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SUBJECT CATEGORIES

- » Population genetics
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Genomic resources for wild populations of the house mouse, *Mus musculus* and its close relative *Mus spretus*

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Wild populations of the house mouse (*Mus musculus*) represent the raw genetic material for the classical inbred strains in biomedical research and are a major model system for evolutionary biology. We provide whole genome sequencing data of individuals representing natural populations of *M. m. domesticus* (24 individuals from 3 populations), *M. m. helgolandicus* (3 individuals), *M. m. musculus* (22 individuals from 3 populations) and *M. spretus* (8 individuals from one population). We use a single pipeline to map and call variants for these individuals and also include 10 additional individuals of *M. m. castaneus* for which genomic data are publically available. In addition, RNAseq data were obtained from 10 tissues of up to eight adult individuals from each of the three *M. m. domesticus* populations for which genomic data were collected. Data and analyses are presented via tracks viewable in the UCSC or IGV genome browsers. We also provide information on available outbred stocks and instructions on how to keep them in the laboratory.

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Design Type(s)	parallel group design • species comparison design
Measurement Type(s)	genetic sequence variation analysis • transcription profiling assay
Technology Type(s)	whole genome sequencing • RNA sequencing
Factor Type(s)	Subspecies • geographic location • tissue
Sample Characteristic(s)	<i>Mus musculus domesticus</i> • <i>Mus musculus musculus</i> • <i>Mus musculus castaneus</i> • <i>Mus spretus</i> • Massif Central • Bonn Urban District • Heligoland Islands • Ahvaz • Municipality of Studenec • Almaty City • Mazar-I-Sharif District • Himachal Pradesh State • Madrid • brain • gut wall • heart • kidney • liver • lung • muscle structure • testis • spleen • thyroid gland

Background & Summary

The house mouse (*Mus musculus*) has a long-standing history as a model system in genetics and biomedical research, with many classical inbred strains available for purchase worldwide. By comparing the genetic make-up of classical inbred strains with those of mice collected in the wild, it became clear that classical inbred strains represent complex genomic mixtures with contributions from different subspecies and species of *Mus*^{1–3}. While some of this genomic mixture stems from captive breeding of mice from different parts of the world during the early establishment of inbred strains, admixture^{4,5} and introgression of genomic material across subspecies and species^{6–9} also occurs in the wild and is thus likely to contribute to the genomic complexity observed in inbred strains. Classical inbred strains were found to exhibit a much reduced amount of genetic variation compared to their wild mice ancestors¹⁰. For example, all classical inbred strains share a single mitochondrial lineage derived from *M. m. domesticus*¹¹, indicating that they all descend from the same female lineage of the wild ancestor.

An appreciation of the genetic diversity found in wild mice came with the advent of molecular mapping techniques that required crosses between lines with informative polymorphisms^{12,13}. This has led to renewed interest in studying the evolutionary history of natural house mouse populations worldwide, with a main focus on clarifying its taxonomy and catalogue genetic variations found in the wild^{14–16}. Currently, three major lineages of *Mus musculus*, classified as subspecies, are distinguished: the western house mouse *Mus musculus domesticus*, the eastern house mouse *Mus musculus musculus* and the southeast-Asian house mouse *Mus musculus castaneus*. All three lineages have their origin in Southern Asia and diverged roughly 0.5 million years ago, but still share haplotypes and appear to exchange genomic material^{6,8,9}. Hybrid zones have been detected at areas of secondary contact between the subspecies^{17–19} and these serve for tracing genes involved in hybrid incompatibility^{20–22} as well as quantitative trait mapping²³.

During the past 10,000 years house mice have developed commensalism with humans, which allowed them to spread across the world. Among the recognized subspecies, *M. m. domesticus* seems to be the most successful colonizer of new continents during the past few hundred years^{12,13}, partly revealing historical shipping routes of humans²⁴, and has repeatedly colonized small islands²⁵. One such recent island colonization occurred about 400 years ago and resulted in the naming of a new subspecies, i.e., *M. m. helgolandicus*²⁶. Because of its molecular proximity to *M. m. domesticus*, we treat the *M. musculus* population from Heligoland, a small German archipelago in the North Sea, as a member of the subspecies *M. m. domesticus* in some further analyses.

One of the closest relatives to the *Mus musculus* subspecies complex is the Algerian mouse *Mus spretus*²⁷, with populations inhabiting areas around the western Mediterranean Sea. With a divergence time of roughly 2 million years^{28,29} the *Mus spretus* lineage serves as an ideal outgroup to *Mus musculus*. Although viable offspring can be produced from crosses between *Mus musculus* and *Mus spretus* in the laboratory, it is morphologically and behaviorally rather distinct, justifying its status as separate species. Nevertheless, some exchange of genomic regions is still possible between these species in the wild^{17,9,30}. A few individuals of *Mus spretus* are also represented among classical inbred strains³¹.

The unique combination of genetic and molecular knowledge derived from the classical inbred strains and the profound knowledge of the evolutionary history of wild mouse populations make *Mus musculus* a prime model for the study of evolution and molecular biology of natural populations^{12,13,32}. We have previously used the *Mus musculus* model system to analyze patterns of positive and negative selection in the genome^{8,33–37}, hybrid sterility^{21,22}, evolution of copy number variation³⁸, mapping of craniofacial traits²³ and the composition and turnover of the microbiota^{39–41}.

Here we describe the genomic resources that we have generated using mice collected in the wild over the past 10 years. These data can serve as a basis for in depth studies at a population level and to inform biomedical research projects on natural polymorphisms that are present in inbred strains. We provide information on a) genomic data for a total of nine populations (Fig. 1), covering the three major house

