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Article:

**Ancient onset of geographic divergence, interpopulation genetic
exchange, and natural selection on the *Mc1r* coat-colour gene in the
house mouse (*Mus musculus*)^β**

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Running head: HYBRID ORIGIN OF HOUSE MICE IN THEIR HOMELAND

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colour – selective sweep.

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We examined genetic variation in house mice from India and Pakistan, a predominant part of the predicted homeland of this species and the territory of the subspecies *Mus musculus castaneus* (CAS), using a nuclear marker for seven tandemly arranged genes (5 *Fanca–Spire2–Tcf25–Mc1r–Def8–Afg3l1–Dbndd1*) and compared them with those previously determined for mice from other parts of Eurasia. Construction of a network with the concatenate sequences yielded three distinct clusters representing the three major subspecies groups: CAS, *M. m. domesticus* (DOM), and *M. m. musculus* (MUS). STRUCTURE analysis provided evidence for further subdivision of CAS into two main 10 haplogroups within the Indian subcontinent. Single-gene networks revealed not only gene-specific architecture for subgrouping in CAS, but also allelic exchange among subspecies. These results suggest the earlier onset of allopatric divergence in the predicted homeland (the Middle East and Indian subcontinent) and subsequent intermittent admixing via gene flow across the CAS haplogroups and among the three 15 subspecies groups. A comparison of the levels of nucleotide diversity among the gene regions revealed a less divergent state in the chromosome region containing *Mc1r* and its adjacent genes, indicative of a selective sweep, suggesting the involvement of natural selection in the *Mc1r* allelic variation.

INTRODUCTION

Mus musculus originated from south-western Asia and extended its range to all of Eurasia in association with prehistoric human movements. The species is now a well-known model organism and a large number of strains with different geographic origins and hence different genetic backgrounds are used for biomedical research (e.g. Takada *et al.*, 2013). Therefore, knowing the genetic background of the species throughout its range would be useful. In addition, tracking the dispersal of mouse lineages should give insight into the prehistoric movements of the humans that the mice accompanied (e.g. Boursot *et al.*, 1993; Bonhomme & Searle, 2012; Cucchi, Auffray & Vigne, 2012; Gabriel, Mathias & Searle, 2013; Jones *et al.*, 2013; Jing *et al.* 2014). The origin of *M. musculus* before its human-mediated dispersal is most commonly identified as either India (Boursot *et al.*, 1993, 1996; Din *et al.*, 1996) or the Middle East (Prager, Orrego & Sage, 1998; Suzuki *et al.*, 2013), based on various molecular markers, including mitochondrial DNA, allozymes, and nuclear intron sequences. Under these scenarios, the ancestral stock gave rise to three dispersing lineages, *M. m. castaneus* (CAS; southern Asia, Southeast Asia, southern China), *M. m. domesticus* (DOM; Western Europe), and *M. m. musculus* (MUS; northern Eurasia excluding Western Europe), which made secondary contacts in Eurasia (Boursot *et al.*, 1993). However, difficulty has arisen in assessing the evolutionary history in this species, even after intensive molecular work, for various reasons, including intra- and interspecies introgression events and intricate human-mediated propagation in both early prehistory and recent times. Regardless, no intensive genetic analyses of mice have been performed in their original range.

Mitochondrial DNA (mtDNA) analyses have identified five distinct lineages that diverged roughly a half million years ago (mya): those representing the three major subspecies CAS, DOM, and MUS and two local lineages confined to Yemen (*M. m. gentilulus*) and Nepal (Prager *et al.*, 1998; Suzuki *et al.*, 2013). In contrast, studies using nuclear markers provide substantially different spatiotemporal views of the evolution of *M. musculus*. Based on isozyme analyses, mice from southern Asia, the hypothesised origin of CAS, are separated into two geographic groups: mice from Iran, Afghanistan, Pakistan, and northern India, called “*Mus bactrianus*” (or *M. m. bactrianus*) and mice from the rest of Eurasia, including southern Asia, called *M. m. castaneus* (Bonhomme *et al.*, 1984). In addition, sequences of nuclear genes such as *Irbp* and *Rag1* suggest ancient divergence dating back to that between *M. musculus* and closely related species such as *M. spretus* and *M. spicilegus* (e.g. 1.7 mya; Suzuki *et al.*, 2004). In comparison,

the mtDNA analyses tend to show lower levels of divergence among the geographic groups, including those from the territory of “*M. bactrianus*” (0.1–0.2 mya; Suzuki *et al.*, 2013). Therefore, the genetic architecture of CAS remains unclear.

Organisms in the wild are subject to a specific evolutionary mode, called
5 reticulate evolution, in which genetic introgression and hybridisation occur
intermittently among closely related species or geographically situated conspecific
populations, as is documented in *M. musculus* (e.g. Bonhomme *et al.*, 2007; Nunome *et al.*, 2010; Rajabi-Maham *et al.*, 2012). Analyses of several different phylogenetic
10 markers are essential to elucidate the evolutionary histories of organisms in which
multiple hybridisation events are suspected over the long term. Haplotype structure
analysis, which monitors recombination events focused on variation in a chromosome
region using representative linked gene markers, is effective for assessing the time span
after detecting recombinant haplotypes and can be used to time ongoing hybridisation
15 events (Martinsen *et al.*, 2001; Koopman *et al.*, 2007; Nunome *et al.*, 2010). Since the
density of recombination points reflects the number of generations since the
hybridisation event, assessment of haplotype structure, which is defined by a
combination of alleles in an appropriate length of chromosome, allows us to consider
spatiotemporal aspects of hybridisation.

Nunome *et al.* (2010) adapted haplotype analyses to wild mice mainly from East
20 Asia using eight linked gene markers. The aligned set of sequences has proven to be
useful as a biparental genealogical marker in inferring population genetic structure.
Moreover, by showing the recombinant haplotypes, it clarified the presence of historical
hybridisation between CAS and MUS in northern Japan, including Hokkaido and
northern Honshu. However, this study drew attention to several unresolved issues
25 related to the evolution of Eurasian wild mice. First, MUS is further subdivided in
northern Eurasia into northern (MUS-I; Eastern Europe, Siberia, and Primorye) and
southern (MUS-II; Kazakhstan, Uzbekistan, northern China, Korea, and Japan)
phylogroups, although little is known about this evolutionary episode. Second, the
recombinant haplotypes from northern Japan involve components of MUS, CAS, and
30 DOM. The presence of a short DOM segment (20–30 kb) was unexpected because it
contradicts the expectation that Japanese wild mice possess ancient MUS and CAS
DNA, as the major and minor components, respectively (Terashima *et al.*, 2006). Third,
the level of variability differs markedly from locus to locus, even within the same
subspecies (Nunome *et al.*, 2010). For example, the gene haplotypes assigned to MUS
35 showed diverse patterns, ranging from one (*Fanca*) or two (*Spire2*) to 10 (*Dbn1d1*) or

23 (*Afg3ll1*) haplotypes. Finally, even CAS and DOM possess their own sequences over the loci examined, except for *Mc1r* (409-bp analysis), for which CAS and DOM share the same *Mc1r* allele, while subspecies-specific sequences are detected in both neighbouring genes.

5 Haplotype structure analysis also offers the prospect of examining the presence of positive natural selection on specific functional genes through an investigation of the reduction in nucleotide diversity, a sign of a selective sweep (*e.g.* Nielsen *et al.*, 2005). This phenomenon is characterised by extensive linkage disequilibrium and limited haplotype diversity and is seen in a variety of organisms, including North American
10 grey wolves (*Canis lupus*; Anderson *et al.*, 2009) and wild mice (*M. spretus* and *M. musculus*; Song *et al.*, 2011). Among rapidly evolving phenotypes, coat colour variation is an ideal character to examine. In Asia, the dorsal coat colour variation in *M. musculus* due to melanin synthesis is significantly correlated with precipitation: dark coats are observed in more humid habitats and pale coats in drier habitats (Lai *et al.*, 2008). This
15 implies a role of concealment as a selective force affecting dorsal coat colour in house mice (Lai *et al.*, 2008). Two genes, agouti signalling protein (*Asip*) and the melanocortin-1 receptor (*Mc1r*), effectively drive the evolution of coat colour via phenotypic change arising from single mutations (*e.g.* Hubbard *et al.*, 2010; Suzuki, 2013). For example, the effect of natural selection on the coat colour gene *Asip* has been
20 documented in the deer mouse *Peromyscus maniculatus* (Linnen *et al.*, 2009; Vignieri, Larson & Hoekstra, 2010). In *Peromyscus polionotus*, another example, amino acid changes in *Mc1r* are associated with an adaptive coat colour phenotype, which functions as camouflage against pale sand dunes (Mullen *et al.*, 2009). The *Mc1r* sequences show different evolutionary rates of amino acid replacement in the phylogeny of *Mus*,
25 possibly associated with coat colour changes for local adaptation (Shimada *et al.*, 2009). The sequence variation in *Mc1r* and its neighbouring genes lying in a 200-kb stretch has been analysed in wild mice collected from throughout Eurasia and a broad area of the Japanese islands (Nunome *et al.*, 2010). Investigation of this chromosome region with sufficient specimens should help us to determine the impact of natural selection on coat
30 colour genes.

In this study, our basic objectives were to explore various evolutionary issues related to *M. musculus*, specifically its genetic state in its homeland before it underwent the long-range dispersal events associated with prehistoric human movements. Knowledge of the genetic architecture of the homeland area would help to identify the
35 exact source areas of the lineages that dispersed with human activities in the past and

recently. Here, we focus on the CAS lineages in Southeast Asia, southern China, and Japan. We also focused on the involvement of natural selection on the *Mclr* gene in the process of the genetic differentiation of the subspecies groups. We determined the sequences of the seven genes located in the chromosome region near *Mclr* in wild mice collected from India and Pakistan and conducted phylogeographic analyses making use of available sequences representing other geographic lineages of Eurasian wild mice (Nunome *et al.*, 2010).

MATERIALS AND METHODS

SPECIMENS

We examined the genetic variation in nearly 40 wild-captured mice from 18 localities in India and Pakistan (Table 1; Fig. 1), most of which were collected by H. Ikeda and K. Tsuchiya in 1989–1992 (Yonekawa *et al.*, 2003; Suzuki *et al.*, 2013). The 54 wild-captured mice previously examined are those from 52 localities throughout Eurasia used in our previous study (Nunome *et al.*, 2010). We determined gene sequences in five closely related species, three from the same species group, *Mus spretus* (strain SEG), *M. spicilegus* (HS3212), and *M. macedonicus* (HS537), and two relatively divergent taxa, *M. caroli* (HS1469) and *M. terricolor* (HS3962) (see Suzuki *et al.*, 2004; Suzuki & Aplin, 2012 for their inferred phylogenetic relationships).

