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2	Evolutionary and dispersal history of Eurasian house mice
3	Mus musculus clarified by more extensive geographic
4	sampling of mitochondrial DNA
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### 1 Abstract

2	We examined sequence variation of mitochondrial DNA control region and cytochrome
3	b gene of the house mouse (Mus musculus sensu lato) drawn from ca. 200 localities,
4	with 286 new samples drawn primarily from previously unsampled portions of their
<b>5</b>	Eurasian distribution and with the objective of further clarifying evolutionary episodes
6	of this species before and after the onset of human-mediated long-distance dispersals.
7	Phylogenetic analysis of the expanded data detected five equally distinct clades, with
8	geographic ranges of northern Eurasia (musculus, MUS), India and Southeast Asia
9	(castaneus, CAS), Nepal (unspecified, NEP), western Europe (domesticus, DOM), and
10	Yemen (gentilulus). Our results confirm previous suggestions of Southwestern Asia as
11	the likely place of origin of <i>M. musculus</i> and the region of Iran, Afghanistan, Pakistan,
12	and northern India, specifically as the ancestral homeland of CAS. The divergence of
13	the subspecies lineages and of internal sublineage differentiation within CAS were
14	estimated to be 0.37-0.47 and 0.14-0.23 million years ago (mya), respectively,
15	assuming a split of <i>M. musculus</i> and <i>Mus spretus</i> at 1.7 mya. Of four CAS sublineages
16	detected, only one extends to eastern parts of India, Southeast Asia, Indonesia,
17	Philippines, South China, Northeast China, Primorye, Sakhalin and Japan, implying a
18	dramatic range expansion of CAS out of its homeland during an evolutionary short
19	time, perhaps associated with the spread of agricultural practices. Multiple and
20	non-coincident eastward dispersal events of MUS sublineages to distant geographic
21	areas, such as northern China, Russia, and Korea, are inferred, with the possibility of
22	several different routes.

- 1 **Key words:** mitochondrial DNA; cytochrome *b*; control region; phylogeography; wild
- 2 house mouse
- 3
- 4 Running head: mtDNA variation in Eurasian House mice
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# 1 Introduction

2	Despite the rapid rise of polygenic and genomic approaches to the analysis of
3	population history (e.g. Abe et al., 2004; Stoneking and Delfin, 2010; Yang et al., 2011),
4	the study of mitochondrial DNA (mtDNA) continues to play a significant role in the
5	investigation of many species. In the case of the house mouse complex (Mus musculus
6	Complex), the availability of large numbers of mtDNA sequences derived from
7	European and other populations has facilitated detailed analysis of both prehistoric and
8	historic range expansions (Rajabi-Maham et al., 2008; Gabriel et al., 2010, 2011;
9	Bonhomme et al., 2011; Jones et al., 2010), often with significant implications for
10	human history. By contrast, the other major lineages of the house mouse are known
11	from far fewer sequences and this has hindered progress on even some of the most
12	basic questions of phylogeography, such as their likely places of origin and the timing
13	and routes of major dispersal episodes.
14	Early investigations of house mouse mtDNA, using the method of Restriction
15	Fragment Length Polymorphism (RFLP; e.g. Yonekawa et al., 1981, 1986), identified
16	three major haplogroups among wild house mice. These appeared to be associated with
17	recognized subspecies and were designated accordingly: a DOM haplogroup in <i>M. m.</i>
18	domesticus from western Europe and North Africa (also southern Africa, Australia and
19	the Americas, all as historical introductions); a MUS haplogroup in M. m. musculus
20	from the northern part of Eurasia excluding western Europe; and a CAS haplogroup in
21	M. m. castaneus from Southeast Asia. Later studies of mtDNA suggested a number of
22	other possible divergent lineages: a BAC haplogroup in M. m. bactrianus from
23	Afghanistan and Pakistan (Boursot et al., 1993, 1996; Yonekawa et al., 1994); a GEN

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1	haplogroup in M. m. gentilulus from Yemen (Prager et al., 1998) and Madagascar
2	(Duplantier et al., 2002); and most recently, another divergent but as yet unnamed
3	haplogroup from Nepal (Terashima et al., 2006). Broader genomic comparisons using
4	microsatellites (Sakai et al., 2005), single nucleotide polymorphic sites (Abe et al.,
5	2004), and whole-genome sequences (Frazer et al., 2007) support the notion that each
6	of the MUS, CAS and DOM mtDNA haplogroups represents a longstanding
7	evolutionary lineage. However, the remaining mtDNA haplogroups (BAC and GEN)
8	have not been subject to the same level of scrutiny, hence their status remains
9	uncertain.
10	In this paper we fill a number of the remaining gaps in geographic mtDNA
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11 12 13 14	coverage for the house mouse, with a particular emphasis on the Indian sub-continent, China, and far eastern Russia. Addition of mtDNA sequences from these key areas sheds light on several issues, including 1) the likely ancestral range of each of the major evolutionary lineages; and 2) the direction and timing of range expansions, with
11 12 13 14 15	coverage for the house mouse, with a particular emphasis on the Indian sub-continent, China, and far eastern Russia. Addition of mtDNA sequences from these key areas sheds light on several issues, including 1) the likely ancestral range of each of the major evolutionary lineages; and 2) the direction and timing of range expansions, with a particular focus on East Asia, China and Japan, where multiple mtDNA lineages are

# 19 Materials and methods

20 Materials

Our new sequencing effort is based chiefly on samples of House mouse genomic DNA
stored in the National Institute of Genetics, Mishima, Japan. These were collected in
China, India, Russia, and a variety of other countries, on expeditions organized by KM

1	during 1983-2003 (MG series, stored in the National Institute of Genetics; and BRC
2	Series, stored in the RIKEN Bio-Resource Center), and by HI and KT during
3	1989-1992 (HI series, stored in Hokkaido University). We also used DNA samples
4	stored at Hokkaido University (HS series), including mice collected by PV (IZEA
5	series, vouchered in the Institut de Zoologie et d'Ecologie Animale of Lausanne
6	University) and KA (ANWC series, vouchered in the Australian National Wildlife
7	Collection). Some of the same samples have been used in previous studies (e.g.
8	Yonekawa et al., 1988, 1994, 2003; Miyashita et al., 1994; Nagamine et al., 1994;
9	Tsuchiya et al., 1994; Munclinger et al., 2002; Spiridonova et al., 2004).
10	New sequences were generated for mtDNA control region (CR) from 212
11	House mouse individuals from 137 localities; the cytochrome $b$ gene (Cytb) was also
12	sequenced from a subset of 167 individuals from 106 localities (Table S1). In our
13	sampling we strived to achieve maximum geographic coverage, at the cost of small
14	samples sizes (frequently just one) for each locality. While this reduced the scope for
15	sophisticated analysis of population expansion scenarios, it increased the likelihood of
16	detecting previously undiscovered components of mtDNA diversity. The geographic
17	distribution of the new sequences is shown in Figure 1.
18	We downloaded a further 571 CR sequences and 41 Cytb sequences of M.
19	musculus from public databases, drawn primarily from the work of Prager et al. (1996,
20	1998), Gündüz et al. (2005), Rajabi-Maham et al. (2008) and Bonhomme et al. (2011),
21	along with representative sequences of closely related species for use as outgroups. Our
22	sequence alignments for each mtDNA region are provided in Appendix A and B.
23	

