A Wide Range of 3243A>G/tRNALeu(UUR) (MELAS) Mutation Loads May Segregate in Offspring through the Female Germline Bottleneck

Francesco Pallotti1,2, Giorgio Binelli3, Raffaella Fabbri4,5, Maria L. Valentino6,7, Rossella Vicenti4,5, Maria Macciocca4,5, Sabina Cevoli6,7, Agostino Baruzzi6,7, Salvatore DiMauro1, Valerio Carelli6,7,*

1 Department of Neurology, Columbia University, New York City, New York, United States of America, 2 Dipartimento di Scienze Chirurgiche e Morfologiche, University of Insubria, Varese, Italy, 3 Dipartimento di Scienze Teoriche e Applicate, University of Insubria, Varese, Italy, 4 Unità Operativa di Ginecologia e Fisiopatologia della Riproduzione Umana, Ospedale S.Orsola-Malpighi, University of Bologna, Bologna, Italy, 5 Dipartimento di Scienze Mediche e Chirurgiche (DIMEC), University of Bologna, Bologna, Italy, 6 IRCCS Istituto delle Scienze Neurologiche di Bologna, Ospedale Bellaria, Bologna, Italy, 7 Dipartimento di Scienze Biomediche e Neuromotorie (DIBINEM), University of Bologna, Bologna, Italy

Abstract

Segregation of mutant mtDNA in human tissues and through the germline is debated, with no consensus about the nature and size of the bottleneck hypothesized to explain rapid generational shifts in mutant loads. We investigated two maternal lineages with an apparently different inheritance pattern of the same pathogenic mtDNA 3243A>G/tRNALeu(UUR) (MELAS) mutation. We collected blood cells, muscle biopsies, urinary epithelium and hair follicles from 20 individuals, as well as oocytes and an ovarian biopsy from one female mutation carrier, all belonging to the two maternal lineages to assess mutant mtDNA load, and calculated the theoretical germline bottleneck size (number of segregating units). We also evaluated “mother-to-offspring” segregations from the literature, for which heteroplasmy assessment was available in at least three siblings besides the proband. Our results showed that mutation load was prevalent in skeletal muscle and urinary epithelium, whereas in blood cells there was an inverse correlation with age, as previously reported. The histoenzymatic staining of the ovarian biopsy failed to show any cytochrome-c-oxidase defective oocyte. Analysis of four oocytes and one offspring from the same unaffected mother of the first family showed intermediate heteroplasmic mutant loads (10% to 75%), whereas very skewed loads of mutant mtDNA (0% or 81%) were detected in five offspring of another unaffected mother from the second family. Bottleneck size was 89 segregating units for the first mother and 84 for the second. This was remarkably close to 88, the number of “segregating units” in the “mother-to-offspring” segregations retrieved from literature. In conclusion, a wide range of mutant loads may be found in offspring tissues and oocytes, resulting from a similar theoretical bottleneck size.

Citation: Pallotti F, Binelli G, Fabbri R, Valentino ML, Vicenti R, et al. (2014) A Wide Range of 3243A>G/tRNALeu(UUR) (MELAS) Mutation Loads May Segregate in Offspring through the Female Germline Bottleneck. PLoS ONE 9(5): e96663. doi:10.1371/journal.pone.0096663

Editor: David C. Samuels, Vanderbilt University Medical Center, United States of America

Received October 24, 2013; Accepted April 10, 2014; Published May 7, 2014

Copyright: © 2014 Pallotti et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: This study was supported by FAR (Fondo di Ateneo per la Ricerca) from University of Bologna and from University of Insubria. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: VC is currently receiving funding for the following clinical trials: clinical trial with EPI-743 ([\(\alpha\)-tocotrienol quinone) in Leber’s hereditary optic neuropathy, Edison Pharmaceuticals, USA, and clinical trial with l-acetyl carnitine in Leber’s hereditary optic neuropathy, Sigma-tau, Italy. This does not alter the authors’ adherence to all PLOS ONE policies on sharing data and materials.

* E-mail: valerio.carelli@unibo.it

Introduction

Human mitochondrial DNA (mtDNA) is assumed to be a clonal multi-copy genome of 16,5 kb that is strictly maternally inherited. In each cell, mtDNA may be present either as identical copies (homoplasmy) or as a mixed population of two or more different sequences (heteroplasmy or polyploidy) [1]. Heteroplasmic mtDNA nucleotide changes, including those causing mitochondrial encephalomyopathies [2], segregate in tissues of the developing embryo as well as in germline cells. Somatic segregation of pathogenic mutations is relevant for clinical expression of mitochondrial diseases by affecting energy-dependent tissues that accumulate high, supra-threshold mutant loads [2,3]. Germline segregation is crucial for maternal transmission of variable mutant loads to the offspring [3].