SEQUENCE ANALYSES

Previously, we examined eight genes, which are arranged at 20–30-kb intervals in a 200-kb portion of mouse chromosome 8 (Nunome *et al.*, 2010). Here, we re-examined the genes, excluding *Tubb3*, which has less pronounced allelic diversity. We amplified segments of *Fanca* (448 bp), *Spire2* (436 bp), *Tcf25* (613 bp), *Mclr* (409 bp), *Def8* (370 bp), *Afg3l1* (510 bp), and *Dbnndl* (550 bp) (see Fig. 2 of Nunome *et al.*, 2010). The primers used in the polymerase chain reaction (PCR) analyses included those used by Nunome *et al.* (2010). The sequence of the entire coding region (948 bp) of mouse *Mclr* was determined using the primers previously described (Shimada *et al.*, 2009). We determined sequences of the non-coding region (*Mclr-NC*) immediately upstream of the *Mclr* coding region, with primers newly designed (Mc1r_ NCF: 5'-TCCCTTCTTCTCCAGAGTCC-3'; Mc1r_ NCR: 5'-GCACGATGCTGACACTTACC-3') using the Ensemble Mouse Genome Database

(NCBI; <http://www.ncbi.nlm.nih.gov/>). The amplifications were carried out for 35 cycles each consisting of 30 s at 96°C for denaturation, 30 s at 55–64°C for annealing, and 30 s at 72°C for extension. The reaction mixtures (20 µl) contained 2.5 mM MgCl₂. Master Mix 360 DNA Polymerase (Applied Biosystems, Foster City, CA, USA) was used in the PCR. The first PCR cycle was preceded by 10 min at 95°C to activate the polymerase. The double-standard PCR product was purified using a 20% polyethylene glycol–2.5 M NaCl precipitation method, sequenced using the PRISM Ready Reaction Dye Deoxy Terminator Cycle Sequencing Kit (Applied Biosystems), and run on an ABI3130 automated sequencer (Applied Biosystems). Both strands were sequenced using either the forward or reverse primer used for PCR.

HAPLOTYPE ASSESSMENT FROM ALLELIC COMBINATIONS OF THE SEVEN LOCI

We used PHASE v2.1 (Stephens, Smith & Donnelly, 2001; Stephens & Scheet, 2005) with the default settings to determine unique allele arrangements in each of the gene regions examined. Networks of the single-gene sequences and concatenated gene sequences for *Fanca*, *Spire2*, *Tcf25*, *Mcl1r*, *Def8*, *Afg31l*, and *Dbnddl* were constructed using the Neighbour-Net (NN) method, as implemented in SPLITS TREE v4.11.3 (Huson & Bryant, 2006). The number of pairwise differences, nucleotide diversity (π), neutrality test, and Tajima's D were calculated using ARLEQUIN (ver. 3.5) (Excoffier & Lischer, 2010).

Population genetic structure was inferred by Bayesian clustering analysis using STRUCTURE, version 2.3 (Pritchard, Stephens & Donnelly, 2000) with concatenated sequences for the seven loci. The dataset used in the clustering analysis consisted of a total of 208 SNPs. K was estimated from 5 000 000 Markov chain Monte Carlo (MCMC) generations, sampled every 10 000 generations after a burn-in period of 2 000 000 generations. To determine an appropriate K , we used Evanno's method in STRUCTURE HARVESTER (Earl & vonHoldt, 2012).

Considering the lengths of exogenous segments, one can predict the date when introgression took place (Nunome *et al.*, 2010). A general indication of the time frames of meiotic recombination events can be inferred from the formula $P = (1 - r)^G$ (Stephens *et al.*, 1998), where $1 - P$ is the probability of recombination, G is generations, and r is the recombination rate (neglecting mutation). Transforming this equation, G can be estimated with $G = -\ln(P)/r$. The average recombination rate within the *Mus* genome was estimated to be 0.52 cM/Mb (Jensen-Seaman *et al.*, 2004). For example, a 10-kb segment carries an estimate of 5.2×10^{-5} for r . The estimate of G is the sum of

generations for exogenous segments in the heterozygous state and does not account for generations in the homozygous state, and is consequently the minimum value for the generations elapsed since the introgression event.

5 ESTIMATION OF TIMES OF DIVERGENCE

BEAUti and BEAST v1.7.5 (Drummond *et al.*, 2012) was used for the Bayesian Markov-chain Monte-Carlo (MCMC) analyses of the nucleotide sequence data (Drummond *et al.*, 2005) to estimate the time to the most recent common ancestor (TMRCA) and 95% highest posterior density (HPD). As outgroup taxa, we used *Mus caroli*, *M. terricolor*, *M. spretus*, *M. spicilegus*, and *M. macedonicus* and the predicted divergence time of 1.7 mya was used as a prior for the node separating *M. spretus* and *M. musculus* (Suzuki *et al.*, 2004). For the MCMC posterior analyses, the chain length was 10,000,000. After a 1000-tree burn-in, 10,000 trees were used for the analyses. The Bayesian tree was generated under the relaxed clock model (HKY substitution) and the best-fit model was inferred in MEGA v5.0 (Tamura *et al.*, 2011). The convergence of the MCMC chains and the effective sample size (ESS) values exceeding 200 for all parameters were assessed using Tracer v1.5 (Rambaut & Drummond, 2009).

DATA ARCHIVING

20 The nucleotide sequences reported in this paper appear in the DDBJ, EMBL, and GenBank nucleotide sequence databases under the accession numbers AB909504–AB909921. Sequence data files in nexus file format, together with Supplementary Information files, will be stored in the Dryad repository.

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RESULTS

PHYLOGENIES OF SINGLE GENE SEQUENCES

We determined the sequences of seven markers used previously (*Fanca*, *Spire2*, *Tcf25*, *Mc1r*, *Def8*, *Afg3l1*, and *Dbnmdl*; Nunome *et al.*, 2010) in 39 mice collected from India and Pakistan covering a major portion of the predicted homeland of CAS (Table 1; Fig. 1) and separated allelic sequences with a Bayesian method using PHASE (Stephens *et al.*, 2001; Stephens & Scheet, 2005). Neighbour-net (NN) networks (Fig. 2) were constructed for the seven markers with the newly determined sequences, together with those previously determined (59 mice) assigned to CAS, DOM, and MUS (Nunome *et*

al., 2010). This exemplified a higher level of allelic diversity in CAS mice from Pakistan and India, integrating the sequences of mice from India and Pakistan into two or more divergent clusters in six gene-sequence sets, excluding that of *Mc1r*. Mice from the DOM territory formed a single cluster for each of the seven genes. For mice from the MUS territory, the sequences formed single clusters for the first five loci (*Fanca*, *Spire2*, *Tcf25*, *Mc1r*, and *Def8*), while they formed three distinct clusters for the last two loci, *Afg3ll* and *Dbn1*.

The NN networks show the genetic exchange between the subspecies. The majority of CAS sequences of *Mc1r* (409 bp) from India and Pakistan (81%) was assigned to a single genotype identical to that of the DOM mice. For *Def8*, several mice from northern India (DNA codes 74, 75) and northern Pakistan (60, 62, 63) possessed a sequence assigned to DOM (Fig. 2A). The allelic sequences of mice from Japan were overwhelmingly derived from MUS, although some from northern Japan (northeast Honshu and Hokkaido) were non-MUS sequences, showing high affinity with CAS sequences (*Spire2*, *Tcf25*, *Mc1r*, and *Dbn1*) and DOM (*Def8* and *Afg3ll*), as previously observed (Nunome *et al.*, 2010). The non-MUS *Def8* sequence from northern Japan was identical to that in DOM mice and the above-mentioned mice from the CAS territory (Fig. 2A, arrowhead). The *Afg3ll* sequence (510 bp) from northern Honshu and Hokkaido differed from those representing DOM by one base (site 445), so that they were near the DOM cluster in the NN network and differed from any allele from the CAS territory (Fig. 2A, arrowhead).

NETWORK OF CONCATENATE SEQUENCES

Combinations of the seven linked gene regions (*Fanca*, *Spire2*, *Tcf25*, *Mc1r*, *Def8*, *Afg3ll*, and *Dbn1*) were assessed with PHASE (Stephens *et al.*, 2001; Stephens & Scheet, 2005). The NN network of the haplotypes of the concatenated sequences formed several distinct clusters (Fig. 3A). We assessed the genetic ancestry of subspecies, based on the clustering patterns perceived in the networks for the single-gene sequences (Fig. 2A). This allowed us to recognise three subspecies clusters representing the three subspecies groups (CAS, DOM, and MUS), while they simultaneously exhibited a number of recombinant haplotypes or haplogroups. Notably, seven divergent clusters were tentatively seen within the CAS cluster. The haplogroups showed rough geographic distributions in northern India and Pakistan (CAS-A), southern India (CAS-B), northern and central India (CAS-C), Southeast Asia (CAS-D), and India and Pakistan (CAS-E) (Fig. 3C). Notably, CAS-D comprised the haplotypes recovered from

a large geographic area of Southeast Asia, South China and Indonesia and can be characterized as the lineage dispersed with the prehistorical human movement.

One of the recombinant haplogroups seen in the concatenate NN network (Fig. 3A), here designated Re-1[C, D, M], was recovered from northern Japan, and possessed three components that could be assigned to CAS, DOM, and MUS. Another recombinant haplogroup (Re-10[C, D, M]) consisted of haplotypes of a mouse (74-HI184 in Fig. 1) from Delhi, India, in which the alleles of *Tcf25* and *Def8* were closely related to MUS and DOM alleles, respectively. The specific CAS haplotypes from India and Pakistan, in which an allele assigned to DOM was embedded, formed a scattered pattern in the CAS cluster.

The MUS mice tended to form two clusters, with the groups corresponding to geographic regions, *i.e.*, a ‘northern’ (MUS-I) subgroup representing mice from Eastern Europe, Siberia, and Primorye, and a ‘southern’ (MUS-II) subgroup from Kazakhstan, Uzbekistan, northern China, Korea, and Japan (Nunome *et al.*, 2010). In contrast, the DOM group formed a tight-knit cluster on the network tree, with much less internal divergence than within either CAS or MUS.

The Bayesian clustering analysis (Drummond *et al.*, 2012; Earl & vonHoldt, 2012) with the combined dataset for the seven gene markers indicated that the wild mouse population is composed of four genetic clusters ($K=4$: CAS-I, CAS-II, DOM, and MUS), the first two of which are represented by the abovementioned haplogroup A and a haplogroup set of B, D, and E (Fig. 3A), with apparent geographic subdivision into northern (Himalayas) and southern parts of the Indian subcontinent, respectively (Fig. 3C). From the visualisation of $K=4$, most of the individuals (codes 57, 73, 83, 84) belonging to haplogroup C were assigned to the DOM cluster in addition to the CAS related clusters CAS-I and CAS-II.