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# 1 Sequence analyses

2	The PCR and direct sequencing of the CR (around 800-bp; Yasuda et al.,
3	2005) and Cytb (1140 bp; Suzuki et al., 2004) were performed according to previously
4	described methods. Two primers were used for sequence determination of CR in M.
<b>5</b>	musculus; CR1: 5'-CATGCCTTGACGGCTATGTT-3' and CR2:
6	5'-ATCGCCCATACGTTCCCCTT-3'. The double-stranded PCR product was
7	sequenced utilizing the PRISM Ready Reaction DyeDeoxy Terminator Cycle
8	Sequencing Kit (ABI) and an ABI3130 automated sequencer. Sequences of M.
9	cypriacus, M. macedonicus, M. spicilegus, and M. spretus were obtained from the
10	databases and used as outgroups in phylogenetic inference.
11	
12	Phylogeny and divergence time estimation
13	Sequences were aligned by eye using MEGA5 (Tamura et al., 2011). Prior to further
14	analyses, we deleted tandem repeat sequences of 75-76 bp in CR of some MUS and
15	some CAS haplotypes (identified by Prager et al., 1996, 1998) and an 11-bp insertion
16	in CR of some DOM sequences, while encoding the occurrence of these repeats into
17	the taxon name to check for conformation with phyletic lineages.
18	To obtain a general impression of clustering topology we constructed
19	Neighbor-Net (NN) networks for reduced datasets of 399 unique CR haplotypes and 98
20	unique Cytb haplotypes, and using the default parameters of uncorrected P distance and
21	the EqualAngle algorithm, as implemented in SplitsTree 4.10 software (Bryant and
22	Moulton, 2004). The principal advantage of this hypothesis-poor method over others
23	that generate dichotomous branching networks or trees is that NN networks illustrates

1	all potentially supported splits among a group of sequences as a reticulation. The
2	potential complexity of a dataset is thus represented rather than reduced by this method,
3	while any predominant network topology remains visible. Further insights into the
4	structure of each of the CAS, MUS and DOM mtDNA lineages was obtained by
5	constructing NN networks, together with Median-Joining (MJ) networks (Bandelt et al.
6	1999), as implemented in SplitsTree 4.10.
7	Maximum likelihood (ML) phylogenies were constructed for each of the CR
8	and Cytb datasets and for a concatenate dataset for 30 individuals. We used the PhylML
9	algorithm (Guindon and Gascuel, 2003) with the HKY substitution model, as
10	implemented on the ATGC website (http://www.atgc-montpellier.fr/). A maximum
11	parsimony (MP) method and the neighbor-joining (NJ; Saitou and Nei, 1987) method
12	were taken for phylogenetic inference with concatenate sequences using PAUP 4.0b10
13	(Swofford, 2001). Bootstrap analysis was carried out with 1000 pseudoreplicates in the
14	ML and NJ analyses, and 100 pseudoreplicates in the MP analysis. Sub-groups are
15	designated within each of the major mtDNA lineages only if there was moderate to
16	good bootstrap support (BSS = $0.7 - 0.9$ ) from the ML analysis, combined with
17	concordant structure in the NN networks.
18	We estimated the age of most recent common ancestors (TMRCAs) for
19	mtDNA clades using the Cytb sequences and a relaxed Bayesian molecular clock with
20	uncorrelated rates (BEAST v1.6.1, Drummond and Rambaut, 2007), as described
21	previously (Nunome et al., 2010b). For this analysis we used <i>M. cypriacus</i> , <i>M.</i>
22	macedonicus, M. spicilegus, and M. spretus, the remaining members of the Mus
23	musculus Species Group, as outgroup taxa. For the root node of the Mus musculus

1	Species Group we assigned a prior value of 1.7 mya (95% HPD: 1.45 - 1.95), which is
2	a based on molecular divergences of single copy nuclear gene sequences (Irbp and
3	Rag1), calibrated against the known fossil record of the genus Mus and other Murinae
4	(Suzuki et al., 2004; Shimada et al., 2010). The monophyletic setting was applied for
5	clades of the lineages of the four subspecies groups (CAS, MUS, DOM, and NEP) and
6	M. m. subspecies. Then TMRCAs were estimated by the Bayesian Markov-chain
7	Monte-Carlo (MCMC) method, using the HKY substitution model as selected under
8	the Akaike Information Criterion in MrModeltest version 2.2 (Nylander, 2004).
9	Analyses were run for 50 million generations from a UPGMA starting tree with
10	sampling at every 5000 generations following 5 million burn-in generations. The
11	convergence of MCMC chains and the effective sample size (ESS) values exceeding
12	200 for all parameters were assessed using the software Tracer version 1.5 (Rambaut
13	and Drummond, 2009). BEAST analysis was not performed with the CR sequences due
14	to the greater inequality in branch lengths observed on the CR ML trees, compared
15	with the Cytb ML trees, suggestive of less regular substitution fixation rates over
16	evolutionary time.
17	

#### 18 Assessment of historical demographical processes

19 The DnaSP programme, version 5.00.7 (Librado and Rozas, 2009), was used to

20 estimate haplotype diversity (*Hd*), nucleotide diversity ( $\pi$ ), mean number of pairwise

- 21 differences among sequences (k), and Tajima's D value. The same software was used
- 22 for the analysis of mtDNA sequence mismatch distributions, measured as substitutional
- 23 differences between pairs of haplotypes. Estimates of the expansion parameter tau  $(\tau)$

1	were calculated using Arlequin version 3.5 (Excoffier and Lischer, 2010). Population
2	expansion times were estimated under the assumption of a constant molecular clock
3	and using mutation rates from 2.5%, 10% and 20% using the online tool developed by
4	Schenekar and Weiss (2011; available at http://www.uni-graz.at/zoowww/mismatchca
5	<u>lc/mmc1.php</u> ). The goodness-of-fit of the observed distribution to the expected
6	distribution under the sudden-expansion model (Rogers, 1995) was tested by
7	computing the sum of squares deviation (SSD).
8	

### 9 **Results**

### 10 Characteristics of major haplogroups

11 The NN network generated from the Cytb dataset features four well-differentiated 12haplogroups (Fig. 2a), three of correspond to the previously identified DOM, CAS and 13MUS lineages. The fourth haplogroup includes two sequences derived from Nepalese 14 mice, one reported previously by Terashima et al. (2006; HS1467) and one new to this 15study (HS1523). For convenience, this haplogroup is herein labeled NEP to indicate its 16 geographic origin. No Cytb sequences are available for the GEN haplogroup. The four 17*Cytb* haplogroups are equally divergent from a common central hub and outgroups 18 either join this central hub or have an affinity with the MUS haplogroup. The CAS 19haplogroup appears to contain deeper lineage diversity than either the MUS or DOM 20haplogroups, both of which have distinctly brush-like terminal segments. Overall, the 21topology of the NN network for the Cytb dataset is suggestive of a more or less 22simultaneous diversification of an ancestral Mus musculus stock into multiple 23evolutionary lineages, and also indicative of much recent diversification in each of the

## 1 DOM and MUS haplogroups.

2	The ML phylogeny generated from the <i>Cytb</i> dataset also features the same
3	four clades with support values between 99% (DOM) and 100% (CAS) (Fig. 3a).
4	Monophyly of Mus musculus (sensu lato) is well-supported relative to the outgroups
5	but there is no support for any special relationships among the four haplogroups.
6	The NN network generated from the CR dataset (Fig. 2b) shows a
7	well-differentiated cluster of DOM sequences but less marked segregation among the
8	other groups which now includes GEN. A network generated without DOM (Fig. 2c)
9	shows well-differentiated haplogroups for GEN and MUS, and a less cohesive cluster
10	of 5-6 haplogroups that includes 4-5 that might be regarded as 'CAS' (CAS-1, CAS-2,
11	CAS-3, CAS-4, and AF074526) and one that includes the two NEP haplotypes
12	(HS1467, Tukuche; HS1523, Kathmandu) as well as 'CAS' types 13 (AF074524,
13	Kathmandu) and 14 (AF074525, Nuwakot) from Prager et al. (1998). The GEN and
14	some CAS haplogroups are more divergent from the central hub than other
15	haplogroups but this may be due in part to missing data in some sequences obtained by
16	Prager et al. (1998) from museum skins.
17	The ML phylogeny generated from the CR dataset features five major clades
18	with support values between 86% (CAS) and 100% (DOM) (Fig. 3b). Monophyly of
19	Mus musculus (sensu lato) is well-supported but there is no strong support for any
20	special relationships among the haplogroups, as well as in the Cytb dataset.
21	To further explore the phylogenetic relationships among the haplogroups, we
22	constructed an ML phylogeny using concatenate sequences (CR+ $Cytb$ ) for the 30
23	individuals represented in both datasets. The resultant trees remain ambiguous for

