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Complete Mitochondrial Genome and Phylogeny of Pleistocene Mammoth *Mammuthus primigenius*

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Phylogenetic relationships between the extinct woolly mammoth (*Mammuthus primigenius*), and the Asian (*Elephas maximus*) and African savanna (*Loxodonta africana*) elephants remain unresolved. Here, we report the sequence of the complete mitochondrial genome (16,842 base pairs) of a woolly mammoth extracted from permafrost-preserved remains from the Pleistocene epoch—the oldest mitochondrial genome sequence determined to date. We demonstrate that well-preserved mitochondrial genome fragments, as long as ~1,600–1,700 base pairs, can be retrieved from pre-Holocene remains of an extinct species. Phylogenetic reconstruction of the Elephantinae clade suggests that *M. primigenius* and *E. maximus* are sister species that diverged soon after their common ancestor split from the *L. africana* lineage. Low nucleotide diversity found between independently determined mitochondrial genomic sequences of woolly mammoths separated geographically and in time suggests that north-eastern Siberia was occupied by a relatively homogeneous population of *M. primigenius* throughout the late Pleistocene.

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Introduction

*Mammuthus, Elephas, and Loxodonta* (family Elephantidae, subfamily Elephantinae) are closely related genera that evolved in the African Pliocene, possibly from the genus *Primelephas* [1–4]. The woolly mammoth *Mammuthus primigenius* became extinct across most of its former range along with other late Pleistocene megafauna, although small, isolated mammoth populations survived into the mid-Holocene [5]. The phylogeny of Elephantinae has not been resolved. Morphological analyses have yielded conflicting phylogenies for *M. primigenius, Elephas maximus,* and *Loxodonta africana* [1–4]. Dental characters suggest a closer relationship between *M. primigenius* and *E. maximus*, trunk tip morphology supports a grouping of *M. primigenius* and *L. africana*, and immunological and hair structure characters could not confidently resolve the phylogeny of these three taxa [1–4]. Molecular analyses have also generated conflicting conclusions [1–4]. Data on short mitochondrial DNA (mtDNA) sequences have variously supported a monophyletic clade of the extant elephant species, have grouped *M. primigenius* with either *E. maximus* [6] or *L. africana* [7–9], or have been inconclusive [10,11]. DNA template modifications caused by oxidation or hydrolysis are a potential source of artificial mutations and may partly explain the high polymorphism initially observed for some short mammoth DNA sequences, although pseudogenes of mitochondrial origin located in the nuclear genome, which are common in elephants, may also be a potential factor [10]. A meta-analysis taking into account potential sequencing and other errors favored the *M. primigenius*–*L. africana* clade [9]. Clearly, a comprehensive analysis of longer genomic sequences is necessary to resolve this phylogeny, but so far only short DNA fragments have been retrieved from *M. primigenius*.

Results/Discussion

The woolly mammoth leg with intact muscle and skin tissue was found in the Enmynveem River valley (Chukotka) in north-eastern Siberia in 1986 (Figures 1 and S1). Since then, the collected muscle specimens of the mammoth (hereafter called Enmyn) were kept frozen. Radiocarbon method dated the specimens to ~33,750–31,950 y BP (before present) [12]. An initial examination suggested that the soft tissue was remarkably well preserved, and no signs of tissue decay were noticed when the specimen was excavated from the permafrost. Because no tissue damage by insects or other animals was observed, the remains presumably were quickly buried and have never been defrosted. The cells with nuclei were observed in epithelia and muscle tissue. The treatment with DAPI efficiently stained the nuclei, indicating the existence of
well-preserved genomic DNA (Figure 2A). To our knowledge, this is the first documented cytogenetically detectable nucleus with genomic DNA in such an ancient tissue (>30,000 y BP).

In accordance with these primary observations, a substantial amount of genomic DNA was extracted and detected on an electrophoresis gel. Although the extracted DNA was apparently degraded, it is yet of remarkable quality and quantity. For example, the major fraction of DNA fragments from one extract (shown in Figure 2B) ranged from 100 base pairs (bp) to 600 bp, with diminishing amounts at higher molecular weights. Microbial DNA may potentially contribute to the high-molecular-weight DNA fraction. Nevertheless, the PCR analysis described below (Figure 2C and 2D) supports the assertion that this mammoth specimen contains preserved DNA. Independent replication in PCR analysis is absolutely imperative for the study of ancient DNA. Therefore, several DNA extracts from the tissue specimens were obtained for the mitochondrial genome analysis.

