The Mitonuclear Dimension of Neanderthal and Denisovan Ancestry in Modern Human Genomes

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Accepted: June 21, 2017

Abstract

Some human populations interbred with Neanderthals and Denisovans, resulting in substantial contributions to modern-human genomes. Therefore, it is now possible to use genomic data to investigate mechanisms that shaped historical gene flow between humans and our closest hominin relatives. More generally, in eukaryotes, mitonuclear interactions have been argued to play a disproportionate role in generating reproductive isolation. There is no evidence of mtDNA introgression into modern-human populations, which means that all introgressed nuclear alleles from archaic hominins must function on a modern-human mitochondrial background. Therefore, mitonuclear interactions are also potentially relevant to hominin evolution. We performed a detailed accounting of mtDNA divergence among hominin lineages and used population-genomic data to test the hypothesis that mitonuclear incompatibilities have preferentially restricted the introgression of nuclear genes with mitochondrial functions. We found a small but significant underrepresentation of introgressed Neanderthal alleles at such nuclear loci. Structural analyses of mitochondrial enzyme complexes revealed that these effects are unlikely to be mediated by physically interacting sites in mitochondrial and nuclear gene products. We did not detect any underrepresentation of introgressed Denisovan alleles at mitochondrial-targeted loci, but this may reflect reduced power because locus-specific estimates of Denisovan introgression are more conservative. Overall, we conclude that genes involved in mitochondrial function may have been subject to distinct selection pressures during the history of introgression from archaic hominins but that mitonuclear incompatibilities have had, at most, a small role in shaping genome-wide introgression patterns, perhaps because of limited functional divergence in mtDNA and interacting nuclear genes.

Key words: coevolution, cytonuclear interactions, hybridization, human evolution, OXPHOS enzymes, speciation.

Introduction

Understanding the mechanisms that lead to speciation is one of the defining challenges in the field of evolutionary biology, and numerous causes of reproductive isolation have been identified (Dobzhansky 1937; Coyne and Orr 2004). Natural curiosity about our own origins has spurred debate about how these mechanisms have shaped the evolution and diversification of hominins (Stringer 2002; Wood and Boyle 2016).

The advent of technologies to extract and sequence ancient DNA samples and the rapid increase in availability of population genomic data have revealed that modern-human populations that expanded out of Africa subsequently underwent hybridization and genetic admixture with related hominin lineages that were already present in Asia and Europe (Green et al. 2010; Reich et al. 2010; Meyer et al. 2012; Prüfer et al. 2014; Sankararaman et al. 2014, 2016; Vernot and Akey 2014; Vernot et al. 2016). Interbreeding with Neanderthals resulted in substantial introgression into modern human populations such that ~2–4% of genome content in non-African individuals is now derived from Neanderthals (Green et al. 2010). Additional interbreeding appears to have occurred between modern humans and Denisovans, an enigmatic lineage of hominins known only from limited fossil remains found in Denisova Cave in the Altai Mountains of Siberia (Krause et al. 2010; Sawyer et al. 2015). Denisovan-derived alleles make up a substantial portion (up to ~5%) of the genomes of modern humans from Melanesia and surrounding regions but appear to have made little or no contribution to other populations (Reich et al. 2010).

There is large variation across the human genome in the extent of introgression from Neanderthals and Denisovans...
(Sankararaman et al. 2014, 2016; Vernot and Akey 2014; Vernot et al. 2016), which is a common theme in hybridizing organisms (Baack and Rieseberg 2007). Substantial variation is expected even under neutral models of introgression, but differences can be magnified by selection for or against certain foreign alleles (Harrison and Larson 2014, 2016; Payseur and Rieseberg 2016; Schumer and Brandvain 2016). Much of the recent interest in introgression from archaic hominins has focused on loci with unexpectedly high frequencies of foreign alleles because they provide evidence that modern human populations acquired beneficial genetic variants from Neanderthals and Denisovans in the process of adapting to novel environments outside of Africa (Racimo et al. 2015).

Even so, regions of the genome that exhibit atypically low rates of introgression are also noteworthy, as they present the opportunity to infer causes of restricted gene flow and to draw parallels with more general mechanisms of reproductive isolation. For example, Neanderthal- and Denisovan-derived alleles are substantially underrepresented on the modern human X-chromosome, suggesting that X-linked loci might have contributed to early stages of reproductive isolation among hominin lineages (Sankararaman et al. 2016). This “large X-effect” on hybrid inviability and sterility is a common pattern from the field of speciation genetics (Presgraves 2008) and is often attributed to the exposure of recessive alleles because of hemizygosity of X-linked genes in males (Qvarnsjö and Bailey 2009; cf. Hu and Filatov 2016). In addition, there is evidence that levels of introgression from archaic hominins are lower for genes near structural rearrangements (Rogers 2015) and for genes that are preferentially expressed in the testes (Sankararaman et al. 2016). These patterns are consistent with findings from other organisms that structural rearrangements can act as a barrier to gene flow (Rieseberg et al. 1999; Rieseberg 2001; Feder et al. 2003; Lowry and Willis 2010) and that tests-specific genes are often involved in hybrid male infertility (Wu and Davis 1993; Presgraves 2008).

Another hypothesis that has emerged from the study of reproductive isolation in eukaryotes is that mitonuclear incompatibilities play a disproportionate role in the creation of species boundaries because hybridization can disrupt coadapted combinations of mitochondrial and nuclear alleles (reviewed in Gershoni et al. 2009; Burton and Barreto 2012; Hill 2016; Sloan et al. 2017). Mitonuclear coadaptations are thought to be driven by the rapid evolution of the mitochondrial genome and subsequent coevolutionary responses in nuclear genes that encode mitochondrially targeted proteins (N-mt genes) (Osada and Akashi 2012; Havird et al. 2017). Although mitochondria retain their own genome, the vast majority of the >1000 proteins that function in mitochondria are encoded in the nucleus (Gray 2015). Coevolutionary dynamics are expected to be the most intense for N-mt proteins that interact directly with mitochondrial gene products in multisubunit complexes such as oxidative phosphorylation (OXPHOS) enzymes and mitochondrial ribosomes (Rand et al. 2004). By incorporating models of molecular structures, various studies have investigated the possibility that physically interacting residues in mitochondrial- and nuclear-encoded subunits can undergo coevolutionary changes that lead to coadapted mitonuclear genotypes (Schmidt et al. 2001; Melvin et al. 2008; Osada and Akashi 2012; Aledo et al. 2014; Havird et al. 2015). However, researchers have only recently begun using genomic data sets to examine the effect of selection on maintaining associations between coadapted combinations of mitochondrial haplotypes and N-mt alleles during genetic admixture/hybridization (Trier et al. 2014; Bar-Yaakov et al. 2015; Beck et al. 2015; Sloan et al. 2015; McKenzie et al. 2016; Baris et al. 2017; Morales et al. 2017; Runemark et al. 2017). These studies have produced mixed results about the extent to which selection can preferentially act on N-mt loci relative to the rest of the nuclear genome. More generally, the hypothesis that mitonuclear interactions play a common or disproportionate role in speciation remains controversial (Eyre-Walker 2017) and seemingly at odds with evidence of widespread introgression of mtDNA in many hybrid zones (Sloan et al. 2017). Therefore, addressing this hypothesis is a pressing biological challenge.