1	branching order among the four major lineages of CAS, DOM, NEP and MUS (Fig. 4).
2	Genetic diversity in each of the main haplogroups is summarized according
3	to a variety of standard parameters in Table 1. Excluding NEP where n=2, for <i>Cytb</i> the
4	highest nucleotide diversity (Pi) is observed in DOM, followed by CAS and MUS;
<b>5</b>	while for CR nucleotide diversity is highest in CAS, followed by DOM and GEN, with
6	MUS once again lowest. The average number of nucleotide differences $(k)$ is highest
7	for Cytb in DOM, followed by CAS and MUS; and highest for CR in DOM, followed
8	by CAS, GEN, and MUS. The number of distinct haplotypes (H) and number of
9	polymorphic sites (S) both are clearly correlated with the total number of samples (N)
10	in each of the <i>Cytb</i> and CR datasets (Table 1).
11	
12	Geographic distribution of major haplogroups
13	The newly determined haplotypes show geographic distributions largely consistent
14	with expectation based on previous findings (Fig. 1a). DOM haplotypes are
15	concentrated around the Mediterranean region but show numerous widely dispersed
16	outliers including localities within MUS territory in western and northeastern Russia
17	and in China, and within CAS territory in the Philippines and Indonesia; MUS
18	haplotypes are predominant in northern part of Eurasia excluding western Europe; and
19	CAS haplotypes are predominant across South and Southeast Asia but with outliers in
20	Japan, the Middle East and eastern Russia.
21	A more detailed mapping of new and previously published sequences from
22	South Asia through to the Middle East illustrates the concentration of mtDNA diversity
23	in southwestern Asia for <i>M. musculus</i> as a whole and additionally for subgroups within

- 1 CAS (Fig. 1b; identity of sub-groups discussed below).
- $\mathbf{2}$

## 3 Genetic and phylogeographic structure of individual haplogroups

## 4 CAS haplogroup

5	Four well-differentiated sub-groups within CAS are clearly depicted in the NN network
6	for the CR dataset (Fig. 2c) and they are also evident in the NN for the smaller Cytb
7	dataset (Fig. 2a) and in the ML tree for the concatenated dataset (Fig. 4); they are
8	herein designated as CAS-1, CAS-2, CAS-3 and CAS-4, as mentioned above. Two of
9	these sub-groups were identified by Terashima et al. (2006) and labeled CAS-II (=
10	CAS-1 of this study) and CAS-I (= CAS-2 of this study). An outlier CR sequence
11	(AF074526 = CAS type 15 of Prager et al., 1998, from Ilam, western Nepal) may
12	represent a fifth sub-group (Fig. 2c) but this requires confirmation as it was obtained
13	from a museum skin and contains several gaps. The CAS sub-groups emerge from a
14	central hub on the NN networks and, with the exception of AF074526, show
15	approximately equivalent degrees of divergence. Each of the main sub-groups also
16	shows relatively deep haplotype diversity; uniquely in CAS-1, this includes a
17	brush-like structure suggestive of recent radiation from a common ancestral haplotype.
18	The ML phylogeny for the concatenated dataset (Fig. 4) recovered
19	monophyletic clades with good support (> 90%) for CAS-1, CAS- 2 and CAS-3,
20	indicated a close relationships between CAS-3 and CAS-4 (BRC3025) with low or
21	moderate support (> 50%), and suggested a basal derivation of CAS-2 with low or
22	moderate support (>50%).
23	The phylogeographic pattern for the CAS haplogroup appears relatively

1	uncomplicated. The greatest haplotype diversity is observed in Pakistan and northern
2	India where all four sub-groups are present but CAS-2 and CAS-3 are dominant (Fig.
3	1b). Approximately half of the sequences of CAS-2 share a 76 bp tandem repeat
4	(reported by Prager et al., 1996, 1998) which further supports the monophyly of this
5	sub-group; these include mice from Taitung in Taiwan and Hanoi in Vietnam (Fig. 2d),
6	constituting the only occurrences of the CAS-2 haplogroup outside of India and
7	Pakistan.
8	Subgroups CAS-1 to CAS-3 are represented in central India but CAS-1 alone
9	is more widely distributed, with representation in southern India and Sri Lanka, and
10	also across southeast Asia, China and eastern Russia to Japan (Fig. 1). A NN analysis
11	of using concatenate sequences (CR+Cytb) from 40 individuals of CAS-1 (Fig. 5a)
12	suggested the presence of a further sub-division that we recognize as CAS-1a and
13	CAS-1b. CAS-1a haplotypes come from two localities in southern China (Guilin and
14	Kunming), from northern Japan (northern Honshu and Hokkaido), and from southern
15	Sakhalin. CAS-1b haplotypes come from a wider geographic area including several
16	parts of India, Bangladesh, Sri Lanka, Myanmar, southern China, Hainan Island,
17	southern Sakhalin and Primorye, eastern Indonesia, and Morocco.
18	
19	MUS haplogroup
20	The MUS haplogroup appears to be comprised of two main sub-groups
21	which are herein designated MUS-1 and MUS-2. These are most clearly expressed in
22	the NN network based on concatenated CR and Cytb data from 38 individuals (Fig. 5b)
23	but they are also evident in the networks generated from the individual data sets (CR,

## 1 Fig. 2e; *Cytb*, Fig. 2f).

2	A total of seven clusters were identified within MUS-1 on the CR NN
3	network (labeled i-vii on Fig. 2e); the majority of these clusters show a high level of
4	geographic fidelity. Based on relationships observed in the NN networks for Cytb (Fig.
5	2f) and the concatenated data set (Fig. 5b), we suggest that these CR phyletic groups
6	can be revolved into three phyletic lineages that we herein designate as MUS-1a (CR
7	clusters i, iii, v, vii), MUS-1b (CR clusters iii, v), and MUS-1c (CR cluster vi).
8	Sub-group MUS-1 as a whole is represented across the entire geographic
9	range of MUS. However, its components show high fidelity to discrete geographic
10	areas: MUS-1a is largely confined to eastern Europe (Ukraine, Moldova, south Siberia,
11	and Primorye in the Russian Far East; see Fig. 1 for the geographic distribution);
12	MUS-1b is predominantly Chinese, being represented at multiple localities spanning
13	the entire breadth of China, from Xanjiang Uyghur Autonomous Region in the
14	northwest to Shandong Province on the eastern seaboard, though there are several
15	occurrences of in Transcaucasia, in Iran, in eastern Europe, and in Russia adjacent to
16	China; and MUS-1c is geographically restricted to far northeast China (Tumen,
17	Qiqihar), Korea and Japan, with one outlier recorded from the coastal city of Kraskino,
18	near the Russian-Korean border (Fig. 1c).
19	The MUS-2 sub-group is distributed across the eastern half of the range of
20	MUS, with representation to the north and east of the Caspian Sea (Kazakhstan and
21	Turkmenistan, respectively), south Siberia in the Altai Mountains, Novosibirsk and
22	Irkutsk, Primorye, and across China including localities in the far northwest (Ili
23	Khazakh Autonomous Prefecture), the central region (Ningxia Hui Autonomous