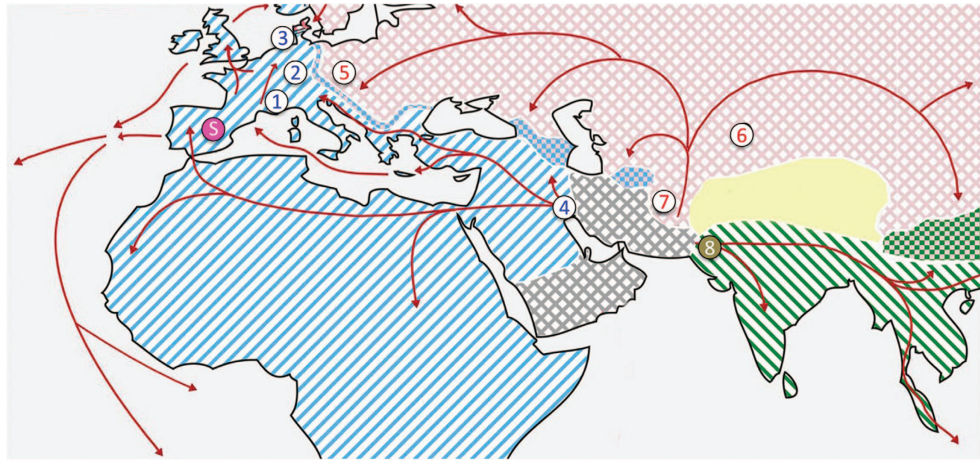


Figure 1. Geographic location of *Mus musculus* (1–8) and *Mus spretus* (S) samples. The map is modified from refs 3,13. The blue area depicts *M. m. domesticus* territory (includes *M. m. helgolandicus* (3), because of its close molecular proximity to *M. m. domesticus*), the red area depicts *M. m. musculus* territory, and the green area depicts *M. m. castaneus* territory. *Mus spretus* co-occurs with *M. m. domesticus* in Spain. The grey area harbors further lineages and possible additional subspecies¹⁶. Red arrows symbolize possible migration routes, mostly in post-glacial times during the spread of agriculture. Locations (year caught): 1, Massif Central/France (2005); 2, Cologne-Bonn/Germany (2006); 3, Heligoland/Germany (2012); 4, Ahvaz/Iran (2006); 5, Studenec/Czech Republic (2003); 6, Almaty/Kazakhstan (2002); 7, Afghanistan (2012); 8, Himachal Pradesh/India (2003); S, Madrid/Spain (2004).

mouse subspecies and one outgroup, b) tissue-specific RNAseq data from three *M. m. domesticus* populations, and c) details on animal husbandry of wild house mice in a Supplementary File and d) some general analyses and browser tracks for visualization. The genomic dataset is summarized using basic descriptive statistics, such as F_{ST} (ref. 42), π (nucleotide diversity⁴³) and Tajima's D ⁴⁴ as measures of population differentiation and selection, as well as a description of copy-number variation based on sequencing read depth. The RNAseq data are summarized as normalized RNAseq read coverage for each base pair of the genome. All statistics are made available as genome browser tracks (bed and bigWig files) to allow close visual inspection of any genomic region of interest in the UCSC browser⁴⁵ or other visualization software such as IGV^{46,47}.

Methods

Sampling procedure and sampling locations

The location of populations used for re-sequencing in this study are depicted in Fig. 1. They include three *M. m. domesticus* populations from Western Europe and Iran, the island subspecies *M. m. helgolandicus*, three *M. m. musculus* populations from the Czech Republic, Kazakhstan and Afghanistan and a *M. spretus* population from Spain. Populations were sampled between 2003 and 2012. The DNA samples used for genome sequencing were obtained either directly from wild caught animals, or from the first or second generation of out-breeding in our animal facility, i.e., they are expected to represent full wild type variation. Some aspects of the genome sequences obtained from the three *M. m. domesticus* populations, as well as the island subspecies *M. m. helgolandicus*, have previously been described^{26,38}.

House mice form naturally extended family groups at a given location including breeding among relatives^{48,49}. To obtain an unbiased population sample from a given region ideally requires sampling mice in a way to avoid catching related animals. Therefore, we aimed to collect only a single mouse per trapping location (or, for the purpose of setting up breeding colonies, one male and one female per location) and selected the next trapping location 500 m to 1 km apart. The whole area sampled ideally comprises a diameter of about 50 km. However, depending on the local conditions, following this sampling regime precisely was not always possible. Moreover, some samples were provided by collaborators who have only recorded the general area for trapping, but not the exact location (e.g., the mice from Kazakhstan). Other trapping locations had regional limitations. For example, the island of Heligoland is only 1.7 km² in size, or the military Camp Marmal, Mazar-e-Sharif (Afghanistan) is only 8.7 km² in size. In both cases, mice were collected in different localities on the island or military base respectively⁵⁰. In Supplementary Table 1 we provide exact location information, as much as it is available for all animals involved in the study, as either directly having been sequenced, or as having been parent to one of the animal facility-born offspring of wild mice (see below).

To complement the genomic resources generated in our laboratory, we also re-analyzed previously published genome sequencing data from *M. m. castaneus* that were collected in the northwest Indian state of Himachal Pradesh⁵¹.

Mice were either caught in snap traps, or live traps ('Mäusewippfalle' No. 3451002, Firma Ehlert & Partner, 53859 Niederkassel, Germany) or were found dead after rodenticide-based pest management. Cervical dislocation was used to sacrifice mice caught in live traps in the field. Transportation of live mice to the animal facility, maintenance and handling were conducted in accordance with German animal welfare law (Tierschutzgesetz) and FELASA guidelines. Permits for keeping mice were obtained from the local veterinary office 'Veterinäramt Kreis Plön' (permit number: 1401-144/PLÖ-004697).

***M. m. domesticus* breeding scheme**

For the *M. m. domesticus* populations our aim was to generate an RNAseq dataset from the same individuals for which we generated the DNAseq data. In order to standardize mice to the same sex, age and environmental conditions, we did not use wild caught mice but bred wild caught mice for one to two generations in our animal facility. Supplementary Table 2 shows the breeding scheme for the *M. m. domesticus* mice in the study. In one case, we caught a pregnant female in the wild and used its male offspring born in the facility for DNA and RNA sequencing. For the Iranian mice, we included two wild caught individuals in the DNAseq study (AH15 and AH23), for which we did not generate RNAseq data. For two additional Iranian individuals in the DNA sequencing study, tissue samples were lost and thus no corresponding RNAseq data exist. To compensate for this, we included four individuals representing male offspring from male AH15 and male AH23 respectively (both individuals are part of the DNAseq dataset; see Table 1 (available online only)). Two male offspring from each of these two males are represented as biological replicates in the RNAseq dataset.

Procedures for wild mouse handling

We established wild-derived outbred populations for *M. m. domesticus* (France, Germany) and *M. m. musculus* (Kazakhstan and Czech Republic). For the first 11–14 generations live mice obtained from the wild were set up in a cyclical breeding scheme aiming at maintaining maximum genetic variability over time. They were then partly refreshed with newly collected mice from the same original area and a HAN rotational breeding scheme⁵² was established. Individuals from each of the 4 outbred populations are maintained at the Max-Planck Institute in Plön, Germany, and are available from the authors upon request.

Wild mice are considerably more agile than classical inbred strain mice. Environmental enrichment is necessary and strongly reduces agitated stereotype behavior in wild mice kept under laboratory conditions. Standard mouse chow is provided *ad libitum* (e.g., Altromin 1,324 from ALTROMIN, 32,791 Lage, Germany). Further details on mouse handling and breeding are provided in Supplementary Material Text 1.

