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1	The First complete Zoroastrian-Parsi Mitochondria Reference Genome: Implications of
2	mitochondrial signatures in an endogamous, non-smoking population
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24 Abstract:

25 The present-day Zoroastrian-Parsis have roots in ancient pastoralist migrations from circumpolar regions¹ leading to their settlement on the Eurasian Steppes² and later, as Indo Iranians in the 26 Fertile Crescent³. From then, the Achaemenids (550 - 331 BC), and later the Sassanids (224 BC -27 642 AD) established the mighty Persian Empires². The Arab invasion of Persia in 642 AD 28 necessitated the migration of Zoroastrians from Pars to India where they settled as Parsis and 29 30 practiced their faith, Zoroastrianism. Endogamy became a dogma, and the community has maintained the practice since their arrival in India. Fire is the medium of worship⁴ as it is 31 32 considered pure and sacrosanct; Social ostracism practiced against smokers resulted in a nonsmoking community, thus forming a unique basis for our study. 33

34 In order to gain a clearer understanding of the historically recorded migration of the Zoroastrian-35 Parsis, decipher their phylogenetic relationships and understand disease association to their 36 individual mitochondrial genomes, we generated the first complete de novo Zoroastrian-Parsi Mitochondrial Reference Genome, AGENOME-ZPMS-HV2a-1. Phylogenetic analysis of 37 38 additional 100 Parsi mitochondrial genome sequences, showed their distribution into 7 major 39 haplogroups and 25 sub-haplogroups and a largely Persian origin for the Parsi community. We 40 have generated individual reference genomes for each major haplogroup and assembled the 41 Zoroastrian Parsi Mitochondrial Consensus Genome (AGENOME-ZPMCG V1.0) for the first

42 time in the world.

We report 420 variants, specifically 12 unique mitochondrial variants in the 100 mitochondrial
 genome sequences compared with the revised Cambridge Reference Sequence (rCRS) standard.

45 Disease association mapping showed 217 unique variants linked to longevity and 41 longevity 46 associated disease phenotypes across most haplogroups. Our results indicate none of the variants

- 47 are linked to lung cancer. Mutational signatures, C>A, G>T transitions³⁶, linked to tobacco
- 48 carcinogens were found at extremely low frequencies in the Zoroastrian-Parsi cohort.
- Our analysis of gene-coding, tRNA and the D-Loop regions revealed haplogroup specific disease
 associations for Parkinson's, Alzheimer's, Cancers, and Rare diseases.

51 These disease signatures investigated in the backdrop of generations of endogamy, in the rapidly

52 declining, endangered Zoroastrian-Parsi community of India, provides exceptional universal

53 opportunity to understand and mitigate disease.

54 Keywords: Mitochondria, Haplotypes, Phylogeny, Human migration, Endogamous, Non-55 smoking, Longevity, Cancers, Neurodegenerative disorders, Rare Diseases, t-RNA, D-loop 56 variants, Population genetics, Unique mitochondrial variants, Zoroastrian Parsis, Persia, Iran, 57 India, Precision healthcare.

- 58 Abbreviations: mt DNA-Mitochondrial DNA ; rCRS-revised Cambridge Reference Sequence ;
- 59 NGS Next Generation Sequencing ; ZPMS- Zoroastrian Parsi Mitochondrial Sequence ; ZPMRG-
- 60 Zoroastrian Parsi Mitochondrial Reference Genome ; ZPMCG- Zoroastrian Parsi Mitochondrial
- 61 Consensus Genome ; AD- Alzheimers Disease ; PD- Parkinsons Disease
- 62

63 Introduction :

64

65 The Burden of History- Travelogue of the Zoroastrian Mitochondrion

- Zoroastrian-Parsis of India are followers of the ancient prophet Zarathushtra, claimed by the Greek
 historian Herodotus to have been born circa 6,450 BC¹. Zarathushtra, advocated the first known
- historian Herodotus to have been born circa $6,450 \text{ BC}^1$. Zarathushtra, advocated the first know monotheistic concept of one supreme intelligence termed Ahura Mazda - 'Majestic Creator'².
- 69 The ancient homeland of the present-day Zoroastrian-Parsis finds mention in their sacred Avestan
- 70 text *Vendidad*, and the location indicated is the North Polar Arctic region³. Sanskrit scholar B G
- 71 Tilak's study Arctic Home in the Vedas is also corroborated by Bennet, suggesting that the Indo-
- 72 European culture originated in the Hyperborean Regions of Northern Siberia and the islands of the
- 73 circumpolar regions⁴.
- Around 12000 years ago, this region suffered a natural calamity and became ice-clad¹ necessitating
- southward migrations of these pastoralist inhabitants, and by 4,000 BC the Indo Europeans took
- 76 over the Eurasian Steppe⁷.

77 From the late second to early first millennium BC, the Indo Europeans, mostly on the basis of

religious worship, split with the Indo Aryans who moved further south and crossed the Hindu

79 Kush, while the Indo Iranians (Medes, Persians, and Parthians) began populating the western

80 portion of the Iranian plateau, close to the Alborz and Zagros Mountains and northern

81 Mesopotamia to Southeast Anatolia, in what is called the Fertile Crescent where significant

82 innovation in agriculture occurred⁸.





84 Image of *Taq-e Bostan* means "Arch of the Garden" or "Arch made by stone", a site with a series of large rock reliefs from the era of the
 85 Sassanid Empire of Persia (Iran), carved around the 4th century CE. Image courtesy of Irandestination.com

86 In 550 BC, Cyrus the Great overthrew the leading Median rule and conquered the Kingdom of

87 Lydia and the Babylonian Empire after which he established the Persian Zoroastrian Achaemenid

88 Empire (the First Persian Empire), while his successors Dariush I (522-485 BC) Xerxes I,

89 Artaxerxes and others extended its borders to encompass several countries across three continents,

90 namely Europe, Africa and Asia. A second Zoroastrian dynasty of Sassanian Kings followed, who

91 ruled Persia starting with Ardashir 1 (224 BC). It was the Golden age of the Persian Empire. During

92 the time of Zoroastrian Achaemenid and Sassanid empires, Persia became a global hub of culture,

93 religion, science, art, gender equality, and technology.

94 The Persians under Yezdezard III were defeated by the Arabs in two decisive battles - (Qadisiyah-

95 636 AD and Nahavand – 642 AD) resulting in the fall of the Zoroastrian Persian Empire.

96 It was almost a hundred years later in the 8th century that a few boatloads of Zoroastrians left Paars

97 and Khorasan from the port of Hormuz to sail south towards India. The boats first touched shore

98 on Diu island on the west coast of India where the refugees stayed for around 19 years. The

99 environment being non-conducive to progress, they once again set sail and arrived in Sanjan,

100 Gujarat. Vijayaditya of the Chalukya dynasty (aka Jadi Rana) the ruler, hesitated to give refuge,

101 but on being explained the principles of Zoroastrianism and observing the similarities with the

102 Vedic religion, the Parsis were given refuge.



103



Map showing early migration of Parsis from Iran. Image courtesy Microsoft Encarta. Reference Library 2005

Endogamy became the norm to preserve their identity, and for the last 1300 years the community has maintained this practice^{9,10}. Fire being the purest of all elements is considered sacred by Zoroastrian-Parsis. Strict measures are employed to maintain the purity of fire, hence the strict social ostracism practiced against smokers in the community.

Today, the Zoroastrian-Parsis, are a small community of <52000 in India (2011 Census, Govt of India). We present the genetic data of the conserved Zoarastrian-Parsi mitochondrion, encapsulated in resilience of thousands of years of magnificent history: of struggles and overcoming them; of building something out of nothingness; of achievement gained with ethical standards; and philanthropy.

114 In recent decades, the analysis of the variability of maternally inherited mitochondrial DNA (mtDNA) has been commonly used to reconstruct the history of ethnic groups, especially with 115 respect to maternal inheritance. The lack of genetic recombination in mtDNA, results in the 116 117 accumulation of maternally inherited single nucleotide polymorphisms (variants). The accumulation of Variants along maternally inherited lineages results in phylogenetically traceable 118 haplotypes¹⁵ which can be used to follow maternal genealogies both historically and 119 120 geographically. This approach has provided insightful findings into the origins and disease etiologies associated with another well documented endogamous European community: the 121 122 Icelandic people (rev in 13).

123

Human mtDNA (mitochondrial DNA) is a double stranded, circular (16,569 kb) genome of bacterial origin¹⁶ primarily encoding vital subunits of the energy generating oxidative phosphorylation and electron transport chain (ETC) pathway that generates Adenosine Tri-Phosphate (ATP), the primary energy substrate of the eukaryotic cell. In addition, 22 tRNAs and 2 rRNAs are also encoded by the mtDNA¹⁷. 129 In this study, our first aim was to gain a clear understanding and genetic impact of the historically

- 130 recorded migration of the Zoroastrian-Parsis from Persia to India, and to link socio-cultural,
- 131 ritualistic practices followed within the community over several millenia manifesting in genetic
- 132 outcomes. To shed further light on the impact of migrations followed by integration into
- 133 communities, where ritual and social practices are strictly followed within communities and
- 134 between communities resulting in specific traceable signatures.
- 135

Secondly, we have attempted to elucidate the genetic basis of commonly occurring diseases in this endogamous community. To address these issues, we generated the first complete *de novo* Zoroastrian-Parsi Mitochondrial Genome (AGENOME-ZPMS-HV2a-1) and used it to arrive at the mitochondrial haplotype specific Reference Genomes from a hundred Zoroastrian-Parsi individuals. Our study for the first time, has assembled the Zoroastrian Parsi Mitochondrial Consensus Genome (AGENOME-ZPMCG V 1.0) thereby creating the first Mitochondrial Consensus Genome for the Zoroastrian-Parsi community.

143

Our phylogenetic analysis confirmed that present day Zoroastrian-Parsis are closely related to Persians, and like most endogamous communities have comparatively lower genetic diversity and tend to be predisposed to several inherited genetic disorders^{5,11}. They also possess longevity as a trait and are a long-living community⁶ with lower incidences of lung cancer¹². The study of the genealogic history of a close-knit community like the Parsis provides an unique opportunity, to understand the link between disease and social behaviour, thus providing the direction for population genetics as a basis for personalized healthcare.

151

152 Materials and Methods

153 Sample collection and ethics statement

154 One hundred healthy nonsmoking Parsi volunteers residing in the cities of Hyderabad-155 Secunderabad and Bangalore, India were invited to attend blood collection camps at the 156 Zoroastrian centers in their respective cities under the auspices of The Avestagenome ProjectTM. Each adult participant (>18 years) underwent height and weight measurements and answered an 157 extensive questionnaire designed to capture their medical, dietary and life history. All subjects 158 159 provided written informed consent for the collection of samples and subsequent analysis. All health-related data collected from the cohort questionnaire were secured in The Avestagenome 160 ProjectTM database to ensure data privacy. All procedures performed in this study involving human 161 participants were in accordance with the ethical standards of the institution (Avesthagen Limited, 162 163 Bangalore, India) and in line with the 1964 Helsinki declaration and its later amendments. This

- 164 study has been approved by the Avesthagen Ethics Committee (BLAG-CSP-033).
- 165

166 Genomic DNA extraction

167 Genomic DNA from the buffy coat of peripheral blood was extracted using the Qiagen Whole

168 Blood and Tissue Genomic DNA Extraction kit (cat. #69504). Extracted DNA samples were

assessed for quality using the Agilent Tape Station and quantified using the QubitTM dsDNA BR

170 Assay kit (cat. #Q32850) with the Qubit 2.0[®] fluorometer (Life TechnologiesTM). Purified DNA

171 was subjected to both long-read (Nanopore GridION-X5 sequencer, Oxford Nanopore

172 Technologies, Oxford, UK) and short-read (Illumina sequencer) for sequencing.

173

174 Library preparation for sequencing on the Nanopore platform

Libraries of long reads from genomic DNA were generated using standard protocols from Oxford
Nanopore Technology (ONT) using the SQK-LSK109 ligation sequencing kit. Briefly, 1.5 μg of
high-molecular-weight genomic DNA was subjected to end repair using the NEBNext Ultra II End
Repair kit (NEB, cat. #E7445) and purified using 1x AmPure beads (Beckman Coulter Life
Sciences, cat. #A63880). Sequencing adaptors were ligated using NEB Quick T4 DNA ligase (cat.
#M0202S) and purified using 0.6x AmPure beads. The final libraries were eluted in 15 μl of elution

181 buffer. Sequencing was performed on a GridION X5 sequencer (Oxford Nanopore Technologies,

- 182 Oxford, UK) using a SpotON R9.4 flow cell (FLO-MIN106) in a 48-hr sequencing protocol.
 183 Nanopore raw reads (fast5 format) were base called (fastq5 format) using Guppy v2.3.4 software.
- 184 Samples were run on two flow cells and generated a dataset of ~14 GB.
- 185