1	Region), the far north (Manasi), the Tibetan Plateau (Lhasa), and Shandong Province in
2	the east (Liyang). Most of the MUS-2 haplotypes recorded to date are known from
3	single localities and many differ by two or more nucleotide substitutions from the
4	closest sequences (Fig. 2e, f). Only two MUS-2 haplotypes were detected at multiple
5	localities and neither appears to be an ancestral haplotype. In each case, the shared
6	haplotypes are recorded from widely separated localities, suggestive of recent
7	long-distance dispersal or translocation.
8	
9	DOM haplogroup
10	The NN network for DOM CR sequences is an explosively radiating structure, likened
11	by Bonhomme et al. (2011: Fig. 1b) to a "multiple-armed sea star" (Fig. 2g). The
12	additional 23 CR sequences added in this study do not disrupt the primary structure of
13	the NN network with eleven haplogroups (HGs), though HGs 1 and 2 appear somewhat
14	more mixed than in the presentation of Bonhomme et al. (2011: Fig. 1b) and the small
15	HG9 appears to have disaggregated into basal positions within HGs 1, 2 and 7. Most of
16	our new sequences fall into HG11 which corresponds with Clade F of Jones et al.
17	(2010), including sequences from the novel (outlier) localities of Somalia (HS3700),
18	central China (MG509, MG566), and Java, Indonesia (HS2322).
19	Most of the HGs are also evident in an ML tree (not shown) though
20	supporting values were low (65% for HG5 and less than 50% for others). However,
21	HG9 occupies a more diffuse central position consistent with its lack of unity in the
22	NN network, and HGs 1 and 2 were not supported, though most members of these
23	putative HGs associated correctly in smaller clades. An aggregation of HG7 with HG8,

1 is also evident in both the NN network and the ML tree, albeit with no substantial

2	support in the ML analysis.
3	The smaller Cytb dataset presents a simpler picture (Fig. 2h). The NN
4	network shows 12 clusters, some of which are represented by single sequences. Six of
5	the clusters can be correlated to CR HGs based on the subset of individuals represented
6	in both datasets. A MJ network (not shown) shows a completely stellar arrangement
7	with minimal reticulation and with all terminal haplotypes similarly divergent (3-6
8	nucleotide substitutions) from a central node. This putative ancestral haplotype has not
9	been detected. HGs 8 and 9 derive from a common primary branch on the MJ network.
10	The various analyses performed on DOM sequences do not suggest any
11	grounds for its formal sub-division, as was suggested above for each of CAS and MUS.
12	Rather, the topology appears to be genuinely explosive, involving differentiation of
13	multiple, regionally-based matrilines, as concluded also by Rajabi-Maham et al. (2008,
14	2012) and Bonhomme et al. (2011).
15	
16	Divergence time among and within the haplogroups
17	Divergence estimates generated by BEAST for each of the four major haplogroups
18	have central values that range between 0.37-0.46 mya (Fig. 6); in each case the 95%
19	highest probability density values have spans of around $\pm 0.16$ mya (Table 2). The
20	TMRCA of subgroup diversification within each of the CAS, MUS, DOM and NEP

21 haplogroups was estimated at 0.22  $\pm$  0.08, 0.15  $\pm$  0.07, 0.13  $\pm$  0.04 and 0.14  $\pm$  0.06

- 22 mya, respectively (Table 2).
- 23 The Tajima's *D* values for all haplogroups and subgroups were significantly

1	negative, indicating various phases of rapid population growth involving mice with
2	matrilines in CAS-1, CAS-1b, MUS-1, MUS-1b, MUS-1c and DOM (Table 3). We
3	estimated the age of population growth in each of the phyletic groups under four
4	different mutation rates of <i>Cytb</i> ; 2.5%, 10% and 20% (Table 3).
5	The mismatch distribution for the CAS-1 Cytb dataset (Supplementary Fig.
6	S1) shows a multi-peaked distribution which is consistent with the notion of CAS-1 as
7	a well-structured haplogroup (Fig. 5a). CAS-1b shows a good mismatch conformation
8	to a model of recent population growth, with further support coming from a statistically
9	significant negative value for Tajima's $D$ (Table 3). Tajima's $D$ was negative but not
10	statistically significant for CAS-1a. In MUS there is support for recent population
11	expansion of MUS-1 as a whole and for each of MUS-1b and MUS-1c, each backed up
12	by statistically significant negative values for Tajima's D, though the SSD value for
13	MUS-1 was significant ( $P < 0.01$ ), as evidence for departure from the estimated model
14	of population expansion (Table 3). In contrast, there is no support for recent population
15	expansion of either MUS-1a and MUS-2. The mismatch distribution for DOM both CR
16	(data not shown) and Cytb datasets (Fig. S1) shows near perfect conformation with the
17	population growth and decline model provided by DNASP. Neutrality test statistics
18	also point to a significant phase of population expansion in the recent history of DOM
19	(Tajima's $D = -2.02$ , P < 0.05), as concluded previously by others (Rajabi-Maham et al.,
20	2008; Bonhomme et al., 2011), though the SSD value was significant ( $P = 0.024$ ).
21	

# **Discussion**

23 Much of our current understanding of *Mus musculus* phylogeography remains little

1	modified from the conclusions of early studies of mtDNA (e.g. Boursot et al., 1993,
2	1996; Yonekawa et al., 1994; Prager et al., 1996, 1998; Boissinot and Boursot, 1997)
3	and of allozymes and other nuclear markers (e.g. Bonhomme et al., 1984; Miyashita et
4	al., 1994; Din et al., 1996). Three of the most persistent notions to emerge from these
5	early studies are: 1) the understanding that the common ancestor of all of the major <i>M</i> .
6	musculus haplogroups arose in the region of western to central Eurasia, either
7	somewhere in the mountainous terrain that extends from Transcaucasia through to
8	northwest India (Boursot et al., 1993, 1996; Din et al., 1996) or possibly in the
9	low-lying region of Mesopotamia (Prager et al., 1993, 1996); 2) the belief that the
10	broader distribution of all major haplogroups is due to range expansions that occurred
11	following the development of commensalism and thus within the last 10,000 years; and
12	3) the conclusion that the CAS lineage is genetically more diverse and probably older
13	than either of DOM or MUS, with MUS probably trailing DOM in this regard.
14	To date these notions have been subject to detailed scrutiny only for the
15	DOM haplogroup (Gündüz et al., 2005; Darvish et al., 2006; Rajabi-Maham et al.,
16	2008; Bonhomme et al., 2011; Duvaux et al., 2011). In this case, the majority of results
17	uphold the general assumptions as outlined above.
18	For each of the MUS and CAS haplogroups the most comprehensive
19	phylogeographic analyses prior to this study were contained in the work of Prager et al.
20	(1996, 1998). For their initial study of the musculus and domesticus lineages
21	geographic sampling was heavily biased toward Europe, with only a smattering of
22	samples derived from the eastern range of <i>musculus</i> . In the later study this was partially
23	rectified through the laborious extraction of DNA from museum skin samples from

1 eastern populations of musculus and castaneus. Despite this remarkable effort, major  $\mathbf{2}$ geographic gaps in sampling remained; and with such large geographic areas to cover, 3 sample sizes were small for all regions. Our sampling has filled many of the gaps in geographic coverage, especially 4  $\mathbf{5}$ for the Indian subcontinent, Indochina and the Far East. However, the issue of small 6 sample sizes remains and will not be solved without further field collecting on a 7multi-regional scale. Nevertheless, our broader sampling produces new insights into 8 the phylogeography of each of the CAS and MUS groups, and allows us to challenge 9 several key aspects of the current understanding.

