The time-dependent evolutionary rate of mitochondrial DNA in small mammals inferred from biogeographic calibration points with reference to the late Quaternary environmental changes in the Japanese archipelago

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Abstract  Mitochondrial DNA (mtDNA) sequences have long been the most popular marker for assessing phylogenetic relationships and uncovering population dynamics. However, the mechanism of the nucleotide substitution rate of mtDNA remains unclear. While the evolutionary rate over tens of thousands of years is thought to be time dependent, the overall picture is not fully understood. This article presents recent achievements related to the time-dependent evolutionary rate of mtDNA in small rodents in the Japanese archipelago. The method focuses on rapid expansion events during the late Quaternary, during which there was a prolonged severe cold period and repeated abrupt warm periods, providing multiple calibration points. The global sea level fluctuation and migration to islands help to specify the calibration points. For calibration points at 11000, 15000, 53000, and 130000 years ago, the evolutionary rates were approximately 0.11, 0.11, 0.047, and 0.029 substitutions/site/million years, respectively, in the mitochondrial cytochrome b gene (Cytb). Applying the higher rate to assess the evolutionary history of the commensal house mouse (Mus musculus) and complete mitochondrial genome sequences (~16000 bp) allowed us to trace prehistoric human culture development based on millet and rice agriculture. The pattern of time-dependent evolutionary rates presented here is likely applicable to other small rodents. The Japanese archipelago is ideal for assessing evolutionary rates with biogeographic calibration points in the late Quaternary in species with multiple genetically distinct local populations.

Key words: land bridge, mitochondrial DNA, murine rodents, Quaternary glacial maximum, time-dependent evolutionary rate

Introduction

Mitochondrial DNA (mtDNA) has long been the phylogenetic marker most commonly used to assess species relationships, genetic structure, and population dynamics (Brown et al., 1979; Avise, 1986; Irwin et al., 1991). The evolutionary rate of this marker proposed more than 40 years ago is 0.01–0.02 substitutions/site/million years (myr) in mammals (Brown et al., 1979; Wilson et al., 1985). Since then, however, no further consensus has been made on more accurate evolutionary rates. Ho and Larson (2006) proposed that the evolutionary rate of mtDNA has changed over time, namely the substitution rate is significantly higher than in older divergences (see also Penny, 2005). However, the rates currently used for the recent divergences differ among researchers, ranging from 0.11 (e.g. Suzuki et al., 2015) to 0.4 (e.g. Rajabi-Maham et al., 2008; Herman and Searle, 2011; Garcia-Rodriguez et al., 2018) substitutions/site/myr in rodents and around 0.05 in humans (protein-coding region; Henn et al., 2009). Here, I propose a substitution rate curve that shows a time-dependent tendency by utilizing the nucleotide substitution rate of mtDNA evaluated from biogeographic evidence.

I focus on historical events during the last 150000 years, when much of the planet was subject to rapid massive climate fluctuations (Brown et al., 2007). Specifically, a prolonged glacial period during the late Quaternary induced extreme population reductions, whereas abrupt rapid warmings triggered sudden population increases. Sudden population expansion can be detected by drawing a star-shaped network of mtDNA sequences and using biogeographic calibration points, which allow estimation of the evolutionary rate. Another way to find biogeographic calibration points is by addressing the dispersal events associated with the construction and destruction of land bridges facilitating migration between land masses separated by deep sea (i.e. at least 120 m depth; Japan Association for Quaternary Research, 1987). While the sea level rose during periods of global warming (Lambeck et al., 2014), the land bridges are
thought to have remained during the initial warming, allowing dispersal among what are now island masses. Bottle-necks are also caused by migration, as a founder effect, providing an opportunity to set a calibration point (e.g. Henn et al., 2009). For example, the house mouse (Mus musculus) migrated to the Japanese archipelago with humans around 3000 years ago along with the introduction of early paddy rice cultivation in the Yayoi period (Crawford and Lee, 2003; Fuller, 2011), yielding a star-shaped network in mtDNA analyses (Kuwayama et al., 2017; Li et al., 2020b).

Characterization of rapid expansion events in small rodents from Japan

Population dynamics in small mammals inferred from mtDNA variation

Phylogenetic analysis, such as the construction of a maximum likelihood tree and bootstrap analysis, helps us to specify clusters of mtDNA sequences. Network construction with mtDNA sequences sometimes exemplifies a cluster with a star-shaped structure, as mentioned above. The distribution of pairwise differences or mismatch distribution gives a unimodal pattern that is also indicative of rapid expansion (Slatkin and Hudson, 1991; Rogers and Harpending, 1992). Significantly negative values in neutrality tests, such as Tajima’s D and Fu’s Fs, confirm rapid expansion events.

