primary oocytes. As the proportion of mitochondria transferred increases towards the estimated rate of $f = 50\%$ [32], the number of mitochondria in primary oocytes increases 4-fold. Pooling therefore cuts the number of rounds of replication needed to reach the extreme ploidy required by mature oocytes, which decreases the input of new mutations from replication errors during oocyte maturation. This benefit accrues whatever the initial mutation load, and more dramatically with a higher mutation rate (Figure S3).

The second benefit arises from selective transfer of mitochondria. Preferential exclusion of mutant mitochondria ($p_{wt} > p_{mut}$), as suggested by experimental evidence [32], lowers the mutation load in primordial oocytes (Figure 4B). The difference between $p_{wt}$ and $p_{mut}$ determines the extent to which the mutation load is reduced, with stronger exclusion of mutant mitochondria (lower $p_{mut}$) reducing the number of mutations when the inherited load is medium or high ($m_0 = 0.01, 0.1$), albeit with a negligible effect at low initial mutation load ($m_0 = 0.001$; Figure 4C). The same effect is seen with lower and higher
mutation rates (Figure S4). Nurse cells retain a higher fraction of mutant mitochondria but undergo apoptosis, removing mutants from the pool of germ cells, and explaining the need for an extreme loss of germ cells during late gestation. This effect acts in concert with pooling leading to a reduction in both the mean and variance of mitochondria mutation load (Figure 4B).

Evolutionary model

The computational model discussed above gives an indication of the effectiveness of selection at the level of individuals, cells or mitochondria in eliminating mitochondrial mutations across a single generation. To address the long-term balance of mutation accumulation versus selection over many generations, we developed an evolutionary model. This assesses the effectiveness of the three representations of germline development in explaining the observed prevalence of mitochondrial mutation load and disease in human populations (see Materials and Methods). This evolutionary model evaluates long-term evolutionary change in an infinite population with non-overlapping generations and is implemented using a number of approximations, which greatly reduce the model complexity (see Materials and Methods).

By iterating the patterns of germline inheritance and selection, the equilibrium mutation distribution was calculated across a range of mutation rates and bottleneck sizes. The accuracy of the three models was then assessed as the likelihood of reproducing the observed levels of mitochondrial mutations in the human population (Fig. 5). Specifically, we used estimated values of 1/5000 for mitochondrial disease (>60% mutant), 1/200 for carriers of mitochondrial mutants (2-60% mutant) and hence 99.5% of individuals are
‘mutation free’ (i.e. carry <2% mutants, the threshold for detection in these estimates of mutation frequency (REFS). Recent deep-sequencing estimates using a mutation detection threshold of >1% [14], show that a minor allele frequency of 1-2% is relatively common in selected human PGCs, but this does not alter earlier population-level estimates of the proportion of carriers not suffering from overt mitochondrial disease, defined as a 2-60% mutation load used here.

Likelihood heatmaps confirm that selection at the level of individuals (bottleneck) or cells (follicular atresia) do not readily approximate the clinical data (Figure 5A-B). Only at very low mutation rates (\( \mu = 10^{-9} \)) do these forms of selection offer a credible explanation of the observed mutation load and disease frequency in humans, especially when using tighter bottlenecks (Figure 5A). These limitations do not apply to the preferential transfer of wildtype mitochondria into the Balbiani body (Figure 5C-D). Even mild selection against the transfer of mutant mitochondria into the Balbiani body (\( p_{mut} = 0.33, p_{wt} = 0.67 \)) generates a high log-likelihood of reproducing the clinical data at realistic mutation rates \( \mu = 10^{-9} / \text{bp} \) and bottleneck sizes (> 100 mitochondria per cell) (Figure 3C). Stronger selection on transfer probabilities (\( p_{mut} = 0.25, p_{wt} = 0.75 \)) can account for the clinical pattern under a wide range of bottleneck sizes and mutation rates (Figure 5D).

**DISCUSSION**

How selection operates on mitochondria has long been controversial. At the heart of this problem is the paradox that mtDNA accumulates mutations faster than nuclear genes, yet is under stronger purifying selection. Mitochondrial mutations accumulate through Muller’s ratchet, as mtDNA is exclusively maternally inherited, and does not undergo recombination.
through meiosis [3]. In addition, mitochondrial genes are highly polyploid, which obscures the relationship between genotype and phenotype, hindering the effectiveness of selection on individuals. Despite these constraints, deleterious mitochondrial mutations seem to be eliminated effectively [6-11], facilitated by female germline processes that have long been mysterious. These include: the excess proliferation of primordial germ cells (PGCs) [51]; the germline mitochondrial bottleneck (when mitochondrial numbers are reduced to a disputed minimum in PGCs) [13-15]; the formation of the Balbiani body (‘mitochondrial cloud’) in primary oocytes [32, 38]; the atretic loss of 70-80% of germ cells during late gestation [26, 30]; the extended oocyte quiescence until puberty or later (during which time mitochondrial activity and replication is suppressed) [43, 52]; and the generation of around half a million copies of mtDNA in mature oocytes [44]. The key question is how do these processes facilitate the maintenance of mitochondrial quality over generations?