Importantly, the replication in this study was not limited to selected mitochondrial genome regions, and the entire mitochondrial genome was sequenced in duplicate. Different DNA extracts from mammoth muscle tissue were used to generate PCR products and reconstruct the complete mitochondrial genome in Moscow (“MOS contig”, completed in 2000) and, independently, at the University of Massachusetts Medical School (UMASS MS) laboratories (“UM contig”, completed in 2005) (see Materials and Methods). No contamination by extraneous DNA was found in multiple PCR experiments. Apart from a variable number of tandem repeats (VNTR) in the control region characterized by high somatic hypervariability, the mtDNA sequences obtained from different extracts and different laboratories matched exactly. This accuracy was achieved with the following protocol. (I) Redundant oligonucleotide primers were designed and tested to yield a sufficient amount of PCR products (Figures S2–S4 and Table S1). Preliminary tests with these primers were conducted to demonstrate the applicability of DNA extracts for PCR. We consistently found efficient recovery of at least ~500- to 700-bp PCR fragments from mammoth DNA extracts. Moreover, PCR of complete gene sequences of ~1,200–1,700 bp was also efficient, but the mtDNA sequence larger than 3,000 bp was not amplifiable (Figure 2C and 2D and Methods and Materials). The data are consistent with excellent preservation coupled with some degradation due to the ancient origin of the DNA. (II) The large amount of DNA enabled us to obtain PCR products sufficient for sequencing after the first round of PCR reaction and minimized the risk that sequence errors arose due to template switching during PCR, because of the low number of original DNA templates [13,14]. (III) Direct sequencing of PCR products gave clean chromatogram reads (Figure S5). In all cases, the replication of sequencing in each nucleotide position was performed from different PCR products using the protocols described in Materials and Methods. In addition, the PCR products were cloned and the independent clones were also sequenced. As expected, the cloned mtDNA fragments occasionally contained random mutations with an average rate of ~6/10,000 bp. No cloned sequences with identical sets of mutations were identified. These sequence modifications were mostly transitions corresponding to type I (A→G/T→C) and type II (C→T/G→A) mutations, which have previously been detected in ancient DNA (Figure S6) [13–15].

![Figure 2](https://example.com/figure2.png)

**Figure 2. The Unusually Well-Preserved Mammoth DNA**

(A) Nuclei with DNA clearly detectable by DAPI staining in muscle cells of ~33,000-y-old *M. primigenius*. (B) Total genomic DNA isolated from the mammoth muscle tissue (lane 1 is 1/10 dilution of the DNA on lane 2); control DNA from fresh human blood samples in lanes 3 and 4. (C) Examples of PCR products (~300–600 bp) for mammoth mitochondrial genome. (D) PCR amplification recovers long sequences for complete mitochondrial genes (1,317 bp *CytB* and 1,613 bp *ATP6* genes), but PCR of larger fragments (3,054 bp *ND5*) is failed.

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As described previously for ancient DNA, type II mutations were predominant in the cloned mammoth mtDNA sequences (>70%). The sequence modifications were not detectable in direct PCR product sequences of the same region (Figure S5). The data indicate that the mutations in cloned DNA were random and rare in each site of individual molecules. No contamination with nuclear mitochondrial DNA (numt) was found for any PCR product of mammoth DNA (Materials and Methods). Direct sequencing of multiple PCR products and independent clones provided evidence of accurate reconstruction of authentic mitochondrial genome sequence of *M. primigenius*.

The sequence of the small region (VNTR) in the control region (D-loop) could not be determined by direct sequencing of PCR products. Thus, the analysis of cloned PCR fragments was undertaken. The sequence analysis demonstrated somatic heteroplasmy and high molecular heterogeneity in this region. The heterogeneity resulted from hypervariability in a number of short hexa-nucleotide tandem repeats (CGCATA)$_n$, resembling VNTR (Figure S4). Similar VNTR was found in the control mtDNA region in *L. africana* and *E. maximus* also.

To determine the evolutionary history of the woolly mammoth, we sequenced complete mitochondrial genomes of *L. africana* and *E. maximus*. To exclude the possibility of DNA contamination, the elephant specimens were obtained only after the primary mitochondrial genome sequence of *M. primigenius* was completed (Materials and Methods). Long-range PCR was initially used for the amplification of the overlapping ~3–5 kb mtDNA fragments of African and Asian elephants. The sequence analysis was also conducted for cloned PCR fragments and short PCR fragments as described in mammoth mtDNA analysis. The *E. maximus* and *L. africana* mitochondrial genomes sequenced in this study were highly similar but not identical to those submitted to GenBank previously.