The potential for mitonuclear interactions among hominins is of interest because, unlike in the nuclear genome, there has not been any detectable mtDNA introgression from Neanderthals or Denisovans into modern human populations (Krings et al. 1997; Serre et al. 2004). Regardless of what has caused this lack of mtDNA introgression, one consequence is that all introgressed Neanderthal and Denisovan nuclear alleles must function on a modern-human mitochondrial background, and it has been proposed that divergence in mitochondrial gene products between modern humans and archaic hominins may have altered physical interactions with N-mt proteins (Green et al. 2008). Accordingly, if coadapted combinations of mitochondrial and nuclear alleles have evolved within hominin lineages, incompatibilities between modern-human mtDNA and Neanderthal/Denisovan nuclear alleles may have preferentially restricted introgression at nuclear loci that encode N-mt proteins and are involved in mitonuclear interactions. Here, we investigate the hypothesis that mitonuclear incompatibilities have contributed to the earliest stages of reproductive isolation between modern humans and related hominins and shaped patterns of genetic admixture following hybridization.

Materials and Methods

Analysis of mtDNA Polymorphism and Divergence

To compare patterns of molecular evolution in mtDNA across hominin lineages, we obtained published mitochondrial genome sequences for 6 Neanderthal samples (NCBI accessions FM865407-FM865411 and AM948965), 3 Denisovan
samples (KT780370, FN673705, FR695060), and 54 samples from globally diverse lineages of modern humans (AF346963-AF347015, NC_012920), which included the revised Cambridge Reference Sequence (Andrews et al. 1999) and the full sampling performed by Ingman et al. (2000). We also included the mitochondrial genome sequence (NC_023100) from an ancient hominin (~400,000 years old) that was recovered from the Sima de los Huesos (SdhH) site in Spain (Meyer et al. 2014). Nuclear, morphological, and geographical data indicate that the SdhH sample is an early representative of the Neanderthal lineage (Meyer et al. 2016), but its mitochondrial haplotype is more closely related to Denisovan mtDNA (Meyer et al. 2014). The SdhH sample is not a primary focus of this study, but we included it in our mtDNA analysis to achieve a thorough sampling of available mitochondrial genomes from archaic hominins.

We aligned all sampled mitochondrial genome sequences with MAFFT v7.305b, using the FFT-NS-i algorithm with a maximum of 1000 iterations (supplementary file S1, Supplementary Material online) (Katoh and Standley 2013). After visual inspection and manual alignment editing, we used a custom Python script (available at https://github.com/jsharbrough/polymorphismsSubstitutions) to identify polymorphisms segregating within Neanderthals, Denisovans, and modern humans as well as fixed differences among the sampled hominin lineages (treating SdhH as a separate lineage; supplementary file S2, Supplementary Material online). To account for sites with multiple hits, we assumed that the number of changes was the minimum required to explain the data. In protein-coding sequences, we further assumed that any codons with multiple hits resulted from the minimum number of nonsynonymous changes needed to explain the data. We generated a pairwise mismatch distribution by quantifying the number of nucleotide substitutions between each modern human mitochondrial haplotype and each other modern human, Neanderthal, or Denisovan sample from the above alignment.

The primary goal of our mtDNA analysis was to distinguish ancestral divergence between modern humans and related hominin lineages from more recent changes segregating within these populations. For the purposes of this analysis, we considered variants to be fixed differences if they were not segregating in our sample of 54 modern human mitochondrial genomes even if they have been identified as low-frequency variants in larger samples of human mitochondrial diversity. To assess the effects of this sampling approach, we repeated the analysis of sequence polymorphism and divergence using two larger modern human mtDNA data sets: 114 mitochondrial genomes from Mishmar et al. (2003) and 9862 mitochondrial genomes from Levin et al. (2013). We also directly compared Neanderthal and Denisovan sequences to a reconstruction of the ancestral modern human mtDNA sequence from Behar et al. (2012).

Neanderthal Introgression Analysis

To test the hypothesis that mitonuclear incompatibilities have preferentially restricted the introgression of N-mt alleles from Neanderthals, we used the “introgression score” calculated by Sankararaman et al. (2014), with raw gene-specific values kindly provided by the authors of the original study (S. Sankararaman, pers. comm.). The introgression score was determined from a conditional random field analysis and based on the marginal probability of Neanderthal ancestry averaged across the entire length of a gene and across all sampled individuals, which were taken from the 1000 Genomes Project (1000 Genomes Project Consortium 2012). Distinct estimates were produced for European populations, Asian populations, and a combined European/Asian data set in the original analysis (Sankararaman et al. 2014). We used the set of high-confidence N-mt genes (“Tmt”), in the MitoCarta 2.0 classification (Pagliarini et al. 2008) to compare rates of introgression against all other nuclear protein-coding genes. We separated these high-confidence N-mt genes into two groups, distinguishing a subset of 150 genes that encode proteins that interact with mitochondrial gene products in multisubunit OXPHOS and ribosomal complexes from the rest of the N-mt gene set (following the classification scheme in Sloan et al. 2015). Differences in rates of introgression among gene classes were assessed with nonparametric Kruskal–Wallis rank sum tests.