186 Library preparation and sequencing on the Illumina platform

Genomic DNA samples were quantified using the Qubit fluorometer. For each sample, 100 ng of
DNA was fragmented to an average size of 350 bp by ultrasonication (Covaris ME220

- 189 ultrasonicator). DNA sequencing libraries were prepared using dual-index adapters with the
- 190 TruSeq Nano DNA Library Prep kit (Illumina) as per the manufacturer's protocol. The amplified
- 191 libraries were checked on Tape Station (Agilent Technologies), quantified by real-time PCR using
- the KAPA Library Quantification kit (Roche) with the QuantStudio-7flex Real-Time PCR system
- 193 (Thermo). Equimolar pools of sequencing libraries were sequenced using S4 flow cells in a
- 194 Novaseq 6000 sequencer (Illumina) to generate 2 x 150-bp sequencing reads for 30x genome 195 coverage per sample.
- 195 196

197 Generation of the *de novo* Parsi mitochondrial genome (AGENOME-ZPMS-HV2a-1)

a) Retrieval of mitochondrial reads from whole-genome sequencing (WGS) data:

199 A total of 16 GB of raw data (.fasta) was generated from a GridION-X5 Nanopore sequencer for

- 200 AGENOME-ZPMS-HV2a-1 from WGS. About 320 million paired-end raw reads were generated
- 201 for AGENOME-ZPMS-HV2a-1 by Illumina sequencing.
- 202 Long Nanopore reads (. fastaq5) were generated from the GridION-X5 samples. The high-quality
- 203 reads were filtered (PHRED score =>20) and trimmed for adapters using Porechop (v0.2.3). The
- high-quality reads were then aligned to the human mitochondrial reference (rCRS) NC_12920.1
- 205 using Minimap2 software. The aligned SAM file was then converted to a BAM file using
- 206 SAMtools. The paired aligned reads from the BAM file were extracted using Picard tools (v1.102).
- 207

208 The short Illumina high-quality reads were filtered (PHRED score =>30). The adapters were

- trimmed using Trimgalore (v0.4.4) for both forward and reverse reads, respectively. The filtered
- 210 reads were then aligned against a human mitochondrial reference (rCRS²¹) using the Bowtie2
- 211 (v2.2.5) aligner with default parameters. The mapped SAM file was converted to a BAM file using
- 212 SAMtools, and the mapped paired reads were extracted using Picard tools (v1.102).
- 213
- b) *De novo* mitochondrial genome assembly
- Mapped reads were used for *de novo* hybrid assembly using the Maryland Super-Read Celera Assembler (MaSuRCA-3.2.8) tool. The configuration file from the MaSuRCA tool was edited by adding appropriate Illumina and Nanopore read files. The MaSuRCA tool uses a hybrid approach that has the computational efficiency of the de Bruijn graph methods and the flexibility of overlap-
- 219 based assembly strategies. It significantly improves assemblies when the original data are
- augmented with long reads. AGENOME-ZPMS-HV2a-1 was generated by realigning the mapped
- 221 mitochondrial reads from Illumina as well as Nanopore data with the initial assembly.
- 222

223 Confirmation of Variants in the *de novo* Parsi mitochondrial genome using Sanger 224 sequencing

- To validate the *de novo* Parsi mitochondrial sequence, AGENOME-ZPMS-HV2a-1, selected variants were identified and subjected to PCR amplification. Genomic DNA (20 ng) was PCR amplified using LongAmpTaq 2X master mix (NEB). The PCR amplicons of select regions were subjected to Sanger sequencing and BLAST analysis to confirm the presence of eight Variants using primers listed in Supplemental Table 1.
- 230

233

243

Generation of the Zoroastrian Parsi mitochondrial consensus genome (AGENOME ZPMCG-V1.0) and Parsi haplogroup-specific consensus sequences

a) Retrieving mitochondrial reads from 100 Parsi whole-genome sequences

234 The whole-genome data from 100 Parsi samples were processed for quality assessment. The 235 adapters were removed using the Trimgalore 0.4.4 tool for paired end reads (R1 and R2), and sites 236 with PHRED scores less than 30 and reads shorter than 20 bp in length were removed. The processed Illumina reads were aligned against a human mitochondrial reference sequence (rCRS¹⁸, 237 238 NC_012920.1) using the Bowtie 2 (version 2.4.1) aligner with default parameters. Mapped reads 239 were further used for the *de novo* assembly using SPAdes (version 3.11.1) and Velvet and IVA 240 (version 1.0.8). Comparison of the assembly and statistics were obtained using Quast (version 241 5.0.2). The assembled scaffolds were subjected to BLASTn against the NCBI non-redundant 242 nucleotide database for validation.

b) Variant calling and haplogroup classification

Sequencing reads were mapped to the human mitochondrial genome (rCRS²¹) assembly using the
 MEM algorithm of the Burrows–Wheeler aligner (version 0.7.17-r1188) with default parameters.

- 246 Variants were called using SAMtools (version 1.3.1) to transpose the mapped data in a sorted
- 247 BAM file and calculate the Bayesian prior probability. Next, Bcftools (version 1.10.2) was used

- to calculate the prior probability distribution to obtain the actual genotype of the variants detected.
- 249 The classification and haplogroup assignment were performed for each of the 100 Parsi mtDNAs
- after variant calling and after mapping reference and alternate alleles to the standard haplogroups
- 251 obtained from MITOMAP (**Appendix 4**).

252 c) Haplogroup-based consensus sequence

Ninety-seven of 100 full-length Parsi mtDNA sequences were segregated based on haplogroups and separately aligned using the MUSCLE program to obtain the multiple sequence alignments. The Zoroastrian Parsi Mitochondrial Reference Genome (ZPMRG) and the Parsi haplogroupspecific consensus sequences were generated after calculation of the ATGC base frequency by comparison of the nucleotides in an alignment column to all other nucleotides in the same column called for other samples at the same position. The highest frequency (%) was taken to build seven

- 259 Parsi haplogroup ZPMRGs and the seven Parsi haplogroup-specific consensus sequences.
- 260

261 **Phylogeny build and analysis**

262 Ninety-seven of 100 full-length Parsi mtDNA sequences generated as described above were compared with 100 randomly chosen Indian mtDNA sequences derived from NCBI Genbank 263 under the accession codes FJ383174.1-FJ 383814.1²², DQ246811.1-DQ246833.1²³, and 264 KY824818.1-KY825084.1²⁴ and from previously published data on 352 complete Iranian mtDNA 265 sequences²⁵. All mtDNA sequences were aligned using MUSCLE software²⁶ using the "maxiters" 266 2" and "diags 1" options, followed by manual verification using BioEdit (version 7.0.0). Following 267 alignment, the neighbor-joining method, implemented in MEGAX²⁷, was employed to reconstruct 268 269 the haplotype-based phylogeny. The neighbor-joining method was used because it is more efficient 270 for large data sets²⁸.

271

272 Variant disease analysis

One hundred Parsi mitochondria sequences extracted from the WGS were uploaded into the VarDiG[®]-R search engine (https://vardigrviz.genomatics.life/vardig-r-viz/) on AmazonWeb Services. VarDiG[®]-R, developed by Genomatics Private Ltd, connects variants, disease, and genes in the human genome. Currently, the VarDiG[®]-R knowledgebase contains manually curated information on 330,000+ variants, >20 K genes covering >4500 phenotypes, including nuclear and mitochondrial regions for 150,000+ published articles from 388+ journals. Variants obtained from Parsi mitochondria were mapped against all the published variants in VarDiG[®]-R. Associations

- 280 with putative diseases was ascertained for each variant through VarDIG[®]-R.
- 281

282 Seventeen tRNA SNP sites were identified in the 100 Parsi mitochondrial SNP data. The PON-283 mt-tRNA database⁴² was downloaded to annotate the tRNA Variants for their impact and disease 284 associations. This database employs a posterior probability-based method for classification of 285 mitochondrial tRNA variations. PON-mt-tRNA integrates the machine learning-based probability 286 of pathogenicity and the evidence-based likelihood of pathogenicity to predict the posterior

of pathogenicity and the evidence-based likelihood of pathogenicity to predict the posterior

probability of pathogenicity. In the absence of evidence, it classifies the variations based on themachine learning-based probability of pathogenicity.

289

For annotation of disease pathways associated with Variants, we employed MitImpact (https://mitimpact.css-mendel.it/) to predict the functional impact of the nonsynonymous Variants on their pathogenicity. This database is a collection of nonsynonymous mitochondrial Variants and their functional impact according to various databases, including SIFT, Polyphen, Clinvar, Mutationtester, dbSNP, APOGEE, and others. The disease associations, functional classifications, and engagement in different pathways were determined using the DAVID and UNIPROT annotation tools.

297

298 Haplogroup and disease linkage

Principal component analysis (PCA) was performed to visualize the linkage of the haplogroup with disease. XLSTAT (Addinsoft 2020, New York, USA. https://www.xlstat.com) was used for statistical and data analysis, including PCA.

302

303 Data Accessibility:

The GenBank (http://www.ncbi.nlm.nih.gov/genbank) accession numbers for the 105 novel complete mtDNA sequences (97 ZPMS, 7 ZPMRG and 1 ZPMCG) reported in this paper are MT506242-MT506346. The raw reads for 97 ZPMS mitochondrial genome sequences have been deposited with BioProject ID: PRJNA636291. The SRA accession numbers for the 97 ZMPS: SRR11888826-SRR11888922.

309

310 **Results**

Assembly of the first complete Zoroastrian Parsi mitochondrial sequence, AGENOME ZPMS-HV2a-1

- 313 The first complete *de novo* non-smoking Zoroastrian Parsi mitochondrial sequence, AGENOME-
- 314 ZPMS-HV2a-1, was assembled from a healthy Parsi female sample by combining the sequence
- 315 data generated from two next-generation sequencing (NGS) platforms using a protocol, as outlined
- 316 in Materials and Methods. Our approach combines the sequencing depth and accuracy of short-
- 317 read technology (Illumina) with the coverage of long-read technology (Nanopore). QC parameters
- 318 for mitochondrial reads, mitochondrial coverage, and X-coverage were found to be optimal, as
- 319 seen in Supplementary Figure 1. The hybrid Parsi mitochondrial genome was assembled as a
- 320 single contig of 16.6 kb (with 99.82% sequence identity), resulting in the consensus sequence for
- 321 the *de novo* Parsi mitochondrial genome with 99.84% sequence identity to the revised Cambridge
- 322 Reference Sequence (rCRS²¹).
- 323

324 Identification of 28 unique Variants in AGENOME-ZPMS-HV2a-1

325 The variants identified from both the Illumina and Nanopore data were considered to be significant

- 326 for this de novo Zoroastrian Parsi mitochondrial genome, henceforth referred to as AGENOME-
- 327 ZPMS-HV2a-1.

328

- A total of 28 significant variants (i.e., variants) were identified by BLAST alignment between the Parsi mitochondrial hybrid assembly and the rCRS²¹ (**Figure 1, Table 1**). To confirm the authenticity of the identified variants, we selected a total of 7 identified variants from the D-loop region and one SNP from the *COI* gene (m.C7028T, A375A) and subjected them to Sanger sequencing using primers. All 8 predicted variants were verified and confirmed for their presence in the consensus Parsi mitochondrial genome (**Figure 2**).
- 335
- The majority (n=11) of the variants identified in the AGENOME-ZPMS-HV2a-1 were found in the hypervariable regions (HVRI and HVRII) of the D-loop. Of the remaining 17 variants, eight were found to represent synonymous variants, while four were in genes for 12S, 16S-rRNA (n=3) and tRNA (n=1) (**Figure 1**). The remaining 5 nonsynonymous variants were located in the genes for *ATPase6* (m8860G>A), *COIII* (m.9336 A>G), *ND4* (m.11016 G>A), and two in the *CytB* gene (m15326 A>G and m15792 T>C, (**Table 1**). Except for the *ATPase6* gene variant, which has been found to be associated with hypertrophic cardiomyopathy in Iranian individuals²⁹, no associations
- were found in the published literature for these gene variants, and they need to be further investigated.
- 345

Given that the Zoroastrian Parsis are known to have originated in Persia and have practiced endogamy since their arrival on the Indian subcontinent, we wished to determine the mitochondrial haplogroup associated with the first complete Zoroastrian Parsi mitochondrial genome. We therefore compared the variants associated with ZPMS-HV2a-1 to standard haplogroups obtained from MITOMAP and determined the haplogroup to be HV2a (**Figure 1**). This haplogroup is known to have originated in Iran²⁵, suggesting Persian origins for this Parsi individual, based on maternal inheritance patterns.