The peak (mode) of the mismatch distribution is defined as \( \tau \), which is a statistical value proportional to the time since expansion. From the \( \tau \) value, the time in generations (\( t \)) since expansion started is calculated from the formula \( t = \tau/2u \), where \( u \) is the mutation rate per generation for the entire haplotype (Rogers and Harpending, 1992). Here, \( u = \mu kg \), where \( \mu \) is the evolutionary rate (substitutions/site/years), \( k \) is the sequence length of the haplotype, and \( g \) is the generation time in years. The time in years (\( T \)) since expansion can be estimated using the formula \( T = t g = \tau/2uk \). The value of \( \tau \) is an average value obtained by comparing a large number of sequences; thus, the value is trustworthy. Assigning historical time points (\( T \)) for the start of expansion, the evolutionary rate in years (\( \mu \)) can be calculated with the formula \( \mu = d/2T \), where \( d \) is the genetic distance (\( d = \tau/k \)).

Our previous study addressed mtDNA cytochrome b gene (Cytb) sequences, which is the most popular mtDNA gene for phylogenetic inference in mammals. Our study of the Cytb gene sequences (1140 bp) of the large Japanese wood mouse Apodemus speciosus (Suzuki et al., 2015) detected an apparent signal for a population expansion event in the sequence clusters for Hokkaido and Honshu/Shikoku/Kyushu with \( \tau = 2.7 \) and 9.2, respectively. Then, we measured the \( \tau \) value excluding the haplotypes from islands associated with the main islands of Honshu and Kyushu and obtained a new \( \tau \) of 8.5 (Honda et al., 2019). In the lesser Japanese wood mouse Apodemus argenteus, the Hokkaido population shows mtDNA diversity that indicated a recent rapid expansion, with a \( \tau \) value of 2.6 (Suzuki et al., 2015). By contrast, the sequences from the three main islands of Honshu, Shikoku, and Kyushu and their adjacent islands (Sado, Oki, Yamashima, etc.; Figure 1A) showed five clusters with expansion signals (Hanazaki et al., 2017), in which the \( \tau \) values could be categorized into three groups based on the values 3.9, 5.7, and 7.8–8.5. The Japanese giant flying squirrel Petaurista leucogenys, a rodent inhabiting broadleaf forests, had two clusters with expansion signals in Cytb sequences with \( \tau \) values of 3.9 and 8.3 (Oshida et al., 2009). Similarly, in a study of grassland-dwelling voles occurring in the Japanese archipelago, we found four clusters with expansion signals from three species: two from Hokkaido (Myodes Rufocanus and Myodes rutilus) and one from Honshu/Shikoku/ Kyushu (Myodes smithii). The \( \tau \) values obtained formed two groups: 4.1 and 5.8–6.3.

Overall, the \( \tau \) values observed in the Cytb sequences of small mammals from the Japanese archipelago can be roughly attributed to three groups of historic time points based on \( \tau \) values (Cytb, 1140 bp): Groups I, II, and III for 2.6–4.1, 5.8–6.3, and 7.8–8.5, respectively. A trend similar to these \( \tau \) values can be seen in the mtDNA sequences of Eurasian populations of M. Rufocanus (Abramson et al., 2012; Honda et al., 2019). The \( \tau \) values representing Group I can be subdivided into groups Ia (\( \tau = 2.6–2.7 \)) and Ib (\( \tau = 3.9–4.1 \)) (Hanazaki et al., 2017; Honda et al., 2019).

Linking population events to turning points in the Quaternary environment

How can one link the four historic expansion events to specific geological events? Since the species considered (wood mice, voles, and flying squirrels) are non-commensal, these rapid expansion events could be attributed to environmental recovery from the harsh glacial periods in the Quaternary, especially after the glacial maxima that occurred every 100000 years, as many researchers have noted (e.g. Hewitt, 2004; Oshida et al., 2009; Herman and Searle, 2011).