In this study, we introduced a computational model that considers these germline processes from the perspective of mitochondrial proliferation, segregation and selection, using realistic estimates of parameter values, drawn from the human literature [39, 40]. Most work to date [13, 15, 45, 53, 54] has focused on the mitochondrial bottleneck as a means of generating variation in mitochondrial content between oocytes and by extension zygotes (Figure 2B), furnishing the opportunity for selection to act on individuals in the following generation. These studies have been unable to reconcile serious differences in experimental estimates of mitochondrial numbers during PGC proliferation, inciting inconclusive debates over the tightness of the bottleneck [13, 15, 45, 53, 54]. More significantly, this earlier work neglects an important germline feature, the introduction of de novo mitochondrial mutations produced by copying errors [55] rather than damage by reactive oxygen species.
These accumulate during PGC proliferation and, equally importantly, during the mass-production of mtDNAs in the mature oocyte. Tighter bottlenecks are disadvantageous as they impose the need for more rounds of mitochondrial replication which means a greater input of de novo mutations. Our modelling shows that for most individuals the mean mutation load shows little meaningful change (Figure 2D), regardless of whether the mutation rate is set low or high (Figure S1) and in fact increases with tighter bottleneck size (Figure 2D). Most individuals have low mutation loads (~99.5% in human populations [4, 5]), and for them, the normal process of repeated segregation during cell division generates sufficient variance in itself. Any marginal increase in variance caused by bottlenecks is more than offset by increased mutational input. Tighter bottlenecks only benefit individuals who already carry high mutation loads (i.e. \( m_0 \geq 0.1 \), Figure 2D). For them, there is benefit in further reductions in bottleneck size as this increases the fraction of mature oocytes with significantly reduced mutation load (Figure 2D).

These results show that the popular idea that a germline mitochondrial bottleneck facilitates selection against mitochondrial mutations is misconstrued. The value of a bottleneck depends on the unforeseen trade-off between increasing genetic variance and mutation input. In fact, the reduction in mitochondrial copy numbers from zygote to primordial germ cells should be thought of as the reestablishment of a typical copy number at the start of cellular differentiation, which commences after multiple cell divisions without mtDNA replication. What counts as a bottleneck are the ‘extra’ rounds of cell division reducing mitochondrial number below the ‘normal’ number, and the incremental increase in variance this induces. Most critically, the bottleneck needs to be understood in relation to oogamy, the massively exaggerated mitochondrial content of the female gamete. This is a
characteristic of metazoan gametogenesis [44]. Previous work has shown it is beneficial in animals with mutually interdependent organ systems [44]. The extreme ploidy in the zygote allows early rounds of cell division to occur without replication, and hence without de novo mutational input. These initial cell divisions generate little between-cell differences, as segregational variance is weak when numbers are high (e.g. Fig 1B before PGC specification). So at the point of cellular differentiation (~12 cell divisions) there is homogeneity in the mutation load among the different organ systems and no one system is likely to fail, which would massively lower the fitness of the whole organism [44]. This contrasts with organisms that have modular growth, such as plants and basal metazoa (sponges, corals, placazoa), which neither sequester a recognizable germline nor have oocytes with massively expanded mitochondrial numbers [44, 57].

Follicular atresia is another female germline feature examined in our modelling, in which there is over-proliferation of PGCs followed by ~80% loss early in development, before oocyte maturation [26, 30]. This massive reduction in germ cell number has long been enigmatic, for it is unlikely to be random, yet does not obviously serve a selective function, as it seems unlikely that such a high proportion of germ cells could have low fitness [23-25]. The model confirms this intuition. Selection among PGCs at the end of the period of proliferation has little effect in significantly reducing mutation load (Figure 3C). Assuming a concave fitness function (Figure 3B), which seems reasonable by extension from the severity of mitochondrial diseases [46, 48], between-cell selection is ineffective, as it only eliminates PGCs with very high mutational numbers. This has little effect in constraining the burgeoning of lower mutation loads. Linear selection does better, even if it seems unrealistic, as it will act against a broader range of mutational states. But as with
bottlenecks, it is only beneficial in individuals already carrying significant mutation loads (i.e. $m_0 \geq 0.1$, Figure 3C). We conclude that cell-level selection produces little measurable reduction in mutation load and so is unlikely to be responsible for follicular atresia.