10

#### 11 A homeland for *M. musculus* in southwestern Asia

12The ancestral homeland of Mus musculus is most likely to coincide with a broad region 13of co-occurrence of the various phylogroups and it should encompass or abut the 14geographic range of the most restricted phylogroups, namely GEN and NEP of the 15Arabian Peninsula and Himalayan region, respectively. Under these criteria, the region 16of southwestern Asia, encompassing modern day Iraq, Iran, Afghanistan, Pakistan, and 17northwestern India stands out as the most likely candidate area. Bonhomme et al. 18(1984) reached the same conclusion on different evidence, namely the higher levels of 19variation in nuclear genes among mice in this area compared with peripheral regions 20(see also Suzuki et al., 1986; Boursot et al., 1996; Boissinot and Boursot, 1997; Prager 21et al., 1998; Darvish et al., 2006; Duvaux et al., 2011). As discussed at length by Prager 22et al. (1998), mtDNA lineage boundaries in this area show general association with major geographic barriers (see also Duvaux et al., 2011). In particular, the Zagros 23

1	Mountains divide DOM in the west from CAS in the east, while the Elburz Mountains
2	divide MUS in the north from CAS in the south. Similarly, the mountain chains of the
3	Hindu Kush separate populations of MUS and CAS in northern Afghanistan, though
4	the present day distribution of the mtDNA haplotypes is not always associated with the
5	mountainous range (e.g., MUS in Kabul, Afghanistan, Fig. 1a). A process of allopatric
6	differentiation is indicated, as suggested also by the lack of overt ecological
7	differentiation among the divergent populations.
8	Our divergence estimates of 0.37-0.47 mya (see Table 2 for confidence
9	interval) for the major mitochondrial phylogroups are in good accord with previous
10	determinations (Rajabi-Maham et al., 2008; Terashima et al., 2006). Initial lineage
11	diversification evidently predates the dispersal of modern humans out of Africa, hence
12	it is likely that initial phases of range expansion were not mediated by human activity,
13	unless of course the impact of early human populations on the environment was much
14	greater than currently understood.
15	An interesting biogeographic observation is that the inferred place of origin of
16	M. musculus southwestern Asia lacks any other co-occurring mouse species belong to
17	subgenus Mus. In this regard, it differs from each of peninsular India, where M.
18	booduga and M. terricolor of the M. booduga Species Groups are both found (Musser
19	and Carleton, 2005); Indochina, that hosts a variety of species in the <i>M. booduga</i> and
20	M. cervicolor Species Groups (Suzuki and Aplin, 2012); and eastern Europe, where
21	other species of the Mus musculus species group are present. It is tempting to speculate
22	that the presence of these ecologically similar native species in surrounding areas
23	formerly served to constrain the geographic distribution of <i>M. musculus</i> .

1	The distribution and ecology of contemporary castaneus populations in Asia
2	provides further clues to its regional history. As summarized by Marshall (1977:
3	205-206) the only 'outdoor commensal' (i.e. agricultural field) populations of CAS
4	mice are found in the semi-arid habitats of Pakistan (the bactrianus morphotype).
5	Elsewhere on the Indian subcontinent and through into Southeast Asia, house mice are
6	found only as 'indoor commensals'; furthermore, across Southeast Asia, they are
7	generally confined to larger towns and absent in rural villages (see also Aplin et al.,
8	2006). Marshall (1977) attributed the absence of house mice from agricultural contexts
9	in these areas to competitive exclusion by other species of Mus, notably members of
10	the Mus booduga Species Group on the Indian subcontinent and members of the Mus
11	cervicolor Species Group in South East Asia; and he attributed the absence of house
12	mice in rural villages in Southeast Asia to the presence of commensal species of Rattus
13	such as <i>R. exulans</i> .

#### 15 **Phylogeography of the CAS lineage**

16The CAS lineage has been subject to two different phylogeographic interpretations. 17Boursot et al. (1993, 1996) proposed that the northern Indian subcontinent was both the 18place of origin of *Mus musculus* and the cradle of genetic diversity within this group. 19This model was based on the discovery in this area of numerous highly divergent 20mtDNA lineages (Boursot et al., 1996) and levels of nuclear diversity (as determined 21by allozyme electrophoresis) that exceeded those found in European populations of 22domesticus and musculus (Din et al., 1996). To explain these dual observations Boursot 23et al. (1993, 1996) proposed a 'centrifugal' model of differentiation in which the

1	ancestors of each of the <i>domesticus</i> , <i>musculus</i> and <i>castaneus</i> lineages dispersed to the
2	west, east and north, each carrying a subset of the mtDNA and nuclear diversity, and
3	subsequently undergoing local differentiation. They referred to populations in the
4	ancestral area as "Mus musculus subsp." and restricted use of the name castaneus to
5	populations in southern India, southern China and Indochina. Boursot et al. (1996)
6	referred to the northern Indian and Pakistani populations as an 'oriental group', while
7	Yonekawa et al. (1994) subsequently applied the existing name bactrianus to these
8	populations.

9 Prager et al.'s (1998) version of CAS phylogeography is based primarily on 10 interpretation of mtDNA phylogeny. While confirming a high diversity of mtDNA 11 types in northern India and Pakistan, they regarded these to be part of a monophyletic 12castaneus lineage distinct from each of domesticus, musculus and the newly recognized 13gentilulus lineage of the Arabian Peninsula. Prager et al. (1998) developed a model of 14'sequential' derivation of the lineages to reflect their phylogenetic branching order -15domesticus being the oldest branch, followed by gentilulus, castaneus and musculus. 16 They preferred to locate the ancestral pre-domesticus stock in the Near East, within the 17current range of *domesticus*, and regarded the progressive derivation of other lineages 18as a consequence of sequential dispersal events that took house mice south onto the 19Arabian Penisula, then east onto the Indian subcontinent, and finally, north through the 20mountains of northwest India and Pakistan to occupy the great Eurasian steppe. Within 21CAS, Prager et al. (1998:858) suggest a relatively long phase of regional 22diversification on the Indian subcontinent, followed by a 'more recent' dispersal into 23the 'humid lowlands of Southeast Asia'.

1	Our sampling for CAS is relatively extensive and the results go far towards
2	illuminating the historical phylogeography of this haplogroup. In keeping with the
3	findings of Prager et al. (1998) and contrary to the predictions of Boursot et al. (1993,
4	1996), we recovered reciprocal monophyly with good to excellent support among all of
5	the major haplogroups, including CAS. While Prager et al. (1998) considered the
6	branching order among the major haplogroups to be resolved, our larger CR and Cytb
7	dataset fails to provide a robust phylogenetic structure at this level, although there is a
8	suggestion of special affinity between CAS and MUS, and between DOM and NEP.
9	Like both groups of previous researchers, we found the highest mtDNA diversity and
10	depth in CAS populations inhabiting the mountainous region of northwest India and
11	Pakistan, with a loss of haplotype lineage diversity from north to south on the Indian
12	subcontinent (Boursot et al., 1996), and from west to east into Southeast Asia (Prager et
13	al., 1998). Despite this general agreement, Boursot et al (1996) clearly regard
14	castaneus in their restricted application of the name to be a long-term resident of
15	Southeast Asia, while Prager et al. (1998) portray this as a relatively recent phase of
16	dispersal of castaneus though without specifying any time frame.
17	Low nucleotide diversity in the widely distributed CAS-1 sub-group is evident
18	in this study. This is consistent with that observed by the recent work on the castaneus
19	subspecies group done by Rajabi-Maham et al. (2012; see also Bonhomme and Searle,
20	2012). Our results are suggestive of a relatively recent range expansion of CAS-1 to a
21	large geographic areas covering the south and east Indian subcontinent, Southeast Asia,
22	Indonesia, South China, Northeast China and the Russian Far East (Fig. 5). On the
23	other hand, the presence of the locally restricted phyletic group, CAS-1a is suggestive

of stepwise historical range expansion of CAS-1. Haplotype diversity within CAS-1a,
the sub-group found in mice from South China (Kunming and Guilin), northern
Honshu, Hokkaido, and South Sakhalin, was most likely produced by subsequent
dispersal and is suggestive of several thousands of years of *in situ* evolution.
Furthermore, the location of the Japanese cluster at the far eastern periphery of the
CAS distribution implies a significantly earlier onset for dispersal onto the Indian
subcontinent and thence through to East Asia.