Similar to other placental mammals, the mammoth mitochondrial genome contains 13 protein coding genes, 22 tRNA and two rRNA genes, and the D-loop control region (Figure 3). The length of the genome varied due to somatic variability of tandem DNA repeats (CGCATA)$_n$ in the control region. The length of the mammoth mitochondrial genome was 16,842 bp, if the longest VNTR tract was included (Figure S4). Start and stop codons were identical in *M. primigenius*, *L. africana*, and *E. maximus*; however, we found a substitution (G→A) in the ND4 gene leading to a premature stop codon in the C-terminus of the protein. Thus, the ND4 protein is four amino acids shorter in *M. primigenius* than in elephants. A multiple alignment of the C-terminal end of the ND4 protein is variable in sequence and in length in other animal species. Substitutions in the mammoth lineage were predominantly transitions (19/339 transitions/transversion) transition ratio in mammoth versus 13/351 in *E. maximus* when polarized with *L. africana*). The total number of nucleotide and amino acid differences between the mammoth and *E. maximus* genomes was 722 and 100, and between the mammoth and *L. africana* genomes 780 and 107, respectively (Table S2).

The phylogeny of Elephantinae was inferred using the newly obtained *M. primigenius*, *E. maximus*, and *L. africana* sequences, and other complete mitochondrial genome sequences of *E. maximus* and *L. africana*, while the mitochondrial genomes of the closest extant taxa [16], the dugong (*Dugong dugon*) and the hyrax (Procavia capensis), were used as outgroups. The same topology was recovered by a variety of tree reconstruction methods (Figure 4). As expected, individuals from the same species, *E. maximus* and *L. africana*, clustered together on the tree. *M. primigenius* was determined to be a sister species to *E. maximus*, i.e., the woolly mammoth shared a common ancestor with the Asian elephant more recently than with the African elephant. A maximum likelihood (ML) ratio test comparing all three possible topologies of the Elephantinae species corroborates this conclusion (*p* < 0.01; see Materials and Methods). We also reconstructed the phylogeny of these species by using only individual protein and rRNA genes (tRNA genes are too short and contain too few substitutions). The majority, but not all (Table S3), of trees reconstructed with the sequence of individual genes supported the topology recovered using the complete genome. Although the topology of this tree appears robust, the unavailability of an outgroup more closely related than the dugong and hyrax calls for the sequencing of nuclear *M. primigenius* genes to further test the results reported here.

The resolution of the Elephantinae phylogeny makes it possible to estimate the time of divergence of *M. primigenius* from its sister species *E. maximus*. Such an estimate usually relies on the existence of a molecular clock, i.e., a uniform rate of evolution of all compared clades, and on having calibration points from the fossil record. However, previous studies suggested that the evolution of the Elephantinae mitochondrial genomes may be inconsistent with the molec-
The analysis of complete mtDNA sequences places *M. primigenius* with *E. maximus* on the tree. The Sirenia (*D. dugon*) and Hyracoidea (*P. capensis*), most closely related species among extant taxa to Elephantinae, were taken as outgroups. Bootstrap values and posterior probabilities were calculated using a Bayesian approach [29,31] assuming a gamma distribution of the rates of evolution across sites with a General Time Reversible model (normal font), HKY model (bold), with a parsimony approach (italic), and by neighbor joining (italic and bold) [28]. The scale is 0.1 substitutions per site. The mitochondrial genomes of *M. primigenius*, *E. maximus* A, *E. maximus* B, *L. africana* A, *L. africana* B, *D. dugon*, and *P. capensis* were used in the analysis.

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Figure 4. Paenungulata Tree and Phylogenetic Relationship of the Woolly Mammoth

The analysis of complete mtDNA sequences places *M. primigenius* with *E. maximus* on the tree. The Sirenia (*D. dugon*) and Hyracoidea (*P. capensis*), most closely related species among extant taxa to Elephantinae, were taken as outgroups. Bootstrap values and posterior probabilities were calculated using a Bayesian approach [29,31] assuming a gamma distribution of the rates of evolution across sites with a General Time Reversible model (normal font), HKY model (bold), with a parsimony approach (italic), and by neighbor joining (italic and bold) [28]. The scale is 0.1 substitutions per site. The mitochondrial genomes of *M. primigenius*, *E. maximus* A, *E. maximus* B, *L. africana* A, *L. africana* B, *D. dugon*, and *P. capensis* were used in the analysis.

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Polarization of the molecular clock [38] and so we tested this possibility using complete genomes. Tajima’s relative rate test showed no difference in the rate of evolution of different Elephantinae lineages (unpublished data). Similarly, a likelihood ratio test did not reject the assumption of equal rate of evolution of the *M. primigenius* versus the *E. maximus* lineage using the *L. africana* genome as an outgroup (Materials and Methods). In contrast, the assumption of a molecular clock was rejected when the likelihood ratio test was performed on simulations involving a comparison of the *M. primigenius*–*E. maximus* clade with the *L. africana* lineage when the dugong and hyrax species were used as outgroups (without molecular clock LnL1 = −56661.45; with molecular clock LnL2 = −56871.63; \( p \ll 0.01 \)).