The turning points in the late Quaternary environment from cold to warm stages are the best occasions to consider as the starting points of population expansions after a bottleneck and subsequent sudden population growth. The transitions from marine isotope stage (MIS) 2 to MIS 1 (10000–15000 years ago) and from MIS 6 to MIS 5 (130000 years ago) are the most likely starts of rapid expansion events (Figure 1B). However, the timing of the expansion event of the remaining group may be difficult to precisely determine. Hanazaki et al. (2017) suggested that the transition from MIS 4 to MIS 3 (the early MIS 3), around 53000 years ago when the climate became substantially warmer in the last ice age (Helmens et al., 2009; Helmens and Engels, 2010; Weber et al., 2018), is a candidate turning point.

The vegetation reconstruction over the past 150000 years provides insight into this assignment (Figure 1C; Igarashi and Oba, 2006). Fossil pollen analyses including Quercus species use the \( Tp \) index, the ratio of the pollen abundance of cool-temperate broadleaf trees to the sum of that of cool-temperate broadleaf plants and subarctic conifers (Igarashi and Oba, 2006). Several peaks of recovery of broadleaf plants have been detected, including those around 150000, 50000–60000, and 130000 years ago (Igarashi and Oba, 2006). These effectively explain the expansion events of acorn-dependent wood mice and large flying squirrels (Hanazaki et al., 2017). By contrast, voles are typical herbivorous animals that were affected by early MIS 3, which corresponds to a period of grassland expansion (Honda et
Figure 1. The possible influence of Paleo-environmental fluctuations on the population dynamics of terrestrial animals. (A) A map of the Japanese archipelago showing the four main islands of Hokkaido, Honshu, Shikoku, and Kyushu, along with the associated islands of Sado, Izu, Tsushima, and Satsunan. These surrounding islands are separated from their respective nearby major islands, as well as from Hokkaido and Honshu, by deep sea (i.e., ~120 m). (B) The marine oxygen isotope curve over the last 150000 years, adapted from Lisiecki and Raymo (2005), showing the marine isotope stages (MIS). The approximate periods of the last glacial maximum (LGM) and penultimate glacial maximum (PGM) are indicated with horizontal bars. The four critical time points for population expansion events are marked with vertical bars, called Ia, Ib, II, and III here. (C) Inferred climatic changes over 144000 years based on high-resolution pollen analysis of a core from the northwest Pacific off the coast of central Japan (Igarashi and Oba, 2006). The $T_p$ value is the ratio of the pollen abundance of cool-temperate broadleaf trees to the sum of that of cool-temperate broadleaf plants and subarctic conifers (defined by Igarashi and Oba, 2006). (D) The sea level change over the last 140000 years (Lambeck et al., 2014), highlighting the two critical time points (Ib and III) when the temperature became warm and the sea levels were very shallow, allowing gene flow between remote islands via land bridges. (E) A global temperature fluctuation since ca. 17000 years ago (source: https://www.climate.gov/sites/default/files/historictemperaturerecord_greenland_large.jpg), indicating the critical time points of Ia and Ib and the period of the Younger Dryas (YD). (F) Global sea level change for the last 35000 years, covering LGM and YD (Lambeck et al., 2014; Harrison et al., 2019).
The importance of the early MIS 3 period for demographic expansion has been noted in grassland voles in North America (Kohli et al., 2015), and in hares in Europe (Fickel et al., 2008). However, an early MIS 3 expansion event has also been posited in a forest-dwelling wood mouse (A. argenteus) from Japan (Hanazaki et al., 2017) and roof rats (Rattus rattus) in Myanmar (Maung Maung Theint et al., 2019).

For the two Group I subgroups of \( \tau \) values, Ia and Ib can be assigned to the expansion events immediately after the termination (c. 11000 years ago) of the Younger Dryas (YD) and the onset of the Bølling–Allerød warming (c. 15000 years ago), respectively, although the environmental recovery is latitude dependent and there are some time differences among regions (Igarashi et al., 2012). YD, a global cold reversal event, is thought to have affected the population dynamics of terrestrial animals (Herman and Searle, 2011; Suzuki et al., 2015).

The assignment of Stages Ib and III to the time periods near the last glacial maximum (LGM) and penultimate (PGM) glacial maximum is congruent with the fact that the haplogroups corresponding to these stages contain members that found refuge on remote islands, such as Sado and Tsushima Islands, currently separated by deep sea. This implies that the expansion events at Stages Ib and III involved migration over land bridges across deep straits at a time when the sea level was low, about ~100 m (Figure 1D, F).