We performed a phylogenetic analysis using the concatenated sequences ($n=34$; 3402 bp) using BEAST v1.7.5 to assess the phylogenetic relationships and the temporal aspects of the divergence of haplotypes representing the geographic groups. We removed the haplotypes that were assigned to be of hybrid origin deduced from the network and STRUCTURE analyses in order to clarify the temporal patterns of ancient divergences. We used five outgroup taxa with a prior of 1.7 mya for the divergence of *M. spretus* from the other members of the *M. musculus* species group. The phylogenetic analysis recovered the four monophyletic groups predicted by the STRUCTURE analysis (Fig. 4). TMRCA for CAS-I and CAS-II, and DOM and MUS were calculated

to be 1.05 mya (95% HPD = 0.85–1.27 mya) and 0.91 mya (95% HPD = 0.70–1.16 mya), respectively.

SEQUENCE VARIABILITY OF *Mclr* AND ITS FLANKING GENES

5 We determined the entire coding region of *Mclr* (948 bp) from 84 mice including 45 from India and Pakistan to get a finer view of the evolutionary dynamics of single-nucleotide polymorphisms (SNPs) across the subspecies, revealing 28 alleles (Fig. 5A). The coding sequences featured a total of 28 variable sites, with 19 nonsynonymous and nine synonymous mutations. The network tree revealed the
10 presence of a predominant allele (allele 5) representing certain portions of the DOM and CAS mice examined. The prevalent allele possessed a C to T mutation at site 302 changing the amino acid at codon position 101 from alanine to valine (Fig. 5B; Table 1). The sequence analysis of the entire *Mclr* coding region revealed another prevalent mutation (A218G) with a predicted amino acid change from tyrosine to cysteine at
15 codon position 73, in mice from Pakistan and northern India (Table 1).

To further address the divergence of *Mclr* sequences in CAS and DOM, we determined the sequences of the 377-bp *Mclr* non-coding region, *Mclr-NC*, immediately upstream from the *Mclr* coding region in 81 mice and constructed a network with the sequences obtained (Fig. 5A). DOM mice were shown to have a single
20 unique NC sequence (red circle). Eleven alleles were identified among the sequences of mice from India and Pakistan (orange circles), but none of them were shared with the allele exclusive to DOM mice. The mice from northern Japan that had the *Mclr* coding sequence (allele 5) shared by CAS and DOM had the predominant CAS allele in *Mclr-NC*, differing from the NC allele unique to DOM by one base, providing a
25 evidence for the subspecies origin of the coding sequence.

To examine the spectrum of nucleotide diversity (π) across the chromosome region examined in each of the three subspecies groups, we choose non-recombinant haplotypes, accounting for the patterns shown in the network and STRUCTURE analyses (Fig. 3). In total, 172 haplotypes that were assigned as CAS (n = 91), DOM (n
30 = 15), or MUS (n = 66) were used to compare π among the seven gene regions, together with those for *Mclr-NC* upstream from the coding region (Fig. 2B; Table S1). In MUS, an apparent reduction in nucleotide diversity was seen in the *Fanca*, *Spire2*, *Tcf25*, *Mclr*, and *Def8* gene loci, while the remaining two gene loci were markedly divergent, suggestive of recent introgression between the haplogroups MUS-I and MUS-II. In

CAS and DOM, a substantial reduction in nucleotide diversity was observed in *Mc1r* and the adjacent loci at the central portion near *Mc1r* and *Mc1r-NC*, while the levels of the nucleotide diversity were relatively high in genes in the flanking regions on both sides. We calculated Tajima's *D* for each of the three subspecies groups using the data for the eight gene regions examined, including *Mc1r-NC*. Negative values were obtained for *Mc1r* for CAS and DOM, although the values were not significant (Table S1).

DISCUSSION

ASSESSING THE ANCIENT GENETIC STRUCTURING OF *M. MUSCULUS* IN ITS HOMELAND
We assessed the ancient state of genetic structure of *M. musculus* in its presumed species homeland and the dispersal events associated with prehistoric human movements, which led to the current occurrence of the three major subspecies groups in northern Eurasia (MUS), southern Asia (CAS), and westernmost Eurasia (DOM). This study highlights the intricate phylogenetic relationships of the subspecies groups and unveiled the previously concealed evolutionary history of *M. musculus*.

The phylogenetic patterns of concatenated sequences give a general view of the genetic structure of *M. musculus* (Fig. 3). The NN network disclosed three genetically distinct haplogroups representing the three major subspecies groups: CAS, DOM, and MUS. STRUCTURE analysis further suggested the subdivision of CAS into two clusters, here designated CAS-I and CAS-II, with apparent geographic affinity in the northern and southern parts of the Indian subcontinent, respectively (Fig. 3). In contrast to the shallow divergence of mtDNA of *M. musculus* (0.35–0.5 mya, Suzuki *et al.*, 2013), the divergence of these geographic groups is likely as deep as those between closely related species (i.e., *M. macedonicus*, *M. spretus*, and *M. spicilegus*; Fig. 4). This is in good accord with molecular phylogenetic studies on nuclear gene sequences that observed higher levels of divergence (e.g. Suzuki *et al.*, 2004; Bonhomme *et al.*, 2007). These considerations thus imply that *M. musculus* has a long evolutionary history with several distinct geographic groups, perhaps arising 1–2 mya in a wide area of its ancient territory, i.e., Iran, Afghanistan, Pakistan, and northern India.

The haplotypes of mice from the CAS territory were substantially diverse (Figs. 3, 4). Our current phylogenetic work portrayed spatial structuring within the Indian subcontinent. A trend in the basal divergences of the lineages recovered from northern India and Pakistan is present along with clear genetic differentiation in other

haplogroups represented by mice from certain geographic areas, such as CAS-B from the southern part of the subcontinent, with an estimated divergence time of ca. 0.5 mya (Fig. 4). This implies the primary occurrence of CAS to be in Pakistan and northern India and an early secondary range expansion to the southern part of the Indian subcontinent. This contradicts the previous view from mtDNA analyses in which mice from the eastern and southern parts of the Indian subcontinent share less divergent mtDNA sequences from a recent expansion within the last 10,000 years (Suzuki *et al.*, 2013; see Fig. 3C). It is thus suggested that mice occurring in eastern and southern India experienced recent intensive mtDNA introgression, while maintaining an ancient divergence in the nuclear genome.

ANCIENT INTROGRESSION AMONG GEOGRAPHIC GROUPS

The genetic exchanges among the original homeland geographic groups before the long-range dispersal events are poorly documented. However, our results shed light on the genetic exchanges among the geographic groups of *M. musculus* occurring in India and Pakistan. The historical genetic exchanges among the geographic groups are evident in the single-gene network (Fig. 2); the DOM-assigned sequence of *Def8* is recovered from mice from the CAS territory, including Islamabad (code 60), Lahore (codes 62, 63), and Delhi (codes 74, 75). Notably, some CAS mice (e.g. codes 60, 89) formed a cluster separate from the rest of the CAS-assigned alleles and localised near the DOM-assigned cluster for *Spire2* (Fig. 2B). These results imply that intersubspecies introgression happened from time to time.

The sizes of present-day DOM segments incorporated in mice from the CAS territory are likely short, *i.e.*, the size of single genes or 10–20 kb. In the case of *Def8*, for example, the mice from Pakistan and northern India carrying the DOM alleles possess CAS alleles at the immediately adjacent gene loci. A parsimonious explanation for the shortened DOM segment is that introgression event(s) happened at some ancient time, followed by long-term backcrossing of the DOM segments against the CAS background genome. Haplotype breakdown arising through recombination in a given 10-kb stretch is expected to be seen with >50% probability after 14,000 generations (1–3 generations per year), assuming that the recombination rate is 0.52 cM/Mb (Jensen-Seaman *et al.*, 2004). These considerations suggest that substantial time has passed since the introgression occurred. Therefore, the gene introgression between the subspecies may have manipulated such gene (or gene segment)-specific incorporation of

exogenous genes in recipient subspecies genomes, facilitating the maintenance of genetic diversity.

The possible occurrence of an introgression event involving genes of DOM origin into the CAS background mice in the homeland area, the Indian subcontinent, is a clue to the enigmatic issue of the CAS segment harbouring DOM-related genes (*Def8* and *Afg311*) in the northern Japanese mouse genome. Given that ancient introgression events involving *Def8* are evident in the homeland area, the presence of a rather long DOM segment (20–30 kb, *i.e.* *Def8* plus *Afg311*) in the northern Japanese mouse population might be attributable to ancient introgression from DOM to CAS and to early isolation by dispersal to a place where conspecific populations were absent, followed by the acquisition of a homogeneous state for the exogenous segment of CAS and DOM in the MUS background mice via a founder effect.

The genetic structuring for the three CAS subgroups in the Indian subcontinent seen in the NN network using the concatenate sequences was not reproduced in the single-gene trees (Fig. 2). This implies that substantial gene flow occurred among the geographic groups during the course of evolution and contributed to generating the diversified haplotypes of the concatenate sequences of the CAS groups. The STRUCTURE analysis supports this view (Fig. 3B). Overall, one can reasonably presume that the evolutionary state of *M. musculus* is a typical example of reticulate evolution.

GENEALOGICAL ASSIGNMENT OF THE HUMAN-RELATED DISPERSAL IN ASIA

Although prehistoric human movements across Eurasia are known to have assisted the long-distance dispersal of house mice from its homeland (Boursot *et al.*, 1993; Prager *et al.*, 1998), the exact source areas are vague. Our previous mtDNA study showed that the eastward movements of CAS mice brought one of the four mtDNA subtypes, CAS-1, to a broad area of southern China, continental Southeast Asia, the Philippines, Indonesia, Japan, and Sakhalin (Suzuki *et al.*, 2013). However, the magnitude of the differences among the CAS-1 sequences from the eastern part of the Indian subcontinent is limited, perhaps due to overlaying of part of the homeland in the Indian subcontinent by the newly expanded mtDNA subtype; therefore, we are unable to identify a dispersal track from the mtDNA data. In this study, the majority of the nuclear haplotypes of the dispersed lineages (CAS-D) was found to be closely related to one another and shown to be nearly identical to the haplotype of the mouse from Pune, central India, while differing from the other haplotypes CAS-A, B, C, and E recovered from India (see Fig.

3). These results suggest that Southeast Asia, southern China, and Indonesia are representative areas to which *M. musculus* expanded efficiently, mediated by the historical human eastward movement, and that its source area is confined to a geographic locality, perhaps somewhere in India with the haplotypes belonging to CAS-D.

The CAS mice appear to have expanded their range to the northern tip of the Japanese islands, where the majority of the present-day genetic component is that of MUS. Accordingly, the 200-kb segments represent recombinant haplotypes between MUS and non-MUS, as seen in mice from northern Japan; this recombinant is

Fanca^{MUS}–*Spire2*^{CAS}–*Tcf25*^{CAS}–*Mc1r*^{CAS}–*Def8*^{DOM}–*Afg3l1*^{DOM}–*Dbn1d1*^{CAS} (Nunome *et al.*, 2010). If we consider that the CAS fragment is 200 kb in size, the recombinant segments comprise components of CAS and DOM and the time of the hybridisation is estimated to be around 700 generations ago (Nunome *et al.*, 2010). The non-MUS *Def8* and *Afg3l1* sequences are assigned as DOM, and the unique combination of CAS and DOM components is thought to have been generated in the source area, as mentioned above. However, no such combination of CAS and DOM has yet been found in the predicted source area, perhaps due to intensive backcrossing of the introgressed DOM segments with CAS mice.