A more recent explanation of PGC loss relates to the formation of the Balbiani body in primary oocytes [32, 34]. In many metazoans, including clams [36], insects [35, 58], mice [50] and probably humans [34], the over-proliferation of PGCs culminates in their organization into germline cysts of multiple oogonia connected by cytoplasmic bridges [32, 50, 58]. These connections allow the transfer of mitochondria and other cytoplasmic constituents by active attachment to microtubules, into what becomes the primary oocyte [32]. The surrounding oogonia that transferred their mitochondria, now termed nurse cells, die by apoptosis [32]. The plethora of terms should not mask the key point that nurse cell death accounts for a considerable fraction of the germ cell loss usually ascribed to follicular atresia. We modelled selective mitochondrial transfer into the Balbiani body, perhaps in part reflecting membrane potential [35, 38]. This achieves two complementary things: it purges mutations and pools high-quality mitochondria in a single cell. If the germline cyst is composed of eight cells that contribute half of their mitochondria to the Balbiani body, then the primary oocyte gains four times as many mitochondria, cutting the need for additional rounds of mtDNA copying, and so reducing de novo mutations during oocyte maturation. Selective transfer and pooling lowers the mutation load across a wide range of mutation rates and inherited loads (Figure 4C, Figure S3-4). This process differs from mitophagy, the main route used in somatic cells for maintaining mitochondrial quality [59, 60], as it not only removes mutant mitochondria, but crucially also increases mitochondrial numbers, a key requirement for prospective gametes. The requirement for pooling of mitochondria to lower the mutation load from
copying errors also aligns with experimental observations of active spindle-associated mitochondrial migration to the generative oocyte in the formation of polar bodies during meiosis I of oogenesis [61]. We predict that selection for mitochondrial quality occurs during this process (i.e. polar bodies retain mutant mitochondria) but have not dealt with that explicitly in the model.

These insights depend in part on the parameter values used in the modelling, many of which are uncertain. We have examined variation around the most representative values drawn from the literature [2, 14, 62, 63], and aimed to be conservative wherever possible. We considered mutation rates across two orders of magnitude, around $10^{-8}$ per bp as the standard [62] and a similar range of bottleneck sizes ($\bar{B} = 8 - 128$). Strong selective pooling of mitochondria into the Balbiani body predicts the observed prevalence of mitochondrial mutations and diseases in human populations [4, 5] under a wide range of mutation rates and bottleneck sizes (Figure 5). Selection at the level of individuals or cells are much more constrained explanations, although we do not rule out some role for these processes (Figure 5). In general, higher mutation rates ($10^{-7}$ per base pair) strengthen the conclusions discussed here (Figures S1-S4) whereas the lowest mutation rates are more commensurate with weaker forms of evolutionary constraint generated by selection on individuals or cells. Plainly, weaker selection approximates best to clinical data when the mutation input tends towards zero (Figure 5). However, such low mutation rates are not consistent with the 10-30-fold faster evolution rates of mtDNA compared with nuclear genes [1, 2], or with the strong signatures of purifying [6] and adaptive [7] selection on mitochondrial genes. In addition, we have ignored the contribution of oxidative damage caused by reactive oxygen species. While this is likely low compared with copying errors [42, 55], oxidative mutations
may accumulate over female reproductive lifespans [56], perhaps contributing to the timing of the menopause [64]. As primary oocytes contain ~6000 mitochondria [64], expansion up to ~500,000 copies in the mature oocyte will amplify any mutations acquired during oocyte arrest at prophase I, potentially over decades [55]. The metabolic quiescence of oocytes can best be understood in light of the need to repress mitochondrial mutation accumulation during the extended period before reproduction [43, 52]. In any case, our assumption that zero mutations are caused by oxidative damage is plainly conservative.

We have addressed here a simple paradox at the heart of mitochondrial inheritance. Like Gibbon’s *Decline and Fall of the Roman Empire*, mitochondrial DNA is often portrayed as being in continuous and implacable decline through Muller’s ratchet [3]; yet like the Empire, which endured for another millennium, mitochondrial DNA has persisted and has been at the heart of eukaryotic cell function for over a billion years [65]. Strong evidence for purifying and adaptive selection implies that the female germline facilitates selection for mitochondrial quality, but the mechanisms have remained elusive. We have modelled segregation and selection of mitochondrial DNA at each stage of germline development, and shown that direct selection for mitochondrial function during transfer into the Balbiani body is the most likely explanation of the observed prevalence of mitochondrial mutations and diseases in human populations. More remarkably, this mitochondria-centric model elucidates the complexities of the female germline. It explains why mature oocytes are crammed with mitochondria [44], whereas sperm mitochondria are typically destroyed, giving rise to two sexes [22]; why germ cells over-proliferate during early germline development; why oogonia organize themselves into germline cysts, forming the Balbiani body; why the majority of germ cells then perish by apoptosis as nurse cells; why primary
oocytes enter metabolic quiescence, sometimes for decades; and even why polar bodies channel most of their mitochondria into a single mature oocyte. Most fundamentally, this perspective challenges the claim that complex multicellularity requires passage through a single-celled, haploid stage to constrain the emergence of lower-level, selfish genetic elements [66, 67]. This is true for nuclear genes in oocytes, whose quality is maintained by sexual exchange and recombination [67], but is not the case for mitochondria, which are transmitted uniparentally, without sexual exchange or recombination. In animals, the oocyte cytoplasm is not derived from a single cell, but instead requires the selective pooling of mitochondrial DNA from clusters of progenitor cells, which together generate high-quality mitochondria at extreme ploidy in mature gametes.
MATERIALS AND METHODS

