

On the subspecific origin of the laboratory mouse

Hyuna Yang¹, Timothy A Bell², Gary A Churchill¹ & Fernando Pardo-Manuel de Villena²

The genome of the laboratory mouse is thought to be a mosaic of regions with distinct subspecific origins. We have developed a high-resolution map of the origin of the laboratory mouse by generating 25,400 phylogenetic trees at 100-kb intervals spanning the genome. On average, 92% of the genome is of *Mus musculus domesticus* origin, and the distribution of diversity is markedly nonrandom among the chromosomes. There are large regions of extremely low diversity, which represent blind spots for studies of natural variation and complex traits, and hot spots of diversity. In contrast with the mosaic model, we found that most of the genome has intermediate levels of variation of intrasubspecific origin. Finally, mouse strains derived from the wild that are supposed to represent different mouse subspecies show substantial intersubspecific introgression, which has strong implications for evolutionary studies that assume these are pure representatives of a given subspecies.

Laboratory mice, the most popular model organism in mammalian genetics^{1,2}, were derived from wild mice belonging to the *Mus musculus* species by an intricate process that included the generation of ‘fancy’ mice in both Asia and Europe and a complex web of relationships between inbred strains³. Early studies showed that the mitochondria and the Y chromosome present in many classical laboratory strains were derived from different subspecies, *M. m. domesticus* for the mitochondria and *M. m. musculus* for the Y chromosome^{4,5}. Furthermore, the Y chromosome was introduced into the laboratory mouse through *M. m. molossinus* males⁶. Based on these findings, it was proposed that the genomes of inbred strains are a mosaic of regions that have different subspecific origins⁷. Recently, the fine structure of such mosaic variation was described⁸. This study reported that strain-to-strain comparisons reveal regions with extremely high variation that span one-third of the genome and regions with extremely low variation that cover the remaining two-thirds of the genome. This distinctively bimodal distribution was assumed to represent regions that have different, or the same, subspecific origins, respectively. This mosaic model has been the driving concept behind mouse association mapping studies and haplotype analysis^{9–12}. However, the origin of a given region of a laboratory strain could not be directly assigned to a subspecies owing to the lack of reference sequences for the three main mouse subspecies. Subsequent studies raised questions about the haplotype structure^{11,13}, the effect of ascertainment biases in subspecific assignment^{14–16} and the contributions of intersubspecific versus intrasubspecific variation¹⁷. Several studies reported the presence of substantial intrasubspecific variation, ancestral polymorphisms and secondary introgression after the divergence of the subspecies^{17–20}, further complicating the interpretation of the data.

In 2004, the National Institute of Environmental Health Sciences (NIEHS) contracted Perlegen Sciences to resequence 15 mouse inbred

strains. This project has released more than 109 million genotypes for 8.3 million SNPs that span the nuclear and mitochondrial genomes²¹. The 15 strains were selected on the basis of their genetic diversity, ease of breeding, inclusion in the Mouse Phenome Project, widespread use in research and background information. This set includes 11 classical strains (129S1/SvImJ, A/J, AKR/J, BALB/cBy, C3H/HeJ, DBA/2J, FVB/NJ, NOD/LtJ, BTBR *T⁺ tfJ*, KK/HIJ and NZW/LacJ) and four strains derived from the wild (hereafter ‘wild-derived’ strains) (WSB/EiJ, PWD/PhJ, CAST/EiJ and MOLF/EiJ), which represent the *M. m. domesticus*, *M. m. musculus*, *M. m. castaneus* and *M. m. molossinus* subspecies, in corresponding order^{22–25} (<http://www.jax.org>). *M. m. molossinus* is a subspecies that arose by natural hybridization between *M. m. musculus* and *M. m. castaneus*. The data are hereafter referred to as the NIEHS data.

We set out to use this resource to examine the ancestral subspecific origin of classical strains, expecting to identify a mosaic of segments that could be assigned to one of three distinct lineages: *M. m. domesticus*, *M. m. musculus* and *M. m. castaneus*^{8,26}. We planned to use the three wild-derived strains as a reference for each subspecies and then assign genomic segments from classical strains to a subspecies, based on the pattern of SNP similarity between the query strain and the reference strains.

RESULTS

Diagnostic SNPs

Evolutionary models suggest that the three main mouse subspecies diverged simultaneously from a common ancestor or, alternatively, that *M. m. musculus* and *M. m. castaneus* diverged from a common ancestor shortly after the divergence of *M. m. domesticus*^{27–29}. This history should be reflected in the distribution of SNPs that are specific to each subspecies. SNPs that have arisen since the divergence of the three subspecies should be equal in number or alternatively, be slightly

¹The Jackson Laboratory, Bar Harbor, Maine 04609, USA. ²Department of Genetics, Carolina Center for Genome Sciences and Lineberger Comprehensive Cancer Center, University of North Carolina, Chapel Hill, North Carolina 27599, USA. Correspondence should be addressed to F.P.-M.d.v. (fernando@med.unc.edu).