From these considerations, calibration points can be set for the four stages: Ia, Ib, II, and III, respectively (see Figure 1B). Previously, we obtained representative \( \tau \) values for the mitochondrial Cytb sequences (1140 bp) corresponding to the four stages (Suzuki et al., 2015; Hanazaki et al., 2017; Honda et al., 2019). If we use \( \tau \) values for the four stages of 2.7, 3.9, 5.7, and 8.5, respectively, the evolutionary rates (\( \mu \)) of Cytb are estimated to be 0.11, 0.11, 0.047, and 0.029 substitution/site/myr (Figure 2); the first two points have identical evolutionary rates. Notably, this curve has two phases, which I call Phases I (high substitution rates) and II (low substitution rates).

Contrary to initial expectations, the evolutionary patterns of the time dependency were similar among small rodent species and other animals. A study of the Eurasian jay (Garrulus glandarius) from Honshu and Kyushu, Japan detected an expansion signal in the Cytb sequences (1143 bp) with a \( \tau \) value of 1.8 (Aoki et al., 2018). If we set the calibration point as 11000 years ago (around the end of YD), the calculated evolutionary rate is 0.072 substitutions/site/myr. In the arthropod Ligidium japonicum, mitochondrial cytochrome c oxidase I gene (COI) sequences (496 bp) from Hokkaido, northern Japan, showed an apparent rapid expansion signal (Harigai et al., 2020). Assuming that the expansion event happened immediately after the end of YD (c. 10000 years ago in Hokkaido; Igarashi et al., 2012), the calculated evolutionary rate of COI is 0.084 substitutions/site/myr, with a \( \tau \) value of 0.83 (Harigai et al., 2020). Therefore, the mtDNA evolutionary rates calculated for divergence on short evolutionarily timescales may not be markedly different among animals including those of vertebrates and invertebrates.

It is possible to estimate the evolutionary rate by considering the divergence of A. speciosus on the remote islands of Hokkaido, Sado, Tsushima, Izu, and Satsunan, which are predicted to have been connected to the main islands of Japan (i.e. Honshu, Shikoku, and Kyushu) via land bridges that arise when the sea level drops every 10000 years (Figure 3A; see Hanazaki et al., 2017). Three calibration points were considered—140000, 250000, and 350000 years ago—allowing the estimation of an evolutionary rate of around 0.03 substitutions/site/myr (Figure 3B, C). The evolutionary rate curve was in good agreement with those derived from the calibration points for the last 130000 years mentioned above (Figure 3C). The pattern of the evolutionary rate curve in rodents is similar to that reported for human mtDNA (Henn et al., 2009).

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**Figure 2.** A plot of genetic distances based on rapid expansion events of the Japanese wood mouse species Apodemus speciosus and Apodemus argenteus, tentatively assigning the historic time points 11000, 15000, 53000, and 130000 years ago. These points were obtained from the assumption of the association of specific rapid cooling and rapid warming events. The plot comprises two phases, designated Phases I and II. Nucleotide substitutions are considered to be of two types: mildly deleterious and non-deleterious mutations, which are expected to be removed and retained from the sequences, respectively. Five straight lines of the genetic distances with the respective evolutionary rates (substitutions/site/year) are indicated for comparison.
Comparison with other mtDNA studies of small rodents

Concern regarding the highly accelerated evolutionary rates

There is still a huge range in the rate of mtDNA evolution considered by researchers. When inferring divergence times on shorter timescales based on the mtDNA D-loop (control region) and Cytb in rodent phylogeographic studies, an accelerated evolutionary rate of around 0.4 substitutions/site/myr (e.g. Rajabi-Maham et al., 2008; Herman and Searle, 2011) is often used; a rate of around 0.1 is used less frequently (e.g. Rajabi-Maham et al., 2008; Suzuki et al., 2013). Here, I would like to point out concerns regarding the higher evolutionary rate.

First, Rajabi-Maham et al. (2008) determined the higher evolutionary rate of mtDNA (0.4 substitutions/site/myr) assuming that the European subspecies of the house mouse, Mus musculus domesticus (DOM), underwent expansion 12000 years ago. However, they presented another option with the expansion starting 60000 years ago using an evolutionary rate of 0.1 substitutions/site/myr. I favor the second option, because a range-expansion event has not always triggered a rapid increase in the genetic diversity of mtDNA; instead, it was often associated with ancient divergence in mtDNA. Relatedly, we revealed that a Japanese rodent (M. smithii) acquired genetic variation in mtDNA in ancient times, and shows no sign of having been affected by the relatively recent rapid expansion event, namely the post-LGM event, although it has extended its distribution range to northern Honshu at present (Honda et al., 2019). This may be because the LGM was insufficiently intense to create a bottleneck.