Heteroplasmay may be theoretically due to coexistence of individual organelles containing either exclusively mutant or exclusively wild-type genomes (inter-mitochondrial heteroplasmy) or to the coexistence in each mitochondrion of both mutant and wild-type genomes in different proportions (intra-mitochondrial heteroplasmy) [4]. The mtDNA molecules are associated with specific coating proteins in discrete nucleoids, physically attached to the inner mitochondrial membrane [5], which may themselves be either homoplasmic or heteroplasmic [6]. Admixture and complementation of heteroplasmic mtDNA genomes may be accomplished by mitochondrial fusion events and exchange of mtDNA between nucleoids [7]. Variable efficiency in complementation has been observed in cellular models harboring different mtDNA mutations [8,9] but inter-mitochondrial complementation has been documented in a mito-mouse model carrying an mtDNA deletion [10]. Recent evidence suggests that
nucleoids do not exchange genetic material frequently and are probably homoplasmic [11,12], and may contain up to only one mtDNA molecule [13].

The load of mutant mtDNA may vary markedly between a mother and each of her children and a bottleneck mechanism has been postulated during the germline segregation of mutant mtDNA to explain rapid shifts of heteroplasmacy observed within one generation [14–16]. However, the nature of the bottleneck mechanism in humans is still under intense debate. Recent studies led to several potential mechanisms that are not necessarily mutually exclusive. These include: i) a marked reduction in the number of mtDNA molecules during the early stages of germline development [17]; ii) aggregation of identical segregating units without a reduction of mtDNA copy number, leading to rapid segregation due to sampling effect [18,19]; iii) preferential replication of a subpopulation of genomes, implying an active selection [20]; iv) rapid mtDNA segregation in preimplantation embryos [21]. Most of the data collected so far have been obtained by studying animal models segregating clusters of mtDNA polymorphic variants [22]. The experimental models provided by animals carrying pathogenic mtDNA mutations (“mutator” mouse) suggested a purifying selection for the most severe mtDNA mutations [23,24]. In humans, the bottleneck model has been tested only in a few studies, using both neutral polymorphisms and pathogenic mtDNA mutations segregating in relatively small pedigrees [25–30]. Apparently, different segregation patterns may operate depending on the mtDNA pathogenic mutation: the 8993TAG mutation associated with neuropathy, ataxia, retinitis pigmentosa (NARP) was characterized by skewed segregation in offspring or oocytes [27], whereas the 3243A>G/tRNALeu(UUR) mutation associated with mitochondrial encephalomyopathy, lactic acidosis, stroke-like episodes (MELAS) followed a random genetic drift model of segregation in a large sample of oocytes from a single woman [30].

We combined quantitative analysis of mtDNA heteroplasmy in both oocytes and somatic tissues to study the germline and somatic segregation of the 3243A>G/tRNALeu(UUR), pathogenic mutation [31] in two Italian pedigrees.

Materials and Methods

Patients

We studied two previously reported [32] Italian maternal lineages (Family A in Figure 1 and Family B in Figure 2) carrying the heteroplasmic 3243A>G/tRNALeu(UUR) mutation. Briefly, the proband from Family A (II-2, Figure 1) was affected with chronic progressive external ophthalmoplegia (CPEO), whereas the proband from Family B (II-4, Figure 2) had the typical MELAS syndrome. Both probands had ragged-red-fibers (RRF) and/or cytochrome c oxidase (COX)-negative fibers in skeletal muscle with different mutation loads.

In family A, the proband’s only daughter (III-2) was asymptomatic except for frequent migraine attacks and she lacked RRF in muscle biopsy. Her son (IV-1) has been treated with growth hormone for short stature.

In family B, the proband’s mother (I-2) was clinically asymptomatic, whereas the proband’s only son (III-3) recently developed the full-blown MELAS syndrome.