The analysis of linked nuclear gene sequences is useful for phylogeographic studies for several reasons. First, a set of concatenated sequences is helpful for reconstructing a reliable population genetic structure without making any particular assumptions. Second, in a species that has experienced dispersal and secondary contact, the method used here can help to identify the source locality and contact areas. By setting the intervals of gene loci properly, one can infer the time span of hybridisation. Moreover, if a gene in the central part of the chromosome region used in the haplotype analyses has undergone adaptive evolution, we can test this possibility by measuring the reduced level of nucleotide diversity, as discussed below.

POSSIBLE INVOLVEMENT OF NATURAL SELECTION ACTING ON THE COAT COLOUR GENE

MC1R

The prominent features of the genetic diversity of *M. musculus* are that both the degree of nucleotide diversity and the phylogenetic patterns differ substantially among the genes examined, despite the fact that they are arranged tightly at approximately 20-kb intervals. This implies that *M. musculus* has experienced gene flow within and between geographic groups and the subsequent geographic allocation of specific alleles across

geographic barriers in a gene-specific manner. Concerning the ancient gene flow across the geographic groups in its homeland, identifying the factors shaping the geographic permeability of specific alleles is of interest. *Mclr* is one of the best-known genes promoting the evolution of pigmentation in birds and mammals (e.g. Klunghand *et al.*, 5 1995; MacDougall-Shackleton, Blanchard & Gibbs, 2003; Hoekstra & Price, 2004; Hoekstra, 2006; Hubbard *et al.*, 2010; Manceau *et al.* 2010; Suzuki, 2013). Our data include evidence for the inclusion of natural selection in shaping the patterns with respect to the coat colour gene *Mclr*.

Comparing the nucleotide diversity across the chromosome region, V- and 10 U-shaped graphs are detected in CAS and DOM, respectively (Fig. 2B; Table S1). This feature, known as selective sweep, is considered a footprint of positive selection (Maynard Smith & Haigh, 1974; Teschke, Büntge & Tautz, 2012) and the cases accompanied by gene flow across different taxa are known as selective gene flow or adaptive introgression (Martinsen *et al.*, 2001; Bull *et al.*, 2006; Anderson *et al.*, 2009; 15 Song *et al.*, 2011). The trend was confirmed in the adjacent upstream non-coding region, *Mclr-NC*, suggesting that the lower diversity of *Mclr* is not due to a comparison of a coding sequence with non-coding (e.g. intron) sequences. The V-shaped pattern in CAS is attributable to long-term processing of the polymorphism. The U-shaped pattern in DOM indicates a greater level of linkage disequilibrium affecting *Mclr* and the adjacent 20 region, indicating the relatively recent involvement of natural selection (Maynard Smith & Haigh, 1974). Accordingly, the values of Tajima's *D* were negative for *Mclr* in CAS and DOM, although not statistically significant (Table S1). These results support the idea that natural selection has acted on the *Mclr* gene or the adjacent region in the homeland of CAS and DOM during the course of evolution.

If natural selection is responsible for the specific population genetic patterns of 25 *Mclr* in CAS and DOM, certain *Mclr* allele(s) should involve a functional change, namely an amino acid replacement(s) altering coat colour. A candidate is allele 5 of the *Mclr*-coding that is shared by CAS (61%) and DOM (30%) (Fig. 5), whose state could be explained either by ancestral polymorphism or introgression. The allele shows two 30 diagnostic derived mutations at sites 51 (C to T) and 302 (C to T) (Fig. 5B), which are synonymous and non-synonymous mutations, respectively. Hence, the latter is a candidate as the responsible mutation, as it is predicted to replace the amino acid alanine with valine at codon position 101 (A101V). The site of the candidate mutation at codon 101 corresponds to the first extracellular loop (LC1), where the reverse agonist 35 of agouti signalling protein (ASIP) makes contact (Patel *et al.*, 2010) and is predicted to be important for the adaptive evolution of coat colour (Eizirik *et al.*, 2003). Considering coat colour (Marshall, 1998), mice occurring in Pakistan and western and central India have a straw- or golden-coloured pelage dorsally, and have been classified as specific

taxa, such as “*bactrianus*” and “*tyleri*”, while mice from adjacent northern (Nepal) and southern (south India) areas have dark pelage dorsally and are traditionally classified as “*homouulus*” and “*castaneus*”, respectively (Marshall, 1998). A future study focusing on the relationships between the *Mc1r* SNP and coat colour variation would illuminate the taxonomic issue of the traditional subspecies grouping.

CONCLUSIONS

Our results demonstrate that *M. musculus* has a long evolutionary history since the establishment of the geographic groups leading to the major subspecies CAS, DOM, and MUS. A deep phylogenetic subdivision within CAS is evident, involving two distinct phylogroups with predicted origins on the Indian subcontinent. Apparent signals of gene flow exist among the three subspecies, which enabled them to mediate allelic exchanges. Notably, natural selection appears to have been involved in the intensive geographic expansion of *Mc1r* alleles during the course of evolution. The wild population of house mice, with its extraordinary genetic diversity and geographically separated groups, is an ideal subject for studies on evolutionary biology, in which natural selection is one of the major concerns (Teschke *et al.*, 2012). The method we adopted in this study (*i.e.* the analysis of linked nuclear multilocus markers) has been proven useful for assessing population genetic structure and genetic interaction in areas of secondary contacts.

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REFERENCES

- Anderson TM, vonHoldt BM, Candille SI, Musiani M, Greco C, Stahler DR, Smith DW, Padhukasahasram B, Randi E, Leonard JA, Bustamante CD, Ostrander EA, Tang H, Wayne RK, Barsh GS. 2009.** Molecular and evolutionary history of melanism in North American gray wolves. *Science* **323**: 1339–1343.
- Bonhomme F, Catalan J, Britton-Davidian J, Chapman VM, Moriwaki K, Nevo E, Thaler L. 1984.** Biochemical diversity and evolution in the genus *Mus*. *Biochemical Genetics* **22**: 275–303.
- Bonhomme F, Rivals E, Orth A, Grant GR, Jeffreys AJ, Bois PR. 2007.** Species-wide distribution of highly polymorphic minisatellite markers suggests past and present genetic exchanges among house mouse subspecies. *Genome Biology* **8**: R80.
- Bonhomme F, Searle JB. 2012.** House mouse phylogeography. In: Macholán M, Baird SJE, Munclinger P, Piálek J, eds. *Evolution of the house mouse (Cambridge studies in morphology and molecules)*, Cambridge: Cambridge University Press, 278–296.
- Boursot P, Auffray JC, Britton-Davidian J, Bonhomme F. 1993.** The evolution of house mice. *Annual Review of Ecology and Systematics* **24**: 119–152.
- Boursot P, Din W, Anand R, Darviche D, Dod B, von Deimling F, Talwar GP, Bonhomme F. 1996.** Origin and radiation of the house mouse: mitochondrial DNA phylogeny. *Journal of Evolutionary Biology* **9**: 391–415.
- Bull V, Beltran M, Jiggins CD, McMillan WO, Bermingham E, Mallet J. 2006.** Polyphyly and gene flow between non-sibling *Heliconius* species. *BMC Biology* **4**: 11.
- Cucchi T, Auffray JC, Vigne JD. 2012.** On the origin of the house mouse synanthropy and dispersal in the Near East and Europe: zooarchaeological review and perspectives. In: Macholán M, Baird SJE, Munclinger P, Piálek J, eds. *Evolution of the house mouse (Cambridge studies in morphology and molecules)*, Cambridge: Cambridge University Press, 65–93.
- Din W, Anand R, Boursot P, Darviche D, Dod B, Jouvin-Marche E, Orth A, Talwar GP, Cazenave P-A, Bonhomme F. 1996.** Origin and radiation of the house mouse: clues from nuclear genes. *Journal of Evolutionary Biology* **9**: 519–539.
- Drummond AJ, Rambaut A, Shapiro B, Pybus OG. 2005.** Bayesian coalescent inference of past population dynamics from molecular sequences. *Molecular Biology and Evolution* **22**: 1185–1192.

- Drummond AJ, Suchard MA, Xie D, Rambaut A. 2012.** Bayesian phylogenetics with BEAUti and the BEAST 1.7. *Molecular Biology and Evolution* **29**: 1969–1973.
- Earl DA, vonHoldt BM. 2012.** STRUCTURE HARVESTER: a website and program for visualizing STRUCTURE output and implementing the Evanno method. *Conservation Genetics Resources* **4**: 359–361.
- 5 **Eizirik E, Yuhki N, Johnson WE, Menotti-Raymond M, Hannah SS, O'Brien SJ. 2003.** Molecular genetics and evolution of melanism in the cat family. *Current Biology* **13**: 448–453.
- Excoffier L, Lischer HEL. 2010.** Arlequin suite ver 3.5: a new series of programs to perform population genetics analyses under Linux and Windows. *Molecular Ecology Resources* **10**: 564–567.
- 10 **Gabriel SI, Mathias MDL, Searle JB. 2013.** Genetic structure of house mouse (*Mus musculus* Linnaeus 1758) populations in the Atlantic archipelago of the Azores: colonization and dispersal. *Biological Journal of the Linnean Society* **108**: 929–940.
- 15 **Hoekstra, H.E. 2006.** Genetics, development and evolution of adaptive pigmentation in vertebrates. *Heredity* **97**: 222–234.
- Hoekstra HE, Price T. 2004.** Parallel evolution is in the genes. *Science* **303**: 1779–1780.
- Huson DH, Bryant D. 2006.** Application of phylogenetic networks in evolutionary studies. *Molecular Biology and Evolution* **23**: 254–267.
- 20 **Hubbard JK, Uy JAC, Hauber ME, Hoekstra HE, Saffran RJ. 2010.** Vertebrate pigmentation: from underlying genes to adaptive function. *Trends in Genetics* **26**: 231–239.
- Jensen-Seaman MI, Furey TS, Payseur BA, Lu Y, Roskin KM, Chen C-F, Thomas MA, Haussler D, Jacob HJ. 2004.** Comparative recombination rates in the rat, mouse, and human genomes. *Genome Research* **14**: 528–538.
- 25 **Jing M, Yu H-T, Bi X, Lai Y-C, Jiang W, Huang L 2014.** Phylogeography of Chinese house mice (*Mus musculus musculus/castaneus*): distribution, routes of colonization and geographic regions of hybridization. *Molecular Ecology* **23**: 4387–4405.
- Jones EP, Eager HM, Gabriel SI, Jóhannesdóttir F, Searle JB. 2013.** Genetic tracking of mice and other bioproxies to infer human history. *Trends in Genetics* **29**: 298–308.
- 30 **Klungland H, Våge DI, Gomez-Raya L, Adalsteinsson S, Lien S. 1995.** The role of melanocyte-stimulating hormone (MSH) receptor in bovine coat color determination. *Mammalian Genome* **6**: 636–639.
- Koopman WJM, Li YH, Coart E, De Weg EV, Vosman B, Roldan-Ruiz I, Smulders MJM. 2007.** Linked vs. unlinked markers: multilocus microsatellite
- 35