Molecular clock analyses are highly sensitive to the molecular distance of the outgroups, which can affect the correct root placement. Clearly, the dugong and hyrax are not ideal outgroups for these analyses, as their approximate nonsynonymous nucleotide divergence (dn) with the Elephantinae species is ~0.15 while synonymous nucleotide divergence is at saturation (ds ~ 1.2). Thus, the change in the rate of evolution implied by the ML analysis must be treated with caution. However, a simple parsimony assay corroborated the possibility that the mitochondrial genomes of different Elephantinae species evolve at different rates (Table S4). A ML estimate of the number of substitutions per lineage yielded quantitatively similar results (unpublished data).

These observations, coupled with previous reports [3,8], imply that a molecular clock assumption may be inappropriate when estimating the time of divergence of Elephantinae species. Thus, we used a heuristic rate smoothing procedure for ML estimates [17], which takes into account different rates of evolution among different branches of the tree. Using the calibration points of 6 million y for the *E. maximus–* *L. africana* split and 65 million y for the Elephantidae–Sirenia split (see [8] and references within), the time of divergence of *M. primigenius* and *E. maximus* was estimated as ~4 million y ago (±0.01 s.e.). This estimate must be treated with caution because it relies on the correct paleontological dating of the *E. maximus–* *L. africana* divergence. Nevertheless, these data clearly indicate that the divergence of the woolly mammoth from the Asian elephant occurred soon after the divergence of their ancestor lineage from African elephants.

To further verify the accuracy of our sequence and to investigate the level of nucleotide diversity in the woolly mammoth population, we compared our sequence to the longest available sequences of the mammoth mitochondrial genome from other individuals. Two CytB gene sequences from mammoth individuals found in north-eastern Siberia (Magadan region, Kirgilyakh Creek near Kolima River) [18] and north-central Siberia (Pyasina River valley in Taimyr Peninsula) [19] show very high similarity, while the 12S RNA gene [19] was identical to the corresponding genes in our mitochondrial genome sequence from Chukotka’s mammoth (Table S5). All these sequences were recovered from muscle specimens enriched in mtDNA and, thus, very likely had no numt contamination. The Chukotka Emmenyveem River valley is ~1,000 km from Kirgilyakh Creek and ~2,900 km from the Pyasina River valley. The data suggest a relatively low genetic diversity of mammoth maternal lineages in mammoth population spanning vast territory in Northern Siberia. Two recent studies have also demonstrated the isolation of multiple fragments of mammoth DNA [20,21]. In one report, a novel approach was applied for direct cloning and shotgun sequencing of random DNA fragments. The set of multiple fragments corresponding to partial mitochondrial genome sequence contained sequences with ancient DNA artifact mutations and potential numts and cannot yet be reliably used for comparative analysis of complete mitochondrial genomes [21]. Another group reported the complete mitochondrial genome sequence [20] of a ~12,170-y-old woolly mammoth that was found in the Beryтекh Yakutia region (70° 35’ N, 145° 00’ E), which is ~900 km from the Emmenyveem River valley (68° 10’ N, 165° 56’ E). A comparison of the two complete genomes revealed a pattern of high similarity with the nucleotide diversity of ~0.3% (Table S5). This diversity may be even lower since sequence errors may be present in the genome sequences recovered from ancient specimens. When the differences between the two mammoth mitochondrial genomes were polarized with *E. maximus* sequence, the number of sequence-specific polymorphisms for the mitochondrial genome sequence determined here was slightly lower (by ~25%) than in the mitochondrial genome sequence described in the other
study [20]. The sequence [20] also showed a higher ratio of nonsynonymous to synonymous substitutions in protein coding genes and a higher number of nucleotide substitutions in the entire genome (C→T/G→A transitions) that are often associated with ancient DNA modifications (Tables S6 and S7). Although these differences were not statistically significant, a large number of nonsynonymous substitutions was observed in the ND2 gene in sequence [20], but not in our sequence (Table S6). The possibility that these changes appeared due to positive selection in one of the mammoth lineages seems highly unlikely, the more reasonable explanation being that the genome sequence from [20] may harbor some post-mortem ancient DNA mutations (most ND2 nonsynonymous mutations were C→T/G→A). However, these artifacts appear to be rare. In sum, the independently determined mitochondrial genome sequences from ∼33,000-y-old (present study) and ∼12,000-y-old [20] animals were highly similar, and, owing to a few potential errors, the true level of mitochondrial nucleotide variability may be slightly lower than 0.3%.

The low nucleotide diversity of mammoth mitochondrial sequence (π = 0.003; Table S5) is an order of magnitude lower than that reported for the overall populations of L. africana (π ~ 0.02) [22] and E. maximus (π ~ 0.017) [23,24], but similar to the values reported for select populations of L. africana (π ~ 0.00084–0.027) and E. maximus (π ~ 0.0024–0.0055) [22–24]. These data suggest that unlike the Asian and African elephants, the mammoth population has not had a complex population structure and has had a relatively low genetic diversity in mitochondrial lineages, at least in the area spanning thousands of kilometers in north-eastern Siberia. Further sequencing of mitochondrial genomes of other mammoth specimens can clarify the diversity of the ancient mammoth population.