353

354 Seven major haplogroups identified in 100 Zoroastrian Parsi individuals

355 Keeping in mind the endogamous nature of the Indian Parsis and to understand the extent of the 356 diversity of the mitochondrial haplogroups in this population, we analyzed mitochondrial genomes from 100 consenting Parsi individuals. Our study had an equal representation of both genders, and 357 358 60% of the subjects were of age 30–59 (mean age 50 ± 1.6) (Figure 3). Complete analysis of the 359 variants in the 100 Parsi samples identified a total of 420 unique Variants (Figure 4, Appendix 360 1). OC analysis of the 100 mitochondrial genomes sequenced were found to be optimal: 361 PHRED>30 (Supplementary Figure 2). Variant distribution in the coding region normalized to 362 gene length showed the *ND6* gene has the highest number of variants (Supplementary Figure 3). 363 The 100 Zoroastrian Parsi mitochondrial genomes were subjected to haplogroup analysis using 364 haplogroup specific variant assignment matrix from MITOMAP (Appendix 4). The haplogroup assignment based on variants classified the genomes into seven principal haplogroups (HV, U, T, 365 366 M, A, F, and Z) and 25 sub-haplogroups were also identified within the principal haplogroups 367 (Figure 5). The variant count across all sub-haplogroups ranged between 14-64 (Figure 6A).

Analysis of the sub-haplogroups demonstrated that HV2a was the single largest representative subhaplogroup within the Parsi population (n=14, n=9 females, n=5 males, (**Figure 6B**), that includes

- 370 the AGENOME-ZPMS-HV2a-1.
- 371

372 The sub-haplogroup HV2a (n=14 subjects) contained 28 variants observed in the AGENOME-373 ZPMS-HV2a-1 are common across all 14 subjects. In total, the HV2a sub-haplogroup had 38 374 variants, with the highest number in the HVR II region (n=8). Coding region mutations constituted 375 20/38 variants, with equal distribution between synonymous (n=10) and non-synonymous 376 substitutions observed for this sub-haplogroup (n=10). Among the coding regions, the largest 377 number of Variants was found in the gene encoding *COI* (n=6, **Supplementary Figure 4A**). Four 378 *COI* Variants distributed across all of the 14 subjects in the HV2a sub-haplogroup (m.6104 C>T, 379 m.6179 G>A, m.7028 C>T, and m.7193 T>C) constitute synonymous mutations (amino acid 380 change: F67F, M92M, A375A, and F430F, respectively). Two Variants (m.7080 T>C and m.7146 381 A>G), found to occur in one subject each in the sub-haplogroup HV2a, were nonsynonymous 382 substitutions (F393L and T415A, respectively). Further analysis of rare Variants (occurring only 383 in single subjects or n<8/14) showed their presence in the 16S-RNR2 gene (m.1883 G>A and 384 m.1888 G>A), as well as the COII, COIII (m.8203 C>T and m.9540 T>C), and HVR I (m.16153 385 G>A and 16274 G>A) genes, which were synonymous substitutions in these coding genes, while 386 we found nonsynonymous substitutions in the COII (m.7650 C>T; T22I), ND5 (m.12358 C>T; 387 T8A), and CYTB (m.14954 A>G; T70A) genes in our analysis. We found a variant in the gene 388 encoding for tRNA[R] at m.10410 T>C (n=14 subjects), but no mutations were observed in the D-389 loop region for the entire group under analysis.

390

391 The sub-haplogroup HV12b (n=1 subject) contained 17 Variants. HVR II harbors four Variants, 392 while the coding genes together contain six Variants that encode three synonymous and three 393 nonsynonymous substitutions. We observed Variants encoding nonsynonymous substitutions in 394 this sub-haplogroup in ATPase6 (m.8860 A>G; T112A), ND5 (m.13889 G>A; C518Y), and CYTB 395 (m.15326 A>G; T194A). Three Variants were found in 12S-RNR1, two Variants in 16S-RNR2. 396 In the non-coding regions 5 variants were observed in HVRII, 1 in HVR I and 1 in the D-loop 397 region (m.16519 T>C). No Variants were observed in the genes coding for tRNAs in the HV12b 398 sub-haplogroup.

399

400 The 21 subjects analyzed that fell into the U haplogroup consisted of four sub-haplogroups U1a 401 (n=1), U4b (n=11), U2e (n=3), and U7a (n=6). The U1a sub-haplogroup contained 44 Variants 402 distributed across 19 positions in the mitochondrial genome. Twenty-one Variants were observed 403 in the coding region (17 synonymous, 4 nonsynonymous). ND5, containing a coding region, 404 contains six Variants, the most for any position within the U1a haplogroup. All ND5 Variants 405 coded for synonymous substitutions, while nonsynonymous substitutions were observed for ND2 406 (m.4659 G>A; A64T), ATPase6 (m.8860 A>G; T112A), and CYTB (m.14766 C>T; T7I and 407 m.15326 A>G; A190T). 21/44 variants fell within coding genes, while the rest were distributed

408 across HVR I (n=4 Variants), HVR II (n=3 Variants), HVR III (n=5 Variants), 12S-RNR1 (n=2
409 Variants), 16S-RNR2 (n=4 Variants), the D-loop region (n=1 SNP), and control regions (n=2
410 Variants). Two Variants were found in regions coding for tRNA[D] and tRNA[L:CUN].

411

412 The U4b sub-haplogroup is the most common sub-haplogroup among the U haplogroup in our 413 analysis. In all, 64 Variants were observed for the U4b sub-haplogroup, with most of the variants 414 (n=20) found in the gene encoding 16S-RNR2 (Supplementary Figure 4B). Twenty-one Variants 415 were found in coding regions (14 synonymous and 7 nonsynonymous substitutions), with the 416 highest number seen in the gene coding for COI (n=6 Variants). Five of six Variants coded for 417 synonymous substitutions, while m.6366 G>A coded for a nonsynonymous substitution (V155I). 418 Three Variants were found in the gene encoding CYTB and were distributed across all subjects 419 (n=11) in the U4b sub-haplogroup. All three encoded nonsynonymous substitutions, m14766 C>T 420 (T7I), m.15326 A>G (T194A), and m.15693 T>C (M316T), and need to be further investigated. 421 Four tRNA mutations were observed in this sub-haplogroup and one mutation in the D-loop region.

422

A total of 52 variants were observed across all samples in the U7a subgroup (Supplementary
Figure 4B). Twenty-seven Variants were found in noncoding regions, 12S-RNR1, 16S-RNR2,
and the D-loop region. Twenty-five Variants were found in the coding region (17 synonymous and
8 nonsynonymous substitutions), with 17/25 distributed among the ND genes coding for *ND1–6*. *ND5* (n=6 Variants) encodes five synonymous mutations, with a nonsynonymous mutation
observed at m.14110 T>C (F592L, in 4/6 subjects).

429

430 A total of 55 Variants was observed for U2e, with the majority (n=33 Variants) falling in the 431 noncoding regions (HVRI-III and D-loop) and the 12S-RNR1, 16S-RNR2, and tRNA genes. 432 Twenty-two Variants fell within the coding region (15 synonymous and 7 nonsynonymous 433 substitutions), of which 8 fell in the ND gene complex (four ND2, four ND5) and four in the CYTB 434 gene. While all the Variants in the ND2 and ND4 genes are synonymous substitutions, all the 435 Variants in the CYTB gene encoded nonsynonymous mutations (m.14766 C>T; T7I in 3/3 subjects, 436 m.15326 A>G; T194A in 3/3 subjects; m.14831 G>A; A29T and m.15479 C>T; F245L, both in 437 1/3 subjects).

438

439 Five subjects in our analysis (n=100) fell within the T haplogroup. We found four sub-haplogroups 440 within this haplogroup (T1a, 2 subjects: T2b, T2i, and T2g, with 1 subject each). Our analysis 441 indicated a total of 39 Variants (Supplementary Figure 4C) for T1a, with 21/39 Variants found 442 in noncoding regions, including 12S-rRNA, 16S-rRNA, tRNAs, and control regions, including the 443 D-loop. Eighteen Variants were observed in the coding region, with the greatest number occurring 444 in the *CYTB* gene (n=5 Variants). Three Variants within the *CYTB* gene coded for nonsynonymous 445 mutations, including m.14776 C>T, m.14905 G>A, and m.15452 C>A, coding for T7I, T194A, 446 and L236I substitutions, respectively.

448 The T2b, T2g, and T2i sub-haplogroups contained 35, 42, and 34 Variants, respectively, in total. 449 We found that CYTB contained the majority of the Variants found in the coding regions in these 450 sub-haplogroups, except for the T2i group in which the CYTB Variants (n=5) constituted the 451 majority of the Variants found in coding and noncoding regions of the genome. Two Variants, 452 m.14766 C>T and m.15326 A>G, seen in all three groups code for nonsynonymous substitutions, 453 and m.15452 C>A was seen in T2g and T2i and codes for a nonsynonymous mutation. Single 454 mutations were seen for m.15497 G>A and m.14798 T>C and code for nonsynonymous 455 substitutions and need further investigation.

456

457 The A haplogroup in our study consists of the sub-haplogroup A2v (n=3 subjects). The subjects in 458 the A2v sub-haplogroup had a total of 17 Variants (Supplementary Figure 4D) distributed across 459 the mitochondrial genome. Twelve of seventeen Variants were found in the noncoding regions 460 (HVR I, II) and in the 12S rRNA and 16S rRNA genes. Five Variants were distributed in the 461 coding region across ND2 (m.4769 A>G and m.6095 A>G), ATPase6 (m.8860 A>G), ND4 462 (m.11881 C>T), and CYTB (m.15326 A>G). Two nonsynonymous substitutions were observed in 463 the ATPase6 and CYTB genes that need further investigation.

464

465 F1g (n=1 subject) is a sub-haplogroup, along with Z1a (n=1 subject). A total of 33 and 32 Variants, 466 respectively, were identified in these groups. Nine CYTB Variants were observed in total for both 467 groups. Two encoded nonsynonymous substitutions, m.14766 C>T (T7I) and m.15326 A>G 468 (T194A), while the seven other Variants resulted in synonymous mutations. Variants for ND4L 469 are seen only across Z1a and F1g, with the m.10609 T>C SNP in F1g resulting in a 470 nonsynonymous shift (M47T), while the Z1a SNP resulted in a synonymous substitution 471 (Supplementary Figure 4D).

472

473 The M haplogroup (n=52 subjects) consists of 12 sub-haplogroups, the most number for a 474 haplogroup in our study (Supplementary Figure 4E). M30d is the sub-haplogroups with the 475 highest number of subjects in the M haplogroup (n=11 subjects). Fifty-one Variants were identified 476 in this sub-haplogroup in total, of which 28 Variants were seen in the noncoding regions (HVR I, 477 II, III), the D-loop region, and the 12S-RNR1 and 16S-RNR2 genes. The remaining 23 Variants 478 were part of the coding region within CYTB (n=8 Variants) and ND4 (n=5 Variants) and formed a 479 majority. Nine of thirteen Variants in CYTB and ND4 code for synonymous substitutions, while 480 four Variants in CYTB resulted in nonsynonymous substitutions (m.14766 C>T; T7I, m.15218 481 A>G; T158A, m.15326 G>A; T194A, and m.15420 G>A; A229T).

482

483 M39b (n=10 subjects) is one of the largest sub-haplogroups, and a total of 59 Variants were seen 484 for this sub-haplolgroup. The noncoding regions, 12S, 16S, and control regions, together constitute

485 33/59 of the Variants. Of the remaining 26 Variants, the 5 Variants in the CYTB complex constitute

2 *ND4*, 3 *ND5*, and 2 *ND6*). Of the nine remaining Variants, six are seen in the *COI*, *II*, and *III*genes (two each), while three Variants are found in the *ATPase6* gene.

489

490 The M2 sub-haplogroup consists of M2a (n=2 subjects) and M2b (n=1 subject). A total of 110 491 Variants was observed in total for M2a and M2b (Supplementary Figure 4E). In M2a, 23/53 492 Variants occurred in noncoding regions (HVR I, II, III), the 12S-RNR1 and 16S-RNR2 genes, the 493 control region (OL), and the D-loop region. Thirty Variants occurred in the coding regions, making 494 this one of the sub-haplogroups in which Variants in the coding region outnumber the Variants in 495 the noncoding region. CYTB harbors seven Variants, followed by three Variants in ND4 and three 496 Variants in ATPase8, ATPase6, and COI. A total of 55 Variants was observed for M2b, in which 497 31/55 Variants occurred in the noncoding regions. Twenty-four Variants were observed in genes 498 coding for COI, III; ND1,2,3,4,5; ATPase6,8; and CYTB. The six Variants in CYTB constitute the 499 greatest number of Variants in the coding region. The M2a/b sub-haplogroup is also conspicuous 500 by the presence of Variants in the ATPase8 gene, which is not observed in any sub-haplogroup 501 besides U4b. The complete distribution of the Variants across all the sub-haplogroups is presented

502 in **Table 2**.503

The M3a sub-haplogroup (n=8 subjects) consists of 38 variants, with 12/38 variants in the HVR I, II, III, D-loop regions (**Supplementary Figure 4E**). 19/38 variants were observed in the protein coding regions, with the most variants in this region occurring in *CYTB* (n=5). We found 15 coding

- for synonymous substitutions and 5 for non-synonymous variants (Supplementary Figure 4E) 508
- 509 M52b sub haplogroup (n=9 subjects) contained a total of 90 variants. 29/90 variants were observed 510 in HVR I, II, III and the D-loop (**Supplementary Figure 4E**). 31 variants were observed for 511 protein coding genes. *CYTB* (n=9 variants) contains the most variants for this region. 2 variants
- 512 were found in t-RNA coding genes. 22 variants coded for synonymous substitutions while 9
- 513 variants coded for non-synonymous substitutions.
- 514

515 M24a subhaplogroup (n=8 subjects) contains a total of 48 variants, 12/48 are seen in HVR I, II, 516 III and D-loop (**Supplementary Figure 4E**). 22/48 are found in protein encoding genes with the 517 most on *CYTB* (n=5 variants). 13 synonymous variants and 7 non-synonymous variants are seen 518 in this sub-haplogroup. The rest of the variants are seen in 12S, 16S-rRNA. No variants for t-RNA 519 genes were observed in this sub-haplogroup.