Molecular methods

DNA and RNA extraction. DNA was extracted from liver, spleen, or ear samples using salt extraction⁵³ or DNeasy kits (Qiagen, Hilden, Germany). RNA was extracted only for *M. m. domesticus* mice from Germany (8 individuals), France (8 individuals) and Iran (8 individuals) using mostly the same individuals, which are included in the whole genome sequencing study (see details above and Supplementary Table 2).

Mice designated for RNA extraction were housed alone after weaning and were routinely visually inspected for health and vigor. Only male mice were included in the study, sacrificed at 12 weeks. All mice were fed standard mouse chow *ad libitum* (Altromin 1,324 from ALTROMIN, 32,791 Lage, Germany). Mice were sacrificed by CO₂ asphyxiation. The coat was sprayed with 75% EtOH to reduce loose hair contaminating the organs during dissection of the animal. Organs were extracted in a specific order to improve comparability. Organs were shock frozen in liquid nitrogen and stored at –80° until RNA was extracted.

The Trizol reagent (Life Technologies, Carlsbad, California, USA) was used according to manufacturers instructions to extract RNA from each organ (Table 2 (available online only)). With the exception of the liver, for which we used only right and left medial lobe, whole organs were processed to minimize heterogeneity of the sample. The extracted RNA was quantified on a Nanodrop and analyzed for integrity on the Agilent Bioanalyzer.

DNA sequencing library preparation. All populations apart of the Afghanistan population were sequenced using the same protocol (see below) in the following batches: Batch 1 included the *M. m. domesticus* populations from France, Germany and Iran (locations 1, 2 and 4 in Fig. 1). Batch 2 included the 3 mice from Heligoland (location 3 in Fig. 1). Batch 3 included the *M. m. musculus* populations from the Czech Republic and Kazakhstan (locations 5 and 6 in Fig. 1) as well as the *M. spretus* population from Spain (location S in Fig. 1). Batch 4 included the *M. m. musculus* mice from the Afghanistan population (location 7 in Fig. 1), which were sequenced using a more recent Illumina technology (see below).

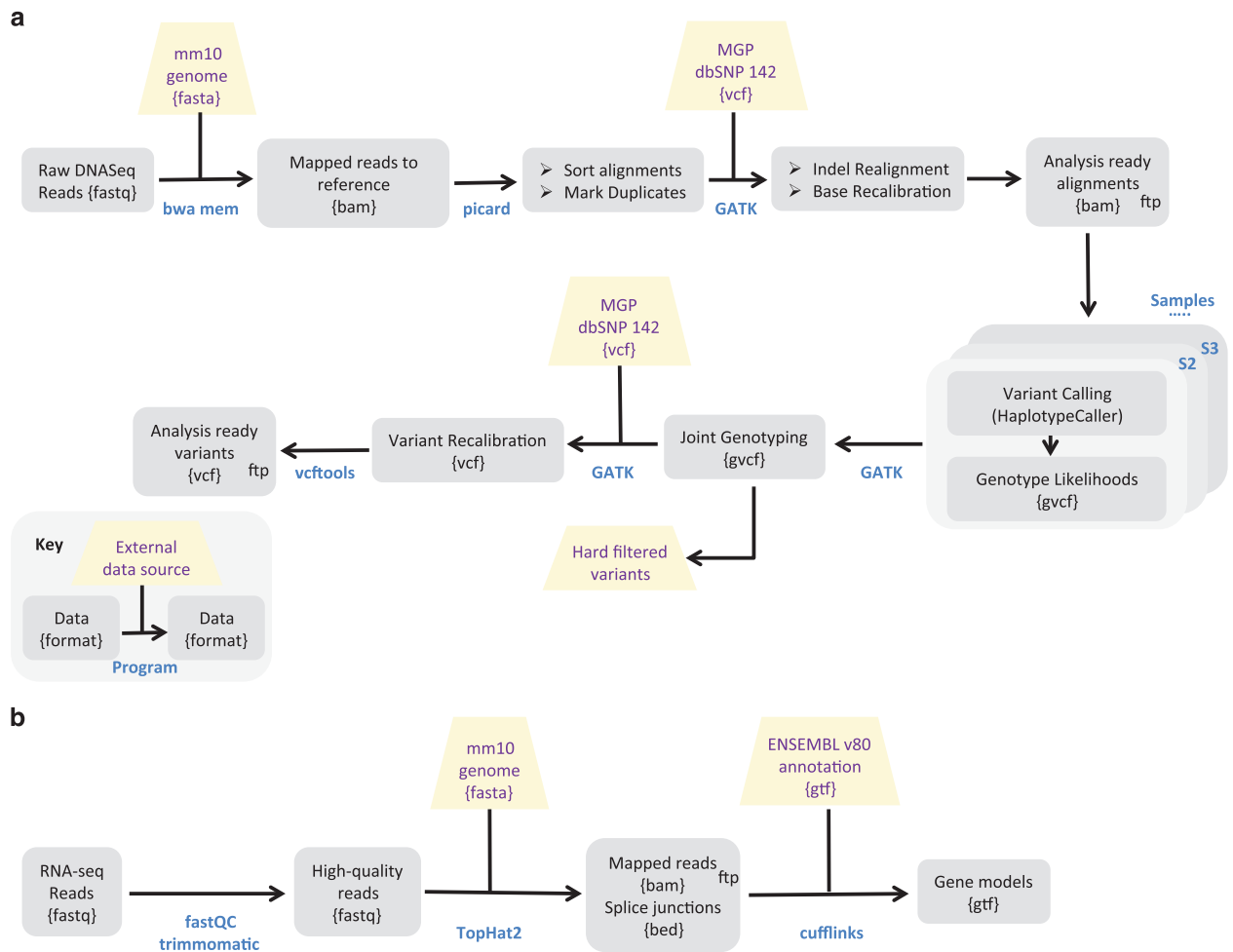


Figure 2. Overview of mapping pipeline for genomic (a) and transcriptomic (b) reads. See Supplementary Material Text 2 for full details. Analysis steps for which files are provided are marked with ‘ftp’.

For whole genome sequencing of batch 1–3 we fragmented 1 μ g of DNA of each individual using the 250 bp sonication protocol (Bioruptor, Diagenode, Liège, Belgium). The fragments were end-repaired and adaptor-ligated, including incorporation of sample index barcodes. The products were then purified and amplified (10 PCR cycles) to create the final libraries. The TruSeq DNA LT Sample Prep Kit v2 was used for all steps. After validation (Agilent 2,200 TapeStation), all libraries were quantified using the Peqlab KAPA Library Quantification Kit and the Applied Biosystems 7900HT Sequence Detection System. One library was loaded on two lanes of a HiSeq2000 sequencer and sequenced with a 2×100 bp v3 protocol.