Computational Model

1. Initial conditions

We use a computational model to follow the distribution of mitochondrial mutations in the female germline over a single generation from zygote to a new set of mature oocytes, as set out in the developmental history given in the main text (Fig. 1A). The initial state of the system is a zygote containing $M_0 = 2^{19} = 524,288$ copies of mtDNA, of which $m_0$ carry a deleterious mutation. Three specific models are considered: bottleneck, follicular atresia and cytoplasmic transfer. A list of terms and parameter values is given in Table 1, which also apply in the evolutionary model considered below.

| Parameters and variables                        | Symbols and values          |
|------------------------------------------------|----------------------------|
| Maximum number of germ cells                   | $N_{max} = 8,388,608$       |
| mtDNA number in mature oocytes                 | $M_0 = 2^{19} = 524,288$   |
| Minimum mtDNA ploidy                           | $B$                        |
| Final number of germ cells                     | $N_{max}/8 = 1,048,576$    |
| Initial mutation load                          | $m_0$                      |
| Mutation rate per bp per cell division         | $\mu$                      |
| Strength of epistatic interactions            | $\xi$                      |
| Transfer probability of mutant mtDNA           | $p_{mut}$                  |
| Transfer probability of wildtype mtDNA         | $p_{wt}$                   |
| Human mitochondrial genome size                | $g = 16,569\text{bp}$      |

Table 1
2. Early embryonic development

During early embryonic development, there is no mtDNA replication. The number of cells doubles at each time step. The existing population of mutant and wildtype mtDNA undergoes random segregation into daughter cells according to a binomial distribution – each mtDNA copy has a 50% probability of being assigned to either daughter cell. During this process, the average number of mtDNA copies per cell halves at each time step. There is no mutational input, as we only consider mutations that arise due to replication errors.

3. PGC proliferation, oogonia cell death and oocyte maturation

The early embryonic period lasts for the first 12 cell divisions. A group of 32 cells is selected at random to form the primordial germ cells (PGC). The PGCs then undergo proliferation for a further 18 rounds of cell division, until the maximum number of germ cells is reached, \( N_{\text{max}} = 32 \times 2^{18} = 8,388,608 \). 

mtDNA replication resumes after cell division 12, at the point of PGC determination. At this point, cells have an average of 128 mtDNA copies. At each following time step, the number of mtDNA copies doubles prior to random segregation into daughter cells. This means that the average number of mtDNA copies per cell is kept constant. New mutations are introduced as errors in mitochondrial replication. During the replication process, the new replica of each wildtype mtDNA copy has a probability of mutation \( \mu/\text{bp} \). The genome wide mutation rate \( U = g \times \mu \) is calculated as genome size \( (g = 16,569 \text{ bp} \text{[68]}) \) multiplied by \( \mu \). Given \( n \) wildtype and \( m \) mutant mtDNAs, the number of new mutants \( \Delta m \) resulting from replication errors is obtained by sampling at random from a binomial distribution with \( n \) trials with probability \( U \). After replication and mutation and prior to segregation the total
number of wildtype and mutant mtDNAs is $2n - \Delta m$ and $2m + \Delta m$ respectively. Back mutation to wildtype is not permitted.

At the end of PGC proliferation, the $N_{\text{max}}$ oogonia undergo random cell death, leaving $N_{\text{max}}/8 = 1,048,576$ primary oocytes. This is achieved by sampling the surviving cells at random with uniform weights (i.e., every cell has an equal probability of survival). The primary oocytes do not undergo further cell division or mitochondrial replication during the quiescent period (this is not explicitly modelled). At puberty, oocyte maturation commences. The number of mitochondria per cell is brought back to the original value $M_0 = 2^{19}$ through 12 rounds of replication without cell division. We assume that the number of mtDNA copies doubles at each time step. This introduces new deleterious mutations, which again are randomly drawn from a binomial distribution (as described above).

4. Specific models of selection

We consider three specific models in the main text with modifications to the base model described above.

The first model adds a bottleneck stage at the time of PGC determination (Fig. 2). As before, 32 cells are selected at cell division 12 to form the PGCs. These go through $b$ extra rounds of cell division without mtDNA replication. This reduces the mean number of mtDNA copies per cell to $\bar{B} = 128 \times (0.5)^b$. The mtDNA replication commences at cell division 14. The PGCs then proliferate as before to produce oogonia that undergo random cell death to produce primary oocytes. The primary oocytes have a reduced number of mtDNA copies, and so must undergo $12 + b$ extra rounds of mtDNA replication in order to regain the
original value $M_0$ mitochondria in mature oocytes. Note that this is an extreme model of the bottleneck, where mtDNA copy number is kept low throughout the period of PGC proliferation, and so maximises the benefit derived from the increase in segregational variation caused by the bottleneck.

For the model of the bottleneck, we allow selection dependent on individual fitness in relation to their mutation load $m$ among mature oocytes, according to the fitness function $f(m) = 1 - \left(\frac{m}{M}\right)^5$ (Fig. 2C). The concave shape of this function accounts for the fact that mitochondrial mutations typically have a detrimental effect on individual fitness only for loads $>60\%$. Changes to the power exponent make little qualitative difference to the outcome of this model (data not shown).