To assess whether differences in levels of introgression may reflect a correlation between mitochondrial targeting and overall functional importance, we used two proxies for the functional importance of each gene. First, we used estimates of transcript abundance based on the premise that highly expressed genes are more functionally constrained (Zhang and Yang 2015). We obtained two separate measures of transcript abundance by averaging across 16 different human tissues from the Illumina Body Map data set (https://www.ebi.ac.uk/gxa/experiments/E-MTAB-513) and by averaging across median RPKM values from each of 53 different human tissues from the GTEx V6p release data set (The GTEx Consortium 2013). Second, we used gene-specific estimates for rates of nonsynonymous substitutions (dN) between humans and chimpanzees (The Chimpanzee Sequencing and Analysis Consortium 2005), with the expectation that slower rates of protein evolution are indicative of greater functional importance. Data sets were log10-transformed after adding 0.001 to introgression scores and dN values and adding 0.01 to mean expression values. After transformation, Pearson correlation coefficients were calculated between introgression score and each of these proxies for functional importance. We also extracted residuals from linear models and used them to repeat the above Kruskal–Wallis tests as a means for controlling for effects of expression or rate of sequence evolution. All statistical analyses were implemented in R v3.1.2 (R Core Team 2014), and all data plots were
generated with the ggplot2 package (Wickham 2009). Introggression data are provided in (supplementary file S3, Supplementary Material online).

Denisovan Introggression Analysis
To perform a similar analysis of introgression from Denisovans, we used haplotype calls from a study of 35 Melanesian genomes (Vernot et al. 2016). The physical locations of individual Denisovan-derived haplotypes were obtained from supporting data sets posted on the personal website of one of the authors of the original study (http://akeylab.gs.washington.edu/vernot_et_al_2016_release_data/). We calculated frequencies of Denisovan-derived alleles for each nucleotide position in the hg19 human genome assembly (i.e., anywhere from 0 to 70 of the haplotypes from the 35 diploid Melanesian genomes). Gene-specific estimates were then calculated by averaging frequencies across the full length of each gene (supplementary file S3, Supplementary Material online). Using the same statistical approach described earlier for Neanderthal introgression, we compared rates of introgression among different gene classes. To assess the consequences of methodological differences between the Neanderthal (Sankararaman et al. 2014) and Denisovan (Vernot et al. 2016) data sets, we repeated our analyses using haplotype calls for Neanderthal-derived alleles in East Asian and European populations that were produced with the same methodology as the Denisovan haplotype predictions (supplementary file S1, Supplementary Material online). We applied the same custom Python package that was used for the mtDNA comparative analysis to identify all homozygous substitutions among the three hominin samples. Identified variants were cross-referenced against dbsNP (Sherry et al. 2001) to determine whether they were segregating within modern human populations, using dbsNP identifiers from the original VCF files. For variants that were unique to the Neanderthal sample, we assessed whether they were fixed within Neanderthals by comparing against additional Neanderthal genomic data sets (Green et al. 2010). UCSF Chimera was used to classify identified variants in N-mt proteins as either contact or noncontact sites as described earlier.

Identification of Contact Residues within the Structures of Mitonuclear Enzyme Complexes
To assess whether fixed differences between modern humans and ancient hominin lineages in mitochondrially encoded proteins and rRNAs may have affected mitonuclear interactions by altering interfaces with N-mt proteins, we mapped substitutions onto published mammalian structural models of OXPHOS complexes and the mitochondrial ribosome. The PDB accessions used for NADH dehydrogenase, cytochrome bc1, cytochrome c oxidase, ATP synthase, and the mitochondrial ribosome were 5LNK, 1BGY, 1V54, 5ARA, and 3J9M, respectively (Iwata et al. 1998; Tsukihara et al. 2003; Amunts et al. 2015; Zhou et al. 2015; Fiedorczuk et al. 2016). We used UCSF Chimera v1.11.2 (Pettersen et al. 2004) to visualize structures and identify mitonuclear contact residues, that is, those that occur at the interfacing surface between proteins or rRNAs encoded by the mitochondrial and nuclear genomes. To identify contact residues, we used the “find clashes/contacts” tool with default parameters, except that the minimum overlap was set to $\geq -1$ Å to increase sensitivity.

Analysis of Sequence Divergence in N-mt Genes
To identify substitutions in N-mt genes encoding proteins that contact variable sites in mitochondrial subunits, we used genotype calls (relative to hg19) from a representative nuclear genome sequence from Neanderthals (Prüfer et al. 2014), Denisovans (Meyer et al. 2012), and modern humans (HuRef; Levy et al. 2007). We obtained RefSeq sequences for each N-mt gene from the UCSC genome browser (http://genome.ucsc.edu) and extracted FASTA sequences from VCF files corresponding to the Neanderthal (http://cdna.eva.mpg.de/neandertal/altai/AltaiNeandertal/VCF/) and Denisovan (http://cdna.eva.mpg.de/denisova/VCF/hg19_1000g/) data sets, using a custom tool called vcf2Fasta in the seqTools_v2.2 package (https://github.com/jsharbrough/seqTools). Variants were filtered based on basecall quality ($\geq Q30$) and depth ($\geq 10x$). We mapped N-mt RefSeq genes to the HuRef alternate genome assembly (ftp://ftp.ncbi.nih.gov/genomes/Homo_sapiens/ARCHIVE/BUILD.37.3/Assembled_chromosomes/seq/) using BWA-MEM v0.7.15-r1140 (Li and Durbin 2010) with pairing and mate-pair rescuing turned off. We used these mapping data to determine N-mt gene positions within the HuRef genome and extracted the specified regions using the samtools fai2x function implemented in SAMtools v1.3.1 (Li et al. 2009).

Protein-coding N-mt gene sequences were extracted from the mapped Neanderthal, Denisovan, and HuRef sequences based on annotated RefSeq exon coordinates. We then used MAFFT v7.305b to align these sequences with the FFT-NS-i algorithm and a maximum of 1000 iterations (supplementary file S1, Supplementary Material online). We applied the same custom Python package that was used for the mtDNA comparative analysis to identify all homozygous substitutions among the three hominin samples. Identified variants were cross-referenced against dbsNP (Sherry et al. 2001) to determine whether they were segregating within modern human populations, using dbsNP identifiers from the original VCF files. For variants that were unique to the Neanderthal sample, we assessed whether they were fixed within Neanderthals by comparing against additional Neanderthal genomic data sets (Green et al. 2010). UCSF Chimera was used to classify identified variants in N-mt proteins as either contact or noncontact sites as described earlier.