The DNA quality of the Afghanistan mice (batch 4) proved problematic. Therefore, to recover high molecular weight genomic DNA, we ran a 0.7% agarose gel over night and extracted the genomic DNA from the gel using the ZymoClean Large Fragment DNA Recovery Kit (Zymo Research Europe, Freiburg im Breisgau, Germany). For whole genome sequencing we used the Nextera DNA library Prep Kit (Illumina) following manufacturer’s instructions and 50 ng of genomic DNA as starting material. Each sample was run on Agilent Bioanalyzer using the Agilent DNA7500 kit to verify that the fragment sizes were in the 500 bp range. To calculate the final concentration for the sequencing run, the samples were measured with the Quant-iT dsDNA BR Assay Kit on a Nanodrop 3,300 fluorometer. The samples were paired-end sequenced (76 bp) independently on a single flow cell on a NextSeq 500 using the NextSeq 500 High Output v2 150 cycles chemistry.

RNA sequencing library preparation. We used the NEBNext Ultra RNA Library Prep Kit for Illumina with the Poly(A) mRNA Magnetic Isolation module to generate the RNAseq libraries. We used a fragmentation time of 15 min (yielding ~ 180 bp fragments) and 15 PCR cycles for library enrichment. After validation (Agilent 2,200 TapeStation), all libraries were quantified using the Peqlab KAPA Library Quantification Kit and the Applied Biosystems 7900HT Sequence Detection System. Samples were pooled such that we generated about 12 million paired reads/sample, which were loaded on one lane each of a HiSeq2000 sequencer and sequenced with a 2×100 bp v3 protocol.

Data analysis

Mapping genomic reads and Single Nucleotide Polymorphism (SNP) calling. All sequencing reads (including those of the 10 previously published *M. m. castaneus*⁵¹ genomes) were processed according to a single standardized pipeline, which is outlined in Fig. 2a and described in detail (including commands) in Supplementary Material Text 2. In brief, reads were mapped against the mouse *mm10* genome reference sequence⁵⁴ (<http://www.ncbi.nlm.nih.gov/projects/genome/assembly/grc/mouse/>) using *bwa-mem*⁵⁵. The Picard tools software suite (<http://broadinstitute.github.io/picard/>) was used for sorting, marking and removing duplicates. Raw SNP and indel calls were obtained from the alignment files following precisely the GATK⁵⁶ 'Best Practice' instructions on joint genotyping of all samples together. The raw .vcf files were subjected to the GATK VSQR SNP filtering step, which uses known variants as training data to predict whether a new variant is likely a true positive, or a false positive. As training data we used the file 'mgp.v5.merged.snps_all.dbSNP142.vcf' downloaded from ftp://ftp-mouse.sanger.ac.uk/current_snps/⁵⁷ which was filtered for 'PASS' SNPs. In addition, we used very stringent hard filtering criteria on our own dataset, and included these SNPs as training sets as well (see details in Supplementary Material Text 2). Due to an absence of a reliable indel reference dataset we did not generate VSQR calls for indels. Thus, all indels called in the .vcf file should be considered 'raw'.

On our ftp website we provide a .vcf file with all SNPs and indels where we flag all SNPs within the 90% VSQR tranche as 'PASS' SNPs (this means that we accept all variants until we reach 90% of our known 'truth' set). This is a rather conservative filter on SNP quality, but users are free to perform their own filtering using their own training set and parameters for filtering on the raw .vcf file.

Transcriptome sequencing of *M. m. domesticus* populations. Transcriptome sequencing reads were processed according to the pipeline depicted in Fig. 2b. In short, they were trimmed according to quality values using *Trimmomatic*⁵⁸, removing bases below Q20 and maintaining an average read quality above Q25. Pairs with one read below the quality thresholds were removed from the analyses. Quality-checked (QC) reads were mapped against the mouse *mm10* genome reference using *TopHat2* (ref. 59) and using the default settings for paired-end samples. The output alignments were sorted and indexed with *samtools*⁶⁰. The sorted alignments were assigned to the version 82 of Ensemble Mouse gtf annotation⁶¹ using *featureCounts* from the *subreads* suite⁶² in paired-end (-p) exon mode (default). The gtf file contained only linear complete chromosomes (no scaffolds, no mitochondria). The unmapped pairs were counted with *featureCounts* from the '<unmapped.bam>' file *TopHat2* generates. Percentages are reported relative to the total number of QC reads. On average (across all tissues and all samples) 93% of the total number of QC reads could be mapped to the *mm10* genome (range 86.5–96%, Table 2 (available online only), Supplementary Table 3). This number also includes spliced reads, which *TopHat2* detects. This number dropped to 57% (range 33–66%) for reads uniquely mapping to ENS 82 annotated features (i.e., exons, Supplementary Table 3).

Copy number variation (CNV) analyses. We used the sequencing read depth approach implemented in the *CNVnator* software⁶³ to predict CNV calls relative to the mouse *mm10* reference assembly. We have previously experimentally confirmed that this is a reliable approach³⁸. Optimal bin size for each individual was chosen such that the ratio of the average read depth signal to its standard deviation was between 4 and 5. Bin size ranged from 100–1,500 bp and was inversely proportional to genome coverage. Only linear complete chromosomes were considered. Calls intersecting annotated gaps in the reference genome were not considered. The CNV detection statistics are provided in Table 3 (available online only). Haploid copy numbers for each detected CNV, either per population or per individual, are included in the bed files for the UCSC browser tracks (available at the ftp site).

Visualization of data within and between populations. For the genomic data we set up UCSC⁴⁵ genome browser tracks for 10 kb windows of nucleotide diversity π , Tajima's D and F_{ST} , calculated using *vcftools*⁶⁴, and CNV tracks. The 90% truth VSQR-filtered 'PASS' data were used for all *vcftools* calculations, allowing only bi-allelic SNPs and a maximum of 20% of missing data per SNP. The output tables were converted to bigWig format using the *BigWig* utilities⁶⁵. For the CNV tracks we used *bedtools*⁶⁶ to intersect calls from all individuals belonging to the same population. Within each population average copy number across individuals was calculated for every given interval and transformed to \log_2 values. The genomic browser tracks provided at the ftp site allow a visualization of differences between the populations and species for genomic regions of interest.