520

M27b (n=1 subject) has a total of 41 variants (**Supplementary Figure 4E**). 16/41 are seen in HVR I, II, III and the D-loop. 22/41 variants are seen in protein encoding genes with the highest variant count in *CYTB* (n=6 variants). 14 synonymous and 8 non-synonymous variants are observed for this sub-haplogroup and 1 variant for t-RNA coding gene.

M4a (n=1 subject) contains a total of 40 variants. 15/40 variants are seen in the non-coding regions
of HVRI, II, III and D-loop (Supplementary Figure 4E). 21 variants are seen in the protein
coding region with *CYTB* gene (n=5 variants) containing the highest variant count. Like M27b,
M4a contains 14 synonymous and 7 non-synonymous variants and 1 variant on the t-RNA coding
gene.

531

A total of 45 variants was seen in M5a sub-haplogroup (n=2 subjects) (**Supplementary Figure** 4E). 19/45 was seen in protein coding genes with *CYTB* (n=7 variants) representing the highest variants in the protein coding region. 13 variants code for synonymous substitutions while 6 code for non-synonymous variants. 1 variant is observed for a t-RNA coding gene.

536

537 M35b sub-haplogroup (1 subject) contains a total of 40 variants (Supplementary Figure 4E).

538 15/40 variants are seen in HVR I, II, III and D-loop and 20/40 variants are found in protein 539 encoding regions with the most variants observed in *CYTB* gene (n=5 variants). 14 code for 540 synonymous substitution while 7 code for non-synonymous substitutions. 1 variant is observed for

- 541 a t-RNA coding gene.
- 542

543 M33a sub-haplogroup (n=1 subject) contains 39 variants (**Supplementary Figure 4E**). 15/39 544 variants are observed in HVR I, II, III and D-loop, 19/39 variants are seen in the protein coding 545 region, with the highest count seen for *CYTB* (n=5 variants) for this region. 12 are synonymous 546 and 7 are non-synonymous substitutions.1 variant for t-RNA coding gene is also observed in this 547 sub-haplogroup. This haplogroup is unique amongst the 25 sub-haplogroups owing to the presence 548 of a variant (m.8562 C>T) at *ATPase6/8* gene.

549

550 Phylogenetic analysis of the Parsi mitochondrial haplotypes with those of Iranians and551 Indians

To further investigate the substructure of the major haplogroups identified in the Parsi cohort, a comparative analysis of haplotypes from 452 complete mtDNA sequences, including 352 Iranians²⁵ and 100 Indian mitochondrial genome sequences, was undertaken. The rationale for selection of these two populations centered around the ancestral migration patterns of the Parsis of India³⁰. This grouping also complements the model of the Parsi origin stemming from the ancient Iranian plateau³¹.

558

Analysis of the haplogroups identified in the Parsis compared with the Iranians, of whom the Persians (n=180) and the Qashqais (n=112) were the most frequent representatives, demonstrated that a) all seven Parsi haplogroups were found within the Iranian haplogroup set and b) a marked lack of haplogroup diversity was observed in the Parsi datatset (n=7 principal haplogroups) compared with the Persians and Qashqais (n=14 principal haplogroups, **Figure 7A, B**). The reason for this lack of haplotype diversity likely lies in the practice of endogamy, which has been strictly adhered to in the Parsi community for centuries, following their arrival from the Iranian plateau. 566 Contemporary populations of Iranians in the Iranian plateau represent diverse haplogroupings, 567 possibly due to admixture following political upheavals in the region after the departure of Parsis 568 from ancient Iran around 745 AD³¹.

569

570 The presence of the predominantly Eurasian mtDNA haplotypes HV, T, and U in our study cohort 571 was remarkable, given that Parsis have resided on the Indian subcontinent for over 1200 years. 572 While the majority of Parsis with M haplogroups can be linked to Persian descent, 2 sub-573 haplogroups (M2a, M2b) and 1 subject from M30d (n=4 subjects in total) were found to be related 574 to relic tribes of Indian origin within the M haplogroups in our analyses.

575

576 A detailed phylogenetic clustering of the Parsis to establish more precise ethnic relationships was 577 next undertaken. Our analysis revealed that the Parsis predominantly clustered with populations 578 from Iran (Persians and people of Persian descent, **Figure 8A**, **8E**), and the most common HV 579 group showed that all Parsis in the HV2a tree (n=14) clustered with Persians and Qashqais 580 (neighbour-joining tree weight > 0.72/72% (**Figure 8A and Table 3**), while the single Parsi in the 581 HV12b (n=1) haplotype demonstrated a strong association with other Iranian ethnicities, including

the Khorasani and Mazandaranis, in addition to the Qashqai and Persians (**Table 3**).

583

584 A total of 20 Parsi individuals in the U macro-haplogroup were found to fall into four subclades,

585 U7a (n=6), U2e (n=3), U4b (n=10), and U1a (n=1), with the highest representation in U4b and

586 U7a (Figure 8B). Phylogentic analysis demonstrated that the Parsis in the U haplogroup cluster

587 with the Persians most frequently, while a few cluster with Kurds, Armenians, Mazandarani,

- 588 Azeris, and Khorasanis, who all claim descent from Mesopotamia and the older Persian empire
- 589 (<u>https://journals.openedition.org/asiecentrale/480</u>). Among the U haplogroup, U4b and U7a (the

dominant branch of U7) haplotypes are distributed throughout the Near East and South $Asia^{24}$ with

591 subclades specific to Central Asia in the Volga-Ural region³³, Mediterranean, and Southeast 592 Europe, with lower frequencies in populations around the Baltic Sea, such as in Latvians and Tver

- 572 Europe, with lower nequencies in populations around the Baltic Sea, such as in Latvians and Tver 593 Karelians³³. Haplogroup U2 harbors frequency and diversity peaks in South Asia, whereas its U2d
- 594 and U2e subclades are confined to the Near East and Europe²⁴.
- 595

596 The T haplogroup in the Parsi cohort was found to consist of T1a, T2g, T2i, and T2b, with an even 597 distribution of samples across the subgroups (n=2, 1, 1, 1, respectively). Similar to the haplogroups 598 HV and U, the Persians and Qashqais form the largest ethnic denomination associated with the 599 Parsis with respect to the T haplogroups (>60%, Figure 8C). Five Parsi individuals of the 600 haplogroups A2v (n=3), F1g (n=1), and Z1a (n=1) were observed to be phylogentically related to 601 Persian, Kurd, Turkmen, and Iranian ethnicities, further attesting to their origin in the Iranian 602 plateau (Figure 8C). The T haplogroup is also well distributed in Eastern and Northern Europe, 603 as well as in the Indus Valley and the Arabian Peninsula. Younger T subclades are reported to have expanded into Europe and Central Asia during the Neolithic transition³⁴ 604

606 Unlike the HV, U, and T haplogroups, within which Parsi's cluster closely with Persians, Parsis

- harboring the M haplogroup appear to demonstrate more diversity in their mitochondrial genomes.
- This study showed the following breakdown: 8/12 M sub-haplogroups of the 29 Parsi M
- 609 haplotypes (M24a [n= 8], M33a [n=1], M5a [n=2], M4a [n=1)], M3a [n=7], M52b [n=8], M27b
- 610 [n=1], and M35b [n=1]) clustered with the Persians, Qashqais, Azeris of Iranian ethnicity, and
- others of Persian descent (Figure 8D, Table 3). Only two sub-haplogroups in our study (M2a and
 M2b [n=21], M30d [n=1], (Figure 8D) clustered more closely with relic tribes of Indian origin.
- 613 Our phylogenetic analyses further showed that 19 Parsi individuals belonging to the M30d (n=10)
- and M39d (n=9) haplogroups did not cluster either with Indian or Iranian ethnic groups (**Figure**
- 615 **8D**) but remained clustered within their own subgroups.
- 616
- 617 Outgroup sampling is of primary importance in phylogenetic analyses, affecting ingroup 618 relationships and, in placing the root, polarizing characters. Accordingly, we used AGENOME-
- 619 OUTGROUP-Y2b to root the phylogenetic tree. AGENOME-OUTGROUP-Y2b did not associate
- 620 with the Zoroastrian-Parsis, Indians and Iranians attesting to the robustness of the method
- 621 employed for phylogenetic analysis (**Figure 8E**, black line)
- 622

Assembly of the Zoroastrian Parsi mitochondrial consensus genome (AGENOME-ZPMRG V1.0) and Parsi haplogroup-specific reference sequences

625 The Parsis of India are a nonsmoking, long-living community despite the prevalence of many 626 genetic disease manifestations. This prompted us to generate a Parsi-specific mitochondrial 627 consensus genome to better understand the nuances of disease and wellness in this unique 628 community. Considering this goal, we classified the Parsi mitochondrial genome based on the 629 seven identified major haplogroups, HV, M, U, T, A, F, and Z. The haplogroup-specific Parsi 630 mitochondrial sequences were aligned, and a consensus call for each nucleotide was made based 631 on the maximal frequency of a base called at each position in the mtDNA genome sequence 632 (Appendix 2).

633

634 Using this approach, we derived the Zoroastrian Parsi mitochondrial reference sequences for each 635 haplogroup: AGENOME-ZPMRG-HV-V1.0 (n=15 sequences), AGENOME-ZPMRG-U-V1.0 636 (n=20 sequences), AGENOME-ZPMRG-T-V1.0 (n=5 sequences), AGENOME-ZPMRG-M-V1.0 637 (n=52 sequences), AGENOME-ZPMRG-A2v-V1.0, AGENOME-ZPMRG-F1a-V1.0, and 638 AGENOME-ZPMRG-Z-V1.0 (Table 4). Additionally, using all 100 Parsi mitochondrial genomes 639 generated in this study (see Materials and Methods), we built the first standard Zoroastrian Parsi 640 mitochondrial consensus genome (AGENOME-ZPMCG-V1.0). The consensus Parsi mtDNA 641 sequence was found to have 31 unique Variants (Table 5), of which five Variants (A263G, A750G, 642 A1438G, A4769G, and A15326G) were found to be common to the reference sequences of all 643 seven haplogroups considered (**Table 5**). While the number of Variants unique to each of the seven 644 haplogroups ranged from 11 to 33, haplogroup M did not appear to have any unique Variants when 645 compared with the overall consensus sequence, AGENOME-ZPMRG-V1.0. The utility of this newly generated reference standard could be found in the accurate mitochondrial-based analyses

- 647 involving the global Zoroastrian Parsi population as well as for individuals of Western Asian,
- 648 Indo-European and Indian origin.
- 649

Disease-specific associations of mtDNA variants predict the prevalence of commonly occurring diseases in the non-smoking Parsi cohort

As demonstrated in this paper (**Figure 7B**), the practice of intermarriage has likely restricted the genetic diversity of the Parsis, as measured by the paucity of haplogroups in our cohort compared with the Persian and Qashqai populations, possibly contributing to a number of autosomal recessive and other genetic diseases. In previous studies, Parsis were found to be disproportionately affected with certain diseases, such as prostate and breast cancers^{5,11}, Parkinsons disease (PD), and Alzheimers disease (AD). However, the Parsis are also considered to be a long-living community⁶ with lower incidences of lung cancer¹².

659

660 In order to determine whether diseases known to be prevalent in the Parsi community could in 661 fact be predicted by association using the collective mitochondrial variants discovered in this 662 study, we first analyzed variants identified in tRNA genes in the mitochondrial genome that have 663 previously been implicated in rare and degenerative diseases. We found a total of 17 tRNA-664 associated variants, with a pathogenic variant (amino acid change: G1644A) implicated 665 significantly in LS/HCM/MELAS, a genetically inherited mitochondrial disease⁴⁷. We also found a total of six tRNA mutations associated with non-syndromic hearing loss, hypertension, 666 breast/prostate cancer risk, and progressive encephalopathies in the analysis of our 100 Parsi 667 668 individuals (Table 6).