Results
mtDNA Polymorphism and Divergence in Modern Human, Neanderthals, and Denisovans
The original publication of Neanderthal and Denisovan mitochondrial genomes revealed that the level of sequence divergence between modern humans and these related hominins greatly exceeds the divergence between any pair of extant human mitochondrial haplotypes (fig. 1; Green et al. 2008; Krause et al. 2010). In addition, these sequencing efforts
revealed that mitochondrial haplotypes of modern humans and Neanderthals share a more recent common ancestor with each other than with Denisovans (Krause et al. 2010), which is reflected in the much greater pairwise divergences with Denisovans (fig. 1 and table 1). Although early efforts were made to describe the functional changes in the mitochondrial genome that separate Neanderthals and modern humans (Green et al. 2008), it is worth revisiting these analyses in light of the subsequent increase in the availability of mitochondrial genomes from Neanderthals and Denisovans.

Despite the rapid rate of sequence evolution in mtDNA, the number of fixed nonsynonymous changes between modern humans and related hominins is small (table 1). Only 22 fixed amino acid substitutions distinguish modern humans and Neanderthals in the set of 13 protein-coding sequences found in the mitochondrial genome. Surprisingly, comparisons between modern humans and Denisovans revealed a nearly identical level of divergence in amino acid sequences (24 fixed differences; table 1) even though overall mtDNA sequence divergence is nearly twice as high (fig. 1). This pattern is consistent with previous observations that the ratio of nonsynonymous to synonymous changes (\(d_\text{N}/d_\text{S}\)) is higher in Neanderthals and modern humans than in other branches of the hominin tree (see supplementary fig. 5 in Krause et al. 2010). Comparisons involving modern humans show smaller numbers of fixed differences (table 1), in part because many variants are polymorphic within modern humans (supplementary table S2, Supplementary Material online).

Individual protein-coding genes have anywhere from zero to six fixed nonsynonymous changes between modern humans and archaic hominins (table 1). Total sequence divergence is even lower in structural RNA genes (tRNAs and rRNAs) than in protein-coding genes (table 1). There was also variation within these structural RNA genes, as the frequency of fixed substitutions was significantly different across the various structural regions within tRNAs (\(P = 3.8 \times 10^{-5}\); Fisher’s Exact Test), with most changes found in the TΨC loop (fig. 2 and supplementary table S3, Supplementary Material online). Not surprisingly, changes were generally not observed at sites known to be crucial for tRNA identity and recognition by the nuclear-encoded aminoacyl-tRNA synthetases (aaRSs) that charge tRNAs with the proper amino acid. However, the identified variant in TRNL2 has been found to affect charging rates (Hao et al. 2005), and TRNW has experienced a change in the acceptor stem, which is typically involved in aaRS interactions (fig. 2). Although none of these tRNA substitutions were segregating in our sample of 54 modern human mitochondrial genomes, some of these variants have been identified at low frequencies in human populations or in heteroplasmic individuals and have been associated with disease phenotypes (Zhu et al. 2009; Lu et al. 2010).

For the purposes of this analysis, variants were classified as fixed differences if they were not segregating within our samples of 54 modern humans, 6 Neanderthals, and 3 Denisovans. We investigated the effects of sampling depth on our estimates by repeating analyses with additional human mtDNA data sets of varying sizes. As expected, performing comparisons directly against the reconstructed ancestral sequence for modern humans (Behar et al. 2012) led to an increased number of fixed differences because some segregating polymorphisms are the result of derived, homoplasious changes within the modern-human genealogy (supplementary tables S4 and S5, Supplementary Material online). In accordance with this pattern, using larger samples of human mtDNA diversity (Mishmar et al. 2003; Levin et al. 2013) led to reclassification of some fixed differences as segregating polymorphisms (supplementary tables S4 and S5, Supplementary Material online). For example, we found 26 fixed nonsynonymous differences between the reconstructed ancestral modern-human sequence and Neanderthals, and this value declined to 22, 17, and 1, when using modern-human data sets consisting of 54, 114, and 9862, respectively.
Underrepresentation of N-mt Genes in Regions of Neanderthal Introgression

If mitonuclear incompatibilities lowered reproductive fitness in hybrids, then the nuclear loci involved in these incompatibilities should have experienced reduced introgression relative to background levels in the genome. We reasoned that such effects should preferentially involve N-mt genes, so we used gene-specific estimates of introgression from Neanderthals (Sankararaman et al. 2014) to compare N-mt genes against all other nuclear genes. We found that N-mt genes have undergone significantly less introgression into both Asian ($P = 5 \times 10^{-5}$) and European populations ($P = 2 \times 10^{-8}$; fig. 3). These results lend some support to the mitonuclear-incompatibility hypothesis and are consistent with findings that certain mitochondrial-related functional classes are significantly enriched among the set of genes with low levels of introgression (see supplementary table 6.1 in Sankararaman et al. 2014). Only a subset of N-mt genes encode proteins that