After approval by the internal review board (1996–1998, Institute of Neurological Clinic, University of Bologna, Director Prof. Elio Lugaresi) and signed informed consent, 20 maternally related individuals from both families agreed to be enrolled in the study aimed at assessing the MELAS mutation loads in somatic tissues. In most cases, we have been able to collect blood samples (leukocytes- and platelet-enriched pellets or whole blood), muscle biopsies, urinary epithelium, and hair follicles. The proband’s only daughter in Family A (III-2), also underwent ovarian stimulation to allow collection of oocytes for genetic analysis and gave informed consent for an ovarian biopsy at the time of oocytes collection. Moreover, two of the proband’s sisters in Family B (II-5 and II-7) became pregnant during the time of our investigation and consented to prenatal diagnosis on cells recovered from amniotic fluid.

mtDNA Analysis

Total DNA has been extracted by standard phenol/chloroform methods from somatic tissues, including amniocytes. Total oocyte DNA was recovered from single oocytes. Briefly, each oocyte was placed in an Eppendorf tube with one drop PBS and 1.5 μL proteinase K 10 mg/ml in ice, and centrifuged 30 sec 3500 rpm in an Eppendorf table-top centrifuge. After adding 50 μL sterile mineral oil, the mixture was centrifuged 30 sec as before, then digested at 37°C for one hour; digestion was blocked by boiling at 95°C for 15 minutes, followed by 80°C for 20 minutes. Each sample was then frozen and maintained at −80°C until the PCR amplification.

Heteroplasmy was determined by restriction fragment length polymorphism (RFLP) analysis after hot-last cycle PCR amplification as previously described [32]. The sensitivity of this method allowed detection of heteroplasmy as low as 1%.

Estimation of Bottleneck Size in Our Families and Review of Previous Reports

We assessed the bottleneck size in two germline segregations from unaffected females carriers of the MELAS mutation to their offspring or oocytes. For the first segregation (Family A; female III-2 in Figure 1), we were able to estimate the proportion $p$ of mutant mtDNA from the heteroplasmic load found in four primary oocytes collected from this woman and in the somatic tissues available from her only son. For the second segregation (Family B; female I-2 in Figure 2), the same estimate was obtained by averaging the loads of mutant mtDNA found in somatic tissues of five offspring. In both cases, $p$ was taken as an estimate of $\pi$, the true proportion in the sample population. Under the binomial distribution, the variance was estimated by $p(1-p)/n$, whereas confidence intervals for $p$ were estimated solving for the equation $z = (p-\pi)/\sqrt{(1-\pi)/n}$. The binomial distribution applies if the levels of mutant mtDNA are solely determined by a sampling error such as may occur during a bottleneck. Confidence intervals were used, in both pedigrees, to test whether the mutant load in a given progeny was compatible with a random sampling event (i.e. the bottleneck in the mother).

The number of “units” undergoing the bottleneck was estimated according to equation (1) in Brown et al. [30] under the assumption that 24 cell divisions are needed to produce the full set of primary oocytes. Each segregating “unit” could be an mtDNA molecule or a nucleoid. We also applied the same statistical approach to a set of “mother-to-offspring” segregations reported in the literature, updating the series reviewed by Chinnery and colleagues [39], and evaluating tissue heteroplasmy in families in which there were at least three siblings besides the proband [34–39].

Oocytes and Ovarian biopsy

The proband’s unaffected daughter in Family A (III-2) underwent surgical laparoscopy during which oocytes were retrieved from both ovaries and a biopsy was taken from the...
right ovary. The oocytes were obtained after ovarian stimulation using a combination of a gonadotrophin-releasing hormone analogue (Triptoreline, Decapeptyl 3.75; Ipsen Biotec, Paris, France) and menotrophins (Metrodin HP, 75 IU; Serono, Milan, Italy) and immediately frozen in liquid nitrogen for DNA analysis [40].

The ovarian biopsy specimen was frozen in liquid nitrogen-cooled isopentane for histological and histoenzymatic staining, following the standard procedure used for muscle biopsies [41]. Ten μM sections were processed for hematoxylin/eosin standard staining and cytochrome c oxidase/succinate dehydrogenase (COX/SDH) double histoenzymatic staining. One age-matched control ovarian biopsy was used for comparison.

**Results**

The heteroplasmic load of MELAS mutation assessed in various somatic tissues of maternally related individuals from Families A and B is summarized in Figures 1 and 2. The mutant mtDNA segregated only in some individuals along the maternal line of both families, as previously reported [32].

In Family A, the mutational event most likely occurred between individual I-2 and the CPEO proband (II-2 in Figure 1), considering that mutant mtDNA was absent in all other siblings.
of II-2, as well as in two maternal descendants in the third generation (individuals III-4 and III-5). We relied on the results in mtDNA from muscle and urinary epithelium, or at least one of the two tissues. The MELAS mutation was transmitted to the proband's daughter, individual III-2 and to her son (IV-1), both currently unaffected. The mutant load slowly increased through these three generations, as shown by all tissues tested. In all individuals, the mutant loads in urinary epithelium and skeletal muscle were remarkably similar, whereas in blood-derived cells they were inversely correlated with age, as reported by others [42–44].

