## Nuclear DNA-encoded fragments of mitochondrial DNA (mtDNA) confound analysis of selection of mtDNA mutations in human primordial germ cells.

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Abstract: The resilience of the mitochondrial genome to a high mutational pressure depends, in part, on purifying selection against detrimental mutations in the germline. It is crucial to understand the mechanisms of this process. Recently, Floros et al. concluded that much of the purifying selection takes place during the proliferation of primordial germ cells (PGCs) because, according to their analysis, the synonymity of mutations in late PGCs was seemingly increased compared to those in early PGCs. We re-analyzed the Floros et al. mutational data and discovered a high proportion of sequence variants that are not true mutations, but originate from NUMTs, the latter of which are segments of mitochondrial DNA (mtDNA) inserted into nuclear DNA, up to millions of years ago. This is a well-known artifact in mtDNA mutational analysis. Removal of these artifacts from the Floros et al. dataset abolishes the reported effect of purifying selection in PGCs. We therefore conclude that the mechanism of germline selection of mtDNA mutations remains open for debate, and more research is needed to fully elucidate the timing and nature of this process.

### Introduction.

Mitochondrial DNA (mtDNA) is known for having a high mutational rate and a high density of crucial genes. Therefore, mtDNA mutations not only underlie a spectrum of mitochondrial diseases (Lander and Lodish, 1990), (Elliott et al., 2008) but also, if left unchecked, could result in long-term detrimental effects on species evolution (Popadin et al., 2007). It is thus vital to understand how mtDNA mutations are kept at a sustainable level through generations. A seminal study from 2008 demonstrated that significant levels of purifying selection against detrimental mutations occur in the germline (Stewart et al., 2008). Mutations are depleted of nonsynonymous variants as early as the second generation after mutations are generated de novo in a mtDNA 'mutator' mouse line. However, the mechanism(s) and precise timing of germline purifying selection remain unknown. More recently, Floros and colleagues published a study of mtDNA mutations in human embryonic primordial germ cells (PGCs), which asserted that purifying selection acted at an early stage in the developing germline during PGC proliferation and migration (Floros et al., 2018). This assertion was based primarily on an observation of a sharp increase of average synonymity of mtDNA mutations in late-stage PGCs compared to early-stage PGCs.

## **Results and Discussion**.

# *Comparison of mutations to known NUMTs reveals significant contamination with NUMT sequences.*

Careful review of the Floros et al. study, revealed that their key result, which was a relative increase in synonymity in late-stage PGCs, was unusual in that it appeared to result not from a depletion of nonsynonymous mutations – as would be expected under negative purifying selection- but instead from an increase in the frequency of synonymous mutations. Further examination of their primary mutational data shown in Table S5 of their supplementary information unexpectedly revealed that the majority (~80%) of sequence variants were found in many unrelated samples. Such recurrence is generally not expected of true nascent mutations, which are individually rare events



**Figure 1.** NUMT contamination in PGC mtDNA mutational dataset: spatial distribution and alignment of sequence variants detected in PGCs .

A. Frequency distribution of Floros et al. variants along the mitochondrial genome. The unexpected dense patch of variants (between ~8kb and 10Kb, dashed vertical blue lines) corresponds to one of the PCR fragments used in Floros et al. (bp:8187-10165, horizontal blue double arrow). The PCR primers of this fragment happen to co-amplify a portion of the nuclear pseudogene of mtDNA (NUMT) on Ch5 (hg38(100045938-100055045), red double arrow).

**B**. A representative section of alignment of reference mtDNA sequence with Ch5 NUMT sequence shows that a great majority of Floros et al. variants in this genome region are identical to the NUMT-derived variants, which corroborates their NUMT origin. 'Floros all muts': a mtDNA sequence where we incorporated **all** variants from Table S3 of Floros et al., including those that are under the 1% cutoff. 'Floros muts >1%': Table S3 variants of Floros et al. were limited to those above the 1% cutoff. Of 10 variants located in the sections of the genomes shown here, one (9506C>T, blue down arrow) exceeded the threshold and thus was erroneously considered a confirmed mutation. This illustrates the mechanism of 'leakage' of NUMT-driven variants into the pool of 'confirmed mutations'.