- haplotype-sharing as a tool to estimate gene flow and introgression. *Molecular Ecology* **16**: 243–256.
- Lai YC, Shiroishi T, Moriwaki K, Motokawa M, Yu HT. 2008.** Variation of coat colour in house mice throughout Asia. *Journal of Zoology* **274**: 270–276.
- 5 **Librado P, Rozas J. 2009.** DnaSP v5: software for comprehensive analysis of DNA polymorphism data. *Bioinformatics* **25**: 1451–1452.
- Linnen CR, Kingsley EP, Jensen JD, Hoekstra HE. 2009.** On the origin and spread of an adaptive allele in deer mice. *Science* **325**: 1095–1098.
- MacDougall-Shackleton EA, Blanchard L, Gibbs HL. 2003.** Unmelanized plumage
10 patterns in old world leaf warblers do not correspond to sequence variation at melanocortin-1 receptor locus (*MC1R*). *Molecular Biology and Evolution* **20**: 1675–1681.
- Manceau, M., V.S Domingues, C.R. Linnen, E.B. Rosenblum and H.E. Hoekstra. 2010.** Convergence in pigmentation at multiple levels: mutations, genes and
15 function. *Philosophical Transactions of the Royal Society* **365**: 2439–2450.
- Martinsen GD, Whitham TG, Turek RJ, Keim P. 2001.** Hybrid populations selectively filter gene introgression between species. *Evolution* **55**: 1325–1335.
- Marshall JT. 1998.** Identification and scientific names of Eurasian house mice and their European allies, subgenus *Mus* (Rodentia: Muridae). Unpublished report, National
20 Museum of Natural History, Washington.
- Maynard Smith J, Haigh J. 1974.** The hitch-hiking effect of a favourable gene. *Genetical Research* **23**: 23–35.
- Mullen LM, Vignieri SN, Gore JA, Hoekstra HE. 2009.** Adaptive basis of geographic variation: genetic, phenotypic and environmental differences among beach mouse
25 populations. *Proceedings of the Royal Society B: Biological Sciences* **276**: 3809–3818.
- Nielsen R, Williamson S, Kim Y, Hubisz MJ, Clark AG, Bustamante C. 2005.** Genomic scans for selective sweeps using SNP data. *Genome Research* **15**: 1566–1575.
- Nunome M, Ishimori C, Aplin KP, Tsuchiya K, Yonekawa H, Moriwaki K, Suzuki
30 H. 2010.** Detection of recombinant haplotypes in wild mice (*Mus musculus*) provides new insights into the origin of Japanese mice. *Molecular Ecology* **19**: 2474–2489.
- Patel MP, Cribb Fabersunne CS, Yang YK, Kaelin CB, Barsh GS, Millhauser GL. 2010.** Loop-swapped chimeras of the agouti-related protein and the agouti signaling

protein identify contacts required for melanocortin 1 receptor selectivity and antagonism. *Journal of Molecular Biology* **404**: 45–55.

- 5 **Prager EM, Orrego C, Sage RD. 1998.** Genetic variation and phylogeography of Central Asian and other house mice, including a major new mitochondrial lineage in Yemen. *Genetics* **150**: 835–861.
- Pritchard JK, Stephens M, Donnelly P. 2000.** Inference of population structure using multilocus genotype data. *Genetics* **155**: 945–959.
- 10 **Rajabi-Maham H, Orth A, Siahsarvie R, Boursot P, Darvish J, Bonhomme F. 2012.** The south-eastern house mouse *Mus musculus castaneus* (Rodentia: Muridae) is a polytypic subspecies. *Biological Journal of the Linnean Society* **107**: 295–306.
- Rambaut A, Drummond AJ. 2009.** Tracer v1.5. Available from <http://beast.bio.ed.ac.uk/Tracer>.
- 15 **Shimada T, Sato JJ, Aplin KP, Suzuki H. 2009.** Comparative analysis of evolutionary modes in *Mc1r* coat color gene in wild mice and mustelids. *Genes & Genetic Systems* **84**: 225–231.
- Song Y, Endepols S, Klemann N, Richter D, Matuschka FR, Shih CH, Nachman M, Kohn MH. 2011.** Adaptive introgression of anticoagulant rodent poison resistance by hybridization between Old World Mice. *Current Biology* **21**: 1296–1301.
- 20 **Stephens JC, Reich DE, Goldstein DB, Shin HD, Smith MW, Carrington M, Winkler C, Huttley GA, Allikmets R, Schriml L, Gerrard B, Malasky M, Ramos MD, Morlot S, Tzetis M, Oddoux C, di Giovine FS, Nasioulas G, Chandler D, Aseev M, Hanson M, Kalaydjieva L, Glavac D, Gasparini P, Kanavakis E, Claustres M, Kambouris M, Ostrer H, Duff G, Baranov V, Sibul H, Metspalu A, Goldman D, Martin N, Duffy D, Schmidtke J, Estivill X, O'Brien SJ, Dean M.**
- 25 **1998.** Dating the origin of the CCR5-Delta32 AIDS-resistance allele by the coalescence of haplotypes. *American Journal of Human Genetics* **62**: 1507–1515.
- Stephens M, Smith NJ, Donnelly P. 2001.** A New Statistical Method for Haplotype Reconstruction from population Data. *American Journal of Human Genetics* **68**: 978–989.
- 30 **Stephens M, Scheet P. 2005.** Accounting for decay of linkage disequilibrium in haplotype inference and missing-data imputation. *American Journal of Human Genetics* **76**: 449–462.
- 35 **Suzuki H, Shimada T, Terashima M, Tsuchiya K, Aplin K. 2004.** Temporal, spatial, and ecological modes of evolution of Eurasian *Mus* based on mitochondrial and nuclear gene sequences. *Molecular Phylogenetics and Evolution* **33**: 626–646.

- Suzuki H, Aplin KP. 2012.** Phylogeny and biogeography of the genus *Mus* in Eurasia. In: Macholán M, Baird SJE, Munclinger P, Piálek J, eds. *Evolution of the house mouse (Cambridge studies in morphology and molecules)*, Cambridge: Cambridge University Press, 35–64.
- 5 **Suzuki H, Nunome M, Kinoshita G, Aplin KP, Vogel P, Kryukov AP, Jin ML, Han SH, Maryanto I, Tsuchiya K, Ikeda H, Shiroishi T, Yonekawa H, Moriwaki K. 2013.** Evolutionary and dispersal history of Eurasian house mice *Mus musculus* clarified by more extensive geographic sampling of mitochondrial DNA. *Heredity* **111**: 375–390.
- 10 **Suzuki H. 2013.** Evolutionary and phylogeographic views on *Mclr* and *Asip* variation in mammals. *Genes & Genetic Systems* **88**: 155–164.
- Takada T, Ebata T, Noguchi H, Keane TM, Adams DJ, Narita T, Shin IT, Fujisawa H, Toyoda A, Abe K, Obata Y, Sakaki Y, Moriwaki K, Fujiyama A, Kohara Y, Shiroishi T. 2013.** The ancestor of extant Japanese fancy mice contributed to the mosaic genomes of classical inbred strains. *Genome Research* **23**: 1329–1338.
- 15 **Tamura K, Peterson D, Peterson N, Stecher G, Nei M, Kumar S. 2011.** MEGA5: molecular evolutionary genetics analysis using maximum likelihood, evolutionary distance, and maximum parsimony methods. *Molecular Biology and Evolution* **28**: 2731–2739.
- 20 **Teschke M, Büntge A, Tautz D. 2012.** Tracing recent adaptations in natural populations of the house mouse. In: Macholán M, Baird SJE, Munclinger P, Piálek J, eds. *Evolution of the house mouse (Cambridge studies in morphology and molecules)*, Cambridge: Cambridge University Press, 335–333.
- 25 **Terashima M, Furusawa S, Hanzawa N, Tsuchiya K, Suyanto A, Moriwaki K, Yonekawa H, Suzuki H. 2006.** Phylogeographic origin of Hokkaido house mice (*Mus musculus*) as indicated by genetic markers with maternal, paternal and biparental inheritance. *Heredity* **96**: 128–138.
- Vignieri SN, Larson JG, Hoekstra HE. 2010.** The selective advantage of crypsis in mice. *Evolution* **64**: 2153–2158.
- 30 **Yonekawa H, Tsuda K, Tsuchiya K, Yakimenko L, Korobitsyna K, Chelomina GN Spiridonova L, Frisman L, Kryukov AP, Moriwaki K. 2003.** Genetic diversity, geographic distribution and evolutionary relationships of *Mus musculus* subspecies based on polymorphisms of mitochondrial DNA. In: Kryukov A, Yakimenko L, eds. *Problems of evolution*, Vladivostok: Dalnauka, Vol 5, 90–108.
- 35

Figure legends

Figure 1. The localities where the wild mice used in this study were collected together with the specimen codes for those previously analysed (1–59; see Nunome *et al.*, 2010 for further details) and those newly determined (60–98). The detailed locality names and sample codes are listed in Table 1. Localities with SNP C→T at site 302 in the *Mclr* coding region are represented by bold circles. Rough geographic ranges for the three major subspecies (CAS: *M. m. castaneus*, DOM: *M. m. domesticus*, and MUS: *M. m. musculus*) and the two previously predicted MUS groups (Nunome *et al.*, 2010) are shown.

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Figure 2. Genetic variation of the seven gene markers. A, Neighbour-Net networks derived from single gene sequence data sets for *Fanca*, *Spire2*, *Tcf25*, *Mclr*, *Def8*, *Afg3l1*, and *Dnadd1* in *Mus musculus* and five species as the outgroup: *M. caroli* (1), *M. terricolor* (2), *M. spretus* (3), *M. spicilegus* (4), and *M. macedonicus* (5). The subspecies groups *M. m. domesticus* (DOM), *M. m. castaneus* (CAS), and *M. m. musculus* (MUS) were assessed by referring to previously obtained data (Nunome *et al.*, 2010). The newly determined allele sequences originating from India and Pakistan were assigned as CAS, although several *Def8* sequences were integrated into the cluster assigned as DOM. The positions of non-MUS sequences from northern Japan are marked with arrowheads. Genetic variation of the six gene markers. B, comparison of the nucleotide diversities (%) using sequences assigned for the three subspecies groups. The non-coding region (*Mclr-NC*) immediately upstream of the *Mclr* (melanocortin-1 receptor gene) coding region was included.