Remarkably, we failed to detect any sign of COX deficiency, neither in the oocyte cytoplasm nor in the other cell types (i.e. granulosa cells of the ovarian follicle and other stromal cells). Figure 4 shows the RFLP analysis in the four oocytes from individual III-2, which revealed mutant loads ranging from 10% to 67%.

In Family B, the female founder (I-2 in Figure 2) showed mutant mtDNA in skeletal muscle and urinary epithelium. This woman segregated mutant mtDNA only in one of her offspring, the proband affected with MELAS (II-4). None of the proband's siblings had mutant mtDNA in any of the tissues investigated, nor did two maternal descendants in the third generation (III-1 and III-2). Furthermore, amniocytes collected during pregnancies of individuals II-5 and II-7 were also negative for the MELAS mutation (data not shown). Mutant mtDNA was transmitted from the proband to her only son, who is affected with MELAS like his mother. The tissue distribution pattern of somatic mutant loads was similar to that described for Family A, except that the female founder of this pedigree had undetectable mutant mtDNA in blood, only traces in urinary epithelium but a relatively high amount in skeletal muscle. Remarkably, this woman had had two...
miscarriages besides the five healthy offspring and the daughter with MELAS.

To investigate the “mother-to-offspring” germ line segregation of the MELAS mutation in these two maternal lineages (from individual III-2 in Family A and from individual I-2 in Family B), we estimated the percentage of mutant mtDNA in somatic tissues of each offspring in Families A and B, and in each oocyte in Family A. The germline segregation was compatible with a bottleneck event in the mother, according to the binomial distribution. Thus, different mutation loads in the progeny have to be ascribed to chance alone. The bottleneck size, based on the assumption that 24 cell divisions are needed to produce primary oocytes, consisted of 89 segregating units for Family A and 84 for Family B, if we consider only the mutant load in skeletal muscle (Table 1). If we take into account the mtDNA heteroplasmy of urinary epithelium in both Family A and B, the segregation units were 108 (Family A, oocytes from subject III-2 plus urinary epithelium from the only son) and 110 (Family B, urinary epithelium from all offspring) (Table 1).

We reviewed previously reported families segregating the MELAS mutation [33–39] and selected those in which the \( p \) of mutant mtDNA was reported for both mother and progeny and included, besides the proband, at least three siblings. We then subjected these “mother-to-offspring” segregations retrieved from the literature to the same test for the binomial distribution that we have used for the analysis of our Italian families. In all included cases (see Table 1) the \( p \) of mutant mtDNA in the progeny was compatible with a random segregation event in the mother. The number \( N \) of “segregating units” was in the range of 59–120, with an average number of \( N = 88 \) (confidence interval at the 0.95 level was 75 \( \leq N \leq 101 \)), remarkably close to the values estimated in our study, \( N = 89 \) for Family A and \( N = 84 \) for Family B. These segregations were calculated using different somatic tissues, such as

---

**Figure 3.** Ovarian follicles. A and C show two ovarian follicles (arrows) of individual III-2 (Family A), stained, respectively, with HE and COX/SDH; B and D, similarly, show three ovarian follicles (arrows), at different stages of maturation, of a control individual (magnification 20x). No evidence of reduced COX stain was observed in any of the tissues from the ovarian biopsy of the individual III-2, in particular the oocytes, as compared to the control (asterisks).

doi:10.1371/journal.pone.0096663.g003

---

**Figure 4.** Quantification of the 3243A>G/tRNALeu mutation loads in four primary oocytes from individual III-2 (Family A) and from five control oocytes.

doi:10.1371/journal.pone.0096663.g004
Table 1. Estimate of bottleneck sizes for the MELAS mutation from a review of the literature.

| Source | Patient's phenotype | Tissue analyzed | p | C.I. | N |
|--------|---------------------|-----------------|----|------|---|
| Liou et al (1994) | Headache, episodic vomiting | Hair follicles | 0.237 | 0.030–0.740 | 120 |
| De Vries et al (1994) | Unaffected | Fibroblasts | 0.282 | 0.065–0.698 | 108 |
| Huang et al (1996) | Unaffected | Skeletal muscle | 0.500 | 0.150–0.850 | 83 |
| Koga et al (2000) | Muscle weakness | Hair follicles | 0.548 | 0.175–0.873 | 84 |
| Dubeau et al (2000) | Deafness, cardio-myopathy, short stature | Urinary epithelium | 0.488 | 0.188–0.793 | 108 |
| Family A (this work) – Unaffected | Four oocytes | Oocytes | 0.423 | 0.113–0.808 | 89 |
| Family A (this work) – Unaffected | Four oocytes + one son | Oocytes | 0.488 | 0.113–0.808 | 110 |
| Family B (this work) – Unaffected | Five, one MELAS | Skeletal muscle | 0.115 | 0.014–0.551 | 110 |

p = frequency of mutant mtDNA in the progeny; C.I. = lower and upper limits of confidence interval for p at the 0.95 level; N = estimated size of the bottleneck.