and therefore have negligible probability to occur simultaneously and independently in multiple samples. We reasoned that this inter-sample repetition of low fraction sequence variants may actually be due to nuclear DNA of mitochondrial origin (NUMT) contamination, a long-known problem of mtDNA mutational analysis (Hu and Thilly, 1994), (Khrapko et al., 1994), (Hirano et al., 1997). NUMTs are nuclear pseudogenes of mtDNA, many of which are millions of years old and have accumulated many sequence differences from the true mitochondrial genome. Our alignment of Floros et al. variants to several known NUMT sequences indeed revealed at least 2 contributing NUMTs. In fact, a single one of these NUMTs located on chromosome 5 was responsible for ~20% of protein coding variants reported by Floros et al. as real mutations. We further found that NUMT contamination was most likely introduced by co-amplification of the NUMT sequences by the PCR approach used in their study (**Figure 1**). Interestingly, we have previously studied this NUMT and its homologs in great apes and demonstrated that it

reveals distant interspecies hybridization event in the evolutionary history of hominids (Popadin et al., 2017).

To further corroborate the NUMT-based explanation of the Floros et al. findings, we searched the raw sequencing data of Floros et al. for typical NUMT-derived reads, which are traditionally recognized by the presence of multiple nucleotide changes within short sequence stretches (Hirano et al., 1997), (Khrapko et al., 1994). This feature reflects the fact that highly diverged ancient NUMT sequences carry a high density of sequence changes (compared to contemporary human mitochondrial genome). For example, the NUMT at chromosome 5, which shows as a part of contamination in Floros study, contains patches of ~10 changes per 100 nucleotides. To determine if the presumably NUMT-derived sequence changes are arranged as characteristic patches, the primary reads were extracted from the Floros et al. dataset. We then performed multiple alignment of the raw read sequences against the sequence of the NUMT at human chromosome 5 (See **Methods** for details). As expected, this analysis revealed a significant proportion of sequence reads characteristic of NUMTs, with patches of multiple mutations (**Figure 2**). Collectively, these results undoubtedly imply that NUMT-derived variants represent a substantial part of the Floros et al. mutation dataset.

There remain a few sequence variants that are repeated in several samples, for which we were not able to identify a known NUMT origin. These variants are likely derived from other, more recent, NUMTs, the presence of which in human nuclear genomes have been confirmed by the authors as well as other groups (Wei et al., 2020), (Lutz-Bonengel et al., 2021). Alternatively, variants repeated in unrelated samples could also be derived from a number of other sources, ranging from contamination by external mtDNA to 'hotspots' associated with instrumental error. In any case, we feel that repetitive variants must be considered errors, unless independent evidence is presented that these events are natural mutational hotspots. Even if the latter proof is obtained, however, such 'hotspot' mutations are of limited use for germline selection analysis, as the dynamics of these mutations is usually strongly affected by the process of mutagenesis itself, which confuses analysis of selection.

## Contamination with NUMT-derived sequence variants affects the estimates of the germline selection.

We next explored if contamination with NUMT-derived sequence variants had significantly affected the key conclusions of Floros et al. study. By comparing mutant fractions of recurrent sequence variants in different samples, we found that NUMT-derived variants are present at systematically and significantly different frequencies. Most of these variants are below the 1% cutoff adopted by Floros et al. and thus are not included in the analysis. However, in some samples, NUMT-derived variants occasionally exceed 1% and thus were deemed 'true' mtDNA mutations. Importantly, a large majority of NUMT-derived variants that exceeded the 1% threshold were found in one particular late-stage PGC sample.

It may look surprising that NUMT-derived contamination has different levels in different samples, because as a nuclear sequence, the NUMTs copy number per cell is expected to be constant. However, the actual level of *NUMT contamination* depends on the nuclear-to-mtDNA ratio, and the copy number of mtDNA can significantly differ between samples of the same tissue type, as was demonstrated in the paper describing NUMT contamination in brain samples (Hirano et al., 1997). These differences are related to the variability of mtDNA levels and/or differences in DNA integrity between samples.