For RNAseq data, the coverage (number of reads at a given base pair) is expected to be proportional to the expression level of the gene from which the read originated. In order to compare RNAseq coverage (and thus gene expression) across individuals with slightly varying total numbers of mapped reads, we normalized the data by proportionally sub-sampling reads from the alignment file (*.bam) for the individuals with higher numbers of total mapped reads. Specifically, for each given tissue, we first determined the individual with the lowest number of total mapped reads. This individual is assigned the normalization factor of 1. For each additional individual we calculated the factor *c*, total number of mapped reads in the individual with the lowest number / total number of reads in sampled individual. Values of *c* range from 0 to < 1. Thus, the individual with the highest number of mapped reads will have a value of *c* closest to zero. This normalization factor is then used with the *samtools view -s x.y* command,

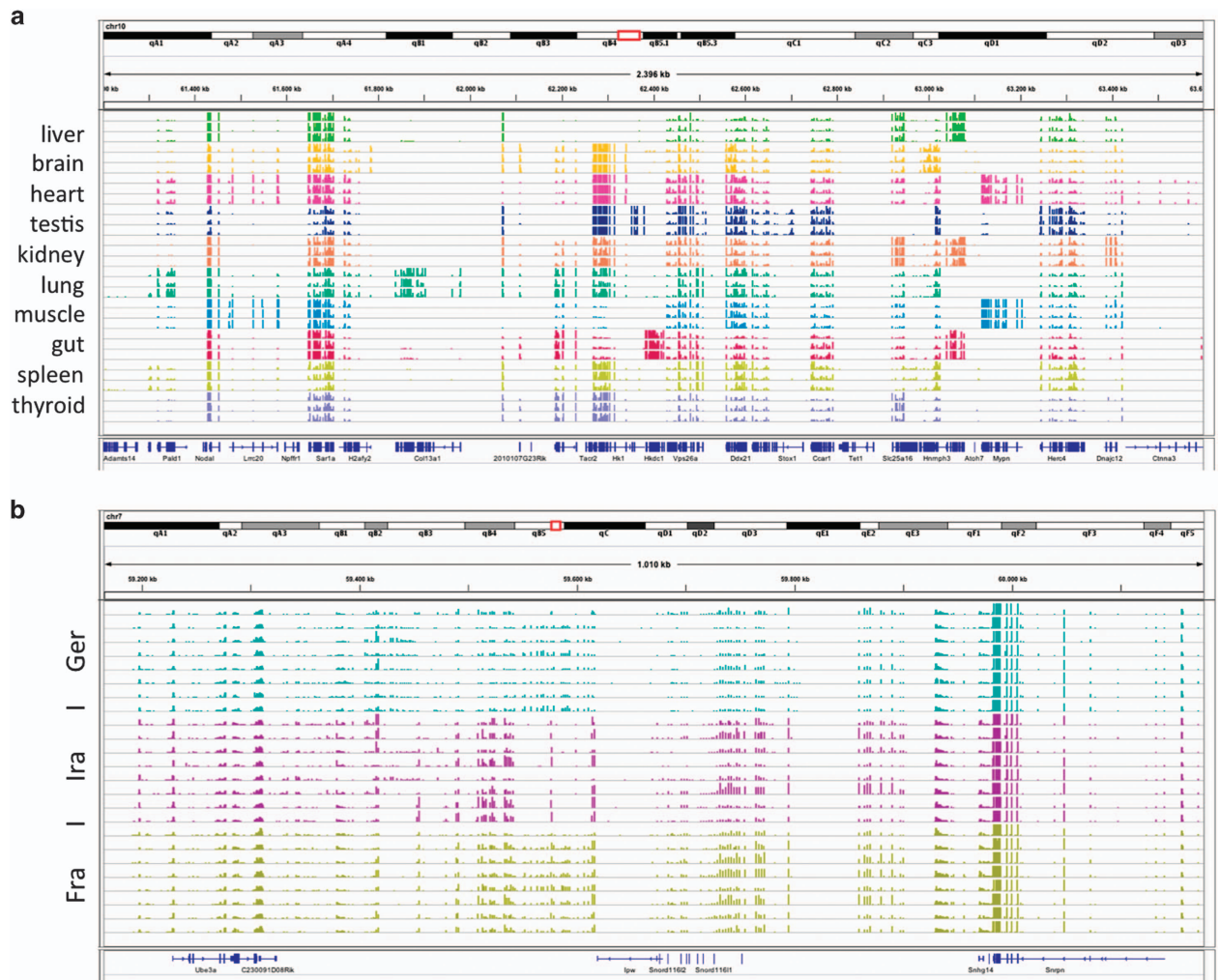


Figure 3. Examples of IGV browser views for the transcriptome data. (a) Region chr10:61,200,000–63,600,000 in the mouse *mm10* genome displaying results for all tissues with the data combined from all individuals of the three *M. m. domesticus* populations. The population order is for each tissue Germany-Iran-France from top. (b) Region chr7:59,165,000–60,177,000 in the *mm10* mouse genome, displaying results for brain normalized RNAseq read coverage with all individuals displayed for each of the three *M. m. domesticus* populations. Note that there is RNAseq read coverage (i.e., ‘gene expression’) between the annotated coding genes, a region corresponding to known non-coding snRNAs located within tandem repeats. This expression is limited to the brain among the tissues sampled. Expression differences between the populations are evident.

where $x=0$ and y =the decimal part of the normalization factor c to generate the normalized alignment file.

To visualize the data as number of reads covering a particular location (i.e., base pair) in the genome, we converted normalized alignment files into bedgraph files, which were further compressed into bigWig format (available on the ftp site). The RNAseq based read coverage for each basepair in the genome can then be visualized in the UCSC (available as public session under ‘wildmouse’) or IGV browsers^{46,47}. Figure 3 shows two examples of screen shots from IGV sessions. The first is a general overview across all tissues from a section of chromosome 10. The second shows only a single tissue (brain), but with read coverage information for each individual.

Data Records

The primary read files for the genome sequences are available at the European Nucleotide Archive (ENA) under project accession number PRJEB9450 (Data Citation 1) for the *M. m. domesticus* genomes, under project accession number PRJEB11742 (Data Citation 2) and PRJEB14167 (Data Citation 3) for the *M. m. musculus* and the *M. spretus* genomes and under project accession number PRJEB2176 (Data Citation 4) for the *M. m. castaneus* genomes processed in this study. All genome samples and their

associated sample designations are listed in Table 1 (available online only). The transcriptome read files are available at ENA under project accession number PRJEB11897 (Data Citation 5). The samples and their sample designation are described in Table 2 (available online only).

The files with the mapped reads (bam), variant calling (vcf) and browser tracks (bigWig) for the genomes and transcriptomes, as well as the IGV session files for the transcriptomes are available at:

<http://wwwuser.gwdg.de/~evolbio/evolgen/wildmouse/> where they can be accessed via ftp.