### Table 1
Fixed Nucleotide Substitutions between Modern Humans (MH), Neanderthals (Nean), and Denisovans (Den)

| Gene/region Type | Length (aligned) | MH—Nean | MH—Den | Nean—Den |
|------------------|------------------|---------|--------|----------|
| ATP6 Protein     | 681              | 4 (2)   | 8 (3)  | 12 (5)   |
| ATP8 Protein     | 207              | 5 (1)   | 7 (1)  | 5 (0)    |
| COI Protein      | 1,542            | 11 (0)  | 18 (1) | 26 (2)   |
| COII Protein     | 684              | 5 (3)   | 18 (4) | 14 (2)   |
| COIII Protein    | 784              | 5 (2)   | 15 (2) | 17 (2)   |
| CYTB Protein     | 1,141            | 13 (4)  | 20 (3) | 26 (6)   |
| ND1 Protein      | 956              | 6 (1)   | 16 (1) | 21 (2)   |
| ND2 Protein      | 1,044            | 7 (1)   | 19 (2) | 18 (4)   |
| ND3 Protein      | 346              | 6 (1)   | 7 (1)  | 9 (0)    |
| ND4 Protein      | 1,378            | 6 (0)   | 14 (0) | 22 (1)   |
| ND4L Protein     | 297              | 2 (1)   | 4 (1)  | 5 (0)    |
| ND5 Protein      | 1,812            | 15 (6)  | 32 (5) | 41 (6)   |
| ND6 Protein      | 525              | 1 (0)   | 6 (0)  | 9 (1)    |
| RNR1 rRNA        | 954              | 1       | 5      | 11       |
| RNR2 rRNA        | 1,561            | 4       | 9      | 9        |
| TRNA tRNA        | 69               | 0       | 0      | 0        |
| TRNC tRNA        | 67               | 0       | 0      | 0        |
| TRND tRNA        | 68               | 0       | 0      | 0        |
| TRNE tRNA        | 69               | 0       | 1      | 1        |
| TRNF tRNA        | 71               | 0       | 1      | 1        |
| TRNG tRNA        | 68               | 0       | 0      | 0        |
| TRNH tRNA        | 69               | 1       | 2      | 1        |
| TRNI tRNA        | 69               | 0       | 1      | 1        |
| TRNK tRNA        | 70               | 0       | 0      | 0        |
| TRNL1 tRNA       | 75               | 0       | 1      | 2        |
| TRNL2 tRNA       | 71               | 0       | 1      | 1        |
| TRNM tRNA        | 68               | 0       | 1      | 1        |
| TRNN tRNA        | 73               | 1       | 0      | 1        |
| TRNP tRNA        | 68               | 0       | 0      | 0        |
| TRNQ tRNA        | 72               | 0       | 0      | 0        |
| TRNR tRNA        | 65               | 0       | 0      | 0        |
| TRNS1 tRNA       | 69               | 0       | 0      | 0        |
| TRNS2 tRNA       | 59               | 0       | 1      | 1        |
| TRNT tRNA        | 66               | 0       | 1      | 1        |
| TRNV tRNA        | 69               | 0       | 0      | 0        |
| TRNW tRNA        | 68               | 0       | 2      | 2        |
| TRNY tRNA        | 66               | 1       | 1      | 0        |
| D-Loop Noncoding | 1,142            | 10      | 20     | 40       |
| Other intergenic Noncoding | 95 | 1 | 0 | 1 |
| Whole genome 16,599 | 105 (22) | 229 (24) | 297 (31)* |

*Note.—For coding regions, nonsynonymous changes are in parentheses. Corresponding divergences with the SdhH sample are reported in supplementary table S1, Supplementary Material online.

*aThe sum of individual gene counts exceeds the whole genome count for comparisons involving Denisovans because of a substitution in a region of overlap between ATP6 and ATP8.

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Denisovans (blue), or both (purple). The primary sequences are from the revised Cambridge Reference Sequence (Andrews et al. 1999), and variants in archaic hominins are indicated next to each variable site. Secondary structures were generated with VARNA v3.93 based on published structural models (Darty et al. 2009).

Because regions of the genome that are subject to greater functional constraint exhibit lower rates of introgression (Sankararaman et al. 2014, 2016), one potential explanation for the lower rates of introgression for N-mt genes is that they are more functionally important on average than other nuclear genes. We used two proxies for functional importance to assess this possibility: 1) levels of transcript abundance and 2) level of amino acid divergence relative to chimpanzees. Using measures of transcript abundance from the Illumina Body Map data set, we found a significant, but very weak, negative association with introgression score in both Asian ($P=0.0073$) and European populations ($P=0.0007$;
Analysis of Denisovan Introggression

If mitonuclear incompatibilities were responsible for the lower rate of introgression of N-mt genes from Neanderthals, we might expect a stronger historical effect on introgression from Denisovans because of the much higher level of mtDNA sequence divergence between modern humans and Denisovans (fig. 1). However, even in Melanesian populations, which have a sizeable amount of Denisovan-derived sequence, obtaining accurate gene-specific estimates of Denisovan introgression poses a technical challenge. It is difficult to confidently distinguish Denisovan alleles in a background that has already acquired alleles via admixture with at least one other archaic hominin lineage (Neanderthals). See Sankararaman et al. (2016) for discussion of these technical challenges.

Nevertheless, efforts have been made to identify Denisovan-derived haplotypes in a set of 35 published Melanesian genome sequences (Vernot et al. 2016). We used these haplotype calls to calculate an average proportion of Denisovan ancestry in the Melanesian population for each nuclear gene. We obtained a global average level of introgression of 0.0063, which is far lower than the estimates of Neanderthal introgression used in our previous analysis (mean introgression scores of 0.031 and 0.026 for Asian and European populations, respectively; Sankararaman et al. 2014). These gene-specific estimates of Denisovan introgression also produced an extremely high proportion of zero values (87.3% of genes). By contrast, zero values were relatively rare for Neanderthal introgression scores in Asian (5.1%) and European (5.6%) populations (Sankararaman et al. 2014). Given the evidence that Denisovan-derived alleles represent ~5% of Melanesian genomes, which is more than the current contribution of Neanderthals to any modern human population (Green et al. 2010), it is likely that the values we used for Denisovan introgression are substantial underestimates, owing to the conservative nature of the published Denisovan haplotype calls (Vernot et al. 2016).

Using these estimates of Denisovan introgression into Melanesian genomes, we performed the same comparison between N-mt genes and other nuclear genes as described earlier for Neanderthal introgression. In this case, however, we did not find any significant differences in levels of introgression for N-mt genes relative to the rest of the genome (P = 0.26, Kruskal-Wallis test; supplementary fig. S2, Supplementary Material online). To assess whether this result represents a true biological difference between the histories of Denisovan versus Neanderthal introgression or a consequence of the more conservative calling of Denisovan-derived alleles, we analyzed a set of Neanderthal-derived haplotypes in Asians and Europeans from Vernot et al. (2016) that were obtained with the same methodology used to analyze

![Figure 3](https://example.com/figure3.png)