Second, if we draw a curve showing the base substitutions with a high evolutionary rate of 0.4 substitutions/site/myr over the short timescale, it is difficult to connect the two straight lines of Phases I and II, as shown in Figure 2. If the evolutionary rate of Phase I is high (e.g. more than 0.2 substitutions/site/myr), it seems unlikely to accommodate the two genetic distance curves of Phases I and II.

Third, phylogeographic studies using a high evolutionary rate of 0.4 substitutions/site/myr inferred very recent expan-
sion, over the past few thousand years, and tend to predict that human activity is a major determinant of expansion events, even in wild rodents (e.g. Herman et al., 2017). However, it is difficult to accept the idea that humans influenced the population dynamics of non-commensal animals such as wood mice and voles.

**Population dynamics of wood and house mice**

For comparison with Japanese wood mouse species, I examined the mtDNA variation of the European wood mouse *Apodemus sylvaticus* from Italy and the western Balkans (Clade Ia; Michaux et al., 2003). Using *Cytb* sequences (949 bp) with no ambiguous sites (*n* = 16) reported in the literature (Michaux et al., 2003; Kryštufek et al., 2012), a median-joining network (Bandelt et al., 1999) was constructed (Figure 4A). This had a star-shaped pattern suggestive of a past rapid expansion event, which was supported by Tajima’s *D* and Fu’s *F* analyses. This was consistent with the results of a mismatch distribution analysis using Arlequin 3.5.1.2 (Excoffier, 2004; Excoffier and Lischer, 2010) showing a unimodal pattern and non-significant sums of the squared deviation and raggedness values (data not shown). The *τ* value obtained was 7.20 and the estimated time (*T*) when the expansion started was 131000 years ago (*T* = 7.2/2/949/0.29), with an evolutionary rate of 0.29 substitutions/site/myr. Notably, the network yielded a small star-shaped cluster of haplotypes from Italy (Cluster Ia-1). Although the number of sequences was small (*n* = 7), the calculated *τ* value was 2.1, implying that the expansion event started 10000 years ago. This suggests that a post-LGM event promoted a rapid expansion somewhere in Italy. Overall, the trend in the occurrence of expansion events was similar to that of the Japanese wood mouse, in which the two expansion events follow the PGM and LGM. Considering this with the results of previous studies (e.g. Michaux et al., 2003; Herman et al., 2017), effort should be made to reconstruct the evolutionary history of the population dynamics of *A. sylvaticus*.

Next, I considered the population dynamics of the European subspecies of *M. musculus*, DOM, in which many researchers, including anthropologists, are interested as a route to better understanding the link between mouse history and prehistoric human activities. I obtained *Cytb* sequences (958 bp) from databases representing western Europe and adjacent Asian countries, including Turkey and Israel (*n* = 107; Suzuki et al., 2013; Fornuskova et al., 2014), and constructed a median-joining network (Figure 5A). The network was star-shaped and comprised 15 distinct lineages, six of which were characterized as haplogroups (denoted as HG), as reported by Fornuskova et al. (2014), with the star-shaped pattern suggestive of past rapid expansion events. The sizes differed among the six haplogroups, suggesting that the expansion events happened at different times during the course of evolution.