We note that NUMT-derived variants, which are in fact differences between mtDNA and the corresponding NUMT sequences, are expected to be (and are)

predominantly highly synonymous. This inference may look surprising because NUMTs are essentially pseudogenes of mtDNA, and mutations in pseudogenes are conventionally thought of as highly nonsynonymous. Explanation for this surprising fact is that from the evolution perspective, a majority of NUMT variants are "reverse mtDNA" mutations". Indeed, on the evolutionary time scale, NUMT sequences, which reside in the 'protective' nuclear DNA environment, mutate at a much slower rate than the corresponding sequences in the far less protected mitochondrial genome. So a great majority of the differences between NUMTs and the contemporary mtDNA sequence are in fact mutations in *mtDNA* (rather than NUMT) that were fixed in the species' mitochondrial genome since insertion of the NUMT into the nuclear genome. Thus, a majority of NUMT/mtDNA differences are historic mtDNA mutations, which ought to be highly synonymous because they are mostly fixed in the population by random drift, and thus need to be fairly neutral to keep selection at zero to allow this to happen. Irrespectively of this theoretical explanation, synonymity of mtDNA mutations is evident from phylogenic analysis (e.g. (Popadin et al., 2017)). We have demonstrated this peculiar feature of NUMTs in humans (Popadin et al., 2017), and this has been demonstrated previously by others in bristletails (Baldo et al., 2011).

Returning to Floros et al., because the most NUMT-rich sample happened to be a late-stage PGC sample, this single sample brought a large number of synonymous NUMT-derived variants into the late-stage PGC mutation pool. This explains the puzzling surge of synonymous variants in late-stage PGCs. More importantly, this resulted in Floros et al. reaching an experimentally unsubstantiated conclusion of negative selection in PGCs. We then asked if removal of the erroneous repetitive variants could yield a 'clean' mtDNA mutational dataset that would allow for a more accurate assessment of the dynamics of true mtDNA mutations during human PGC development. Unfortunately, once these repetitive variants were removed, there remain only a few protein-coding mtDNA mutations in PGCs, which are not sufficient in number to make statistically significant inferences.

In summary, our results indicate that contamination of the Floros et al. mtDNA mutational data with NUMT sequences created bias in favor of their assertion of negative selection during PGC proliferation and migration. When these contaminating variants are removed, the data become statistically underpowered for analysis. We therefore conclude that the mechanism and timing of germline selection of mtDNA mutations requires more research to be fully elucidated.

#### Methods.

### Synonymity and repeatability analysis of sequence variants of mtDNA.

The data pertaining to the Floros et al. study has been imported from the supplemental table 5 at <a href="https://doi.org/10.1038/s41556-017-0017-8">https://doi.org/10.1038/s41556-017-0017-8</a>, (Floros et al., 2018). Synonymity was analyzed using in house excel-based spreadsheets based on lookup function in excel. Identical mutations in different samples were identified using 'countif' functions.

### Extraction and multiple alignment of raw reads from the NGS dataset.

We extracted raw read NGS data from NCBI Sequence Read Archive record SRR6288291. We used TrimGalore software to remove adapters and trim low quality reads. We filtered out 3.7% of bases from raw data due to low quality, removed 0.1% of

sequences, which become shorter than 20bp. Adapters were removed from 29.4% of reads. Next, we aligned reads against two references - chr5 NUMT reference and mtDNA reference with Smith-Waterman alignment algorithm implemented in bwa utility (bwasw). For mtDNA alignment we filtered out alignment reads leaving in only reads a between positions 8750 and 10250 of mtDNA, in order to isolate region, which is homologous to NUMT at chromosome 5 (hg38(ch5100045938-100055045). For each read, we derived quality measure from CIGAR as a percentage of M symbols - we calculate this from SAM aligned files and do this procedure independently for each read in both mtDNA and NUMT alignment. Next we stratified reads out as aligned onto both references with quality measures greater than 0.9 for both NUMT and MTDNA references, NUMT -aligned reads as reads with mtDNA quality less than 0.8 and NUMT quality greater than 0.99, and mtDNA-aligned reads with mtDNA quality greater than 0.99 and NUMT quality less than 0.8. Last, we convert each set of reads from fastq into fasta format, leaving only nucleotide strings before BLAST operation.

*Multiple alignment of raw reads* was performed using the NCBI BLAST server with default parameters, with the sequence of the human NUMT at chromosome 5 (hg38(ch5100045938-100055045) as a query.

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