A second model considers non-random death during the cull of oogonia as these cells transition to being primary follicles (Fig. 3). Selection in this case is applied at the cell level. Cell fitness is expressed as $f(m) = 1 - \left(\frac{m}{M}\right)^\xi$, where $m$ is the number of mutant mitochondria. The parameter $\xi$ determines the strength of epistatic interactions (Fig. 3B).

As in other models, the number of cells is reduced from $N_{\text{max}} = 8,388,608$ to $N_{\text{max}}/8 = 1,048,576$. This is achieved by sampling without replacement the surviving cells at random, with weights proportional to cell fitness (i.e., every cell has a probability of survival proportional to its fitness).

The third model considers that the oogonia are organised in cysts of 8 cells each. These are the descendants of a single cell (i.e. three cell divisions prior). One cell is randomly designated as the primary oocyte using the MATLAB function randsample. The Balbiani body
of the primary oocyte contains a proportion $f$ of the mtDNA copies of all cells in the cyst.

The mitochondria that join the Balbiani body are sampled at random without replacement from each cell with different weights for wildtype ($p_{wt}$) and mutant ($p_{mut}$). After mitochondrial transfer to the Balbiani body, nurse cells undergo apoptosis (i.e. all cells except the one designated as the primary oocyte), reducing the total number of oocytes to $N_{max}/8 = 1,048,576$.

**Evolutionary Model**

In order to calculate the equilibrium distribution of a population undergoing the developmental dynamics mentioned in the previous section, we develop an analytical model for the distribution of mitochondrial mutations in an infinite population, with non-overlapping generations. As it was not possible to find an analytical solution, we solved the equations through numerical iterations. The system converges to a unique equilibrium state, independent of the initial conditions.

The state of the system is described by the vector $p(t) = \{p_0(t), ..., p_M(t)\}$, where $M(t)$ is the number of mtDNA copies per cell at time $t$. The elements $p_m(t)$ are the frequency of mutation load $m(t)/M(t)$ at time $t$. The evolution of the system is determined by a set of transition matrices whose elements are the transition probabilities between states. To avoid unnecessary complexity in the evolutionary model, we assume that fluctuations in mitochondrial number per cell due to segregation are negligible (i.e. in contrast to the computational model which allows binomial segregation at each division). Therefore, the mtDNA number per cell is constant across the whole population of cells at every time step. That is, during early embryonic development (when there is no mtDNA
replication), after $t$ cell divisions, the total number of mitochondria per cell is $M^{(t)} = 2^{-t} M_0$. Then, during PGC proliferation, the total number of mitochondria per cell is constant. Finally, during oocyte maturation, the number of mitochondria per cell exactly doubles with each mtDNA replication cycle. To aid in calculations, we also set the initial number of mtDNA copies to be proportional to the bottleneck size, i.e. $M_0 = 2^{12} \times B$. As the mtDNA number per cell halves at each cell division during early embryonic development, setting $M_0$ this way allows the mtDNA number to remain an integer. This is important for the modelling procedure, because the dimension of the transition matrixes (which is determined by the mtDNA number) must be an integer.

1. Early embryonic development

During early embryonic development, when mitochondrial replication is not active, changes in frequency arise purely from the process of segregation. Let $W^{(t)}$ be a $M^{(t)} + 1 \times M^{(t)} + 1$ square matrix, whose elements $W_{mn}$ represent the transition probabilities from a state with $m$ mutants to a state with $n$ mutants:

$$W_{mn}^{(t)} = \binom{m}{n} \left( \frac{M^{(t-1)} - m}{M^{(t-1)} - n} \right) / \binom{M^{(t-1)}}{M^{(t)}}$$

These matrix elements model the probability of transitioning from a state with $m$ mutants and $M - m$ wildtype to a state with $n$ mutants and $M - n$ wild type via the segregation of $2M$ mitochondria into two daughter cells with $M$ mitochondria each.
After $t$ cell divisions, the average number of mutants per cell is $\bar{m} = 2^{-t} m_0$, and the variance is $\text{Var}(t) = \frac{1}{4} [\text{Var}(t - 1) + 2^{-t} m_0]$. The state of the system is updated as $\vec{p}^{(1)} = (\Pi_t W^{(t)}) \times \vec{p}^{(0)}$.