The phylogenetic reconstructions based on complete mitochondrial sequence analysis is a powerful method to determine the relationship between taxa or even closely related extinct and extant species as demonstrated here and elsewhere [25]. However, mitochondrial and nuclear DNA may have a different coalescent history, and cytonuclear-genomic dissociation has been described recently in African forest and savanna elephants [26]. Given the unique quality of the specimens from the Enmyn mammoth, nuclear DNA may potentially be recovered [21] and used for further confirmation of the topology reconstructed in this study. Our data demonstrate that very high-quality genomic DNA may be recovered from ancient remains of the Pleistocene age and that sequencing of complete mitochondrial genomes can lead to reliable phylogenetic reconstructions and population studies not only for extant, but also for extinct species.

Materials and Methods

Extraction of genomic DNA. Microscopy analysis and DAPI staining of cells demonstrated the preservation of DNA in the mammoth specimens. Genomic DNA was extracted from the muscle mammoth specimens using silica-based and phenol-chloroform extraction methods, but the phenol-chloroform method was more efficient when extracting large amounts of DNA. The following protocol was used for phenol-chloroform DNA isolation. DNA was extracted from ∼0.1–0.4 g of muscle tissue in a solution containing 50 mM Tris-HCl (pH 8.0), 100 mM NaCl, 10–25 mM EDTA, 1%–2% SDS, and at least 100 µg/ml proteinase K for 16–24 h at 37 °C, followed by phenol-chloroform extraction. DNA purification was completed by precipitation with 75% ethanol or by concentrating with Centricon YM centrifugal filter devices (with Amicon filter). DNA quality was analyzed by agarose electrophoresis and PCR.

Mammoth DNA. First, DNA prepared from different samples and by different methods (silica-based and phenol-chloroform extraction methods) was used for PCR of the CytB gene. The analysis demonstrated that the sequences from various DNA extracts were identical and corresponded previously reported references for mammoth CytB gene. After the preliminary experiments, an average length of PCR product of 500- to 600-bp range was chosen to determine the complete mtDNA sequence of M. primigenius. To design the PCR primers, we used the sequences of mitochondrial genomes in the GenBank database available at the time. (Note: this project was launched in January 2000). Each primer pair was placed over the alignment of mitochondrial genomes of L. africana, hippopotamus, rhinoceros, and donkey. We gave preference to the primers located in the most conservative regions to keep the number of ambiguous nucleotides in designed oligonucleotide primers as small as possible. Ultimately, to amplify the whole mitochondrial genome of the mammoth, 35 pairs of redundant primers producing overlapping PCR fragments were designed.

On average, the length of overlapped sequences between PCR fragments produced from different primers was ∼30 bp (excluding primer sequences), but were as long as ∼200 bp in some cases. The anticipated size of the PCR fragments ranged from 325 bp to 650 bp and was 929 bp for the VNTR region. To determine the complete mtDNA sequence in UMSS MS, additional oligonucleotides were designed as shown in Table S1. On average, the mtDNA sequence was covered ∼ nine times in MOS contig and ∼ five times in UM contig by sequencing from different PCR products and clones. The total number of sequenced nucleotides determined from two complementary strands was 240,094. The total length of overlapped sequence obtained from different fragments was 12,000 bp. In addition, we were able to produce several large PCR fragments (the sequence length between direct and reverse primers and corresponding mitochondrial genome region are indicated): L-H5 968 bp (16S rRNA; tRNA Leu; ND3), L-H10 1,642 bp (for 1,683 bp with primers) (ND1; tRNAs Ile, Glu, Met; ND2; tRNAs Trp, Ala, Asn), L16-H18 1,572 bp (COX2; tRNA Lys; ATP6; COX3), L24-H26 1,502 bp (tRNA Leu; ND5), L29-H31 1,271 bp (tRNA Gln; CytB; tRNAs Thr, Pro), L35-H35 1,131 bp (D-loop; tRNA Phe; 12S rRNA) that covered the shorter PCR fragments. No mismatches in the overlapped sequences or in the independent PCR products were found.