669

670 Further analysis of the nucleotide transitions and transversions that constitute the 420 variants

- 671 revealed that the mutational signatures (C>A and G>T) found in tobacco smoke-derived cancers³⁶ 672 were found at an extremely low frequency (<6% compared to other mutational signatures) on both
- the H and L strands of the mitochondrial genomes of the Parsi population (**Figure 9**), who refrain
- 675 the frame is built of the international and easiel habits
- 674 from smoking due to their religious and social habits.
- 675

676 Variant analysis

Furthermore, we found that the 420 variants analysed were associated with 41 diseases. SNP disease-association analysis revealed that Parkinson's disease is highly associated with our variants (178 Variants, **Supplementary Figure 5**). Other neurodegenerative diseases, rare diseases of mitochondrial origin, and cardiovascular and metabolic diseases associated with the variants in our study were also predicted (**Supplementary Figure 5**).

682

683 While a predisposition to 41 diseases were spread across 25 sub-haplogroups, many diseases were

- found to be recurring across haplogroups, totalling 188 diseases (**Figure 10A**). Haplogroup U4b
- harbored 15 diseases associations, while the majority of M and T groups had five diseases (Figure

686 6B). Some of the mitochondrial rare diseases, such as mitochondrial encephalomyopathies,

- 687 MELAS syndrome and cytochrome c oxidase deficiency were found to be associated with M2a 688 and U1a, U4b and M2b sub-haplogroups respectively (**Figure 10B**).
- 689

690 Haplogroup and disease linkage

Since the 420 variants identified fell into 25 sub-haplogroups contributing to 41 diseases and conditions, Principal component analysis (PCA) showed the grouping of variants and haplogroups (**Figure 11**). Alzheimers disease, breast cancer, cardiomyopathies, and Parkinsons disease were represented in all the 25 sub-haplogroups (**Appendix 3**), and longevity was represented in 23 subhaplogroups, with the exception of HV12b and U1a groups. Our tRNA pathogenicity analysis showed that the variability in tRNA was highest in the U, T, and M haplogroups compared with other haplogroups (**Table 6**).

698

699 Analysis of Variants in tRNA genes and the D-loop region in the mitochondrial genome

700 While most of the variants in mtDNA genome sequences do not affect mitochondrial function, 701 unlike synonymous/neutral variants, nonsynonymous/non-neutral variants may have functional 702 consequences, and their effect on mitochondrial metabolism may be strongly deleterious, mildly 703 deleterious, or even beneficial. We thus analysed, a SNP dataset obtained from 100 Parsi subjects 704 for nonsynonymous mutations and identified 63 such Variants located within different 705 mitochondrial genes (Figure 12). Twenty of sixty-three variants were found in genes encoding 706 CYTB (n=13) and ND2 (n=7), followed by ND5 and ND1. Disease-association analysis showed 707 that these genes were implicated in the onset of neurodegenerative conditions like AD, PD, cancers 708 of colorectal and prostate origin, metabolic diseases such as type 2 diabetes, and rare diseases such 709 as LHON (CYTB and ND2), (Figure 13, Figure 14). Variants implicated in longevity were 710 observed in our study and distributed across the *ND2* gene (Figure 10B). As observed earlier, we 711 found no association of the nonsynonymous variants in our data set that linked to lung cancer or a 712 risk of lung cancer.

713

714 To understand the mitochondrial pathways affected by the variants in our study, we annotated the pathways associated with Variants with DAVID and UNIPROT and found that the major genes 715 716 CYTB and ND2 were implicated in pathways that include the mitochondrial respiratory complex 717 (COI/COII/COIII/COIV), OXPHOS, and metabolic pathways implicated in mitochondrial 718 bioenergetics. Critical disease-related pathways in Parkinsons disease, Alzheimers disease, and 719 cardiac muscle contraction were also associated with CYTB- and ND2-specific Variants, which 720 possibly explains the high incidence of these disease in the Zoroastrian-Parsi population (Figure 721 15).

722

A total of 87 variants, including 6 unique variants, were observed in the D-loop region across all

- 724 25 sub-haplogroups (n=100 subjects, **Table 2**). 74/100 Parsis in our study, were found to have the
- polymorphism m.16519 T>C that is associated with chronic kidney disease⁴³, an increased risk

for Huntingtons disease, migraine headache, and cyclic vomiting syndrome⁴⁴ and schizophrenia and bipolar disorder⁴⁸ a. While six subjects of the M52 sub-haplogroup were found to have m.16525 A>G. The rest of the variants were found at m.16390 G>A (n=4 subjects) and m.16399

- A>G, m.16401 C>T, and m.16497 A>G (all with n=1 subject each). Taken together, these results
 warrant a deeper investigation into the D-loop variants in the Zoroastrian-Parsi community.
- 731

732 Identification of unique, unreported variants from the 100 Parsi-Zoroastrian mitogenome 733 analysis

- 734 We performed a comparative analysis of the 420 variants in the Zoroastrian-Parsi community with MITOMASTER⁴⁵, a database that contains all known pathogenic mtDNA mutations and common 735 736 haplogroup polymorphisms, to identify unique Variants in our population, that are not reported 737 previously. Our analysis showed the presence of 12 unique Variants distributed across 27 subjects 738 that were not observed in MITOMASTER and additionally in the VarDIG[®]-R disease association dataset (Figure 16). These unique variants were observed across different gene loci. 12S-rRNA 739 740 (2 variants), 16S-rRNA (5 Variants), 1 each at ND1, COII, COIII, ND4 and ND6. The SNP 741 haplogroup association showed that they fell into 4 major haplogroups and 13 sub haplogroups; 742 HV2a=1, M24a=4, M2a=1, M30d=3, M35b=1, M39b=2, M3a=1, M4a=1, M52b=4, M5a=1, 743 T2b=1, U4b=6, U7a=1. Of the 12 variants identified, no disease associations were observed for on
- analysis with MITOMASTER and VarDIG[®]-R and needs to be further investigated.
- 745

746 **Discussion**

747

The first *de novo* Parsi mitochondrial genome, AGENOME-ZPMS-HV2a-1 (Genbank accession:
 MT506314) from a healthy, non-smoking female of haplogroup HV2a when compared with the
 revised Cambridge Reference Standard (rCRS) showed 28 unique variants. Upon extending our

mitochondrial genome analyses to an additional 99 Parsi individuals, we found that 94 individuals

- r52 separate into four major mitochondrial haplogroups, HV, U, T, and M, while 5 individuals belong
- to the rarer haplogroups A, F, and Z. The largest sub-haplogroup was found to be HV2a (n=14).
- 754

Due to the strict endogamy practiced by the Zoroastrian Parsis, their maternally inherited mitochondrial lineage seems to have remained aligned with those of their ancestors in Old Persia, prior to 642 AD. On comparison of the major mitochondrial haplogroups in our Parsi cohort with 352 Iranian²⁵ and 100 Indian mitochondrial genomes, we observed that the Zoroastrian-Parsi genomes are phylogenetically related to the Persians and Qashqais²⁵ in HV, T, U, F, A and Z haplogroups, those associated with peopling of western Europe, Central Asia and the Iranian plateau.

762

The haplogroup HV2, dated at 36–42 kya, most likely arose in Persia between the time of the first settlement by modern humans and the last glacial melt, and the subclade HV2a has a demonstrated

765 Persian ancestry. HV12b, a branch of the HV12 clade, is one of the oldest HV subclades and has

been found in western Iran, India, and sporadically as far as Central and Southeast Asia. It has
strong associations with the Qashqais, who are Turkic-speaking nomadic pastoralists of southern
Iran and who previously resided in the Iranian region of the South Caucasus^{32,35}. The presence of
these predominantly Eurasian mtDNA haplotypes, HV, T, U, F, A and Z in our Parsi cohort attests
to their practice of endogamy, given that Parsis have resided on the Indian subcontinent for over
1300 years.

772

Despite the large grouping of the M haplogroup (the largest haplogroup in the Indian subcontinent ³⁴) in our Parsi cohort, phylogenetic analysis showed that 47/51 Parsis belonging to the M haplogroups in our study, cluster with the Persians, suggesting Persian descent, with a small minority of Parsis found to be related to relic tribes of India. This observation suggests minimal gene flow from indigenous Indian females into the Parsi gene pool, as had been previously proposed³⁰.

779

Phylogenetic analysis also revealed that two Parsi M sub-haplogroups, M30d and M39b formed aunique cluster that needs further resolution.

782

We further present the first complete Zoroastrian Parsi Mitochondrial Consensus Genome (AGENOME-ZPMCG V1.0), built from the mitochondrial genomes of 100 non-smoking, Parsi individuals representing seven mitochondrial haplogroups. The need for the generation of such an ethnic-specific consensus genome, specifically for the Parsis, is self-evident for studies involving comparative analyses, designed to precisely understand patterns of maternally inherited mitochondrial DNA and aid in reconstructing the history and prevalent disease associations in this unique community.

790

791 We found that *CYTB* gene contained the maximum number of variants ($n \ge 5$) in the coding region 792 of haplogroup M, besides having maximal representation in F1g, T, and HV12b. Haplogroups U, 793 A2v, and Z1a showed dominance for the ND complex genes ND5 and ND2, while the COI genes 794 were the most highly represented in HV2a and U4b. Variants in the CYTB gene are associated with 795 Alzheimers disease, diabetes mellitus, cognitive ability, breast cancer, hearing loss, and 796 asthenozoospermia and associated with changes in metabolic pathways, cardiac contraction and 797 rare diseases such as Huntington's disease, whereas the ND2 and ND5 variants were associated 798 with prostate, ovarian cancer, rare mitochondrial neuronal diseases, such as LHON, 799 cardiomyopathy, Alzheimers disease and Parkinsons disease.

800

801 tRNA disease-association analysis in our study showed that these genes were implicated in the 802 onset of neurodegenerative conditions, such as Alzheimers disease, Parkinsons disease, cancers of 803 colorectal and prostate origin, metabolic diseases, such as type 2 diabetes, and rare diseases, such

as LHON (*CYTB* and *ND2*). The D-loop SNP analysis showed the prevalence (74/100 subjects) of

the m.16519 T>C polymorphism, which has been implicated in chronic kidney disease⁴³, an

806 increased risk for Huntingtons disease, migraine headache, and cyclic vomiting syndrome⁴⁴. Taken

- together, these results warrant a deeper investigation into the tRNA and the D-loop variants in the
- 808 Zoroastrian-Parsi community.
- 809 Consanguineous marriages amongst the Parsis has given rise to longevity¹¹ and number of associated diseases, including colon, prostate, breast cancers^{9,10}, Parkinson's disease (PD), 810 Alzheimer's disease (AD), and lower incidences of lung cancer¹². Interrogation of the 420 811 variants across seven haplogroups in the Parsi cohort using the VarDIG[®]-R database revealed that 812 Parkinson's disease, known to be prevalent in the Parsi community³⁶, was predicted to have the 813 814 highest prevalence with 178 of the 420 variants represented. Not surprisingly, longevity, which 815 often co-occurs with Parkinsons disease, was also predicted to be highly prevalent in the Parsi 816 cohort, but with a notable absence in the U1 sub-haplogroup, an interesting observation that 817 warrants further investigation.
- 818

Analysis of additional disease associations revealed that Alzheimer's disease (also related to ageing), breast cancer, and cardiomyopathies^{38,39,40}, were predicted to be associated with all the 25 Parsi sub-haplogroups. Additionally, the observed low birth rate among Parsi could be predicted from the presence of variants associated with asthenozoospermia³⁷; a condition associated with reduced sperm motility.

824

825 It is noteworthy that previously published studies demonstrating lower rates of lung cancer amongst the Parsis⁴¹, appears to hold true, given that no haplogroup in the Parsi cohort 826 827 demonstrated a predicted predisposition to lung cancer. The lack of mitochondrial signatures for 828 lung cancer in all haplogroups examined in this non-smoking Parsi population coupled with the 829 low frequency of mutational signatures for tobacco smoke-derived cancers was not surprising, 830 particularly since the Zoroastrian-Parsi venerate fire, and smoking would lie in gross violation of 831 that sacred tenet. Other diseases predicted to occur at high frequencies in our analyses await further 832 investigation in the Parsi community.

833

The Parsi haplogroup specific variant-disease association analysis has shed predictive light on the association of mitochondrial variants linked to longevity, neurodegenerative diseases, cancers of the colon, breast and prostate and low birth rate, among others; diseases that have been well documented to occur in the Parsi community. The Parsis thus represent a small but unique, nonsmoking community where genomic disease signatures, both mitochondrial and nuclear, can be investigated in the backdrop of generations of endogamy thus providing exceptional opportunities to understand and mitigate disease.