**Fig. 3.** A violin plot summarizing the distribution of gene-specific Neanderthal introgression scores in Asian (left) and European (right) populations of modern humans from Sankararaman et al. (2014). Separate distributions are shown for N-mt genes that contribute to mitonuclear OXPHOS and ribosomal complexes (orange), other N-mt genes (blue), and all other nuclear genes (gray). The median value for each distribution is indicated with a red bar. Overall statistical significance was assessed with a Kruskal-Wallis test (see reported P values), and lower case letters summarize significant differences that emerged from post hoc pairwise comparisons between each distribution. For the purposes of plotting on a log-scale, a minimum introgression score was set at 0.001.
Denisovan introgression in Melanesians. These gene-specific estimates of Neanderthal introgression exhibited a strongly significant positive correlation with estimates from Sankararaman et al. (2014) that were analyzed above, demonstrating a general consistency between the methods (\(P < 0.001\); \(r = 0.61\) and 0.57 for Asian and European data sets, respectively; supplementary fig. S3, Supplementary Material online). However, the data from Vernot et al. (2016) yielded lower estimates and a much larger proportion of zero values even though they were derived from the same Asian and European populations as the estimates from Sankararaman et al. (2014). When we used these new estimates to compare levels of Neanderthal introgression for N-mt genes to background rates in the genome, the differences between N-mt genes and other nuclear genes were marginally significant at best (\(P = 0.03\) and 0.07 for European and Asian data sets, respectively; supplementary fig. S3, Supplementary Material online). Therefore, the more conservative haplotype calling methods appear to result in a substantial reduction in statistical power to detect differences in rates of introgression, so we must be cautious in interpreting the biological significance of the difference in N-mt introgression results from Neanderthals (fig. 3) versus Denisovans (supplementary fig. S2, Supplementary Material online).

Molecular Interactions between Nuclear and Mitochondrial Gene Products

To test for sequence changes that potentially alter interactions between nuclear and mitochondrial gene products, we analyzed fixed differences in mitochondrially encoded proteins and structural RNAs to determine which sites physically contact nuclear-encoded proteins (fig. 4 and supplementary fig. S4, Supplementary Material online). Ten of the 22 amino acid substitutions in mitochondrially encoded proteins that distinguish humans and Neanderthals were identified as occurring at contact sites (supplementary table S6, Supplementary Material online). In total, these residues physically interact with nine different N-mt subunits within OXPHOS complexes (table 2). Similarly, 8 of the 24 amino acid substitutions that distinguish modern humans and Denisovans occurred at contact residues (supplementary
Neanderthals and/or Denisovans (table 1 and fig. 2). In addition, there are a total of 11 tRNAs with fixed nucleotide substitutions distinguishing modern humans and Denisovans (table 2 and supplementary table S6, Supplementary Material online). About 10 of the 14 rRNA substitutions that have fixed differences distinguishing modern humans from Neanderthals and/or Denisovans (table 1 and fig. 2). Although we have not specifically modeled the many protein–tRNA interactions that occur during mitochondrial tRNA maturation and charging (Pett and Lavrov 2015), we included the corresponding aaRS genes for these 11 tRNAs in our subsequent analysis of N-mt proteins.

Table 2
N-mt Genes That Directly Contact Sites with Fixed Substitutions in Mitochondrially Encoded Protein or Structural RNA Sequences between Modern Humans and Neanderthals

| N-mt Gene | Complex | Mito Contact(s) | Introgression |
|-----------|---------|----------------|--------------|
| NDUFA13   | I       | ND1 (186)      | 2.1          |
| NUFB5     | I       | NDS (24)       | 65.2         |
| NUFB9     | I       | NDS (515)      | 41.2         |
| NUFC2     | I       | ND2 (346)      | 34.2         |
| CYC1      | III     | CYTB (233, 245)| 53.1         |
| COX5a     | IV      | COI (54)       | 51.0         |
| COX6B1    | IV      | COI (95)       | 79.3         |
| COX6B2    | IV      | COI (95)       | 70.3         |
| COX6C     | IV      | COI (146)      | 12.5         |
| ATP5F1    | V       | ATP6 (176)     | 7.3          |
| MRPL2     | Ribosome| RNR2 (855)     | 69.1         |
| MRPL27    | Ribosome| RNR2 (1,163)   | 19.5         |
| MRPL37    | Ribosome| RNR2 (39)      | 46.1         |
| MRPL47    | Ribosome| RNR2 (39)      | 62.3         |
| MRPL48    | Ribosome| RNR2 (1,163)   | 41.1         |
| MRPL57    | Ribosome| RNR2 (386)     | 78.5         |
| HARS2     | aaRS    | TRNH (52)      | 35.5         |
| NARS2     | aaRS    | TRNH (57)      | 5.6          |
| YARS2     | aaRS    | TRNY (52)      | 33.9         |

Note.—The mitochondrial gene products and contact residue positions based on the reference structures (in parentheses) are also indicated. An introgression ranking scaled between 0 and 100 is reported based on gene-specific introgression scores (in parentheses) are also indicated. Asterisks indicate genes with identified Denisovan-derived alleles in Melanesian population based on analysis of Vernot et al. (2016). Note that both tissue-specific paralogs for COX6B and for COX4I2 are included.

Table 3
N-mt Genes That Directly Contact Sites with Fixed Substitutions in Mitochondrially Encoded Protein or Structural RNA Sequences between Modern Humans and Denisovans

| N-mt Gene | Complex | Mito Contact(s) |
|-----------|---------|-----------------|
| NUFB5     | I       | NDS (24)        |
| NUFB9     | I       | NDS (515)       |
| CYC1      | III     | CYTB (233)      |
| UQORC1    | III     | CYTB (84)       |
| COX4I1    | IV      | COI (456)       |
| COX4I2    | IV      | COI (456)       |
| COX6B1    | IV      | COI (95)        |
| COX6B2    | IV      | COI (95)        |
| COX6C     | IV      | COI (22, 146)   |
| COX7B     | IV      | COI (456)       |
| MRPL2     | Ribosome| RNR2 (734, 855)|
| MRPL15    | Ribosome| RNR2 (616)      |
| MRPL16    | Ribosome| RNR2 (1,287)    |
| MRPL27    | Ribosome| RNR2 (1,163)    |
| MRPL37    | Ribosome| RNR2 (39, 736)  |
| MRPL43    | Ribosome| RNR2 (616)      |
| MRPL44    | Ribosome| RNR2 (616)      |
| MRPL47    | Ribosome| RNR2 (39)       |
| MRPL48    | Ribosome| RNR2 (1,163)    |
| MRPL57    | Ribosome| RNR2 (386)      |
| MRPS5     | Ribosome| RNR1 (281)      |
| MRPS6     | Ribosome| RNR1 (362)      |
| MRPS21    | Ribosome| RNR1 (362)      |
| EARS2*    | aaRS    | TRNE (50)       |
| FARS2*    | aaRS    | TRNF (20)       |
| HARS2     | aaRS    | TRNH (52, 56)   |
| IARS2     | aaRS    | TRNI (56)       |
| LARS2*    | aaRS    | TRNL1 (48), TRNL2 (46) |
| MAR2      | aaRS    | TRNM (55)       |
| TARS2     | aaRS    | TRNT (54)       |
| WARS2     | aaRS    | TRNW (5, 28)    |
| YARS2*    | aaRS    | TRNY (52)       |