Discussion

This study shows that germline segregation of the 3243A>G/tRNA^Leu^ MELAS mutation may lead to a wide range of mutational loads in offspring through a similar bottleneck size. Its estimation in the two Italian families here investigated was remarkably close to the average number of segregating units calculated for other “mother-to-offspring” germline segregations retrieved from the literature. In Family A, individual III-2 transmitted intermediate, largely distributed loads of heteroplasmic mutant mtDNA (10% to 75% mutant; Figure 5), as measured in four of her oocytes and in her only son. This resembled most of the other segregations retrieved from the literature (Figure 5). In contrast, in Family B we observed a sharply skewed transmission of mutant from individual I-2 to only one of her offspring (81% mutant; Figure 5). All other siblings had only wild-type mtDNA in the tissues analyzed (0% mutant; Figure 5), including amniocytes from two pregnancies of individuals II-5 and II-7. This was paralleled by only one family previously reported by Huang et al. [37], which had an essentially identical distribution of mutant loads in skeletal muscle of offspring (Figure 5).

The number of “segregating units” (bottleneck size), calculated in these two Italian families and in the several cases retrieved from the literature [28–30] was substantially lower than the 173 segregating units estimated by Brown et al. in the only study that sampled a large set of oocytes (N = 82) from a female carrier of the same MELAS mutation [30]. An important limitation of the current study and those retrieved from the literature is the large error associated with the variance estimated from a very low sample number (≥4) [45]. This is an obvious drawback by working with living patients from human pedigrees. A recent study [46] on the segregation of the MELAS mutation through the human embryofetal development concluded that random drift drives germline segregation, similar to Brown’s and colleagues conclusions [30], but with some appreciable individual-dependent differences in bottleneck size. Interestingly, in a study based on a large cohort of individuals carrying the MELAS mutation, the mothers with a mutation load greater that 50% tended to have offspring with lower or equal heteroplasmy, whereas the opposite was true for mothers with less than or equal to 50% mutation load [47]. These authors concluded that the random genetic drift model could not fully explain the transmission of the MELAS mutation [47]. Ascertainment bias has also to be considered. The recent finding that one in 200 healthy humans harbors a pathogenic mtDNA mutation out of the ten most frequent, indicates that there is a large pool of maternal lineages were probably these mutations segregate silently, and are possibly skeletal muscle, hair follicles, fibroblasts and urinary epithelium (Table 1).

Overall, these cumulative data show a close relationship between the tissues analyzed and the relative calculation for bottleneck size (N); for both a postmitotic tissue, such as skeletal muscle and oocytes, N resulted similar, despite the resulting mutation load in offspring was largely distributed in Family A and skewed to the extremes in Family B. Our literature revision revealed that in most cases the “mother-to-offspring” transmission resembled Family A [34,35,36,38,39], whereas only one family was essentially identical to Family B [37], still with very similar estimated bottleneck sizes. The overview of the relationship between mother and offspring mutant loads from our Families A and B, and those retrieved from literature are graphically represented in Figure 5, including the theoretical bottleneck calculated for each of these segregations.
selected out, missing to express any pathology and not being ascertained at all [48]. Thus, investigation of single pedigrees identified by an affected proband introduces a bias that may be resolved by pooling large cohort of families through multi-centric studies, or by meta-analyses of reported pedigrees.

The analysis of somatic segregation of the MELAS mutation in our two families confirmed that the mutant load is inversely correlated with age in blood cells, whereas skeletal muscle is the tissue of choice, followed by urinary epithelial cells, for detection of the mutation [42–44]. The pattern of mutational load in somatic tissues distinguished the two families, which also differed for the clinical phenotype. In Family A (CPEO), the mutational load in the unaffected female individual III-2 appeared to be similar in skeletal muscle (44%) and urinary epithelium (29%–46%), whereas in the female individual I-2 of Family B (MELAS) the mutational load in skeletal muscle (40%) was much higher than in urinary epithelium (1%). This latter observation might be related to the skewed transmission of mutant mtDNA in the offspring of this woman, resulting in one MELAS patient (81% mutant mtDNAs in skeletal muscle), two miscarriages conceivably due to very high mutant loads, and all remaining unaffected individuals with wild-type mtDNA.