![Figure 4. Median-joining networks constructed from (A) Cytb sequences (*n* = 16; 949 bp) of *Apodemus sylvaticus* from Italy and the Balkans. Michaux et al. (2003) called the cluster ‘Subclade Ia.’ The number of mutations between haplotypes is indicated on the branches. The size of the circles is proportional to the number of samples. The compositions of sample localities are also reflected in each haplotype. Mismatch distributions of the mitochondrial Cytb sequences of Clusters Ia (B) and Ia-1 (C) indicated significant evidence of expansion. Bars indicate the observed frequencies of mutations between haplotypes and the line is the expected frequency under the sudden expansion model.](image-url)
Figure 5. Median-joining networks constructed from (A) Cytb sequences (n = 107; 958 bp) of Mus musculus domesticus. The number of mutations greater than ten between haplotypes is indicated on the branches. The size of the circles is proportional to the number of samples. The compositions of sample localities are also reflected in each haplotype. Six blocks (T1–T6) forming the ‘torso’ are indicated. Mismatch distributions of the mitochondrial Cytb sequences using data for Mus musculus (B), haplotype groups (HG; Fornuskova et al., 2014) HG-A (C), and HG-F (D) show significant evidence of expansion. Bars indicate the observed frequencies of mutations between haplotypes and the line denotes the expected frequency under the sudden expansion model. Haplogroup HG-F addresses structure T1 (E), which can be explained by either parallel mutation (F) or back (reverse) mutation (G).
The DOM cluster (Figure 5B) and haplogroups HG-A (Figure 5C) and HG-F (Figure 5D) were subjected to a mismatch distribution analysis and neutrality tests using Arlequin 3.5.1.2 (Excoffier, 2004; Excoffier and Lischer, 2010), confirming the notion of rapid overall expansion. The $r$ value of the DOM cluster ($n = 107$) was $5.00$ (95% confidence interval (CI): 4.40–5.52) and the estimated time when the expansion started ($T$) was $53900$ (95% CI: 47440–59600) years ago, with an evolutionary rate of 0.047 substitutions/site/myr, attributing the expansion event to the abrupt warming in early MIS 3. This is in good agreement with the estimate of Rajabi-Maham et al. (2008) of 60000 years ago for the DOM D-loop sequence data.

Haplogroup HG-A ($n = 12$) comprised haplotypes from western Europe and experienced a rapid expansion event, with a $r$ value of 1.82 (95% CI: 0.95–3.00). With an evolutionary rate of 0.11 substitutions/site/myr, the calculated start of the expansion was $8400$ (95% CI: 4400–13800) years ago. By contrast, HG-F comprising haplotypes mainly from northwestern Europe had a $r$ value of 1.60 (95% CI 0.57–3.06), with an estimated expansion start time of $7600$ (95% CI: 2700–14500) years ago. Conceivably, the rapid population growth of the European house mouse is linked to the early introduction of agriculture to Europe roughly 7000–8000 years ago (Shennan et al., 2013).

**Application of the evolutionary rate to evolutionary events in the Holocene**

**Inference of the evolutionary history of the house mouse from whole mitochondrial genome sequences**

We applied the time-dependent evolutionary rate to assess the natural history of the house mouse, addressing the factors shaping its long-distance dispersal with human agriculture in prehistoric times (Li et al., 2020b). We determined whole genome sequences of 98 *M. musculus* collected mainly from Asia, constructed a phylogenetic tree, and estimated time using BEAST software (Drummond and Rambaut, 2007). Figure 6 shows a schematic representation of the early prehistoric eastward movements of two *M. musculus* subspecies, *M. m. musculus* (MUS) and *M. m. castaneus* (CAS), as predicted from the whole mitochondrial genome sequences (Li et al., 2020b). The movement of MUS was achieved mainly by sublineage MUS-1 and involved five processes: (1) the initial spread of MUS-1 to northern Eurasia, including westernmost China; (2) movement to the eastern part of western China; (3) expansion to northern China; (4) introduction to the Korean peninsula; and (5) colonization of Japan by a descendant of the Korean haplotype group. The eastward movement of the CAS lineage was undertaken by sublineage CAS-1, resulting in coverage of northernmost China by 9000 years ago. The second CAS-1 dispersal event was movement either from the east coast of India to southern China or vice versa. The next step is simultaneous dispersal from southern China to several peripheral regions, including the eastern coast of India, Sri Lanka, and Bangladesh. The final step is dispersal from southern China to the Japanese archipelago, Far Eastern Russia, and Yunnan, China.

The history of the house mouse inferred from phylogenetic analysis with 0.11 substitutions/site/myr matches the spatiotemporal dynamics of prehistoric humans based on agriculture, particularly the introductions of millet farming in the Korean peninsula around 5300 years ago (Crawford and Lee, 2003; Li et al., 2020a) and paddy rice culture to the Japanese islands in the early Yayoi period 3000 years ago (Crawford and Lee, 2003; Fuller, 2011). The Austronesian language peoples dispersed to distant locations, including Indonesia, 4000–5000 years (e.g. Lipson et al., 2014; Deng et al., 2020) and this may have caused the dispersal of the house mouse to Indonesia. Recent work on the population dynamics of *M. musculus* from Myanmar showed evident population expansion in the Cyth sequence data that began about 5400 (95% CI: 2300–8000) years ago (Maung Maung Theint et al., 2020). In addition, the migration from southern China to the Japanese archipelago was deduced to have occurred 3500 years ago (Li et al., 2020b), which may be correlated with the introduction of cold-tolerant temperate japonica rice strains developed in response to the extreme environmental degradation 4200 years ago (Mao et al., 2019; Guo et al., 2020; Gutaker et al., 2020).