Note.—The mitochondrial gene products and contact residue positions based on the reference structures (in parentheses) are also indicated. Asterisks indicate genes with identified Denisovan-derived alleles in Melanesian population based on analysis of Vernot et al. (2016). Note that both tissue-specific paralogs for COX6B and for COX4I2 are included.
Because of the short divergence time between modern humans and Neanderthals/Denisovans, we are expected to share genetic variants with these related hominins, and all but two of these pairwise differences were amino acid polymorphisms that are segregating within modern human populations (supplementary table S8, Supplementary Material online). One of the two exceptions was a substitution (Y91C) in the complex III subunit UQCRC1 in Denisovans, which is a residue that does not contact any mitochondrially encoded subunits (fig. 5). The other exception is a premature stop codon in the ribosomal protein MRPL43 that appears to have been polymorphic within Neanderthals. This nonsense mutation truncates 86 amino acids from the longest predicted human isoform, but much of the truncated region is absent from some other reported human isoforms, and it is not found within the solved structure of the human mitochondrial ribosome (Amunts et al. 2015). Overall, we observed extremely little variation in the sequences of N-mt proteins. In particular, there were no cases of structurally correlated changes between directly interacting residues.

An additional predicted effect of structural interactions is that incompatibilities and restricted introgression would be most pronounced for N-mt genes encoding proteins that directly contact mitochondrially encoded sites with recent substitutions. However, we found little evidence to support this prediction. In the Neanderthal data set, only one such N-mt gene (NDUFA13) had an introgression score of 0 across both Asian and European population, and most of the other examples had introgression scores that are consistent with background levels in the genome (table 2). In the Denisovan data set, 28 of the 32 N-mt genes involved in interactions with altered mitochondrial sites had no detectable introgression (table 3), but this is consistent with the random expectation because most genes in the genome (87%) are in this category.

**Discussion**

The Role of Mitonuclear Incompatibilities during Hominin Introgression

The hypothesis that motivated our analysis was that incompatibilities (i.e., deleterious epistatic interactions) between Neanderthal/Denisovan nuclear alleles and modern-human mtDNA may have shaped patterns of introgression. In this light, the significant underrepresentation of N-mt loci in genomic regions of Neanderthal introgression suggests that mitonuclear interactions might have had an effect (fig. 3). However, it is possible that mitochondrial function per se has not affected introgression and is only correlated with some other important feature. For example, it has been shown that genomic regions with high levels of functional constraint tend to have lower frequencies of introgressed alleles in modern humans (Sankararaman et al. 2016), so it is possible that N-mt genes are simply more functionally important on average. However, the underrepresentation of introgressed alleles at N-mt loci remained significant even when we controlled for levels of expression and rates of sequence evolution (supplementary fig. S1, Supplementary Material online), which we used as proxies for functional importance.

It is also conceivable that N-mt genes are enriched in genomic regions that happen to be less prone to introgress for reasons such as low recombination rate (Baack and Rieseberg 2007). We consider this possibility unlikely, however, because the lone evidence for a nonrandom distribution of N-mt genes in the human genome is that they are underrepresented on the X-chromosome relative to autosomes (Drown et al. 2015). Because autosomes exhibit higher rates of introgression than the X-chromosome, the overrepresentation of N-mt genes on autosomes would tend to introduce a bias that is opposite the pattern that we observed in our data. We are also skeptical of any interpretation that N-mt alleles from archaic hominins suffered from a genotype × environment interaction. Mitochondrial biology has often been implicated in local adaptation and thermal tolerance, including in humans.
Mitochondrial responses to a greater amount of functional divergence. mtDNA divergence in Denisovans, it is not clear that this cor-
denisovan analysis. First, despite the higher overall level of methodological issues that potentially complicate the

effects would have been even stronger during introgression for reduced introgression of Neanderthal N-mt alleles, such

tions than those of Neanderthal and Denisovan populations that already resided there.

Therefore, although the cause of lower rates of introgression for N-mt genes is not entirely clear, we conclude that the most likely explanation involves deleterious epistatic interactions with modern-human nuclear and/or mitochondrial genetic backgrounds. However, it is very unlikely that such epistatic effects were mediated by substitutions at physically interacting sites even though direct structural interactions have been implicated in other studies of mitonuclear coevolution (Osada and Akashi 2012; Meiklejohn et al. 2013; Havird et al. 2015). A role for changes at directly interacting sites is inconsistent with the extremely low levels of primary sequence divergence in hominin N-mt proteins (table 3; supplementary tables S6–S8, Supplementary Material online). Moreover, rates of introgression appear to be lower for N-mt loci regardless of whether they contribute to direct molecular interactions in complexes with mitochondrially encoded gene products (fig. 3). It is possible that the rapid evolution of mtDNA has effects on diverse nuclear genes involved in mitochondrial function because epistasis does not necessarily require direct physical interaction between gene products (Moore and Williams 2005; Clark et al. 2012; Baris et al. 2017). Given the limited amounts of protein sequence divergence, it is likely that any functional and fitness effects that distinguish alleles from modern humans and related hominins largely reflect differences in gene expression and regulation.

We reasoned that, if changes in mtDNA were responsible for reduced introgression of Neanderthal N-mt alleles, such effects would have been even stronger during introgression from Denisovans because of their much more divergent mitochondrial haplotypes (fig. 1; Krause et al. 2010). Therefore, it was somewhat surprising that there was no detectable difference in rates of Denisovan introgression between N-mt genes and other nuclear loci (supplementary fig. S3, Supplementary Material online). However, there are both biological and methodological issues that potentially complicate the Denisovan analysis. First, despite the higher overall level of mtDNA divergence in Denisovans, it is not clear that this corresponds to a greater amount of functional divergence. Mitochondrial $d_\theta$/$d_s$ values for modern humans and Neanderthals are substantially higher than those for Denisovans and internal branches in the hominin tree (table 1; Krause et al. 2010), suggesting a history of more efficient purifying selection in Denisovans against deleterious functional changes in mtDNA. Second, as we have already noted, the power to detect variation in rates of Denisovan introgression appears to be limited by the conservative methodology used to identify Denisovan-derived alleles (Vernot et al. 2016). Although methods based on conventional $D$-statistics can be used to estimate the total contribution of introgressed alleles from a given population, it is far more challenging to assign a probability of introgression for individual loci. Great progress has been made in addressing this challenge (Sankararaman et al. 2014; Vernot and Akey 2014), but it remains difficult to confidently identify Denisovan-derived alleles in Melanesian genomes because they already contain a large contribution of introgressed alleles from another source (Neanderthals).