Many recent studies have tried to tackle the issue of mtDNA germline segregation testing the bottleneck hypothesis [17–22]. These studies have employed murine and primate heteroplasmic models and there is no consensus on whether the bottleneck exists, whether there are one or more bottlenecks, and at what stage of development the bottleneck[s] operate. These models do not closely recapitulate the situation of a single mtDNA pathogenic mutation segregating along the maternal line of human pedigrees because most heteroplasmic animals were generated by mixing two mitochondrial genomes that differed for a cluster of polymorphic variants, which may have no or small functional relevance [49]. This condition is different from the case of a single pathogenic mtDNA mutation arising on a clonal mitochondrial genome, which is typical of humans with mitochondrial disorders. Important differences between the two situations may include the nucleoid composition and the level of mtDNA exchange, if any, between nucleoids. Nucleoids seem to follow the faithful replication model, without consistent genome exchange [6,11,13]. Furthermore, it has been demonstrated that mtDNA molecules may recombine within mitochondria [50–52], a phenomenon that is not relevant when mtDNA is clonal as in most humans, but that may become important in the case of different coexisting genomes with clusters of distinct variants, as in the heteroplasmic animal models or sometimes in humans with multiple heteroplasmies [17–21,53]. No studies address how frequently mtDNA recombination may occur, in which cell type, or during which stage of germ line segregation. Neither heteroplasmic animal models [17–21,49] nor the few available pathologic mito-mouse models [10,23,24] have been fully exploited yet to answer all these questions.

One final question concerns the possible selective pressure on mtDNA pathogenic mutations. The currently available mito-mice clearly indicated that severe mtDNA mutations undergo purifying selection over a few generations [23,24]. The segregation of the MELAS mutation in human tissues has been proposed to be non-random [54], and in vitro studies using cybrids with different nuclear backgrounds showed that segregation of the mutant mtDNA could be driven in opposite directions depending on the nuclear genome [55–57]. Thus, selection of mutant mtDNA may occur differently in different somatic tissues, impinging on the phenotypic expression. Whether such a genotypic selection may also operate during the germ line segregation for “mild” changes, including the MELAS mutation, is currently debated, casting doubts on the random genetic drift mechanism [47]. Staining the ovarian tissue for the histoenzymatic COX/SDH activities failed

---

**Figure 5.** Graphical representation of mother-to-offspring transmission of the MELAS mutation in the two Italian families and the seven other pedigrees retrieved from literature (see Table 1). The mutant load of MELAS mutation (%) is on the y axis. In all panels, the leftmost point is the mother’s mutant load, connected to each of the offspring mutant load. The reference, the tissues from which mtDNA mutant load has been assessed and the bottleneck size are indicated.

doi:10.1371/journal.pone.0096663.g005
to reveal any COX-deficient oocyte, nor other cell types. This may indicate that in this particular case there was no oocyte with suprathreshold loads of MELAS mutation or that a very efficient complementation occurs within oocytes, which may escape in the case of MELAS mutation any selection along the germline.

In conclusion, the mechanisms governing the germline segregation and the subsequent somatic distribution of single pathogenic mtDNA mutations in humans remain far from being elucidated. Our study of mother-to-oocytes/offspring tissues transmission of the same pathogenic MELAS mutation shows how wide may be the range of mutant loads segregating through the same bottleneck size.