Technical Validation

We used the software *angsd*⁶⁷ and its `-doDepth 1` command (with options `-minMapQ 30 -minQ 20`) to assess the quality of each DNA sequencing library with respect to good quality coverage (both mapping and base quality) of the genome. The *angsd* depth analysis is based on all sequenced and mapped bases, rather than only on called genotypes at variable sites. Thus, this is the most comprehensive way to assess coverage across the whole genome. The average per-base coverage for each sequenced genome is given in Table 1 (available online only) for autosomes, the X-chromosome and the Y chromosome separately. It was calculated as:

$\frac{\sum_{i=1}^{60} n_i \times i}{\text{genome size}}$ where n_i is number of bases sequenced at depth i and genome size is 2,395,908,738 for autosomes, 163,487,995 for the X chromosome and 88,124,698 for the Y chromosome.

The autosomal coverage is variable across individuals (both within and between sequencing batch) but should be high enough to obtain good quality SNP calls. For males, the X-chromosome coverage is expected to be half of the autosomal coverage, while for females, the coverage should be similar for X chromosome and autosomes. We can use this fact to obtain independent confirmation of the animal's sex recorded in the field. For all but one case, the genomic sex (based on X/autosome coverage ratio) matched the sex determined in the field. Individual AL42 was recorded as male in the field, but its genomic data clearly suggest it was a female. For juvenile wild mice it can sometimes be difficult to accurately determine their sex. The lower X-chromosomal coverage for males indicates that some caution should be taken when using called genotypes for this chromosome for population genetic inferences. Approaches that take the genotype likelihoods into account to estimate parameters may be better suited for the X-chromosomal data (i.e., ref. 67). The Y chromosome is extremely poorly covered (see below).

We also assessed the uniformity of coverage across the genome, aiming at identifying specific regions, where few or no reads could be mapped. We ran the *angsd* `-doDepth 1` command for non-overlapping 100 kb windows recording the average (across individuals within subspecies) % bases covered at >10 reads/individual in 100 kb windows. As shown in Supplementary Fig. 4 using all *M. m. domesticus* individuals as an example, there are some regions in the genome where coverage is low or absent. This pattern was highly correlated in the other subspecies/species (data not shown), suggesting that features of the reference genome (possibly presence of repetitive elements or un-sequenced parts of the reference genome) limit mapping of reads in these regions. Regional variation in coverage is especially striking on the Y-chromosome, where coverage is limited to several short regions within the proximal 10 Mb of the chromosome. This region roughly corresponds to the male-specific short arm of the Y chromosome and its centromere.

Confirmation that mice are naturally inbred

The natural inbreeding status of wild caught mice is expected to be influenced by population history, as well as their tendency to form extended family structures with breeding among relatives^{48,49}. The degree of inbreeding is expected to be higher for small populations and also for populations that recently colonized new habitats, a process which often involves a bottleneck. For the *M. musculus* mice included in this dataset, Iranian, Afghanistan and Indian mice are closest to the center of origin of this species and thus are expected to be least inbred, as such populations are expected to have been large and stable over time. On the other hand, *M. m. helgolandicus* inhabits a very small island and experienced a strong founder event during colonization²⁶. For *Mus spretus*, we do not have a good expectation, as wild populations of this species have never been studied. It is generally assumed that *M. spretus* does not form extended family structures, as these mice are not human commensals but live in fields, hedges and boundaries to forests.

We used a combination of *angsd*⁶⁷ to calculate genotype likelihoods and *ngsF*⁶⁸ to calculate inbreeding coefficients for each individual based on randomly selected 1,000 autosomal 10 kb fragments (see Supplementary Material for scripts). As shown in Fig. 4, our expectations are mostly met, with ancestral populations from India and Iran showing the lowest estimated inbreeding coefficient, while Heligoland individuals are close to representing inbred lines. Surprisingly, the *Mus spretus* individuals from Spain are highly inbred, suggesting that they may have experienced a recent bottleneck.

Confirmation that animals cluster with their respective population

The aim of this analysis was to confirm that each sequenced individual is assigned to its respective population, based on its SNP genotypes. We used the software *NgsAdmix*⁶⁹ on the same randomly selected 1,000 autosomal 10 kb fragments as used in the previous analysis. As before, genotype likelihoods were generated by *angsd*. We ran the *NgsAdmix* software for $K=1$ to $K=9$ (Fig. 5). The likelihood of the

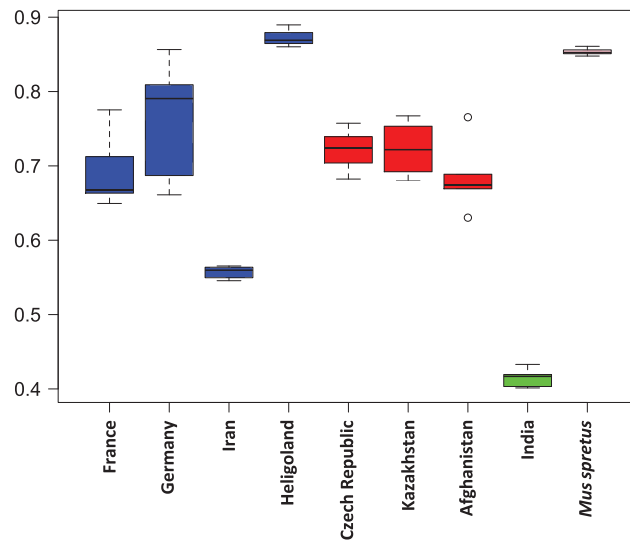


Figure 4. Distribution of inbreeding coefficients within populations. *M. m. domesticus* and *M. m. helgolandicus* populations are highlighted in blue, *M. m. musculus* populations are highlighted in red, the *M. m. castaneus* population is highlighted in green and *Mus spretus* is highlighted in purple.