2. PGC proliferation, oogonia cell death and oocyte maturation

During PGC proliferation, new mutations are introduced at a rate $U = \mu \times g$. The transition coefficient $Q_{mn}$ from a state with $m$ to a state with $n$ mutants results from the combined effects of replication, mutation and segregation:

$$Q_{mn} = \sum_k \binom{M - n}{k - n} U^{k-n} (1 - U)^{M-k} \binom{k}{m} \binom{M - k}{M - m} / \binom{2M}{M}$$

$$= \sum_k \binom{M - n}{k - n} U^{k-n} (1 - U)^{M-k} a_{k,m}$$

The coefficient $a_{k,m} = \binom{k}{m} \binom{M - k}{M - m} / \binom{2M}{M}$ models the probability of transitioning from a state with $k$ mutants and $M - k$ wildtype to a state with $n$ mutants and $M - n$ wild type via the segregation of $2M$ mitochondria into two daughter cells with $M$ mitochondria each; the remaining part of the equation models the probability of reaching a state with $k$ mutant mitochondria through replication and mutation of $M$ mitochondria, of which $m$ are mutant (this corresponds to the probability of introducing $k - m$ new mutations). The system is updated $\vec{p}^{(2)} = Q^q \times \vec{p}^{(1)}$, across $q$ rounds of PGC cell division. We then apply particular processes to capture the effects of the bottleneck, follicular atresia and cytoplasmic transfer.
As before, we model the bottleneck as $b$ extra rounds of segregation before the onset of mtDNA replication, following Eq(1) with $q + b$ cell divisions. This has no effect on the mean mutational number but increases mutational variance between the resulting PGCs. The transition between oogonia and primary oocytes occurs at random, and so does not alter the frequency distribution of mutants. Finally, during oocyte maturation, the mtDNA content of each cell doubles at every time step until the initial ploidy $M_0$ is restored. The transition matrix $G_{mn}$ is analogous to the first term of Eq(2), incorporating replication and mutation, but without segregation (last term of Eq(2)):

$$G_{mn}^{(t)} = \binom{M^{(t)} - m}{n - m} U^{n-m} (1 - U)^{M-k}$$  \hspace{1cm} (3)

Eq(3) models the probability of transitioning from a state with $m$ to a state with $n$ mutants, which is equivalent to the probability that exactly $n - m$ out of $M^{(t)} - m$ wildtype acquire a deleterious mutation. As the bottleneck reduces mtDNA copy number per cell, there is the need for $b$ extra rounds of replication of mtDNA during oocyte maturation. Hence, the transition coefficient $G$ is applied $b + 12$ times in the bottleneck model, to restore the number of mtDNA copies per oocytes to the original ploidy level $M_0$:

$$\bar{p}^{(3)} = (\prod_t G^{(t)}) \times \bar{p}^{(2)}.$$

At the end of the maturation phase, for the bottleneck model, selection is applied on individual fitness using a vector $w$ whose elements $w_m$ are equal to the corresponding
fitness: \( w_m = f(m) = 1 - \left( \frac{m}{M} \right)^5 \). This causes a change in the population mutation load as the system is updated to:

\[
\vec{p}^{(3)} = (I\vec{w}) \vec{p}^{(2)} / \vec{w}^T \vec{p}^{(2)}
\]

(4)

where \( I \) is the identity matrix.

In the model of follicular atresia, an extra step is included to reflect selection that operates when the population of oogonia are culled to produce the primary oocytes. This causes a change in the population mutation load analogous to that described in \textbf{Eq(4)}, but using the cell fitness function \( w_m = f(m) = 1 - \left( \frac{m}{M} \right)^\xi \) instead. This determines the shift in mutation loads that arises from fitness-dependent culling of oogonia. The transition coefficient \textbf{Eq(3)} for oocyte maturation is then applied 12 times in the follicular atresia model, to restore the original level of ploidy.

Finally, in order to model cytoplasmic transfer, a different process is used in the production of primary oocytes. A set of 8 clonally derived cells is selected. The mutation levels of each cell in the cyst is obtained by applying \textbf{Eq(3)} three times. Then, 50\% of the mitochondria in each cell are pooled into the Balbiani body of the primary oocyte. The probability for a cell with \( m \) mutants to contribute \( n \) mutants to the Balbiani body is given by:

\[
C_{nm} = \binom{m}{n} p_{\text{mut}}^n \left( \frac{M - m}{M/2 - \frac{n}{2}} \right) p_{\text{wt}}^{M/2 - n} / N
\]

(5)

Which gives the number of permutations of \( n \) mutant and \( M/2 - n \) wildtype mtDNA copies, weighted by the probability of transfer \( p_{\text{mut}} \) and \( p_t \) respectively, and divided by a
normalisation constant $N$. As the primary oocyte contains half of mitochondria from 8 cells, it needs to undergo 2 fewer rounds of replication during oocyte maturation. Hence only 10 rounds of replication following Eq(3) are carried out in this case to restore the original level of ploidy.

For all three models (bottleneck, follicular atresia and cytoplasmic transfer), the frequency distribution of mutation loads after these steps is used as the starting point for the next generation.

3. Evolutionary dynamics and model accuracy

The processes described above are iterated until the Kullback-Leibler divergence (a theoretical measure of how two probability distributions differ from each other [69]) between the new and the old distribution is smaller than a threshold $\eta = 10^{-9}$. We then assume that the system has reached a stationary state, e.g. without significant changes in the overall distribution of mutation loads between generations (mutation-selection balance).