References

1. Chinnery PF, Thorburn DR, Samuels DC, White SL, Dahl H-HM, et al. (2000) The inheritance of mitochondrial DNA heteroplasmy: random drift, selection or both? Trends Genet 16: 500–505.
2. Schon EA (2000) Mitochondrial genetics and disease. Trends Biochem Sci 25: 555–560.
3. Carling PJ, Cree LM, Chinnery PF (2011) The implications of mitochondrial DNA copy number regulation during oogenesis. Mitochondrion 11: 686–692.
4. Lightowler RN, Chinnery PF, Turnbull DM, Howell N (1997) Mammalian mitochondrial genetics: heterology, heteroplasmy and disease. Trends Genet 13: 450–455.
5. Sato M, Kuroiwa T (1991) Organization of multiple nucleoids and DNA molecules in mitochondria of a human cell. Exp Cell Res 196: 137–140.
6. Jacobs HT, Lehitten SK, Sperlhick JN (2000) No sex please, we’re mitochondria: a hypothesis on the somatic unit of inheritance of mammalian mtDNA. Bioessays 22: 564–572.
7. Legros F, Malka F, Frachon P, Lombes A, Rojo M (2004) Organization and dynamics of human mitochondrial DNA. J Cell Sci 117: 2653–2662.
8. Takai D, Inoue K, Goto YI, Nonaka I, Hayashi JI (1997) The interorganellar interaction between distinct human mitochondria with deletion mutant mtDNA from a patient with mitochondrial disease and with HeLa mtDNA. J Biol Chem 272: 6028–6033.
9. Enríquez JA, Cabezas-Herrera J, Bountour-Rafalou MP, Attali G (2000) Very rare complementation between mitochondria carrying different mitochondrial DNA mutations points to intrinsic genetic autonomy of the organelles in cultured human cells. J Cell Biol 275: 11207–11215.
10. Nakada K, Inoue K, Ogo K, Ogiura A, et al. (2001) Inter-mitochondrial complementation: Mitochondria-specific system preventing mice from expression of disease phenotypes by mutant mtDNA. Nat Med 7: 934–940.
11. Gilkerson RW, Schon EA, Hernandez E, Davidson MM (2000) Mitochondrial nucleoids maintain genetic autonomy but allow for functional complementation. J Cell Biol 151: 1117–1128.
12. Poe BG et al. Daffy CF, Greminger, MA, Nelson BJ, Arriaga EA (2010) Detection of heteroplasmy in individual mitochondrial particles. Anal Bioanal Chem 397: 239–247.
13. Nakada K, Wurm CA, Spa ˚hr H, Falkenberg M, Larsson NG, et al. (2011) Super-resolution microscopy reveals that mammalian mitochondrial nucleoids have a uniform size and frequently contain a single copy of mtDNA. Proc Natl Acad Sci U S A 108: 33534–33539.
14. Olivio P, Van De Walle MJ, Laips P, Hansohr WW (1983) Nucleotide sequence evidence for rapid genotypic shifts in the bovine mitochondrial DNA D-loop. Nature 306: 400–402.
15. Ashley MV, Laips P, Hansohr WW (1989) Rapid segregation of heteroplasmic bovine mitochondria. Nucleic Acids Res 17: 7325–7331.
16. Koehler CM, Lindberg GL, Brown DR, Beitz DC, Freeman AE, et al. (1991) Replacement of bovine mitochondrial DNA by a sequence variant within one D-loop heteroplasmy in mtDNA from tissues and single oocytes: support for a genetic bottleneck. Am J Hum Genet 60: 408–416.
17. Marchington DR, Macalay V, Hartshorne GM, Barlow D, Poulton J (1998) Evidence from human oocytes for a genetic bottleneck in an mtDNA disease. Am J Hum Genet 63: 769–775.
18. Brown DT, Samuels DC, Michael EM, Turnbull DM, Chinnery PF (2001) Random genetic drift determines the level of mutant mtDNA in human primary oocytes. Am J Hum Genet 69: 535–536.
19. Goto Y, Nonaka I, Horai S (1990) A mutation in the tRNA(Leu(UUR)) gene associated with the MELAS subgroup of mitochondrial encephalomyopathies. Nature 348: 631–633.
20. Cevoli S, Pallotti F, La Morgia C, Valentinii ML, Pierangeli G, et al. (2010) High frequency of migraine-only patients negative for the 3243 A>G rRNA mutation in two MELAS families. Cephalalgia 30: 919–927.
21. Chinnery PF, Howell N, Lightowler RN, Turnbull DM (1997) Molecular pathology of MELAS and MERRF: The relationship between mutation load and clinical phenotypes. Brain 120: 1713–1721.
22. Matrinuzzu A, Bartolonei L, Carrozzi R, Mostacciuolo M, Carboni C, et al. (1991) Correlation between clinical and molecular features in two MELAS families. J Neurol Sci 113: 222–229.
23. Liao CW, Huang CC, Chieh EC, Jeng YP, Tsai JL, et al. (1994) MELAS syndrome: correlation between clinical features and molecular genetic analysis. Acta Neurol Scand 93: 190–202.
24. Koga Y, Akiya T, Takane N, Sato Y, Kato H (2006) Heterogeneous presentation in a MELAS134G mutation in the mitochondrial tRNA-Leu(UUR) gene. Arch Dis Child 91: 402–407.
25. D’Alessandro F, De Stefano N, Zifkin BG, Arnold DL, Shoubridge EA (2000) Oxidative phosphorylation defect in the brains of carriers of the tRNA-Leu(UUR) gene. Annu Rev Med 51: 179–185.
26. Fabbrini R, Venturoli S, D’Errico A, Iannasci C, Gobbi E, et al. (2005) Ovarian tissue banking and fertility preservation in cancer patients: histological and immunohistochemical evaluation. Gynecol Oncol 89: 259–266.
27. Dubowitz V, Sevry CA, Lane RMJ (2007) Muscle Biopsy: A practical Approach (3rd Ed), Saunders Elsevier, London, UK.
28. Shanke S, Pancredu J, Kaufmann P, Engwall J, Juhann S, et al. (2004) Varying loads of the mitochondrial DNA A3243G mutation in different tissues: implications for diagnosis. Am J Med Genet 130A: 134–137.
29. McDonnell MT, Schaefer AM, Blakely EL, McFarland R, Chinnery PF, et al. (2004) Mitochondrial mutations in the mtDNA A3243G mutation using urinary epithelial cells. Eur J Hum Genet 12: 778–781.