841

842 Conclusion

843 We have generated the first *de novo* Zoroastrian Parsi Mitochondrial Reference Sequence 844 (AGENOME- ZPMS-HV2a-1) and the Zoroastrian Parsi Mitochondrial Consensus Genome

845 (AGENOME-ZPMCG V1.0). This newly generated reference standards will contribute in the

analysis of mitochondrial genomes of not only the Zoroastrian Parsi population but also other

- 847 populations. We have also provided evidence that the Zoroastrian Parsis of India, through centuries
- 848 of endogamy, have retained their Persian genetic heritage, distinct traits of longevity and associated
- diseases. We have shown the rôle of social habits in genetic signatures exemplified by the lack of
- 850 mitochondrial variants associated with lung cancer.
- 851

In sum, The Parsi haplogroup specific variant-disease association analysis has shed predictive light on the association of mitochondrial variants linked to longevity, neurodegenerative diseases, cancers of the colon, breast and prostate and low birth rate, among others; diseases that have been well documented to occur in the Parsi community. The Parsis thus represent a small but unique, non-smoking community where genomic disease signatures, both mitochondrial and nuclear, can be investigated in the backdrop of generations of endogamy thus providing exceptional opportunities to understand and mitigate disease.

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860 **References**

- 1. Mistry, R. K. Glimpses of Parsi history, Insights Into The Zarathustrian Religion, p.20.
- 862 2. Nariman, R. F. *The Inner Fire Faith, Choice, and Modern Day Living in*
- 863 Zoroastrianism, p. 20-21
- 3. The Vendidad: The Zoroastrian Book Of The Law Paperback September 10, 2010. I, 1-2
 & II, 5. Charles. F. Horne. ISBN-10: 1162910089; ISBN-13: 978-1162910086. Kessinger
 Publishing, LLC (September 10, 2010)
- 867 4. Bennet, J. G. The Hyperborean Origin of the Indo-European Culture, Journal Systematics.
 868 *J Syst.* 1, (1963).
- Jussawalla, D. J., Yeole, B. B. & Natekar, M. V. Histological and epidemiological
 features of breast cancer in different religious groups in greater bombay. *J. Surg. Oncol.*(1981) doi:10.1002/jso.2930180309.
- B72 6. Jussawalla, D. J. The persistance of differences in cancer incidence at various anatomical
 B73 sites 1300 years after immigration. *Recent Results Cancer Res.* (1975) doi:10.1007/978-3B74 642-80880-7_22.
- 875 7. Anthony, DW, (2007), The Horse, The Wheel, And Language. How Bronze-Age Riders
 876 from the Eurasian Steppes Shaped the Modern World, Princeton University Press. p. 9.
- 8. Alizadeh, A. The Rise of the Highland Elamite State in Southwestern Iran. Current. *Curr*878 *Anthropol.* 51, 353–383 (2010).
- 879 9. Shroff Z, C. M. The potential impact of intermarriage on the population decline of the
 880 Parsis of Mumbai, India. *Demogr Res.* 25, 545–564 (2011).
- 881 10. Karkal, M. Marriage among Parsis. *Demogr. India* 4, 128 (1975).
- Barnabas-Sohi, N. *et al.* Breast carcinoma in a high-risk population: Structural alterations
 in neu, int-2, and p-53 genes. *Breast Dis.* (1993).
- 12. Jussawalla, D. J. & Jain, D. K. Lung cancer in Greater Bombay: Correlations with religion

885 and smoking habits. Br. J. Cancer (1979) doi:10.1038/bjc.1979.199. 886 13. Helgason, A., Sigurŏardóttir, S., Gulcher, J. R., Ward, R. & Stefánsson, K. mtDNA and 887 the origin of the Icelanders: Deciphering signals of recent population history. Am. J. Hum. 888 Genet. (2000) doi:10.1086/302816. 889 14. Wallace, D. C. Mitochondrial DNA Variation in Human Radiation and Disease. Cell 890 (2015) doi:10.1016/j.cell.2015.08.067. 891 Wallace, D. C., Brown, M. D. & Lott, M. T. Mitochondrial DNA variation in human 15. 892 evolution and disease. Gene (1999) doi:10.1016/S0378-1119(99)00295-4. 893 Roger, A. J., Muñoz-Gómez, S. A. & Kamikawa, R. The Origin and Diversification of 16. 894 Mitochondria. Current Biology (2017) doi:10.1016/j.cub.2017.09.015. 895 17. Garcia, I., Jones, E., Ramos, M., Innis-Whitehouse, W. & Gilkerson, R. The little big 896 genome: The organization of mitochondrial DNA. Front. Biosci. - Landmark (2017) 897 doi:10.2741/4511. 898 18. Stewart, J. B. & Chinnery, P. F. The dynamics of mitochondrial DNA heteroplasmy: 899 Implications for human health and disease. Nature Reviews Genetics (2015) 900 doi:10.1038/nrg3966. 901 19. Bussard, K. M. & Siracusa, L. D. Understanding mitochondrial polymorphisms in 902 cancer.Cancer Research (2017) doi:10.1158/0008-5472.CAN-17-1939. 903 Alston, C. L., Rocha, M. C., Lax, N. Z., Turnbull, D. M. & Taylor, R. W. The genetics 20. 904 and pathology of mitochondrial disease. J. Pathol. 241, 236–250 (2017). 905 21. Andrews, R. M. et al. Reanalysis and revision of the cambridge reference sequence for 906 human mitochondrial DNA [5]. Nature Genetics (1999) doi:10.1038/13779. 907 22. Chandrasekar, A. et al. Updating phylogeny of mitochondrial DNA macrohaplogroup m 908 in India: dispersal of modern human in South Asian corridor. PLoS One 4, e7447-e7447 909 (2009).910 23. Rajkumar, R., Banerjee, J., Gunturi, H. B., Trivedi, R. & Kashyap, V. K. Phylogeny and 911 antiquity of M macrohaplogroup inferred from complete mt DNA sequence of Indian 912 specific lineages. BMC Evol. Biol. 5, 26 (2005). 913 24. Sahakyan, H. et al. Origin and spread of human mitochondrial DNA haplogroup U7. Sci. 914 *Rep.* 7, 46044 (2017). 915 25. Derenko, M. et al. Complete mitochondrial DNA diversity in Iranians. PLoS One (2013) 916 doi:10.1371/journal.pone.0080673. 917 Edgar, R. C. MUSCLE: multiple sequence alignment with high accuracy and high 26. 918 throughput. Nucleic Acids Res. 32, 1792–1797 (2004). 919 27. Kumar, S., Stecher, G., Li, M., Knyaz, C. & Tamura, K. MEGA X: Molecular 920 Evolutionary Genetics Analysis across Computing Platforms. Mol. Biol. Evol. 35, 1547-921 1549 (2018). 922 Tamura, K., Nei, M. & Kumar, S. Prospects for inferring very large phylogenies by using 28. 923 the neighbor-joining method. Proc. Natl. Acad. Sci. U. S. A. 101, 11030–11035 (2004). 924 29. Houshmand, M. et al. Is 8860 variation a rare polymorphism or associated as a secondary

925		effect in HCM disease? Arch. Med. Sci. (2011) doi:10.5114/aoms.2011.22074.
926	30.	Chaubey, G. et al. 'Like sugar in milk': Reconstructing the genetic history of the Parsi
927		population. Genome Biol. (2017) doi:10.1186/s13059-017-1244-9.
928	31.	López, S. et al. The Genetic Legacy of Zoroastrianism in Iran and India: Insights into
929		Population Structure, Gene Flow, and Selection. Am. J. Hum. Genet. (2017)
930		doi:10.1016/j.ajhg.2017.07.013.
931	32.	Quintana-Murci, L. et al. Where west meets east: the complex mtDNA landscape of the
932		southwest and Central Asian corridor. Am. J. Hum. Genet. 74, 827-845 (2004).
933	33.	Shamoon-Pour, M., Li, M. & Merriwether, D. A. Rare human mitochondrial HV lineages
934		spread from the Near East and Caucasus during post-LGM and Neolithic expansions. Sci.
935		<i>Rep.</i> 9 , 14751 (2019).
936	34.	Farjadian, S. et al. Discordant Patterns of mtDNA and Ethno-Linguistic Variation in 14
937		Iranian Ethnic Groups. Hum. Hered. 72, 73-84 (2011).
938	35.	Thangaraj, K. et al. In situ origin of deep rooting lineages of mitochondrial
939		Macrohaplogroup 'M' in India. BMC Genomics 7, 151 (2006).
940	36.	Alexandrov LB, Ju YS, Haase K, et al. Mutational signatures associated with tobacco
941		smoking in human cancer. Science. 2016;354(6312):618 - 622.
942	37.	E. Ruiz-Pesini, A.C. Lapeña, C. Díez, E. Alvarez, J.A. Enríquez, M.J. López-Pérez
943		Seminal quality correlates with mitochondrial functionality. Clin. Chim.
944		Acta., 300 (2000), p. 97 105.
945	38.	Fang, H., Shen, L., Chen, T. et al. Cancer type-specific modulation of mitochondrial
946		haplogroups in breast, colorectal and thyroid cancer. BMC Cancer 10, 421 (2010).
947	39.	Van der Walt JM, Dementieva YA, Martin ER, Scott WK, Nicodemus KK, Kroner CC,
948		Welsh-Bohmer KA, Saunders AM, Roses AD, Small GW, Schmechel DE, Murali
949		Doraiswamy P, Gilbert JR, Haines JL, Vance JM, Pericak-Vance MA. Analysis of
950		European mitochondrial haplogroups with Alzheimer disease risk. Neurosci Lett. 2004
951		Jul 15; 365(1):28-32.
952	40.	van Oven M, Kayser M Hum Mutat. Updated comprehensive phylogenetic tree of global
953		human mitochondrial DNA variation. 2009 Feb; 30(2): E386-94.
954	41.	Balkrishna Bhika Yeole, AP Kurkure, SH Advani, Sunny Lizzy; An Assessment of
955		Cancer Incidence Patterns in Parsi and Non Parsi Populations, Greater Mumbai. Asian
956		Pacific Journal of Cancer Prevention, Vol 2, 2001; 293-298
957	42.	Niroula A, Vihinen M. PON-mt-tRNA: a multifactorial probability-based method for
958		classification of mitochondrial tRNA variations. Nucleic Acids Res. 2016;44(5):2020-
959		2027. doi:10.1093/nar/gkw046
960	43.	Chen JB, Yang YH, Lee WC, et al. Sequence-based polymorphisms in the mitochondrial
961		D-loop and potential SNP predictors for chronic dialysis. PLoS One. 2012;7(7):e41125.
962		doi:10.1371/journal.pone.0041125
963	44.	Zaki EA, Freilinger T, Klopstock T, et al. Two common mitochondrial DNA
964		polymorphisms are highly associated with migraine headache and cyclic vomiting

syndrome. Cephalalgia. 2009;29(7):719-728. doi:10.1111/j.1468-2982.2008.01793.x

966 967	45.	Brandon MC, Ruiz-Pesini E, Mishmar D, et al. MITOMASTER: a bioinformatics tool for the analysis of mitochondrial DNA sequences. Hum Mutat. 2009;30(1):1-6.							
968		doi:10.1002/humu.20801.							
969	46.	Narasimhan VM, Patterson N, Moorjani P, et al. The formation of human populations in							
970		South and Central Asia. Science. 2019;365(6457):eaat7487. doi:10.1126/science.aat7487.							
971	47.	47. Menotti F, Brega A, Diegoli M, Grasso M, Modena MG, Arbustini E. A novel mtDNA							
972		point mutation in tRNA(Val) is associated with hypertrophic cardiomyopathy and							
973		MELAS. Ital Heart J. 2004;5(6):460-465.							
974	48.	Schulmann A, Ryu E, Goncalves V, et al. Novel Complex Interactions between							
975		Mitochondrial and Nuclear DNA in Schizophrenia and Bipolar Disorder. Mol							
976		Neuropsychiatry. 2019;5(1):13 - 27. doi:10.1159/000495658							
977									
978	Cont	ributions							
979	VMP	conceptualized and guided the experiments; NP, CG analysed the sequences, bioinformatics							
980	analys	sis, interpreted the results; RM, SR, NS co-ordinated wet-lab work flows and data analysis; NP,							
981	BM, I	KK analysed data, plotted graphs and figures; VMP, AKG, PB, KK, RJ drafted the manuscript							
982	with i	nputs from RM, SR, NS, CG. All authors reviewed the manuscript							
983									
984	All at	thors researched data for the article, made substantial contribution to discussion of content,							
985	and w	prote, reviewed, and edited the manuscript before submission.							
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1008 Main Figures

1009

1010 Figure 1 : Identification of 28 variants in the de novo Parsi mitochondrial

- 1011 genome, AGENOME-ZPMS-HV2a-1
- 1012



Variant distribution in de novo assembly

- 1013
- 1014

Number of variants



1016 Representative histogram showing the base change, variant type, type of loci and distribution of

- 1017 variants across genes in the *de novo* mitochondrial genome AGENOME-ZPMS-HV2a-1
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- 1022

Figure 2 : Validation of variants in the AGENOME-ZPMS-HV2a-1 by Sanger sequencing