In order to compare the prediction of the model with the clinical data, we use the equilibrium distribution to calculate the fraction of the population which carries a detectable load of mitochondrial mutations but does not manifest any detrimental phenotype ($\alpha_1$) and the fraction of individuals affected by mitochondrial disease ($\alpha_2$) using a threshold of 60% mutation load to discriminate between carrier and disease status. Individuals are assumed to be mutation free beyond the detection threshold of 2% [4].
The accuracy of the model is evaluated as the logarithm of the probability of reproducing clinical data by sampling the theoretical distribution at random. This is calculated as follows:

Let $X_1$ be the number of healthy individuals with detectable mutation load, and $X_2$ be the number of individuals affected by mitochondrial diseases; $N_1$ and $N_2$ the total number of individuals in the two trials; $\alpha_1$ and $\alpha_2$ the probability of observing, respectively, a healthy individual with detectable mutation load and an individual affected by mitochondrial disease, according to the prediction of the model. The log-likelihood of observing $X_1$ and $X_2$ by random sampling the theoretical distribution is given by

$$
\log (\lambda_{tot}(\mu, M, ...)) = \log \left[ \prod_{i=1}^{2} p(X_i | \alpha_i(\mu, M, ...)) \right] \\
= \sum_{i=1}^{2} \log \left[ \binom{N_i}{X_i} \alpha_i^{X_i}(1 - \alpha_i)^{N_i-X_i} \right] \\
= \sum_{i=1}^{2} \log \left( \binom{N_i}{X_i} \right) + N_i \alpha_i + (N_i - X_i)(1 - \alpha_i)
$$

(6)

**Estimation of the deleterious mutation rate**

The parameter values for the deleterious mutation rate we investigate reflect data collected from a number of species. Estimates of mtDNA point mutation rates in the crustacean *Daphnia pulex* range between $1.37 \times 10^{-7}$ and $2.28 \times 10^{-7}$ per site per generation [70].

Assuming this rate applies to humans and there are ~20 cell divisions before oocyte maturation, leads to a range between $0.68 \times 10^{-8}$ and $1.14 \times 10^{-8}$ per site, per cell division. Analysis of *Caenorhabditis elegans* mtDNA leads to a similar estimate of

$\sim 1.6 \times 10^{-7}$ per site, per generation [71], which corresponds to a rate of $0.8 \times 10^{-8}$ per
site, per cell division. For *Drosophila melanogaster*, the mtDNA mutation rate yields an estimate of $6.2 \times 10^{-8}$ per site, per generation, and hence $\sim 0.31 \times 10^{-8}$ per site, per cell division [72]. Finally, analysis of human mtDNA point mutation rates give a mutation rate of 0.0043 per genome per generation [62], corresponding to $\sim 1.3 \times 10^{-8}$ mutations per site, per cell division.

These values do not take into account the presence of a number of processes likely to remove mutants and is therefore a conservative estimate. The loss of mutations would mean that the actual mutation rate is higher than the estimates above. But unlike nuclear rates, the compact structure of mtDNA where intergenic sequences are absent or limited to a few bases, means that the rate of point mutations is probably not much higher than the rate of deleterious mutations. Therefore, for this study we consider a broad interval of possible deleterious mutation rates, ranging between $10^{-9}$ and $10^{-7}$. 
**FIGURES**

**Figure 1. Stages in female germline development.** (A) Timeline of oocyte development showing the main stages modelled. (B) Numerical simulation of the base model. Top panel: number of germ cells from specification of the 32 primordial germ cells (PGCs) after 12 cell divisions; proliferation to form 8 million oogonia; random cell death reducing to 1 million primary oocytes; quiescent period (not shown) and finally oocyte maturation at puberty. Middle panel: copy number of mitochondria (i.e. mtDNA); from zygote with ~500,000 copies, which are partitioned at cell division during early embryo development until replication begins (first vertical line) during PGC proliferation; copy number is amplified during oocyte maturation back to ~500,000 copies; dotted line shows the mean.
mitochondria copy number, with the distribution across oocytes shown in yellow. Note, skew reflects the log-scale. Bottom panel: mean (dotted line) and distribution of mutation load through development. The yellow shaded area shows the 90% quantile. Other parameter values $\mu = 10^{-8}$, $m_0 = 0.1$. 
Figure 2. Model of germline bottleneck and individual selection. (A) A bottleneck with two extra rounds of cell division without replication (cell division 13 and 14; first two vertical lines), reducing mitochondria copy number per PGC (by a quarter on average). Two extra
rounds of mitochondrial replication are required to regenerate the copy number in mature oocytes. Compared to the base model (Figure 1), mean mutation load (dotted blue line) is slightly higher and variation in load is substantially greater (yellow shaded area, 90% quantile). Parameter values $\mu = 10^{-8}$, $m_0 = 0.1$. (B) Violin plots of the distribution of mutations (mean ± SD shown in red) at two developmental stages, PGC specification and mature oocytes, given 5 mean bottleneck sizes ($\bar{B}$) when $m_0 = 0.1$. (C) Strength of selection on individual fitness, with a concave fitness function based on clinical data from mitochondrial diseases$^{27,28}$. (D) Change in mutation load ($\Delta m$) across a single generation for three initial mutation loads ($m_0$), given 5 mean bottleneck sizes ($\bar{B}$), showing the median (red line) and distribution (box plot IQR with min/max whiskers and outliers).
Figure 3. Model of follicular atresia and cell selection. (A) After PGC proliferation, follicular atresia occurs through selective apoptosis of oogonia. (B) Cell fitness is assumed to be linear ($\xi = 1$) or follow negative epistasis ($\xi = 2, 5$) in which mutations are more deleterious in combination. (C) Change in mutation load, $\Delta m$, across a single generation after cell selection, at an intermediate mutation rate ($\mu = 10^{-8}$), for individuals with low ($m_0 = 0.001$), medium ($m_0 = 0.01$) and high ($m_0 = 0.1$) initial mutation loads, for variable levels of epistasis (median (red line) and distribution (box plot IQR with min/max whiskers and outliers)).
Figure 4. Model of cytoplasmic transfer and mitochondria selection. (A) Cytoplasmic bridges form among oogonia in the germline cyst, leading to selective transfer of wild-type mitochondria (blue) to the primary oocyte, leaving mutant mitochondria (red) in nurse cells that then undergo apoptosis. (B) Cytoplasmic transfer which selectively pools $f = 50\%$ of...
mtDNA from 8 germline cyst cells into a single primary oocyte causes a large reduction in mean (dotted line) and distribution of mutation load (yellow shaded area shows the 90% quantile), which persists during oocyte maturation. Pooling of mtDNA requires two fewer rounds of mtDNA replication to regenerate copy number in mature oocytes. Parameter values $\mu = 10^{-8}$, $m_0 = 0.1$. 