Even if mitonuclear interactions were responsible for reduced introgression rates of Neanderthal-derived N-mt alleles, the magnitude of the effect appears to have been relatively small (fig. 3). Therefore, it is likely that these interactions have had limited consequences for the genome-wide landscape of introgression, at least in the European and Asian populations that were included in this analysis.

These limited effects could indicate that there is little functional divergence in mitochondrial haplotypes among hominin lineages, which would be consistent with recent inferences that mtDNA has introgressed between some hominin lineages. Even though there appears to have been no mtDNA introgression into modern human populations (Krings et al. 1997; Serre et al. 2004), the discordance between mitochondrial and nuclear genealogies in hominins has been interpreted as evidence for a mitochondrial replacement event late in the evolution of Neanderthals (Meyer et al. 2016). These findings are also relevant to recent debates about the extent of mitonuclear incompatibilities within modern humans, which have garnered attention with the advent of mitochondrial replacement therapy (Reinhardt et al. 2013; Chinnery et al. 2014; Morrow et al. 2015; Sloan et al. 2015; Rishishwar and Jordan 2017) and led to arguments that mitonuclear incompatibilities may be less widespread and slower to emerge than previously proposed (Eyre-Walker 2017). Of course, the lack of strong genome-wide signatures of mitonuclear incompatibilities does not preclude the possibility of strong effects on individual loci, and the existence of large introgression “deserts” within the modern human genome (Sankararaman et al. 2014) may be the result of linkage with loci involved in strongly deleterious epistatic interactions, including some N-mt genes (Runemark et al. 2017). Indeed, one of the pressing questions in the field of mitonuclear coevolution is whether incompatibilities emerge from individual changes of large effect or the more gradual and stepwise accumulation at many loci (McKenzie et al. 2003; Barnard-Kubow et al. 2016).

Epistatic versus Additive Effects on Patterns of Gene Flow

Two general models have been put forth to explain the evidence for purifying selection against Neanderthal and Denisovan alleles in modern human populations. The first is based on incompatibilities between introgressing alleles from archaic hominins and the genetic background in modern humans (Currat and Excoffier 2011; Sankararaman et al. 2014; Sharbrough et al. 2015; Vernot et al. 2016).
fits with a broader view that epistatic interactions have pervasive effects (Breen et al. 2012; Levin and Mishmar 2017). The second model is based on additive effects and does not require reproductive isolation or genetic incompatibilities among loci. Specifically, it has been argued that Neanderthals had lower effective population sizes than modern humans and, therefore, harbored larger numbers of weakly deleterious alleles because of inefficient selection (Harris and Nielsen 2016; Juric et al. 2016). Although this second model provides a cogent argument for why Neanderthal-derived alleles may have been generally disfavored in gene-rich regions (Sankararaman et al. 2014) and declined in frequency over time (Fu et al. 2016), it is less clear that it can explain the genome-wide underrepresentation of introgressed alleles in specific functional classes such as testis-expressed genes (Sankararaman et al. 2016) or the N-mt genes that were analyzed in our study. These functional gene classes would have to somehow be enriched for variants with deleterious effects that were weak enough to accumulate by drift in archaic hominin populations but strong enough to be effectively selected against in modern humans. While it is certainly possible that such variants are overrepresented in genes with mitochondrial- or testis-specific functions relative to the rest of the genome (Harris and Nielsen 2016), we are not aware of any a priori rationale for why this would be the case.

One prediction associated with an incompatibility model is that genes exhibiting limited introgression into modern humans would also have experienced lower levels of introgression moving in the opposite direction (i.e., from modern humans into Neanderthals and Denisovans). By contrast, if the historical accumulation of harmful mutations in Neanderthal and Denisovan populations is responsible for the underrepresentation of introgressed alleles at these loci in modern humans, we might expect that affected loci would have been subject to higher levels of introgression into archaic hominins as a form of adaptive replacement. We suggest that this offers a general framework for distinguishing between additive and epistatic effects in cases of bidirectional gene flow. Sign epistasis should contribute to a positive correlation between locus-specific rates of introgression in one direction versus the other because loci involved in incompatibilities would be restricted from moving in both directions. By contrast, differences in additive effects on fitness should lead to a negative correlation between locus-specific rates of introgression because the alleles that carry the heaviest deleterious mutation load will be least likely to introgress into a hybridizing population but most likely to be replaced themselves by introgression from that population. There is potential for future research to test this prediction in hominins because introgression appears to have occurred from modern humans into some eastern Neanderthal populations (Kuhlwilm et al. 2016). There is already evidence from this work that introgression has been restricted in both directions for genomics regions of greater functional importance, as might be expected in cases of epistatic incompatibilities.

**Supplementary Material**

Supplementary data are available at Genome Biology and Evolution online.

**Acknowledgments**

We thank two anonymous reviewers and the lab groups of D.B.S. and Rachel Mueller for helpful comments on an earlier version of this manuscript and BZ 460 students at Colorado State University for stimulating discussion. We also thank Sriram Sankararaman for providing introgression score data and Joshua Akey for directing us to posted haplotype-call data. We are grateful to Thierry Chambert for helpful discussion and statistical advice. Our research on mitonuclear interactions is supported by a Division of Molecular and Cellular Biosciences grant at the National Science Foundation (MCB 1412260) and a National Institutes of Health Ruth L. Kirschstein National Research Service Award (F32GM116361).

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Mitonuclear Dimension of Neanderthal and Denisovan Ancestry in Modern Human Genomes

Genome Biol. Evol. 9(6): 1567–1581 doi:10.1093/gbe/evx114 1581

Associate editor: Bill F. Martin