Received 13 February; accepted 31 May; published online 29 July 2007; doi:10.1038/ng2087

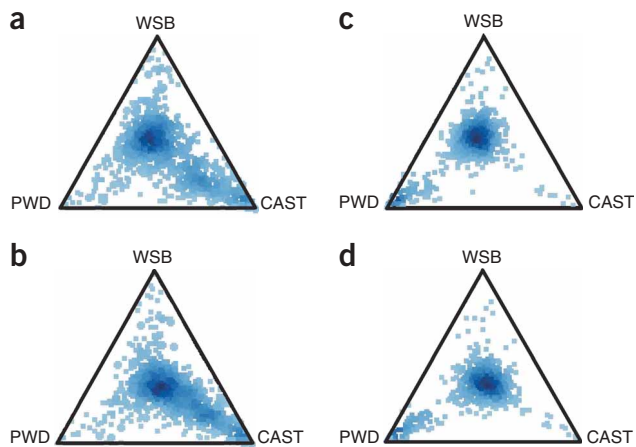


Figure 1 Frequency distribution of diagnostic subspecific SNPs. The relative frequency of diagnostic SNPs in 100-kb intervals is represented as a density plot over the simplex. In each plot, the three reference strains are indicated at the vertices of a triangle, and the relative proportions of diagnostic SNPs in each interval are represented as blue dots. Darker areas represent regions with a higher density of intervals. (**a,c**) Data for chromosome 14. (**b,d**) Data for chromosome X. **a** and **b** show the original data; **c** and **d** show the predicted proportions of diagnostic SNPs after correcting for the frequency-dependent SNP discovery rate.

enriched for *M. m. domesticus* SNPs, and these SNPs should be distributed evenly throughout the genome.

For the purpose of interpreting the NIEHS data, we define diagnostic SNPs as those that are completely genotyped and polymorphic among the three reference strains, WSB, PWD and CAST. Note that the diagnostic allele at some of these SNPs may not be shared by all individuals of that subspecies if it arose recently. Furthermore, because of incomplete sorting or homoplasy, the allele can also be present in individuals of other subspecies. Despite the limitations of using a single reference strain to define diagnostic SNPs, it remains the simplest method to test our expectations on the basis of phylogenetic history. We identified 4,373,427 diagnostic SNPs: 1,481,373 (33.9%) are *M. m. domesticus* SNPs that distinguish WSB from CAST and PWD; 1,280,328 (29.3%) are *M. m. castaneus* SNPs that distinguish CAST from WSB and PWD; and 1,611,726 (36.9%) are *M. m. musculus* SNPs that distinguish PWD from WSB and CAST.

We divided the genome into nonoverlapping 100-kb intervals and determined the proportion of diagnostic SNPs for each subspecies in each interval. These proportions can be represented in a simplex, a triangular region of three-dimensional space that represents the proportions of the three types of diagnostic SNP. In this representation, an interval that contains equal numbers of the three types of diagnostic SNP is located at the center, whereas an interval that contains only one type of diagnostic SNP is located at the corresponding vertex (Fig. 1). In contrast to our expectations, we found that most intervals are not located at the center but consistently deviate away from the CAST vertex (Fig. 1a,b). The degree of distortion varies among chromosomes, but this observation holds true for all autosomes and the X chromosome (Supplementary Fig. 1 online). Although a slight deviation toward WSB (*M. m. domesticus*) is predicted by one evolutionary model²⁹, a genome-wide deficit of diagnostic *M. m. castaneus* SNPs can be explained only by either a differential mutation rate in that subspecies, or a systemic undercounting of diagnostic CAST SNPs across the genome.

We also found many intervals with extremely distorted frequencies of diagnostic SNPs (Fig. 1). The pattern of extreme distortion varies among chromosomes (Supplementary Fig. 1). The two most common patterns are intervals that have an excess of *M. m. castaneus*

SNPs (intervals that are located close to the CAST vertex; Fig. 1a) and intervals that have an excess of *M. m. musculus* SNPs (intervals that are located close to the PWD vertex; Fig. 1b). Remarkably, the low-level distortion against *M. m. castaneus* SNPs in many intervals exists on the same chromosome with extreme distortion in favor of *M. m. castaneus* SNPs in other intervals. This inconsistency suggests that low-level distortion and extreme distortion have different origins.

SNP ascertainment bias

The basic properties of the NIEHS data are provided in a **Supplementary Note** online. We determined the false-positive rate (FPR) and false-negative rate (FNR) in the NIEHS data set by direct resequencing of selected genomic fragments in the 15 NIEHS strains and the C57BL/6 strain (Methods). The FPR is 1.3%, which is similar to previously reported results^{11,30}. We found six discordant genotypes between our data and the NIEHS data (0.3%) among the 2,089 genotypes compared. Therefore, the FPR is low and should have little impact on the distorted frequency of diagnostic SNPs that are observed in the reference strains. By contrast, the FNR is significantly higher than previously reported in humans³⁰. Because Perlegen's SNP discovery algorithm was designed to minimize the FPR, a high FNR is expected.