Acknowledgments

We thank Prof. Eric Schon for supporting this project and for his helpful and constructive discussions. We are also indebted with the family members of the two Italian families investigated for their participation to this study.

Author Contributions

Conceived and designed the experiments: FP AB SDM VC. Performed the experiments: FP GB RF MLV RV MM SDM VC. Contributed reagents/materials/analysis tools: RF MLV SC. Wrote the paper: FP GB RF MLV VC.
44. Rahman S, Poulton J, Marchington D, Suomalainen A (2001) Decrease of 3243 A→G mtDNA mutation from blood in MELAS syndrome: a longitudinal study. Am J Hum Genet 68: 238–240.
45. Wonnapinij P, Chinnery PF, Samuels DC (2010) Previous estimates of mitochondrial DNA mutation level variance did not account for sampling error: comparing the mtDNA genetic bottleneck in mice and humans. Am J Hum Genet 86: 540–550.
46. Monnot S, Gigaret N, Samuels DC, Barlet P, Hesters L, et al. (2011) Segregation of mtDNA throughout human embryofetal development: m.3243A→G as a model system. Hum Mutat 32: 116–123.
47. Uusimaa J, Moilanen JS, Vainionpää L, Tapanainen P, Lindholm P, et al. (2007) Prevalence, segregation, and phenotype of the mitochondrial DNA 3243A→G mutation in children. Ann Neurol 62: 278–287.
48. Elliott HR, Samuels DC, Eden JA, Relton CL, Chinnery PF (2006) Pathogenic mitochondrial DNA mutations are common in the general population. Am J Hum Genet 83: 254–260.
49. Sharpley MS, Marciniaik C, Erkel-Mahan K, McManus M, Crimi M, et al. (2012) Heteroplasmy of mouse mtDNA is genetically unstable and results in altered behavior and cognition. Cell 151: 333–43.
50. Yoshida M, Miyazaki T, Attardi G (1994) Complementation of mutant and wild-type human mitochondrial DNAs coexisting since the mutation event and lack of complementation of DNAs introduced separately into a cell within distinct organelles. Mol Cell Biol 14: 2699–2712.
51. Kraytsberg Y, Schwartz M, Brown TA, Ehralishe K, Kunz WS, et al. (2004) Recombination of human mitochondrial DNA. Science 304: 981.
52. D’Aurelio M, Gaziewski CD, Lin MT, Maack WM, Shao LZ, et al. (2004). Heterologous mitochondrial DNA recombination in human cells. Hum Mol Genet 13: 3171–3179.
53. Zourka G, Hannel KG, Kudina T, Kornblum G, Kraytsberg Y, et al. (2007) Inheritance of mitochondrial DNA recombinants in double-heteroplasmic families: potential implications for phylogenetic analysis. Am J Hum Genet 80: 290–305.
54. Chinnery PF, Zwijnenburg P, Walker M, Howell N, Taylor RW, et al. (1999) Nonrandom tissue distribution of mutant mtDNA. Am J Med Genet 83: 498–501.
55. Yoshida M, Chomyn A, Martinuzzi A, Hurko O, Attardi G (1992) Marked replicative advantage of human mtDNA carrying a point mutation that causes the MELAS encephalomyopathy. Proc Natl Acad Sci U S A 89: 11164–11168.
56. Dunbar DR, Moonie PA, Jacobs HT, Holt JF (1995) Different cellular backgrounds confer a marked advantage to either mutant or wild-type mitochondrial genomes. Proc Natl Acad Sci U S A 92: 6562–6566.
57. Lehtinen SK, Hance N, El Meziane A, Juohla MK, Juohla KM, et al. (2000) Genotypic stability, segregation and selection in heteroplasmic human cell lines containing np 3243 mutant mtDNA. Genetics 154: 363–380.