(C) Change in mutation load ($\Delta m$) across a single generation (median (red line) and distribution (box plot IQR with min/max whiskers and outliers), for individuals with low ($m_0 = 0.001$), medium ($m_0 = 0.01$) and high ($m_0 = 0.1$) initial mutation loads, with variable strengths of selective transfer ($p_{mut}$). Parameter value $\mu = 10^{-8}$. 
**Figure 5.** Heatmaps showing log-likelihood of reproducing the observed mutation load and disease frequency in humans, for equilibrium conditions under the evolutionary model with **(A)** bottleneck and selection on individuals, **(B)** follicular atresia and selection on cells \(\xi = 5\), **(C)** cytoplasmic transfer with intermediate \(p_{mut} = 0.33, p_{wt} = 0.67\) or **(D)** strong \(p_{mut} = 0.25, p_{wt} = 0.75\) selective transfer of wildtype mitochondria. Yellow depicts high likelihood; blue, low likelihood. All models are shown for variable bottleneck size (the minimum mitochondria population size at which replication commences) and variable mutation rates.
Figure S1. Change in mutation load ($\Delta m$) across a single generation after individual selection with variable mean bottleneck size ($\bar{B}$). This is shown with (A) low ($\mu = 10^{-9}$) and (B) high ($\mu = 10^{-7}$) mutation rate, for individuals with low ($m_0 = 0.001$), medium ($m_0 = 0.01$) and high ($m_0 = 0.1$) initial mutation loads. Box plots show the median (red line) and distribution (box plot IQR with min/max whiskers and outliers).
Figure S2. Change in mutation load ($\Delta m$) across a single generation after cell selection with variable levels of epistasis ($\xi$). This is shown with (A) low ($\mu = 10^{-9}$) and (B) high ($\mu = 10^{-7}$) mutation rate, for individuals with low ($m_0 = 0.001$), medium ($m_0 = 0.01$) and high ($m_0 = 0.1$) initial mutation loads. Box plots show the median (red line) and distribution (box plot IQR with min/max whiskers and outliers).
**Figure S3.** Change in mutation load ($\Delta m$) across a single generation, given a variable proportion of transferred mitochondria ($f$) to the Balbiani body when transfer is non-selective ($p_{mut} = p_{wt} = 0.5$). This is shown with (A) low ($\mu = 10^{-9}$), (B), intermediate ($\mu = 10^{-8}$) and (C) high ($\mu = 10^{-7}$) mutation rate, for individuals with low ($m_0 = 0.001$), medium ($m_0 = 0.01$) and high ($m_0 = 0.1$) initial mutation loads. Box plots show the median (red line) and distribution (box plot IQR with min/max whiskers and outliers).
Figure S4. Change in mutation load ($\Delta m$) across a single generation, for individuals undergoing cytoplasmic transfer the Balbiani body with variable strength of selection, given a fixed probability of transfer of wildtype mitochondria ($p_{wt} = 0.5$) and a decreasing probability of transfer of mutant mitochondria ($p_{mut}$). Note the null case is when ($p_{mut} = p_{wt} = 0.5$). This is shown with (A) low ($\mu = 10^{-9}$) and (B) high ($\mu = 10^{-7}$) mutation rate, for individuals with low ($m_0 = 0.001$), medium ($m_0 = 0.01$) and high ($m_0 = 0.1$) initial mutation loads, and a fixed proportion of transferred mitochondria ($f = 0.5$). Box plots show the median (red line) and distribution (box plot IQR with min/max whiskers and outliers).
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