8 We suspect that the dispersal of CAS-1 mice occurred in response to 9 ecological transformation of the landscape by early agriculturalists and the emergence 10 of urban centers and trade networks. As has been postulated for the Middle East 11 (Auffray et al., 1990; Cucchi and Vigne, 2006), South Asian populations of Mus 12musculus are likely to have benefited from the creation of new agricultural landscapes, 13and the common practice of storing harvested grain inside villages and even inside 14houses provided the context for development of commensalism. Long-distance 15dispersal is part and parcel of commensalism, with mice being carried as stowaways 16during transport of grain, building materials, clothing and bedding (Pocock et al. 2005). Although the archaeological record of agriculture is less comprehensive for 1718Asia than for the Middle East and Europe, there is good evidence for domestication of 19cereal crops including rice and millet by about 9,000 years ago in several parts of 20South and East Asia (Khush 1997; Londo et al., 2006; Zheng et al., 2009; Molina et al., 212011) and even earlier evidence for long distance overland and maritime trade (Oka 22and Kusimba, 2008). Assuming that populations experienced a sudden or exponential 23growth, we calculated  $\tau$  values from the *Cyt-b* sequences and estimated times since the

1	onset of population expansions. Higher rates of mutation (e.g. 10% or 20% per million
2	years per lineage) rather than lower rates (e.g. 2.5%) are considered to be realistic for
3	assessing rather recent diversifying events (Ho et al., 2005). We obtained a $\tau$ value of
4	1.7 for CAS-1b (n=17) which under mutation rates (per million years per lineage) of
<b>5</b>	10% and 20%, gives expansion times of 7,600 and 3,800 years, respectively (Table 3).
6	A $\tau$ value was not calculated for CAS-1a but it too is likely to have commenced its
7	dispersal and diversification in China, the Russian Far East and Japan in prehistoric
8	times. In this regard, it is of interest to note archaeological evidence for rice cultivation
9	along the upper Yangtze river (e.g. Yunnan province, here represented by mice from
10	Kunming) at 4500 years ago (Fuller et al., 2010) and recent genetic evidence from an
11	intensive genome survey on wild and cultivated rice, suggesting the Pearl River
12	(Guangxi province, here represented by mice from Guilin) in southern China is the
13	place of the first development of cultivated rice (Bonhomme and Searle, 2012; Huang
14	et al., 2012).

15Comparatively recent long-distance dispersal of CAS mice most likely 16explains the detection of CAS-2 haplotypes at isolated localities in Taiwan and 17Vietnam, though we could not exclude out the possibility that these are relictual 18haplotypes, either rare survivors of an earlier dispersal of CAS-2 mice out of India that 19was swamped by a later CAS-1 dispersal, or the last remnants of incomplete lineage 20sorting of an immigrant population with a mixture of CAS-1 and CAS-2 haplotypes. In the case of the individual from Taiwan, the fact that its nuclear genetic profile is fully 2122consistent with other East Asian populations of CAS and differ from mice from India

1 and Pakistan with the CAS-2 mtDNA haplotypes (Nunome et al., 2010a; Kodama et al.,

2 unpublished) suggests that we are not dealing with a novel invader but perhaps with a

- 3 product of mtDNA introgression.
- 4

#### 5 Phylogeography of the MUS lineage

6 Previous phylogeographic interpretations of the house mouse group do not vary much 7in regard to the geographic origin of the MUS haplogroup. Boursot et al. (1993: 406) 8 speculated that "the cradle of M. m. musculus could be in Transcaucasia or east of the 9 Caspian Sea", while Prager et al. (1993, 1996) saw the origin of MUS as the product of 10 northward dispersal from a proto-CAS population occupying the region east of the 11 Caspian Sea, followed by range expansion. Both groups of researchers also agree that 12MUS populations subsequently dispersed west into central Europe and east into China 13and Japan, and this scenario has been adopted as paradigmatic by Japanese researchers 14interested in the origin of the indigenous molossinus population (Yonekawa et al., 151988; Terashima et al., 2006; Nunome et al., 2010a). Yonekawa et al. (1988) postulated 16that MUS populations relatively recently expanded into China where CAS populations 17had already colonized but few other workers have expressed an opinion on the earlier 18timing of the remarkable eastward expansion of MUS. Nunome et al. (2010a) suggested a latitudinal division within MUS between northern (MUS-I) and southern 1920(MUS-II) groups, based on phylogeographic analyses of nuclear gene sequences, and 21posited that range expansion of the MUS haplogroup from west to east across 22continental Eurasia followed separate northern and southern dispersal routes, with 23separate expansion again into eastern Europe.

1	Much of the interest in the geographic distribution of MUS has focused on its
2	genetic interaction with mice of other haplogroups. In the European context numerous
3	studies have examined the evolutionary dynamics of a narrow hybrid zone with DOM
4	that runs from Norway through Denmark, Germany and Austria to eastern Bulgaria
<b>5</b>	(Hunt and Selander, 1973; Sage et al., 1993; Boursot et al., 1993; Jones et al., 2010).
6	There are grounds to believe that initial contact may have occurred further west in
7	Europe with the current position stabilizing after a period of eastward retreat of
8	musculus (Gyllensten and Wilson, 1987). Whatever the case, the age of the contact
9	zone is constrained by the timing of the DOM migrations along the shores of the
10	Mediterranean, an event that is thought to date to within the last 2-3,000 years (Cucchi
11	et al., 2005).
12	In Transcaucasia, gene flow between complexly parapatric populations of
12 13	In Transcaucasia, gene flow between complexly parapatric populations of MUS and DOM is thought to explain a 300 km wide zone of genetic admixture
13	MUS and DOM is thought to explain a 300 km wide zone of genetic admixture
13 14	MUS and DOM is thought to explain a 300 km wide zone of genetic admixture (Mezhzherin et al., 1989; Frisman et al., 1990; Milishnikov et al., 1990); however, an
13 14 15	MUS and DOM is thought to explain a 300 km wide zone of genetic admixture (Mezhzherin et al., 1989; Frisman et al., 1990; Milishnikov et al., 1990); however, an alternative interpretation attributes the genetic diversity to a high level of ancestral
13 14 15 16	MUS and DOM is thought to explain a 300 km wide zone of genetic admixture (Mezhzherin et al., 1989; Frisman et al., 1990; Milishnikov et al., 1990); however, an alternative interpretation attributes the genetic diversity to a high level of ancestral polymorphism in the regional MUS population (Milishnikov et al., 2004), equivalent to
13 14 15 16 17	MUS and DOM is thought to explain a 300 km wide zone of genetic admixture (Mezhzherin et al., 1989; Frisman et al., 1990; Milishnikov et al., 1990); however, an alternative interpretation attributes the genetic diversity to a high level of ancestral polymorphism in the regional MUS population (Milishnikov et al., 2004), equivalent to that observed among the 'oriental group' of mice in northern India and Pakistan
<ol> <li>13</li> <li>14</li> <li>15</li> <li>16</li> <li>17</li> <li>18</li> </ol>	MUS and DOM is thought to explain a 300 km wide zone of genetic admixture (Mezhzherin et al., 1989; Frisman et al., 1990; Milishnikov et al., 1990); however, an alternative interpretation attributes the genetic diversity to a high level of ancestral polymorphism in the regional MUS population (Milishnikov et al., 2004), equivalent to that observed among the 'oriental group' of mice in northern India and Pakistan (Boursot et al., 1993, 1996; Din et al., 1996). This would be consistent with long
<ol> <li>13</li> <li>14</li> <li>15</li> <li>16</li> <li>17</li> <li>18</li> <li>19</li> </ol>	MUS and DOM is thought to explain a 300 km wide zone of genetic admixture (Mezhzherin et al., 1989; Frisman et al., 1990; Milishnikov et al., 1990); however, an alternative interpretation attributes the genetic diversity to a high level of ancestral polymorphism in the regional MUS population (Milishnikov et al., 2004), equivalent to that observed among the 'oriental group' of mice in northern India and Pakistan (Boursot et al., 1993, 1996; Din et al., 1996). This would be consistent with long residency of the MUS population in this area. MUS and CAS populations also come