**A presumed evolutionary trend in the time-dependent evolutionary rate of mtDNA**

One concern is the mechanisms underlying the time dependency of the mtDNA evolutionary rate. A plausible hypothesis is that a weakly harmful mutation is removed over time (e.g. Rand, 2001; Penny, 2005; Henn et al., 2009; Ho et al., 2011), so the evolutionary rate will reflect the rate of weak deleterious mutations, in addition to the rate of neutral mutations that are not harmful (Figure 2). From this perspective, the initial stage (Phase I) is one in which new mutations are created, some of which are subjected to selection pressure and removed in the subsequent stage (Phase II). This probably occurs because, while plants and other organisms have a specific mtDNA repair system, animals lack this system (Wu et al., 2020). In the early stage of divergence in animal mtDNA, the rate of evolution is inevitably high because the proportion of mildly deleterious mutations is relatively high. As the divergence time becomes older, the proportion of the latter increases and the former portion becomes negligible.

The removal of weakly deleterious mutations is known as compensatory evolution and can be explained by the removal of the weakly deleterious mutations from each sequence (Meer et al., 2010; Osada, 2016). One of the mechanisms is called back (reverse) mutation. In relation to this issue, homoplasic mutations appear as a ‘torso’ in a network (Bandel et al., 1999). The torso structure can be either explained by parallel or back mutations, because mtDNA does not experience recombination (Figure 5E–G).

The occurrence of homoplasic mtDNA mutations is well documented (Rand, 2001; Henn et al., 2009). This may due to their particular mode of evolution, namely a severe germline bottleneck (Rand, 2001; Zaidi et al., 2019), which may contribute to the relatively high frequency of homoplasies given the high incidence of competition among haplotypes during oocyte development. The trends to a high AT base composition, high transition-to-transversion ratio, high
Comparison of the time dependency among mtDNA genes

Does the time dependency of mtDNA genes differ? Here, I examined the Cytb (1140 bp), COI (1463 bp), and ND2 (929 bp) genes from 98 house mouse whole genome sequences (16038 bp; Li et al., 2020b). I identified two phases of haplotype diversity with distinct genetic distances (d): one for divergence within the subspecies groups \(0 < d < 0.015\), and the other for divergence between subspecies groups \(0.02 < d < 0.03\). In the relatively shallow comparison of Cytb, COI, and ND2, the ratios to that of the whole mitochondrial genome sequences are 1.3, 1.1, and 1.0, respectively (Figure 7A–C), indicating that the three genes are somewhat similar. By contrast, on comparing the relatively deeper divergence, the ratios are 1.3, 0.96, and 1.45, respectively (Figure 7D–F), and the differences among genes are larger, suggesting that the evolutionary rates are similar in the initial accumulation of mutations and differ with the removal of moderately deleterious mutations. In other words, the difference in evolutionary rates among gene regions is due to the difference in the extent of purifying selection among gene regions; the deleterious effects are greater in ND2 than Cytb. This provides evidence of the time dependency of the evolutionary rate of mtDNA.
Conclusion

Considering the global environmental changes during the late Quaternary (i.e. the last 150,000 years), the time periods responsible for the rapid expansion events were estimated to be around 110,000, 150,000, 530,000, and 130,000 years ago. These give evolutionary rates of 0.11, 0.11, 0.047, and 0.029 substitutions/site/myr, respectively, showing apparent time dependency (Figure 3C). The evolutionary history of the house mouse *M. musculus* inferred from the higher evolutionary rate effectively explains its association with regional agricultural development in prehistoric times, implying that the rate is applicable for assessing divergence events in the Holocene. The higher evolutionary rate might apply to a short timescale, i.e. the last 20,000 years; lengthening the timescale would give unrealistic results (Figure 3C). I would like to emphasize that the Japanese archipelago, which consists of many islands extending from subarctic to subtropical and separated by deep sea, makes it possible to utilize biogeographic calibration points to draw evolutionary rate curves in other faunal groups as well.
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