The standard PCR was conducted in total volume of 25 µl containing 1–2 U Taq1 DNA polymerase in a buffer with 10mM Tris-HCl (pH 8.3), 0.25 mM dNTPs (Gibco BRL, Gaithersburg, Maryland, United States), 0.2–2 mM BSA, 1.5–2.5mM MgCl2, 20 pmol of each primer or in total volume 25 or 50 µl containing 1–2 U of PicoMaxx High Fidelity PCR System (Taq2000 DNA polymerase, cloned Phu DNA polymerase, and ArchaeMaxx polymerase enhancing factor) in 1x PicoMaxx reaction buffer (Stratagene, La Jolla, California, United States), and 0 or 90 pmol of PCR primers. The PCR conditions were 94 °C for 1–2 min, followed by 35 cycles of 94 °C denaturation for 30–40 s, 53–58 °C annealing for 30–40 s, and 72 °C extension for 30 s to 1 min. The PCRs were accompanied by negative controls containing the reaction solutions with no DNA (Figures 2 and S2). We found that some DNA extracts taken in high concentration may have inhibitory effects on PCR. The series of dilutions of the extracts were undertaken to identify the optimal concentration of DNA sufficient to produce PCR products after 32–35 rounds. Re-amplification by secondary PCR was not necessary. The PCR fragments were analyzed by 1.7%–2 % agarose gel stained by ethidium bromide. The PCR fragments were excised from agarose gels and purified via Qiagen columns. The PCR fragments were cloned into pGEM-T vectors (Promega, Madison, Wisconsin, United States), and the PCR fragments and clones with inserts were subjected to sequencing from mtDNA primer or vector SP6 and T4 primers using ABI PRISM BigDye kit for ABI analyzers following manufacturer’s instructions (Applied Biosystems, Foster City, California, United States).

Asian and African elephant DNA. The blood samples of the L. africana and E. maximus (originating from Burma) were used for DNA extraction by Qiagen (Valencia, California, United States) kits for blood DNA. We initially used long-range PCR to generate mitochondrial genome segments of ~3–5 kb. In addition, to produce longer and shorter PCR fragments, the series of primers were designed on the PCR conditions described above. The total PCR fragments and some cloned PCR fragments were sequenced. In both L. africana and E. maximus, rare mismatched sequences obviously corresponding to nuclear insertions of mito-
Chondroidal sequences were found. These diverged units were found when some pairs of primers were used for long-range PCR using elephant DNA isolated by QIAGEN kits. These pairs of primers had never been used for mammoth DNA PCR analysis. The mammoth DNA was also isolated from muscle tissue that has a relatively high proportion of mtDNA to nuclear DNA. The elephant unit number sequences were compared in the alignment analysis, and full-length sequences were sequenced using other sets of primers and PCR products. Finally, the complete mitochondrial genome sequences were determined from multiple independent PCR products. Comparison of the sequences determined in this study (animal A) with the corresponding *L. africana* B and *E. maximus* B sequences from the GenBank revealed their high similarity. The divergence of the sequences of these animals is likely due to polymorphisms. The potential sequence errors in the GenBank *E. maximus* mtDNA and *L. africana* mtDNA sequences might be of some concern. However, the results of phylogenetic reconstructions using these two species are the same regardless of which sequences were used (ours or GenBank’s) in comparative analysis with mammoth mitochondrial genome sequence.

**Sequence authentication.** The work was performed in accordance with commonly accepted standards for ancient DNA analysis. Multiple measures were undertaken to exclude contamination, potential artifact DNA changes specific to post-mortem specimens, or the inclusion of nonts in the mitochondrial sequence. The measures are summarized as follows. Large amounts of DNA were extracted from well-preserved specimens. The sequencing of PCR products obtained after the first round of amplification guaranteed that no sequence errors arise due to the low number of original DNA templates and template switching during PCR. The mtDNA sequences obtained from different DNA extracts were identical. The complete mitochondrial genome sequence fragments were determined in replicates with PCR, by different laboratories in different countries at different times. Apart from the VNTR region, the two mitochondrial genome sequences (Mos and UMA contigs) were identical. Stringent precautions were undertaken for PCR work with ancient DNA. The work in the Moscow laboratory was conducted in a special sterile room that included two rooms designed for overnight UV irradiation. The Pcr where the work with ancient DNA was conducted was separated from other laboratory rooms and equipment. The PCR and electrophoresis were always performed in separate rooms. No work with any animal DNA had ever been conducted in this laboratory prior to this project. The work at UMASS MS was performed in a new laboratory using novel equipment, pipettes, PCR station, and space that had never been used before for DNA analysis. No DNA work with elephants had taken place in the laboratories before the primary complete mammoth mtDNA sequence was determined. We have not found a single case of contamination by nonmammal mtDNA (human or animal) in PCR products or clones in Mos or in UMASS MS PCR products. Direct sequencing of PCR products from the mammoth specimen showed a high quality of chromatograms comparable to sequencing of cloned DNA (Figure S5). In all cases, the replication of sequence was performed from different PCR products from the PCR boxes. As expected, the cloned mtDNA fragments occasionally contained random mutations, transitions (A→T/G→C or C→T/G→A) corresponding to those documented previously as common type I and type II mutations in ancient DNA. The type II mutations (C→T/G→A) were predominant (75%). These mutations were easily discriminated since they were absent in sequences of total PCR products and other clones. As a final step, the mammoth mitochondrial genome sequences were compared with GenBank sequences of other mitochondrial genomes. As we report in our phylogenetic analysis section, the closest similar sequences to *L. africana* and *E. maximus* mtDNA sequences, even when the closest relative (*Sirenia*) was included in the comparison. The distribution and ratio of synonymous to nonsynonymous substitutions in the mammal genome is similar to that found in modern and nonmammal lineages and showed a high prevalence of synonymous substitutions. No nucleotide mismatches were found in multiple overlapped regions determined from independent PCR fragments (refer to earlier section “Mammoth DNA”). A muscle specimen was used for DNA extraction. This type of cell tissue is particular in that it is enriched with mitochondria, which provides a relatively high proportion of mitochondrial versus nuclear DNA.