23 significant new light on the evolutionary history of the MUS haplogroup. We identify

1	two major sub-groups within MUS – MUS-1 and MUS-2 – and a total of three
2	phylogeographic components within MUS-1: MUS-1a in Moldova, Ukraine, N
3	Caspian Sea and Russian Siberia; MUS-1b in East Europe, Kazakhstan and China; and
4	MUS-1c in Korea and Japan. The origin of the MUS-1 and MUS-2 sub-groups is
5	ancient, with a divergence estimate from BEAST of 150,000 $\pm$ 13,000 years (Fig. 6,
6	Table 2). Both sub-groups are represented in the area around the Caspian Sea and it
7	seems likely that both matrilines originated within this ancestral geographic area.
8	Rapid population expansion was inferred for each of MUS-1b and MUS-1c
9	(Table 3). Estimates of expansion times for these lineages (Table 3) suggest an early
10	expansion of MUS-1b in northern China ( $\tau = 4.9$ CI: 2.9 – 6.5; e.g., 21,000 and 10,800
11	years ago, with an assumption of the mutation rate of 10% and 20% per million years
12	per lineage, respectively), followed by a later expansion of MUS-1c in northeastern
13	Russia, Korea and Japan at ( $\tau = 1.5$ CI: 0.7 – 2.5; e.g. 6,600 and 3,300 years ago).
14	The notion of ancient population expansions in eastern Eurasia is clearly at
15	odds with the conventional notion of a recent west to east dispersal of the MUS
16	haplogroup. However, other lines of genetic evidence similarly point to a long
17	residency of the MUS haplogroup in central Russia and the Far East. For example, the
18	beta-hemoglobin gene (Hbb) shows contrasting predominant alleles in the lower
19	Yellow River basin and in the remaining western portion of northern China $(Hbb^{p})$ and
20	<i>Hbb<sup>w1</sup></i> , respectively; Miyashita et al., 1994; see also Moriwaki, 1994); and mice from
21	the eastern part of China are known to have relatively longer tails (tail ratio: ~93%)
22	than those from the rest of MUS territory in China (81%; Tsuchiya et al., 1994).
23	Finally, we note that the area in which MUS-1c is predominant – the Korean

1	Peninsula and nearby continental area – harbors unique genetic components in both
2	Y-specific gene sequences and nuclear gene sequences (e.g. Nagamine et al., 1994;
3	Terashima et al., 2006; Nunome et al., 2010a). Under the existing paradigm of west to
4	east dispersal, these phylogeographic patterns might be attributed either to genetic drift
5	following migration of ancestral populations with diverse genetic components or to
6	multiple migration events by mice carrying different genetic components, perhaps by
7	different routes. However, neither of these scenarios can readily account for the
8	evidence of ancient population expansions within geographically restricted matrilines.
9	Accordingly, we favor the alternative model of regional differentiation within a
10	long-term resident population.
11	The fossil record should be able to arbitrate this issue and it is of great interest
12	to note that paleontologists have long recognized Mus musculus as a component of the
13	Chinese mammal fauna since the middle part of the Middle Pleistocene (i.e. c. 500,000
14	years ago); e.g. Zheng et al. (1997 and references cited therein). While the taxonomic
15	identification of the fossils might be challenged, the determination is at least plausible
16	given the molecular evidence for early diversification among Chinese musculus
17	populations. However, there is a risk of circularity in such arguments and an urgent
18	need for critical appraisal of the relevant fossils.
19	The European sub-group MUS-1a contains substantial haplotype diversity
20	including persistent ancestral haplotypes and two deeply divided haplotype series, each
21	of which contains relatively shallow stellar clusters derived from populations near the
22	western limit of the MUS geographic range. This pattern is suggestive of a broad
23	westward expansion of a MUS population into eastern Europe, with limited filtering of

1	haplotype diversity. As summarized by Auffray et al. (1990), the long history of Mus
2	musculus in Europe is dominated by large expansions and contractions of range driven
3	by glacial cycles. At the height of the last glaciation <i>M. musculus</i> was rare or absent
4	across most of eastern Europe which supported a mosaic of periglacial forest-steppe,
<b>5</b>	steppe and semi-desert habitats (Markova et al., 2009). Refugial forest habitats were
6	restricted to small patches in the Crimea, in the Transcarpathian region, and in the
7	Caucasus (Markova et al., 2009) and it is of interest to note fossil occurrences of <i>M</i> .
8	musculus in the Carpathian-Balkan region during the warm interval (33-24,000 years
9	ago) immediately prior to the last glacial maximum (Markova, 2010). However, in
10	view of the high level of genetic diversity within MUS-1a and the lack of a strong
11	signal of recent population expansion, it seems likely that mice persisted in multiple
12	localities, perhaps including both forest and semi-desert habitats. This issue warrants
13	further consideration.
14	MUS-1a contains a discrete lineage characterized by a 75-bp duplication, first
15	detected by Prager et al. (1998) in a mouse from Kishinev in Moldova. We found
16	closely related haplotypes at low frequency in mice from eastern Europe (e.g. Donetsk,
17	Ukraine) and also from Khasan in Primorye, Russia (Fig. 5b). Given the other evidence
18	of regional differentiation of mtDNA within MUS, we are inclined to view MUS-1a as
19	originally restricted to eastern Europe (Ukraine), with its more easterly occurrences
20	being a product of long-range transport by modern means. The locality of Novosibirsk,
21	for example, is sited on the Turkestan-Siberia Railway that was built in the early 20th
22	Century (in 1930) and connects the Caspian Sea to localities in Central Asia. The link
23	to the Primorye region of the Russian Far East is less readily accounted for by overland

transportation but might be explained by the activities of the Russian government to
introduce kazak and peasants to the Russian Far East in the late 19<sup>th</sup> Century; upwards
of 90,000 people (and perhaps a few mice) from Odessa in the Ukraine settled in the
Ussuri Region of Primorye (<u>http://www.fegi.ru/prim/geografy/etap.htm</u>).

 $\mathbf{5}$ 

#### 6 Phylogeography of the DOM lineage

7Our small number of new DOM sequences contributes only a few insights into the 8 history of this well-studied lineage (Gabriel et al., 2011; Bonhomme et al., 2011; Jones 9 et al., 2010). The onset of the expansion of is estimated to be 12,000 years ago as the 10 youngest timing, assuming the mutation rate of 20% per million years per lineage 11 (Table 3), which is harmonious with the recent arguments based on zooarcheological 12records (Cucchi et al., 2005; Rajabi-Maham et al., 2008; Bonhomme and Searle, 2012). 13We recovered the expected "Clade F" haplotypes from mice collected in North 14America, Australia and Africa (Senegal, Somalia) but also detected them in mice from 15several localities in Asia, namely Lanzhou and Xining in China, and Bogor on Java in 16Indonesia. At Bogor, CAS and DOM mtDNA haplotypes were found to co-occur in one 17population. 18A high frequency of DOM haplotypes was also detected in the Russian Far East, thereby supporting previous claims of DOM-MUS-CAS interactions in this area 1920based on studies of chromosomes, allozymes and RAPDs (Frisman et al., 2011; 21Spiridonova et al., 2011). Interestingly though, the DOM haplotypes recovered at 22Primorye (HS1466) and Sakhalin (HS3606, HS3607) are not "Clade F" but are related

23 specifically to haplotypes from Cameroon (e.g. AFWCMR41; Bonhomme et al., 2011).