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Figure 3. Assessment of population genetic structure using the dataset of the seven nuclear genes. A, Neighbour-Net network with the concatenated sequences from 98 *Mus musculus* with subspecies groups *M. m. domesticus* (DOM), *M. m. castaneus* (CAS), and *M. m. musculus* (MUS) and five outgroup species. Seven haplogroups seen in the CAS cluster (CAS- A–E) are tentatively indicated. Those apparently recognised as recombinant haplotypes (Re- 1~ 8: Nunome *et al.*, 2010; Re-9~11: this study) are indicated by subspecies components (C, *castaneus*; D, *domesticus*; M, *musculus*). B, Population genetic structure was assessed using STRUCTURE, version 2.3. Genetic component proportions of individuals are represented by thin horizontal bars composed of four clusters. C, Distribution of the seven CAS haplogroups tentatively observed in

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the network analysis. Representative haplotypes related to the seven gene markers are schematically shown, exhibiting DOM components embedded in CAS chromosome segments in wild mice from northern Japan. Mitochondrial DNA types are shown (Suzuki *et al.*, 2013), emphasizing the CAS subtypes (CAS 1-4).

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Figure 4. Phylogenetic tree of *Mus musculus* constructed using the Bayesian method with concatenate sequences (seven genes; 3402 bp). Statistical support is indicated by Bayesian posterior probabilities. The times of the most common recent ancestors and 95% highest posterior density estimated by the Bayesian analysis are shown for each major calibration point. The numbers below the lines at the nodes are the Bayesian posterior probabilities. See Figure 3 for the four CAS haplogroups.

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Figure 5. A, Median Joining (MJ) networks constructed with the entire coding region of the melanocortin-1 receptor gene (*Mclr*, 948 bp; right) and the immediate upstream non-coding region (*Mclr-NC*, 377 bp; left), showing the subspecies groups *M. m. domesticus* (DOM), *M. m. castaneus* (CAS), and *M. m. musculus* (MUS) based on geographic origin. Each circle represents an allele and its size reflects the allele frequency. Numbers along branches and adjacent to alleles indicate nucleotide positions and codes of alleles (alleles 1- 5) of interest. Allele 1 is predicted to be the prototype from the comparison with outgroup sequences. Allele B, Schematic representation of the gene structure of *Mclr*. Presumed derived mutations of nucleotides and amino acids, differing from the predicted original haplotype (hap 1), are shown in haplotypes representing the subspecies groups (hap 2-6).

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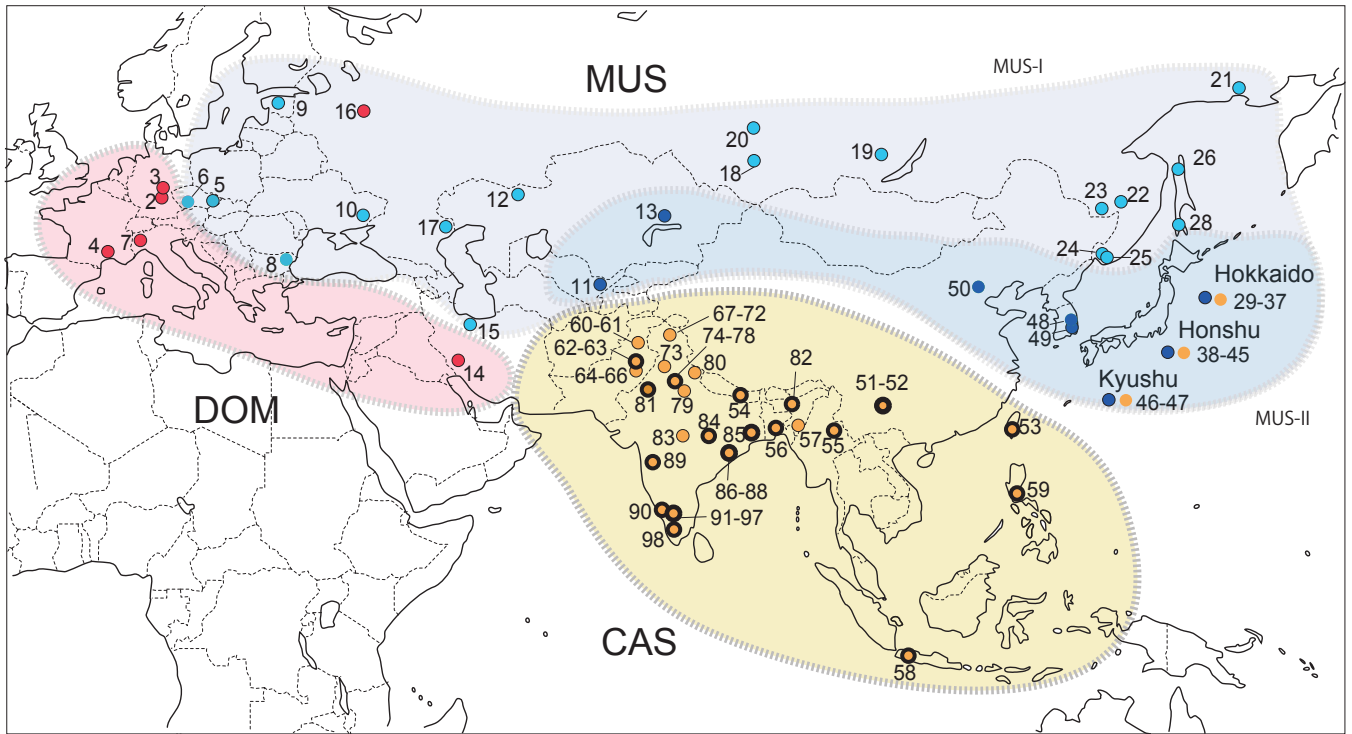
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SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article at the publisher's website:

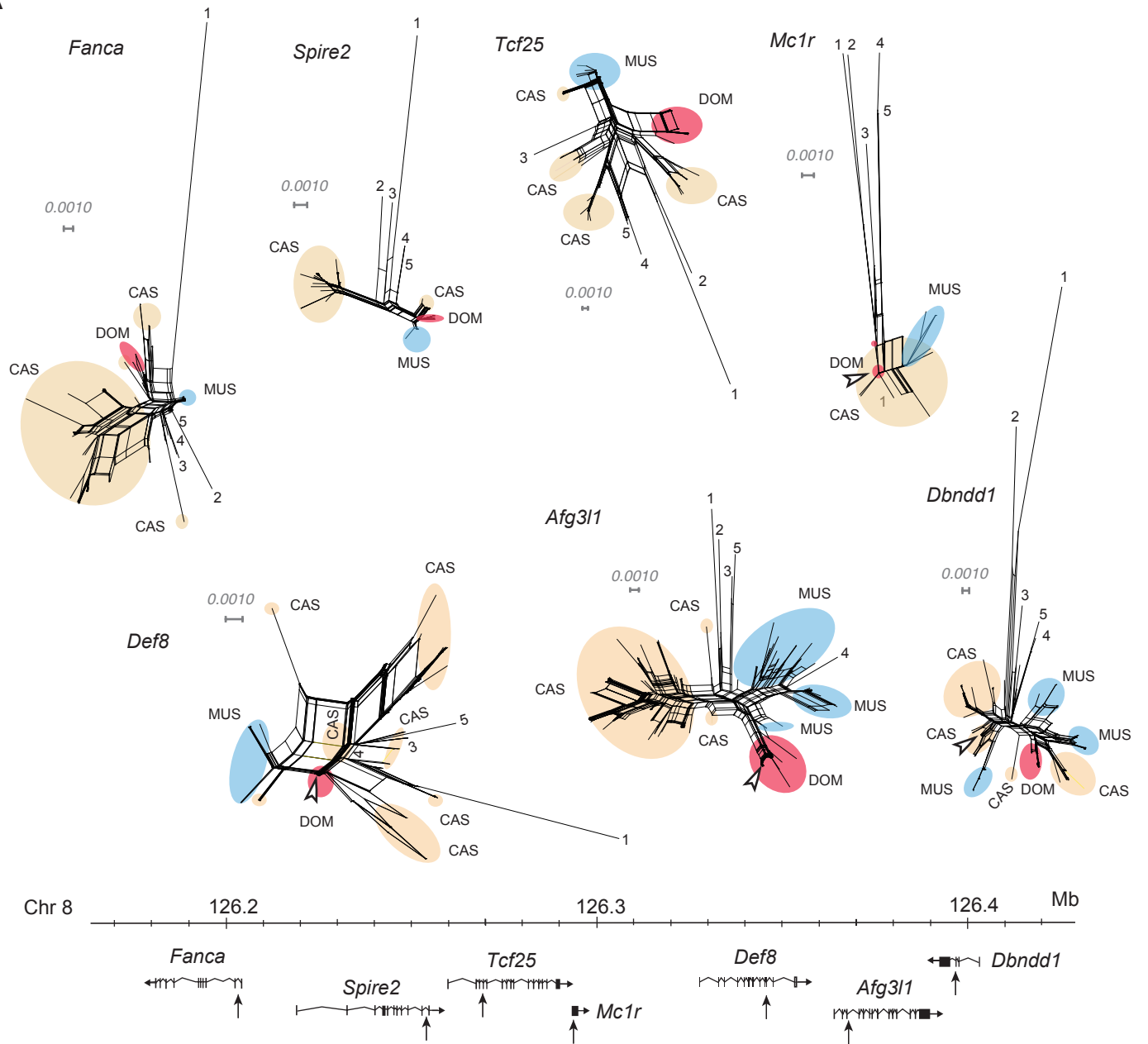
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Table S1. Genetic diversity indices of the eight nuclear gene markers. The number of segregating sites (S), mean number of pairwise differences (π), standard error (SE), and Tajima's *D* were calculated using ARLEQUIN 3.1(Excoffier & Lischer, 2010).