Taken together, the methods and results indicate that contamination by exogenous DNA, nonts sequences, or sequence errors are unlikely to be a factor in our study of mammoth DNA and that the mammoth mtDNA sequence is authentic.

**Phylogenetic analysis.** A multiple alignment of the seven complete mitochondrial genomes was made using the MUSCLE program [27] and checked against amino acid sequence and RNA secondary structure whenever possible. Due to an uncertain alignment in the variable region of the control region, this part of the genome (~500 bp) was excluded from all analyses. The resulting alignment (referred to as the complete mitochondrial alignment throughout the text) was used for all of our phylogenetic and comparative analyses. To estimate synonymous and nonsynonymous nucleotide divergence, a concatenated alignment of all protein-coding genes was used, and dN and dS values were estimated with the PAMO-Bianchi-Li method as implemented in MEGA [28]. All analyses performed with the complete mitochondrial alignment were repeated using the concatenated alignment of all protein and RNA genes (discarding ~1,000 bp of nontranscribed sequence), and all of the results remained qualitatively unchanged.

To reconstruct the tree topology, parsimony and neighbor-joining methods were used as implemented in MEGA [28] with default parameters and 10,000 bootstrap replicates [28–31]. Bayesian inference of phylogenies was done with MrBayes [29] with two different prior models: the General Time Reversible model [28,29] and the HKY model [28,29], which estimates fewer rate parameters. Both models were run assuming a gamma-distribution of substitution rates across sites for 1 million iterations (mcmc ngen = 1000000 in MrBayes). Phylogenetic trees using individual gene sequences were tested with Bayesian inference set to the HKY model (Table S3). An ML ratio test was performed for ML values obtained for the possible topologies of the three Elephantinae species. The topology with *M. primigenius* and *E. maximus* as sister species had a log ML of ~4240.629, compared with ~4245.549 and ~4256.866 for the topologies linking *E. maximus* with *L. africana* and *M. primigenius* with *L. africana*, respectively (p < 0.01; ML ratio test). The log ML scores reported here are averages of log ML scores obtained for 10,000,000 iterations of the BEAST [30] program and run under the HKY [28,29] model with set tree topologies.

The molecular clock was tested with the likelihood ratio test, using the log likelihood scores obtained with the basal program in PAML [32] when the dataset was analyzed with the assumption of no molecular clock (clock = 0 in baseml) and a local clock that tested the difference between the species in question (clock = 2 in baseml). To estimate the branch length for synonymous and nonsynonymous substitutions, the branches of interest were assumed to have different dn/ds ratios (model = 2 in codeml) and an absence of a molecular clock (clock = 0 in codeml). To estimate the divergence time of *M. primigenius*, a heuristic rate smoothing method for ML estimates [16] was used as implemented in PAML [32]. The number of synonymous and nonsynonymous substitutions (Table S4) was obtained by parsimony using the African elephant as the outgroup for the mammmoth–Asian elephant comparison and dugong as the outgroup for the mammmoth–African elephant and Asian–African elephant comparisons without correcting for multiple substitutions and with the assumption that the outgroup accurately resembles the ancestral state. These results yielded were quantitatively very similar to those obtained with a ML approach that corrected for multiple substitutions (unpublished). Control files for all stand-alone programs ran here and other methodological materials are available on request.

**Supporting Information**

**Figure S1.** Mammoth Specimen Found in Siberia
(A) Restored exhibit of mammoth Ennym leg found in the Ennymveev River valley, Chukotka, northeastern Siberia. (B) The geographic location of the find is designated by the red dot on the map.
Found at DOI: 10.1371/journal.pbio.0040073.sg001 (2.3 MB PDF).

**Figure S2.** Electrophoresis Analysis of the PCR Products Used for Sequencing of *M. primigenius* Mitochondrial Genome (“MOS Mammoth mtDNA”)
A and B agarose gel panels. The negative controls are shown in (A). *designates the area of oligonucleotide primer dimers formed in PCR.
Found at DOI: 10.1371/journal.pbio.0040073.sg002 (631 KB PDF).