1025

1. rCRS	CCTGACTGGCATTGTATTAGCAAACTCATCACTAGACATCGTACTACACGACACGTACTA	7018
2. AGENOME-ZPMS-HV2a-1	CCTGACTGGCATTGTATTAGCAAACTCATCACTAGACATCGTACTACACGACACGTACTA	7018
3. SANGER-SEQUENCED	CCTGACTGGCATTGTATTAGCAAACTCATCACTAGACATCGTACTACACGACACGTACTA	434
1. rCRS 2. AGENOME-ZPMS-HV2a-1 3. SANGER-SEQUENCED	CGTTGTAGCCACTTCCACTATGTCCTATCAATAGGAGCTGTATTTGCCATCATAGGAGG CGTTGTAGCTCACTTCCACTATGTCCTATCAATAGGAGCTGTATTTGCCATCATAGGAGG CGTTGTAGCTCACTTCCACTATGTCCTATCAATAGGAGCTGTATTTGCCATCATAGGAGG *******	7078 7078 494
1. rCRS	CAACCGCTATGTATTTCGTACATTACTGCCAGCCACCATGAATATTGTACGGTACCATAA	16138
2. AGENOME-ZPMS-HV2a-1	CAACCGCTATGTATTTCGTACATTACTGCCAGCCACCATGAATATTGTACGGTACCATAA	16138
3. SANGER-SEQUENCED	CAACCGCTATGTATTTCGTACATTACTGCCAGCCACCATGAATATTGTACGGTACCATAA	126
1. rCRS	ATACTTGACCACCIGTAGTACATAAAAACCCAATCCACATCAAAACCCCCTCCCCATGCT	16198
2. AGENOME-ZPMS-HV2a-1	ATACTTGACCACCIATAGTACATAAAAACCCAATCCACATCAAAACCCCCTCCCCATGCT	16198
3. SANGER-SEQUENCED	ATACTTGACCACCIATAGTACATAAAAACCCAATCCACATCAAAACCCCCTCCCCATGCT	186
1. rCRS	TACAAGCAAGTACAGCAATCI ACCCTCAACTATCACACATCAACTGCAACTCCAAAGCCA	16258
2. AGENOME-ZPMS-HV2a-1	TACAAGCAAGTACAGCAACCI ACCCTCAACTATCACACATCAACTGCAACTCCAAAGCCA	16258
3. SANGER-SEQUENCED	TACAAGCAAGTACAGCAACCI ACCCTCAACTATCACACATCAACTGCAACTCCAAAGCCA	246
1. rCRS	CCCCTCACCCACTAGGATACCAACAAACCTACCACCCTTAACAGTACA <mark>FAC</mark> TACATAAA	16318
2. AGENOME-ZPMS-HV2a-1	CCCCTCACCCACTAGGATACCAACAAACCTACCCACCCTTAACAGTACA <mark>FGC</mark> TACATAAA	16318
3. SANGER-SEQUENCED	CCCCTCACCCACTAGGATACCAACAAACCTACCACCCTTAACAGTACAFGCTACATAAA	306
	GCCATTTACCGTACATAGCACATTACAGTCAAATCCCTTCTCGTCCCCATGGATGACCCC GCCATTTACCGTACATAGCACATTACAGTCAAATCCCTTCTCGTCCCCATGGATGACCCC GCCATTTACCGTACATAGCACATTACAGTCAAATCCCTTCTCGTCCCCATGGATGACCCC	16378 16378 366

1. SANGER-SEQUENCED	CAGGACATCCCGATGGTGCAGCCGCTATTAAAGGTTCGTTTGTTCAACGATTAAAGTCCT	181
2. AGENOME-ZPMS-HV2a-1	CAGGACATCCCGATGGTGCAGCCGCTATTAAAGGTTCGTTTGTTCAACGATTAAAGTCCT	3060
3. rCRS	CAGGACATCCCGATGGTGCAGCCGCTATTAAAGGTTCGTTTGTTCAACGATTAAAGTCCT	3058

1. SANGER-SEQUENCED	ACGTGATCTGAGTTCAGACCGGAGTAATCCAGGTCGGTTTCTATCTA	240
2. AGENOME-ZPMS-HV2a-1	ACGTGATCTGAGTTCAGACCGGAGTAATCCAGGTCGGTTTCTATCTA	3119
3. rCRS	ACGTGATCTGAGTTCAGACCGGAGTAATCCAGGTCGGTTTCTATCTA	3118

	CCCTGTACGAAAGGACAAGAGAAATAAGGCCTACTTCACAAAGCGCCTTCCCCCGTAAAT	300
1. SANGER-SEQUENCED	CCCTGTACGAAAGGACAAGAGAAATAAGGCCTACTTCACAAAGCGCCTTCCCCCGTAAAT	3179
2. AGENOWE-ZPINS-HV2a-1	CCCTGTACGAAAGGACAAGAGAAATAAGGCCTACTTCACAAAGCGCCTTCCCCCGTAAAT	3178
3. rCRS	****	
1. SANGER-SEQUENCED	GATATCATCTCAACTTAGTATTATACCCACACCCACCCAAGAACAGGGTTTGTTAAGATG	360
2. AGENOME-ZPMS-HV2a-1	GATATCATCTCAACTTAGTATTATACCCACACCCACCCAAGAACAGGGTTTGTTAAGATG	3239
3. rCRS	GATATCATCTCAACTTAGTATTATACCCACACCCACCCAAGAACAGGGTTTGTTAAGATG	3238

1. SANGER-SEQUENCED	CCCCA	5
2. AGENOME-ZPMS-HV2a-1	ACAATTGAATGTCTGCACAGCCGCTTTCCACACAGACATCATAACAAAAAATTTCCACCA	300
3. rCRS	ACAATTGAATGTCTGCACAGCCACTTTCCACACAGACATCATAACAAAAAATTTCCACCA	300
	* ***	
1. SANGER-SEQUENCED	AACCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	65
2. AGENOME-ZPMS-HV2a-1	AACCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	360
3. rCRS	AACCCCCCCCTCCCCCGCTTCTGGCCACAGCACTTAAACACATCTCTGCCAAACCCCAA	358

1. SANGER-SEQUENCED	AAACAAAGAACCCTAACACCAGCCTAACCAGATTCAAATTTATCTTTTGGCGGTAGCACT	125
2. AGENOME-ZPMS-HV2a-1		395
3. rCRS	AAAUAAAGAAUUUTAACACCAGCCTAACCAGATTT	393
	* * * * * * * * * * * * * * * * * * * *	555

1029	Figure 2 : Confirmation of variants identified with next-generation sequencing (NGS) data
1030	and confirmation by Sanger sequencing. Sequences obtained from desired regions were
1031	analyzed for presence of variants/Variants. Low quality bases were trimmed from both ends of the
1032	sequences and used for alignment with the reference Mitochondrial Genome (rCRS). A total of 13
1033	variants/Variants from D-loop and internal region of mitochondrial genome were verified.
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1040 Figure 3 : Representation of Males and Females in the 100 Zoroastrian-Parsi 1041 whole mitogeneme study

1041 whole mitogenome study



Figure 3: Distribution of 100 Parsi subjects. Distribution of the subjects classified based on
 gender and age. The bars on the histogram depict further segmentation of the total number of
 subjects, Male and Female numbers according to their age range.

1057 Figure 4 : Distribution of 420 variants across gene loci in the 100 Zoroastrian-

1058 Parsi whole mitogenomes

1059



Figure 4 : Annotation and distribution of 420 variants across 100 Parsi complete
 mitogenomes

Figure 5 : Identification of 25 sub-haplogroups in the 100 Zoroastrian-Parsi study group

Table: Haplogroup and Haplotype count in Parsis Parsis Number of Parsis Major haplogroup Sub-haplotypes HV2a 14 HV2a HV HV12b 1 HV12b U7a 6 U7a U2e 3 U2e υ U4b U4b 11 U1a U1a 1 2 T1a T1a T2g T2g 1 т T2i T2i 1 T2b T2b 1 Sub-haplotypes M5a M5a 2 M39b M39b 9 M33a МЗЗа 1 M52b M52b 9 M24a 8 M24a МЗа МЗа 8 M30d M M30d 11 M2a M2a 2 M4a M4a 1 M2b M2b M35b 1 M27b M35b 1 A2v M27b 1 F1g A2v 3 A Z1a F F1g 1 0 2 4 6 8 10 12 14 z Z1a 1 Sample count

1070

1071Figure 5 : Distribution of Parsis across major haplogroups and sub-haplogroups. The table1072and the histogram shows the distribution of 100 Parsi subjects across 7 major haplogroups and 25

1073 sub-haplogroups

1075Figure 6 : Distribution of variants across haplogroups and demographic1076classification of the 100 Parsi study group

- 1077
- 1078 A





- 1085
- 1086

1087Figure 7 : Lack of haplogroup diversity in the Parsi cohort suggesting1088endogamy







B

1092 Figure 7: Comparative analysis of Major haplogroup distribution in the Parsis and

populations of Iranian ethnicities (Persians, Qashqais) (A) Histogram depicting the analysis of

1094 The 7 Major haplogroups across Parsis (n=100), Persians (n=180) and Qashqais (n=112) (B)

1095 Representative figure showing the diversity of major haplogroups in the Parsis and the Persians

-



Figure 8: Phylogenetic analysis depicting individual sub-haplogroup clusters of 97 Parsis, 352 Iranian and 100 relic tribes of Indian origin (A) Representative cladograms of the HV sub-haplogroup (B) Representative cladograms of the U sub-haplogroup



С

Figure 8: Phylogenetic analysis depicting individual sub-haplogroup clusters of 97 Parsis, 352 Iranian and 100 relic tribes of Indian origin (C) Representative cladograms of the T, F and A sub-haplogroup



Figure 8: Phylogenetic analysis depicting individual sub-haplogroup clusters of 97 Parsis, 352 Iranian and 100 relic tribes of Indian origin (A-D) Representative cladograms of the each sub-haplogroup

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Ε



Figure 8E: Table indicates the number of Zoroastrian-Parsis who cluster with Persians or people of Persian origin, relic tribes of Indian origin. Pie chart indicates the percentage of clustering of the HV2a Zoroastrian-Parsis in the phylogenetic clustering analysis. Circular dendrogram of the complete Phylogenetic clustering analysis of Parsis (Blue clades) with Iranian mitogenomes (Green clades) and Indian mitogenomes (Brown clades). Outgroup is indicated by the black line.

Figure 9: Lack of smoking induced mutational sign24atures in the Parsi cohort



Figure 9: Mutational signatures observed in the 100 mitochondrial genomes of Parsis. Graph depicts the quantification of both transitions and transversions on both H&L strands of the 100 mitochondrial genomes of Parsis.

Figure 10: Observation of Longevity variants across all sub-haplogroups and predisposition of U and M haplogroups to diseases

А



В



Figure 10: Haplogroup specific distribution of diseases. (A) Distribution of 188 diseases across 25 sub-haplogroups of the 100 Parsi subjects analyzed in this study (B) Histogram depicting longevity and disease prevalence across U1a, M52b, M35b, M27b





Figure 11: Principal Component Analysis of disease associations with sub-haplogroups in the Parsi-Zoroastrian group under study



Figure 12: CYTB gene has the highest occurrence of non-synonymous variants in this study

Figure 12: Analysis of the non-synonymous variants within 420 variants in the 100 Parsi mitochondrial genome sequences. The histogram and the table show the location of the non-synonymous variants in the coding gene loci in the mitochondrial genome analysed with MitImpact database





Figure 13: Analysis of the non-synonymous variants within 420 variants in the 100 Parsi mitochondrial genome sequences for and their disease associations.





Figure 14 : Distribution of non-synonymous Variants across coding genes. Analysis was performed on the 420 Variants linked to the 100 Parsi mitochondrial genomes.

Figure 15: Gene ontology associated with non-synonymous variants among 420 variants



Figure 15: Analysis of non-synonymous mutations and their functional classification, engagement in different pathways respectively using DAVID and UNIPROT annotation tools.