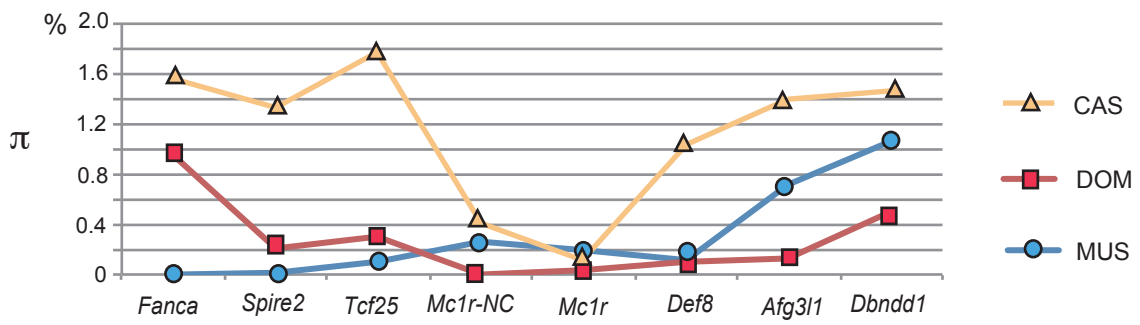


Kodama et al. Fig. 1

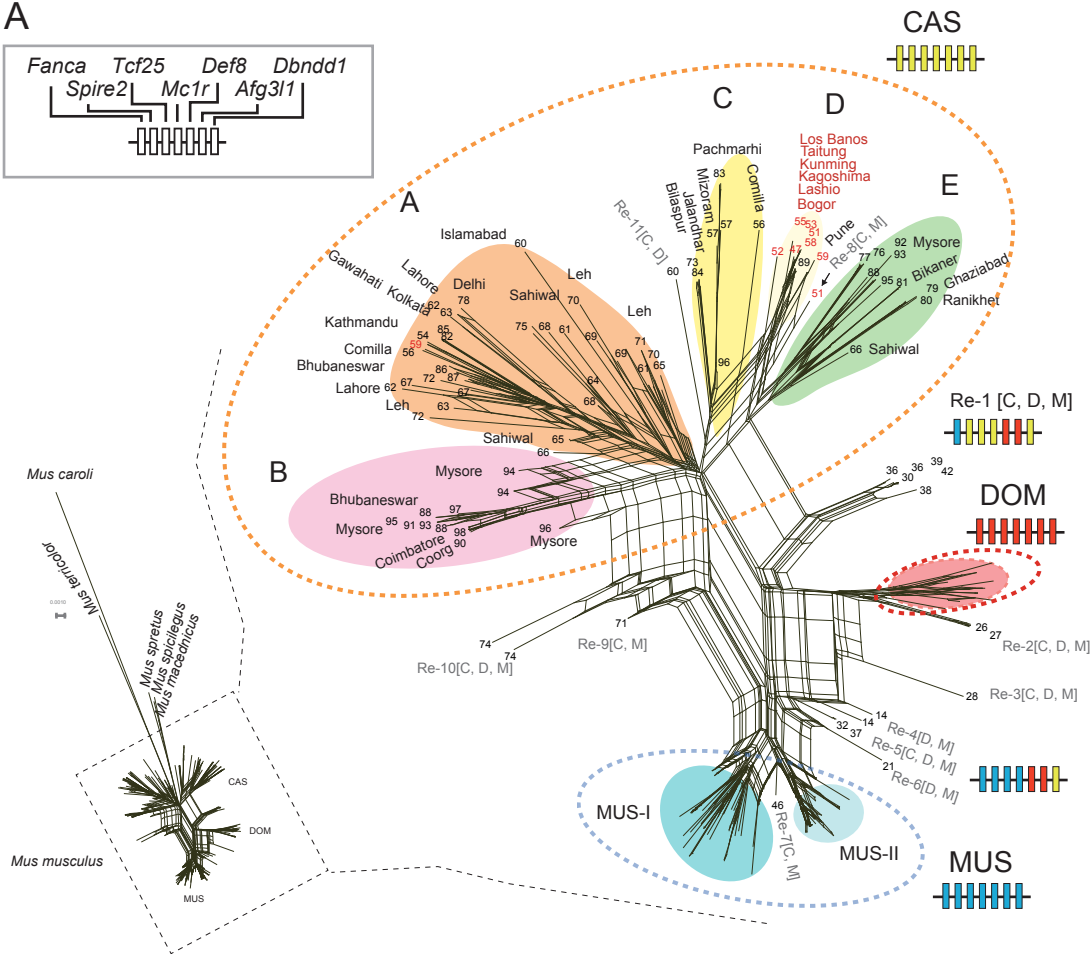
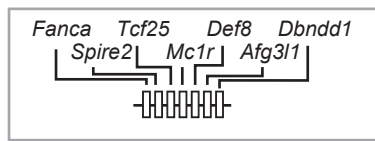
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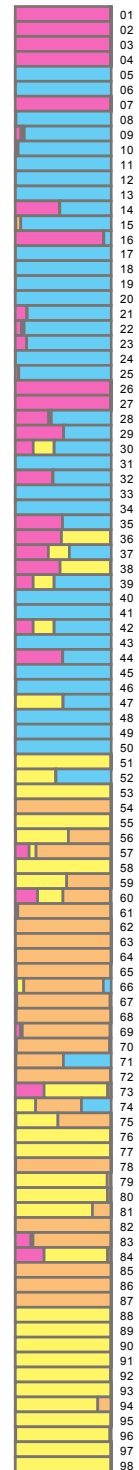
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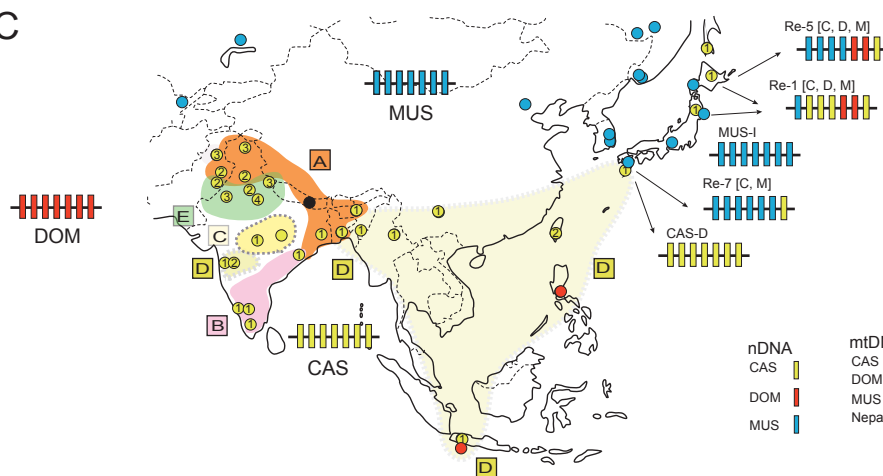
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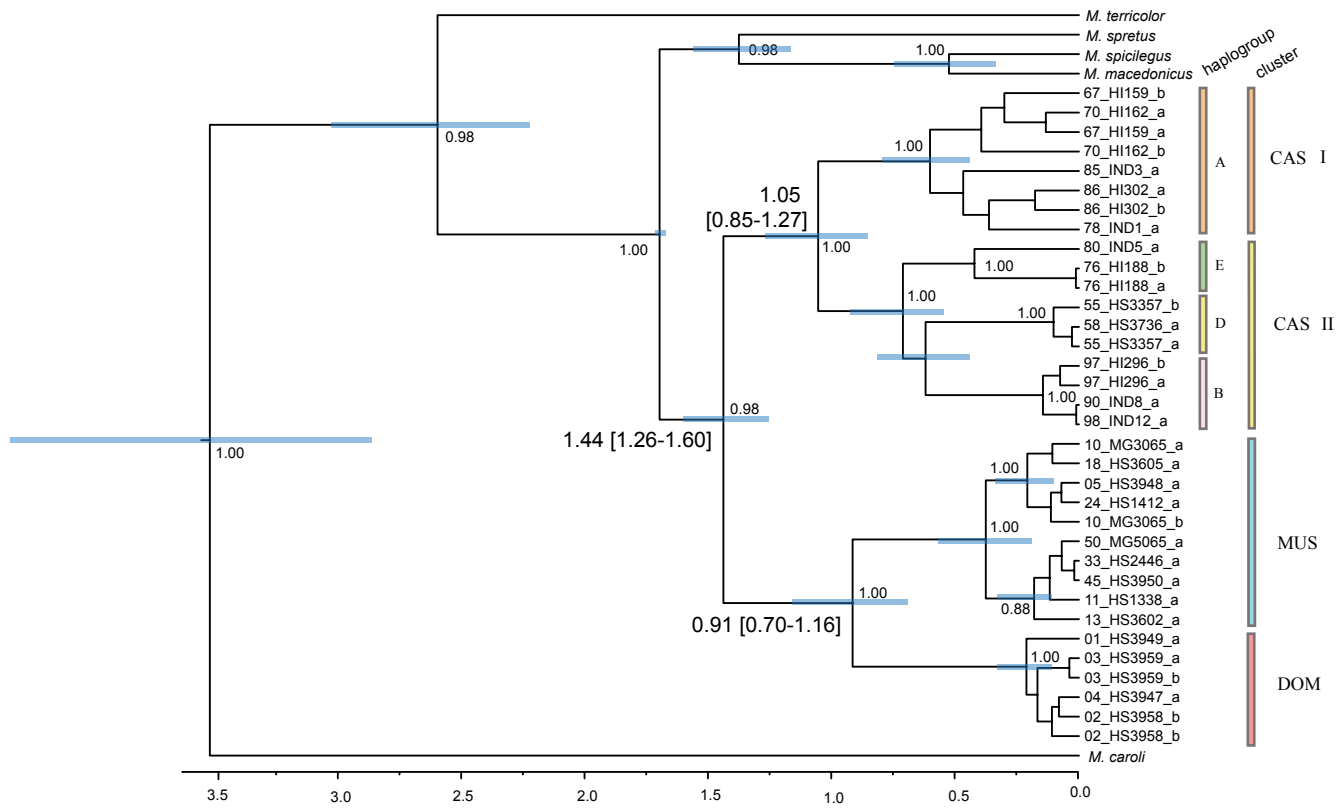