**Figure S3.** Electrophoresis Analysis of the PCR Products Used for Sequencing of *M. primigenius* Mitochondrial Genome (“UM Mammoth mtDNA”)
A–D agarose gel panels. *designates lines with low amount of PCR products.
Found at DOI: 10.1371/journal.pbio.0040073.sg003 (522 KB PDF).
Figure S4. Extreme Heteroplasmy in Hypervariable Region HVR (VNTR Region) in the Control Region of Mitochondrial Genome of M. primigenius
(A) Electrophoresis of PCR products from two DNA extracts (m1 and m22) and (B) corresponding PCR clones with VNTR region mtDNA insertions showed variability in their length. (C) Nucleotide sequences of cloned PCR products with VNTR. The sequence for the longest VNTR was incorporated in a reported complete sequence of M. primigenius.

Found at DOI: 10.1371/journal.pbio.0040073.sg004 (455 KB PDF).

Figure S5. Sequencing Electrophoresis Data for the Mitochondrial Genome of M. primigenius with High-Quality Chromatograms of Direct Sequencing of PCR Products, in Comparison with the Cloned PCR Fragments

Three mtDNA regions are shown: (A) L11-H11, (B) L17-H17, and (C) L22-H22. Lines 1 and 2 (A) and line 1 (B and C) from the bottom are the PCR products. Other lines are cloned sequences determined from opposite strands. The rare mutations are observed in cloned sequences but not in direct PCR product sequences.

Found at DOI: 10.1371/journal.pbio.0040073.sg005 (503 KB PDF).

Table S1. Primer Oligonucleotides and Predicted PCR Fragments Used for Sequencing of Complete Mitochondrial Genome of M. primigenius, E. maximus, and L. africana

Redundant primer oligonucleotides were used to amplify mammoth DNA. The sequences of mitochondrial genomes of E. maximus and L. africana were determined using both long-range PCR fragments and short PCR products.

Found at DOI: 10.1371/journal.pbio.0040073.s0001 (30 KB DOC).

Table S2. Different Types of Substitutions Revealed by Pairwise Comparisons of Elephantinae Genomes

Number of synonymous and nonsynonymous substitutions calculated with the modified Nei-Gojobori method as implemented in MEGA [28]; the results were independent of the transition/transversion ratio.

Found at DOI: 10.1371/journal.pbio.0040073.s0002 (25 KB DOC).

Table S3. Bayesian Phylogeny Inference [29] of Elephantinae Using Single Mitochondrial Protein Coding and rRNA Genes

Posterior probabilities are shown in parentheses.

Found at DOI: 10.1371/journal.pbio.0040073.s0003 (26 KB DOC).

Table S4. Lineage-Specific Substitutions Inferred by Pairwise Comparisons of Elephantinae Genomes and Polarized with One Outgroup

We used the sequences of E. maximus and L. africana determined in this study for these comparisons. Pairwise comparisons of Elephantinae species using a single outgroup to polarize substitutions suggested that the M. primigenius–E. maximus clad evolves faster than the L. africana lineage in both synonymous and nonsynonymous sites.

Found at DOI: 10.1371/journal.pbio.0040073.s0004 (26 KB DOC).

Table S5. Nucleotide Diversity Inferred by Pairwise Comparisons of Different M. primigenius mtDNA Sequences

Sequences are reported in the following studies: DQ316067 (present study), NC_007596 [20], D83047 [18], and D50841–D50842 [19].

Found at DOI: 10.1371/journal.pbio.0040073.s0005 (31 KB DOC).

Table S6. Genome-Specific Nonsynonymous and Synonymous Substitutions Inferred by Pairwise Comparisons of Protein-Coding Genes from Two M. primigenius Genomes (DQ316067 versus NC_007596) and Polarized with E. maximus (DQ316068) Sequence

A large number of nonsynonymous substitutions was observed in the ND2 gene (seven nonsynonymous versus two synonymous mutations in NC_007596 accumulated in a short ~360-bp region; whereas the DQ316067 sequence determined in this study had zero nonsynonymous and one synonymous substitution in the same gene).

Found at DOI: 10.1371/journal.pbio.0040073.s0006 (38 KB DOC).

Table S7. Genome-Specific Substitutions Inferred by Pairwise Comparison of Two M. primigenius Genomes (DQ316067 versus NC_007596) and Polarized with E. maximus (DQ316068) Sequence

While the C→T and G→A substitutions may be a sign of ancient DNA modification (most frequent type II mutations), they represent also real polymorphisms. The NC_007596 mammoth genome shows more of these substitutions, suggesting that a few of these substitutions may have occurred after death of the sampled mammoth; however, this difference is not statistically significant (Fisher’s exact test p > 0.05).

Found at DOI: 10.1371/journal.pbio.0040073.s0007 (25 KB DOC).

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