1 This connection is very likely explained by long-distance dispersals associated with

 $\mathbf{2}$ human activities in modern times. 3 The detection of DOM haplotypes in numerous corners of the world is testimony to the ongoing dispersal of M. musculus, and encourages further study of the 4  $\mathbf{5}$ impact of occasional arrival of 'exotic' mice on the genetic constitution of 6 pre-established mouse populations (Rajabi-Maham et al., 2008; Searle et al., 2009a, b; 7Gabriel et al., 2010; Bonhomme et al., 2011). To further illustrate this point, mice with 8 both DOM and CAS mtDNA haplotypes have been captured in Japanese international 9 ports (Tsuda et al., 2007) and Nunome et al. (2010a) provided robust evidence from 10 their nuclear haplotype analysis of genetic introgression by DOM components of 11 Japanese house mice. The extent to which genetic introgression may now be shaping 12the future evolution of the house mouse is an interesting topic - one that has bearing on 13other commensal mammals including the black rat *Rattus rattus* which also displays 14comparable signals of former geographic subdivision and recent intermingling as a 15consequence of commensalism and human-assisted dispersal (Chinen et al., 2005; 16Aplin et al., 2011; Bastos et al., 2011; Lack et al., 2012). 17

#### 18 Concluding remarks

The expanded mtDNA dataset raises a number of important new issues regarding the prehistory of the house mouse. Most significantly, it has identified one particular CAS sub-group (CAS-1) that has expanded into southern India, Southeast and East Asia, and raised the possibility that this expansion is linked to the emergence of agricultural lifestyles and of Asian civilizations. Also of significance is our suggestion that MUS

1	populations have a long history of residency in eastern Russia and China, contrary to
2	the existing paradigm of recent expansion from west to east. Finally, our results
3	emphasize the role of long-distance dispersal in shaping contemporary pattern of
4	distribution and opportunities for interaction between each of the major lineages within
<b>5</b>	Mus musculus.
6	Our study also demonstrates the value of continuing efforts to fill gaps in
7	geographic coverage of Mus musculus mtDNA. Moreover, it highlights the need for
8	ongoing field collecting to increase local sampling and the need for more
9	comprehensive assessments of population genetic history using nuclear markers. From
10	our preliminary work with nuclear genes on this group, it is clear that much deeper
11	divergence between subspecies groups is observed in some regions of the genome than
12	in others (e.g., Suzuki et al., 2004; Nunome et al., 2010a), and also evident that
13	different markers can yield strongly contrasting phylogeographic structure, such as in
14	southern China, where CAS mtDNA dominates but both CAS and MUS components
15	are detected in nuclear genes (e.g., Nunome et al., 2010a). Finally, it is worth
16	mentioning the as yet unexplored potential for detailed study of Central and East Asian
17	house mouse populations to reveal important new aspects of human history, including
18	the emergence of agricultural lifestyles and of regional trade networks.
19	

### 20 DATA ARCHIVING

The nucleotide sequences reported in this paper appear in the DDBJ, EMBL, and
GenBank nucleotide sequence databases under the following accession numbers
AB649455–AB649770, AB819902-AB819920 and AB820897-AB820942 (Table S1).

- 1 Sequence data files in nexus file format, together with Supplementary Information files
- 2 are stored at Dryad repository: doi:10.5061/dryad.rf161.
- 3

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## 1 Figure legends

 $\mathbf{2}$ 

3 Figure 1. Collection localities and mitochondrial genotypes in Eurasia of Mus 4 musculus samples examined in this study (a). New samples genotyped for this study  $\mathbf{5}$ are shown. Detailed locality names and sample codes are listed in Supplementary Table 6 1. Five major mitochondrial groups representing five subspecies groups, M. m. 7musculus (blue: MUS), M. m. domesticus (red: DOM), and M. m. castaneus (yellow: 8 CAS), M. m. gentilulus (white: GEN), and the divergent lineage occurring in Nepal 9 (orange: NEP) are differentially shown. The specific haplotype group of DOM that 10 broadly dispersed to a variety of countries (Australia, Canada, China, Germany, 11 Indonesia, Senegal, Somalia) are marked with arrowheads. Together with those from 12Prager et al. (1998), spatial patterns for the mitochondrial genotypes are shown for 13mice from Central Asia based on combination of new and previously published 14sequences (sources) (b), where further subdivision of the CAS lineage into four 15(CAS-1, CAS-2, CAS-3, CAS-4) are detected. The types of the four subgroups of CAS 16are shown in circle with numerical numbers (black, Prager et al., 1998; red, in this 17study). Further subdivision of the MUS lineages into two, MUS-1 (light blue) and 18MUS-2 (dark blue), and the MUS-1 sublineage into three (MUS-1a, MUS-1b, MUS-1c) is suggested in this study (**a**, **c**). 19

20

Figure 2. Neighbor-Net networks tree based on the cytochrome *b* gene (*Cytb*; **a**, **f**, **h**) and control region (CR; **b**, **c**, **d**, **e**, **g**) of the mitochondrial DNA, with tip labels for the three major subspecies groups, *M. m. musculus* (MUS), *M. m. castaneus* (CAS) and *M.* 

1	m. domesticus (DOM) and two rather geographically confined groups of M. m.
2	gentilulus (GEN) and Nepalease mice (NEP). The portion of the CR network was
3	enlarged to show the details of the branching patterns for CAS-2, in which most of
4	members possess a 75-bp repeat (d). The codes for the haplogroups (HGs) in the CR
5	(g) and Cytb (h) network for DOM were taken from those used in Bonhomme et al.
6	(2011).
7	
8	<b>Figure 3.</b> ML trees for mitochondrial DNA sequences of the cytochrome $b$ gene ( <b>a</b> )
9	and control region (b). The PhylML algorithm (Guindon and Gascuel, 2003) was used
10	for the tree reconstruction and bootstrap analysis (100 replications). Bootstrap values
11	(>50%) are shown under basal branches.
12	
13	Figure 4. ML tree for concatenated mitochondrial DNA haplotypes (control region
14	and cytochrome $b$ gene) using representatives for the four major haplogroups of $Mus$
15	musculus and M. macedonicus as outgroup. Bootstrap values (>50%) are shown under
16	basal branches (ML/MP/NJ).
17	
18	Figure 5. Neighbor-Net networks of concatenate sequences of control region and
19	cytochrome $b$ gene (ca. 2020 bp) from individuals representing the sublineage of CAS,
20	CAS-1 (a) and MUS (b). Prominent subgroups appeared in the networks are indicated.
21	
22	Figure 6. Divergence time estimates (million years ago, mya) of Mus musculus
23	phylogroups and its closely related species, based on a Bayesian relaxed molecular

clock applied to the mitochondrial cytochrome *b* sequences (1140 bp). The posterior probability and 95% HPD intervals of node ages in mya (gray bars) are shown in particular nodes with ancient divergent. The time estimates of 1.7 mya for the root node of the divergence of *M. spretus* and the other species of *M. musculus* Species Group (Suzuki et al., 2004) was used as calibration point. Sequences obtained from the databases are marked with their accession numbers and asterisks.