B



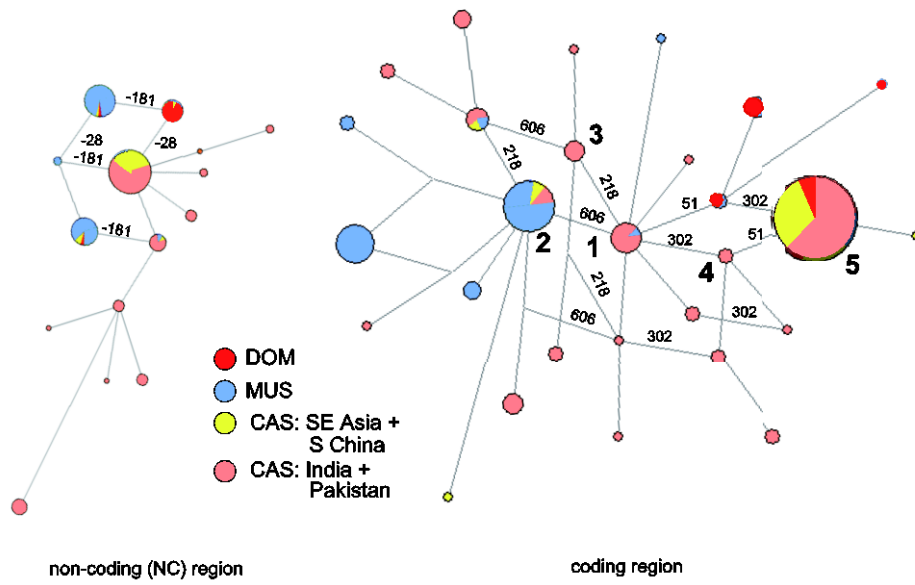
C



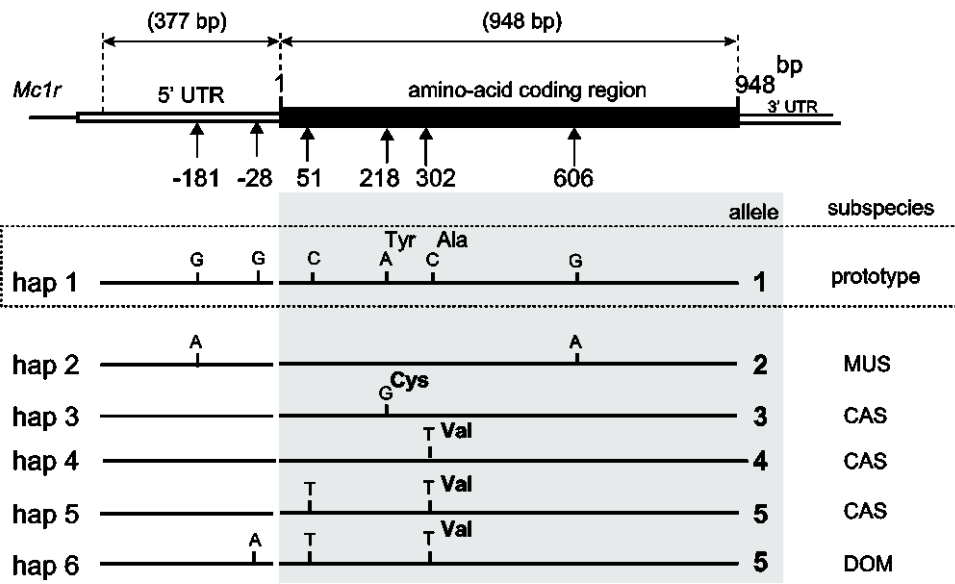


Kodama et al. Fig. 4

A



B



Kodama et al. Fig. 5

Table 1
List of specimens used in this study and phylogroup assignment

Specimen Code	Locality	Haplogroup	Specimen Code	Locality	Haplogroup
1 PGN2:HS3949	Canada: Pegion	DOM	51 HS506	China: Kunming	CAS-D ^a , Re-8[C, M] ^a
2 MB8/HS3958	Germany: Weidesgrun	DOM	52 HS507	Kunming	CAS-D ^a , MUS-II
3 MB9/HS3959	Kubelhof	DOM	53 HS2400	Taiwan: Taitung	CAS-D ^a
4 BFM/2:HS3947	France: Montpellier	DOM	54 HS1467	Nepal: Kathmandu	CAS-A ^a , CAS-A ^b
5 PWK:HS3948	Czech Rep.: Lhotka	MUS-I	55 HS3357	Myanmar: Lashio	CAS-D ^a
6 BLG2:HS3951	Blugaria: Toshevo	MUS-I	56 HS3701	Bangladesh: Comilla	CAS-A ^a , CAS-D ^a
7 HS589	Italy: Aosta	DOM	57 HS3441	India: Mizoram	CAS-C
8 HZ788	Czech Rep.: Liborezy	MUS-I	58 HS3736	Indonesia: Bogor	CAS-D ^a
9 MG3044	Estonia: Tallinn	MUS-I	59 HS3882	Philippines: Los Banos	CAS-A ^c , CAS-D ^a
10 MG3065	Ukraine: Donetsk	MUS-I, MUS-II	60 HI178	Pakistan: Islamabad	CAS-A ^c , Re-11[C, D]
11 HS1338	Uzbekistan: Tashkent	MUS-II	61 HI179	Islamabad	CAS-A ^a
12 HS1464	Kazakhstan: Aktubinsk	MUS-I	62 HI266	Lahore	CAS-A ^{a, b, d} , CAS-A ^{b, c}
13 HS3602	Balkhash	MUS-II	63 HI267	Lahore	CAS-A ^c , CAS-A ^{b, c}
14 NIG934	Iran: Ahvaz	Re-4[D, M]	64 HI260	Sahiwal	CAS-A
15 NIG935	Nowshahr	MUS-I	65 HI261	Sahiwal	CAS-A
16 MG3055	Russia: Moscow	DOM ^a	66 HI262	Sahiwal	CAS-A ^d , CAS-E
17 HS3612	Astrakhan	MUS-I	67 HI159	India: Leh	CAS-A ^d
18 HS3605	Gorno-Altaysk	MUS-I	68 HI160	Leh	CAS-A, CAS-A ^d
19 HS3608	Irkutsk	MUS-I	69 HI161	Leh	CAS-A
20 HS3604	Tomsk	MUS-I	70 HI162	Leh	CAS-A, CAS-A ^d
21 HS1845	Magadan	MUS-I, Re-6[D, M]	71 HI163	Leh	CAS-A, Re-9[C, M]
22 HS1461	Khabarovsk	MUS-I, MUS-I	72 HI164	Leh	CAS-A ^d
23 HS1466	Amurskii	MUS-I	73 BRC3024/Ind10	Jalandhar	CAS-C ^c
24 HS1412	Kraskino	MUS-I	74 HI184	Delhi	Re-10[C, D, M] ^c
25 HS1335	Khasan	MUS-I	75 HI187	Delhi	CAS-A ^c
26 HS3606	Okha	Re-2[C, D, M]	76 HI188	Delhi	CAS-E ^a
27 HS3845	Okha	Re-2[C, D, M]	77 HI189	Delhi	CAS-E ^a
28 HS945	Tomari	MUS-I, Re-3[C, D, M]	78 BRC3015/Ind1	Delhi	CAS-A ^d
29 HS1946	Japan: Kushiro	DOM, MUS-II	79 BRC3025/Ind11	Ghaziabad	CAS-E
30 HS1947	Asahikawa	MUS-II, Re-1[C, D, M]	80 BRC3019/Ind5	Ranikhet	CAS-E
31 HS2781	Obihiro	MUS-II	81 BRC3021/Ind7	Bikaner	CAS-E ^a
32 HS2340	Otaru	Re-5[C, D, M]	82 BRC3023/Ind9	Gawahati	CAS-A ^a
33 HS2446	Naganuma	MUS-II	83 BRC3018/Ind4	Pachmarhi	CAS-C ^d
34 HS2451	Date	MUS-II	84 BRC3020/Ind6	Bilaspur	CAS-C ^a
35 HS2779	Kyowa	DOM, MUS-II	85 BRC3017/Ind3	Kolkata	CAS-A ^a
36 HS3834	Rankoshi	Re-1[C, D, M]	86 HI302	Bhubaneswar	CAS-A ^a , CAS-A ^d
37 HS394	Onuma	MUS-II, Re-5[C, D, M]	87 HI303	Bhubaneswar	CAS-A ^a
38 HS2461	Sakekawa	Re-1[C, D, M]	88 HI305	Bhubaneswar	CAS-E ^a , CAS-B ^a
39 HS2458	Tendo	MUS-II, Re-1[C, D, M]	89 BRC3016/Ind2	Pune	CAS-D ^a
40 HS2454	Otsuti	MUS-II	90 BRC3022/Ind8	Coorg	CAS-B ^a
41 HS2456	Sendai	MUS-II	91 HI273	Mysore	CAS-B ^a
42 HS2457	Sakata	MUS-II, Re-1[C, D, M]	92 HI274	Mysore	CAS-E ^a
43 HS2468	Chiba	MUS-II	93 HI275	Mysore	CAS-E ^a , CAS-B ^a
44 HS3839	Atsugi	DOM, MUS-II	94 HI280	Mysore	CAS-B ^a
45 MSM:HS3950	Mishima	MUS-II	95 HI281	Mysore	CAS-E ^a , CAS-B ^a
46 HS2472	Miyazaki	MUS-II, Re-7[C, M]	96 HI295	Mysore	CAS-C ^d , CAS-B ^a
47 HS3603	Kagoshima	CAS-D, MUS-II	97 HI296	Mysore	CAS-B ^a
48 HS1368	Korea: Gyeongju	MUS-II	98 BRC3026/Ind12	Coimbatore	CAS-B ^a
49 HS3540	Busan	MUS-II			
50 MG5065	China: Beijing	MUS-II			

Individuals with codes 1- 59 were those analyzed in previous study (Nunome *et al.*, 2010). These DNA samples are stored in National Institute of Genetics (NIG) and the RIKEN Bio-Resource Center (BRC), and personally by HS at Hokkaido University (HS; HI, originally collected by H. Ikeda). Phylogroups are taken from Fig. 3 and recombinant haplogroups are indicated with its subspecies component (C, *castaneus*; D, *domesticus*; M, *musculus*).
^aHaplotypes containing the *Mc1r* alleles with the 302T (a) and 218G (b) mutations are indicated in mice from the CAS territory (codes 51-98).

^cHaplotype from India and Pakistan, incorporating the *Def8* alleles assigned to DOM.

Table S1

Genetic diversity indices of the eight nuclear gene markers. The number of segregating sites (S), mean number of pairwise difference (π), its standard error, and Tajima's D were calculated using ARLEQUIN 3.1 (Excoffier and Lischer, 2010).

Subspecies	<i>Fanca</i>	<i>Spire2</i>	<i>Tcf25</i>	<i>Mc1r upst.</i>	<i>Mc1r</i>	<i>Def8</i>	<i>Afg3l1</i>	<i>Dbn1d1</i>
CAS								
n	85	91	73	92	91	89	87	76
S	25	21	36	10	5	14	20	23
π	0.015624	0.013251	0.017766	0.004099	0.001137	0.010405	0.013933	0.014696
SE	+/- 0.008213	+/- 0.007089	+/- 0.009058	+/- 0.002743	+/- 0.001098	+/- 0.005920	+/- 0.001098	+/- 0.007645
Tajima's D	-0.09297	0.85586	1.30983	-0.60743	-1.10120	-0.09957	0.90854	0.29922
DOM								
n	15	15	15	14	15	15	15	15
S	8	2	5	0	1	1	4	10
π	0.009418	0.002058	0.003016	0	0.000314	0.001008	0.001272	0.005022
SE	+/- 0.00555	+/- 0.0017	+/- 0.002078	0	+/- 0.000554	+/- 0.00118	+/- 0.00117	+/- 0.003156
Tajima's D	0.65091	1.19844	0.4709	0	-1.15945	0.23502	-1.51811*	-0.38804
MUS								
n	65	66	66	54	66	66	66	66
S	0	2	7	2	6	5	16	16
π	0	0.000139	0.00101	0.002658	0.001926	0.001146	0.007099	0.010664
SE	0	+/- 0.000338	+/- 0.000887	+/- 0.002016	+/- 0.001926	+/- 0.001208	+/- 0.004042	+/- 0.005721
Tajima's D	0	-1.43204*	-1.4308	2.25873	-0.61065	-1.44214*	0.22811	2.20778

n: the number of sequences examined (n). The number of segregating sites (S), mean number of pairwise difference (π), its standard error, and Tajima's D

* $P < 0.05$.