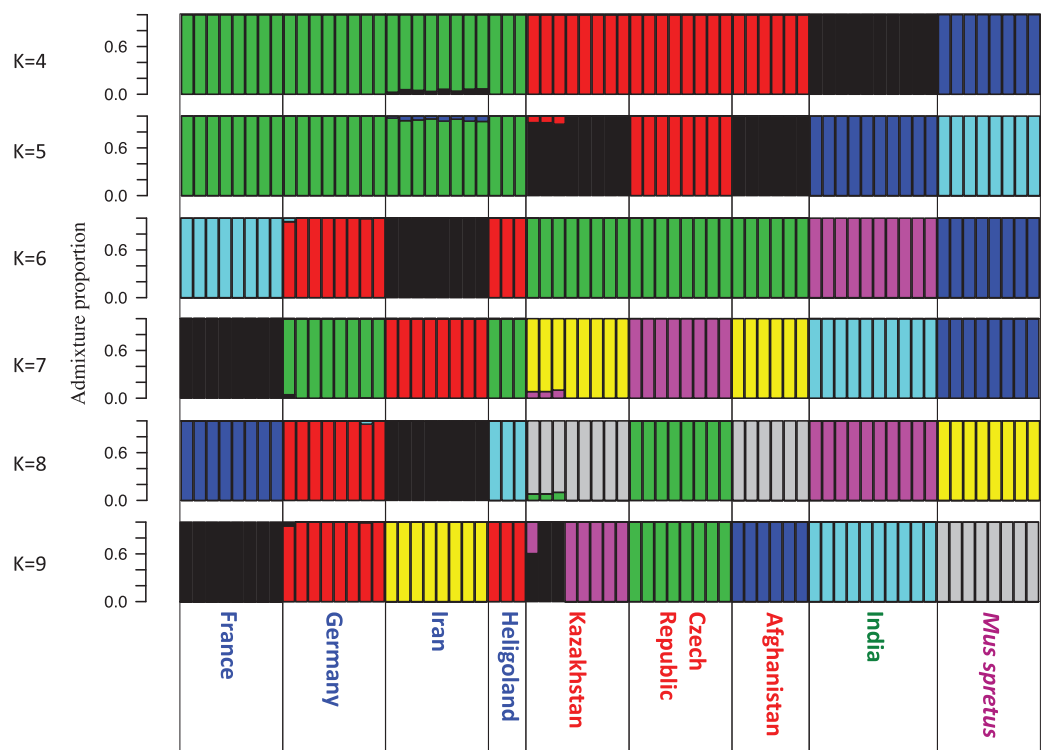


Figure 5. Estimated cluster membership and admixture proportions. Plots for each individual in the sequencing study, for $K=4$ to $K=9$ (number of assumed populations). Individuals are sorted by population and subspecies. *M. m. domesticus* populations are highlighted in blue, *M. m. musculus* populations are highlighted in red, the *M. m. castaneus* population is highlighted in green and *Mus spretus* is highlighted in purple.

data increases dramatically from $K=1$ to $K=4$ and then plateaus (data not shown). Such a pattern (see ref. 70) is usually taken as evidence that $K=4$ fits the data best. $K=4$ clusters all individuals with their correct subspecies (*M. m. helgolandicus* correctly clusters with *M. m. domesticus*²⁶) and species respectively. At $K=7$, we can subdivide the *M. m. domesticus* subspecies into its respective populations

(with *M. m. helgolandicus* clustering with the German individuals, confirming the previous results based on microsatellites²⁶). For the *M. m. musculus* subspecies we find the individuals from the Czech Republic to split off from those from Afghanistan and Kazakhstan. The latter two populations seem to be more closely related. Generally, the genetic clustering analysis suggests that the populations are well defined and differentiated and that there are no recent immigrants from other areas among the sequenced individuals (note that the seeming admixture in $K=9$ is an artifact of too high K).

Confirmation that VSQR 90 tranche filtering yields expected levels of polymorphism

We assessed the GATK VSQR 90% tranche PASS-filtered SNPs for levels of polymorphism (Watterson's θ^{71}) within each population and compared the estimates to several reference data sets, which have previously been generated using Sanger sequencing on smaller number of loci. The Indian population is especially informative, as it has been extensively sequenced^{51,72}. For each chromosome and each population we determined the number of segregating sites using the software *PopGenome*⁷³. Values were summarized over the autosomal genome and converted into Watterson's θ in % by dividing by a_i^{71} and the number of sites sequenced (Supplementary Table 4). The resulting θ_w/bp in % for the Indian population (0.74) lies in between the value obtained by ref. 51 (0.91, for 4-fold degenerated sites and 0.83 for intronic sites) and the one obtained by ref. 72 for the same Indian population (0.664). θ_w for the Western European *M. m. domesticus* individuals was 0.213 in ref. 72, 0.18 for our German population and 0.2 for our French population. Thus, overall, the VSQR 90% PASS-filtered SNPs dataset seem to reflect the previously inferred levels of polymorphism of house mouse populations quite well.

Analysis of relatedness in the sample

We used the `—relatedness2` option of *vcftools* to assess pairwise individual relatedness among all mice in the dataset, using the KING method⁷⁴. This analysis is based on GATK called genotypes and the 90% tranche PASS-filtered SNPs. We restricted the dataset to only include autosomal SNPs, thinned to 1 SNP every 1 Mb. We also removed sites that had more than 20% missing data and only included bi-allelic markers in the analysis. As described in Table 1 of ref. 74, expected ranges of kinship coefficients ('Phi') are >0.354 for duplicate samples/monozygotic twins, $[0.177-0.354]$ for 1st degree relatives, $[0.0884-0.177]$ for 2nd degree relative, $[0.0442-0.0884]$ for 3rd degree relatives and <0.0442 for unrelated samples. Out of 2,211 pairwise individual relatedness estimates, 35 indicated first (10 pairwise comparisons), second (three pairwise comparisons) and third degree (22 pairwise comparisons) relatives (Supplementary Table 5). However, since we detected third degree relationship also among animals that were unequivocally caught far apart (e.g., SP39-SP68), we only consider first and second degree relatedness relevant here. No duplicate samples were detected (expected $\text{Phi} = 0.5$). Relatedness was only detected within populations, and was absent between them. No first or second-degree relatedness was found for the German *M. m. domesticus*, the Afghanistan *M. m. musculus* and the Indian *M. m. castaneus* populations. Most related animals were found in the populations from Iran and Kazakhstan. In the case of the Iranian population the increased relatedness within the sample can be explained by the fact that some breeding adults were used in multiple crosses (see Supplementary Table 2). The relatedness observed in the population from Kazakhstan is best explained by the fact that mice were collected in close proximity, rather than over a larger regional scale.

We can use the known breeding setup of the Iranian mice to confirm the inferred relatedness categories in this population. The KING method detected 2 first-degree relationships in the Iranian population. Male AH15 is indeed the father of JR5-F1C. However, JR-7F1C is the brother of the mother (i.e., uncle) of JR11 and thus a second-degree relative. The second-degree relative identified by the KING method in the Iranian sample is consistent with the breeding scheme, with JR11 and JR15 being half siblings (they have the same father). Two third degree relationships are incorrectly identified and should be second-degree relationships instead (i.e., JR5-F1C is the uncle of JR15 and AH15 is grandfather of JR15) and one third-degree relationship does not have any known breeding history confirming it. The KING method assumes Hardy-Weinberg equilibrium among SNPs with the same underlying allele frequencies. This assumption is most likely being violated in our dataset (given that wild mice are generally inbred, see above), and could explain some of the miss-assignments between categories.