Figure 16: 12 unique variants found in the current study

Figure 16: Comparative analysis of the 420 variants in the AVESTAMITOME[™] Zoroastrian-Parsi community dataset with common and disease associated polymorphisms in MITOMASTER database and VarDiG®-R

Main Tables

Reference_position	72	73	152	195	263	309.1	309.2	310	750	1438	2706	4769	5075	6104	6179	7028	7193
Reference_base	т	Α	т	т	Α			Т	Α	А	А	А	Т	С	G	С	т
AGENOME-ZPMS-HV2a-1	С	G	С	С	G	С	Т	С	G	G	G	G	С	Т	А	Т	С
Mitochondrial genome loci		HVR-II							12S-rRNA:RNR1		16S-rRNA:RNR2	ND2		СОІ			
Amino Acid change	nc	nc	nc	nc	nc	nc	nc	nc	rRNA	rRNA	rRNA	M100M	12021	F67F	M92M	A375A	F430F
Conservation index									98%	87%	84%	24%	44%	100%	100%	100%	100%
Protein Position												100	202	67	92	375	430
Variant Type												syn	syn	syn	syn	syn	syn
Type of base change	trans	trans	trans	trans	trans	ins	ins	trans	trans	trans	trans	trans	trans	trans	trans	trans	trans

Table 1: Annotation of 28 variants in the AGENOME-ZPMS-HV2a-1

Reference_position	8860	9336	10410	11016	11935	12061	15326	15792	16153	16217	16309	Haplogroup
Reference_base	Α	Α	т	G	т	С	Α	т	G	т	А	
ZPMS-HV-1	G	G	С	А	С	Т	G	С	А	С	G	HV2a
Mitochondrial genome loci	ATPase6	COIII	tRNA [R]		ND-4		C١	тв		HVR-I		
Amino Acid change	T112A	M44V	tRNA	S86N	T392T	N434N	T194A	I349T	nc	nc	nc]
Conservation index	71%	16%	22%	7%	89%	69%	18%	58%				
Protein Position	112	44		86	392	434	194	349				
Variant Type	n-syn	n-syn		n-syn	syn	syn	n-syn	n-syn				
Type of base change	trans	trans	trans	trans	trans	trans	trans	trans	trans	trans	trans]

Table 1: Annotation of the de novo Parsi mitochondrial genome AGENOME-ZPMS-HV2a-1. B) The table indicates the Variants (n=28) found in the AGENOME-ZPMS-HV2a-1 in relation to the revised Cambridge Reference Sequence (rCRS, Reference bases

Table 2: Distribution of 420 variants for each sub-haplogroup for protein coding regions, D-loop of 100 Parsi mitogenomes

30

40

Sub-haplogroup	Coding gene SNPs	Gene with max SNPs	D-loop			ig gene s	JNPS
HV2a	20	6 COI	1	18/0-	_	_	
HV12b	6	2 CYTB	0	HV2a			
U4b	21	6 COI	4				
U2e	22	4 CYTB, 4 ND2, 4 ND5	2	U2e		- Ing	
U7a	25	6 ND5	2	U7a		1	1
U1a	21	6 ND5	2	U1a			
T1a	18	5 CYTB	1	T1a			
T2b	17	3 CYTB	0	T2b			
T2g	24	6 CYTB	1	T2g			
T2i	17	5 CYTB	1	T2i			
A2v	5	2 ND2	0	A2v			
F1g	20	7 CYTB	0	F1g			
Z1a	17	3 ND5	1	Zia			
M30d	24	8 CYTB	1	M33a			
M33a	19	5 CYTB	1	M35b			
M35b	21	5 CYTB	1	M39b			
M39b	22	5 CYTB	0	M3a			
M3a	19	5 CYTB	1	M4a			
M4a	20	5 CYTB	1	M52b			
M52b	31	9 CYTB	2	M5a			
M5a	21	6 CYTB	1	M2a		1	
M2a	29	7 CYTB	2	M2b			L
M2b	25	6 CYTB	2	M2/D		-	
M27b	22	6 CYTB	1	M24a			- 6
M24a	20	5 CVTB		0	10	20	3

Association of coding region, D-loop with sub-haplogroup



Table 3: Phylogenetic clustering of complete mitogenomes of Parsis with 352Iranian and 100 relic tribes of Indian origin

Major haplogroup	Sub- haplogroups	People of Persian origin (PO)	People of Indian & Relic tribal origin (IO)	Max BS value to nearest PO	Max BS value to nearest IO
	HV2a	Persian	N.A	0.7270	0
HV	HV12b	Persian, Qashqai, Mazandarani	N.A	0.6550	0
	U7a	Persian, Kurd, Tajik	N.A	0.8980	0
	U2e	Persian, Qashqai, Azeri	N.A	1.000	0
U	U4b	Persian, Khorasani, Qashqai	N.A	0.5100	0
	U1a	Persian, Armenian	N.A	0.6850	0
	T1a	Persian	N.A	0.7320	0
-	T2g	Persian	N.A	0.4880	0
1	T2i	Persian	N.A	0.4480	0
	T2b	Persian	N.A	0.4320	0
	M5a	Persian	Munda, Mahali	0.9860	0.6270
	M39b	Unique cluster			
	M33a	Azeri	Jenu Kuruba	0.2250	0.0960
	M52b	Indian Shia Muslim	Mathakur, Dirang Monpa	0.7950	0.1170
	M24a	Persian, Qashqai	Pauri Bhaiya, Nihal	0.8560	0.0200
	M3a	Persian	N.A	0.9380	0
М	M30d	Unique cluster	1 M30d with Brahmin Iyengar, Bhovi	0	0.4020
	M2a	N.A	Lambadi, Hill Kolam, Katkari, Dongri Bhil	0	0.6110
	M4a	Persian	N.A	0.8560	0
	M2b	N.R	Korku, Hill Kolam	0	0.9400
	M35b	Persian	N.A	0.3860	0
	M27b	Indian Shia Muslim	N.A	0.4220	0
Α	A2v	Persian	N.A	0.4690	0
F	F1g	Kurd, Turkmen	N.A	0.9970	0
z	Z1a	Qashqai, Persian	N.A	0.2470	0

			The second s
Table: Clustering	of Parsis with	population of Persian	and Indian descent

Table 3 : Results of the Phylogenetic clustering of the 100 Parsis mitochondrial genomes with 352 mitochondrial genomes of Iranian origin and 100 mitochondrial genomes of relic tribes of Indian origin through Neighbour Joining method. BS indicates Boot-Strap values between each sample. *N.A. indicates *No Association*, indicating a lack of representation of samples in the specific sub-haplogroup

Table 4: Variants associated with haplogroup specific Zoroastrian Parsi Mitochondrial Reference Genome (n=7) and Zoroastrian Parsi Mitochondrial Consensus Genome (n=1) mitochondrial genome sequences

Consensus Sequence	Number of Variants	Variants
AGENOME-ZPMCG-V1.0	31	T65TT, A73G, A263G, C309CCCT, T310C, T489C, G513GCA, A567ACCCCCC, A750G, A1438G, A2706G, A3158AT, A4769G, C7028T, A8701G, A8860G, T9540C, A10398G, C10400T, T10873C, G11719A, C12705T, C14766T, T14783C, G15043A, G15301A, A15326G, C16169CC, A16182AC, C16223T, T16519C
AGENOME-ZPMRG-A2v-V1.0	11	A263G, C309CCT, T310C, A750G, A1438G, A4769G, A8860G, C11881T, A15326G, C16168T, C16239T
AGENOME-ZPMRG-HV-V1.0	26	T72C, A73G, T152C, T195C, A263G, C309CCT, T310C, A750G, A1438G, A2706G, A4769G, T5075C, C6104T, G6179A, C7028T, T7193C, A8860G, A9336G, T10410C, G11016A, T11935C, C12061T, A15326G, T15792C, T16217C, A16309G
AGENOME-ZPMRG-M-V1.0	29	T65TT, A73G, A263G, C309CCCT, T310C, T489C, A567ACCCC, A750G, A1438G, A2706G, A4769G, C7028T, A8701G, A8860G, T9540C, A10398G, C10400T, T10873C, G11719A, C12705T, C14766T, T14783C, G15043A, G15301A, A15326G, C16169CC, A16182AC, C16223T, T16519C
AGENOME-ZPMRG-U-V1.0	25	A73G, A263G, C309CCCT, T310C, G499A, G513GCA, A567ACCCCCCC, A750G, A1438G, A1811G, A2706G, A3158AT, A4769G, C7028T, A8860G, C11332T, A11467G, G11719A, A12308G, G12372A, C14620T, C14766T, A15326G, T16189TT, T16519C
AGENOME-ZPMRG-T-V1.0	28	A73G, A263G, C309CCT, T310C, G709A, A750G, A1438G, G1888A, A2706G, T4216C, A4769G, A4917G, C7028T, G8697A, A8860G, T10463C, A11251G, G11719A, G13368A, C14766T, G14905A, A15326G, C15452A, A15607G, G15928A, T16126C, C16294T, T16519C
AGENOME-ZPMRG-F1g-V1.0	32	A73G, A248d, A263G, C315CC, CA514d, A750G, A1438G, C2389T, A2706G, T3398C, C3970T, T3999C, A4769G, T6392C, G6962A, C7028T, A8589G, A8860G, G10310A, T10609C, G11719A, G12406A, C12882T, G13928C, C14766T, A15326G, T15916C, A16183C, T16189C, C16193CC, T16304C, T16519C
AGENOME-ZPMRG-Z-V1.0	33	A73G, C151T, T152C, A263G, C315CC, T489C, A750G, A1438G, A2072d, A2706G, A4769G, C7028T, A8701G, A8860G, T9540C, A10149T, A10398G, C10400T, C10556T, T10873C, G11719A, G12007A, C12705T, C14766T, T14783C, G15043A, G15301A, A15326G, G15346A, T15784C, C16223T, T16311C, T16519C

Table 4: List of unique variants associated with the Haplogroup specific Zoroastrian Parsi Mitochondrial Reference Genomes (ZPMRG) for A2v, HV, M, U, T, F1g, Z and overall unique variants in the Zoroastrian Parsi Mitochondrial Consensus Genome (ZPMCG)

 Table 5: Variants associated with Zoroastrian Parsi Mitochondrial Reference Genome (ZPMRG) and unique variants of each ZPMRG compared to Zoroastrian Parsi Mitochondrial Consensus Genome (ZPMCG)

AGENOME-							
ZPMRG-	ZPMRG-	ZPMRG-M-	ZPMRG-T-	ZPMRG-U-	ZPMRG-F-	ZPMRG-F-	
A2v-V1.0	HV-V1.0	V1.0	V1.0	V1.0	V1.0	V1.0	
C11881T	A16G		C6G	A21G	A248d	C151T	
C16168T	T72C		G709A	G499A	CA514d	A2072d	
C16239T	T195C		G1888A	A1811G	C2389T	C10556T	
	T5075C		T4216C	C11332T	T3398C	G12007A	
	C6104T		A4917G	A11467G	C3970T	G15346A	
	G6179A		G8697A	A12308G	T3999C	T15784C	
	T7193C		T10463C	G12372A	T6392C	T16311C	
	A9336G		A11251G	C14620T	G6962A		
	T10410C		G13368A		A8589G		
	G11016A		G14905A		G10310A		MAPLOGROUPS
	T11935C		C15452A		T10609C		ER 15
	C12061T		A15607G		G12406A		
	T15792C		G15928A		C12882T		
	T16217C		T16126C		G13928C		
	A16309G		C16294T		T15916C		
					A16183C		Phi P H P I V I V
					C16193CC		
					T16304C		HAPLOGROUPS

Table 5: (A) Unique Variants found in the haplogroup specific Reference Genomes (ZPMRG) compared to the Zoroastrian-Parsi Consensus Genome (AGENOME-ZPMCG-V1). The histogram (right) lists the exact number of variants in each ZPMRG compared to ZPMCG

mt-tRNA	Variation	Probability_of_ pathogenicity	Classification	Frequency %	Haplogroup	Disease association
Phe	T593C	0.16	Neutral	0.06	M52b	Non-syndromic hearing loss (Reported)
Val	G1644A	0.67	Pathogenic	0.01	U4b	LS/HCM/MELAS (Reported)
Val	T1654C	0.12	Neutral	0.01	M3a	
Met	T4454C	0.13	Neutral	0.02	M5a	Possible contributor to mito dysfuntion / Hypertension (Reported)
Asp	G7521A	0.46	Likely neutral	0.01	U4b	
Asp	T7561C	0.33	Neutral	0.01	U7a	
Asp	T7581C	0.42	Likely neutral	0.01	U1a	
Arg	T10410C	0.17	Neutral	0.14	Hv2a	
Arg	T10463C	0.31	Neutral	0.04	T1a,T2g,T2i	
His	A12172G	0.53	Likely pathogenic	0.01	U4b	
His	C12191G	0.11	Neutral	0.01	M27b	
Leu(CUN)	A12279G	0.37	Likely neutral	0.06	M52b	
Leu(CUN)	A12308G	0.41	Likely neutral	0.21	U4b,U7a	Stroke, CM, CPEO, Breast/Renal/Prostate cancer risk, Altered brain pH(Reported)
Glu	A14696G	0.26	Neutral	0.01	A2v	Progressive Encephalopathy (Reported)
Thr	A15907G	0.23	Neutral	0.03	U2e	
Thr	T15908C	0.5	Likely pathogenic	0.01	M33a	Deaf Helper mutation (Reported)
Thr	T15916C	0.33	Likely neutral	0.01	F1g	

Table 6: mt-t-RNA variants in our study and their disease association

Table 6: Analysis of the occurrence of the 420 variants in the tRNA and their disease associations annotated with the PON-mt-tRNA database. A frequency score ≥ 0.5 – pathogenic, =0.5 – likely pathogenic, <0.5 – neutral

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