Confirmation of known t-haplotype carriers and identifying t-haplotype carriers in the total dataset

The t-haplotype is complex set of 4 inversions, comprising a 20 cM (30–40 Mbp) region of the proximal third of chromosome 17 in house mice⁷⁵. It is a selfish genetic element, which causes transmission ratio distortion, with heterozygous t-haplotype carriers predominantly (sometimes up to 99% of times) transmitting the t-haplotype carrying chromosome to their offspring. Homozygous individuals for the t-haplotype, however, die *in utero*. Despite their massive transmission advantage, t-haplotype carrying individuals are rare in natural populations of mice, but have been found in all recognized subspecies.

We have previously used two published primer pairs⁷⁶ to genotype individuals in our collections of mouse samples for presence/absence of the t-haplotype. Both primer pairs span t-haplotype diagnostic indels that can be analyzed on an agarose gel (the proximal locus *Tcp1* spans a 175 bp indel and the distal locus *Hba-4ps* spans a 16 bp indel). Three individuals typed with those primers are identical to samples included in the whole genome sequencing study described here: male AL41, male CR16 and female H14.

Of those, AL41 and H14 were found to be carriers (heterozygous) of the t-haplotype, while CR16 was wildtype. We use ENSEMBL Blast to determine the location of those primers in the *mm10* reference sequence. All primers yielded unique hits. We then extracted all indels spanning the region between the primer locations for all samples from the .vcf-file, and searched for (combinations of) indels matching the sizes above. For the distal locus, we found a single 17 bp indel at position chr17:26,286,509 ('TACTACTATGCACTGAA'). For the proximal locus, we found one indel at position chr17:12,921,682 that was identified as 'GTTTTTTTTTTTTT'. Illumina sequencing is not capable to sequence through long homo-polymer stretches which is likely the reason why the identified indel is reported shorter than the expected 175 bp. However, it is the only indel >3 bp in the region and moreover, genotypes at this indel are in almost perfect linkage disequilibrium with genotypes at the distal locus (see Table 1 (available online only)), which is highly unexpected over a distance of 13.3 Mb. The two individuals that we previously found to be positive for the t-haplotype using the PCR primers are also heterozygous for the respective indels in the whole genome dataset, while CR16 was wildtype based on the whole genome sequence. Thus, indels at chr17:12,921,682 and chr17:26,286,509 were used to genotype the remaining individuals in the dataset for the presence/absence of t-haplotypes. All *M. musculus* individuals positive for the t-haplotype indels were heterozygous for that t-allele. *Mus spretus*, on the other hand, was homozygous for the proximal t-specific indel, which is consistent with a rather old origin of the t-allele (1–3 Myr⁷⁷), despite not having any obvious transmission ratio distortion properties in a species outside the *M. musculus* complex. *t*⁺/*wt* individuals were found in every population apart from the Iranian population and among the three individuals from Heligoland. T-haplotype carriers reached frequencies of 50% in two *M. musculus* populations (Afghanistan and Czech Republic). 37.5% of the French individuals carried the t-haplotype, as did 30% of Indian individuals, 25% of individuals from Kazakhstan and 12.5% of the German population. The frequency of t-haplotypes in our data is somewhat higher than reported previously (reviewed in ref. 78), however well below the theoretical expectations based on transmission ratio distortion (see ref. 79).

K-mer distributions to determine complexity of RNAseq libraries

We analyzed the *k*-mer (DNA sequence stretch of length *k*) frequency spectrum to identify potential problems with the RNAseq libraries, such as DNA contamination and overabundance (potentially due to PCR amplification) of particular sequence stretches. Ideally, libraries are complex, meaning they exhibit great diversity in unique sequences and repeated structures⁸⁰ and there should be no sign of DNA contamination among the RNA based reads generated. In total, we generated 224 RNAseq libraries from up to 10 tissues in 24 *M. m. domesticus* individuals. For each library, we ran the software *jellyfish*⁸¹ on all forward reads with a *k*-mer length of *k*=12. Using reverse reads yielded the same results (data not shown). For visualization in Supplementary Fig. 5 we randomly choose three individuals from each population and three tissues (brain, testis and liver). The complete set of all 224 RNAseq *k*-mer distributions are available on our ftp server. The *k*-mer distribution of the mouse *mm10* DNA sequence (red in Supplementary Fig. 5) produces a characteristic line with 2 prominent humps when plotted on a log₁₀–log₁₀ scale. The *k*-mer distribution for annotated cDNAs in the mouse genome (ENSEMBL version 83 (ref. 61)) does not produce such humps and compared to the DNA *mm10* sequence is more 'complex', i.e., shows proportionally more unique sequences (with low 12-mer occurrence, but high frequency, left side of plots in Supplementary Fig. 5). The *k*-mer distribution of the RNAseq forward reads (grey in Supplementary Fig. 5) falls in between the cDNA and genomic DNA profile. Since we generated our RNAseq libraries from poly-adenylated RNAs, comparing the RNAseq profile to annotated cDNA seems appropriate. Most notably, the RNAseq *k*-mer profile lacks the characteristic humps of the genomic DNA *k*-mer profile, suggesting that the RNA libraries are not contaminated with DNA. Moreover, the diversity of unique sequences matched the cDNA profile much better than the DNA profile. Very similar patterns have been observed for human RNAseq data, that have been deemed good quality (see Supplementary Fig. 6A in ref. 80).

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Data Citations

1. *European Nucleotide Archive* PRJEB9450 (2016).
2. *European Nucleotide Archive* PRJEB11742 (2016).
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Author Contributions

B.H. contributed to study design, collected mice, prepared samples, established the genomic mapping pipeline, analyzed DNaseq data, and wrote the manuscript. E.K. contributed to and ran the genomic mapping pipelines. R.N. prepared samples, generated and ran the RNAseq mapping pipelines and analyzed RNAseq data. M.T. contributed to study design, collected mice and prepared samples. C.P. collected mice and developed the wild mouse keeping procedures. Z.P. conducted the copy number

analyses. H.B. collected mice and prepared samples. M.L. collected mice and prepared samples. I.M. collected mice and prepared samples. R.S. collected mice and prepared samples. M.A. collected mice. M.P.M. collected mice. M.S. collected mice. A.B. prepared RNA samples. R.G.U. collected mice. J.A. conducted the sequencing. M.F. conducted the sequencing. S.K. conducted the sequencing. D.T. organized the study, wrote the manuscript, with input from the co-authors.

Additional Information

Tables 1–3 are only available in the online version of this paper.

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