The FNR is strongly correlated with the minor allele frequency (MAF). The number of undetected SNPs decreases as the MAF increases from 76% for singletons (SNPs in which the minor allele is present in a single strain) to 42% for SNPs in which the minor allele is shared by seven strains (Fig. 2a). Among singletons, the FNR is constant with respect to the genomic position and strain (Fig. 2b,c), which suggests that the MAF is directly responsible for the pronounced differences in FNR. The local FNR varies across the genome, depending on the MAFs of SNPs that are present in a given region, which in turn depends on the phylogenetic relationships between the strains in that region. We estimate that the average genome-wide FNR in the 15 resequenced strains is 67%, based on the distribution of MAFs among the 3.8 million completely genotyped SNPs and the experimental FNR for each MAF. Based on that FNR and the genome

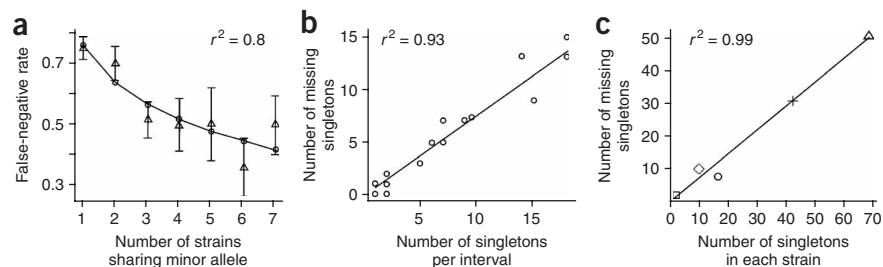


Figure 2 SNP discovery bias. (**a**) Effect of the minor allele frequency (MAF) on the false-negative rate (FNR). Triangles and vertical lines represent observed values that are ± 1 s.e.m. Circles represent the best fit of the data to the regression of $\log(\text{FNR})$ on MAF. (**b**) The FNR for singletons in 24 resequenced intervals distributed across the genome. (**c**) The FNR for singletons from different strains (square, classical strains; open diamond, WSB; circle, PWD; cross, MOLF; triangle, CAST).

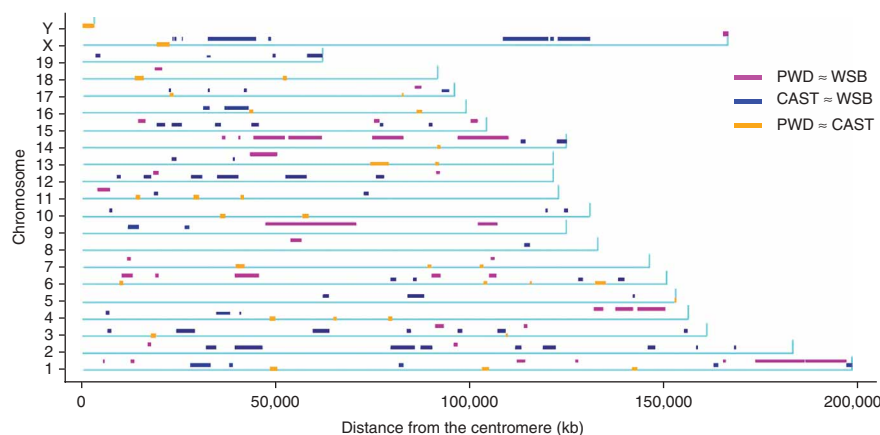


Figure 3 Regions of intersubspecific introgression in the reference strains. Inferred regions of intersubspecific introgression after smoothing the intervals with a hidden Markov model (HMM). Purple denotes regions with an excess of CAST diagnostic SNPs and a deficit of both WSB and PWD diagnostic SNPs. Blue denotes regions with an excess of PWD diagnostic SNPs and a deficit of both WSB and CAST diagnostic SNPs. Orange denotes regions with an excess of WSB diagnostic SNPs and a deficit of both CAST and PWD diagnostic SNPs.

coverage, we estimate that there are 45 million SNPs among the 15 resequenced strains. This number is within the range predicted by direct resequencing studies¹⁸. In conclusion, despite its exceptional size, density and quality, the NIEHS data capture only a fraction of the variation that is present in the laboratory mouse.

The finding that the FNR depends on the MAF implies that the probability of observing each type of diagnostic SNP depends on the local phylogenetic relationships between the 15 NIEHS strains. Furthermore, the MAFs of diagnostic SNPs vary among subspecies: singletons are predominantly CAST SNPs, doubletons are predominantly PWD and SNPs with higher frequencies are predominantly WSB (Supplementary Fig. 2 online). To account for the allele-frequency-dependent FNR, we applied a branch-length correction (Methods) to phylogenetic trees. We then plotted the corrected length of the branches that represent each type of diagnostic SNP in each interval (the distance between the common node and the three reference strains CAST, PWD and WSB) and found that most intervals shifted toward the center of the simplex (Fig. 1 and Supplementary Fig. 1). Therefore, the genome-wide low-grade distortion is due to a frequency-dependent SNP discovery rate that undercounts SNPs from lineages that are locally underrepresented among the NIEHS strains.

Intersubspecific introgression

Correcting for a frequency-dependent FNR had little effect on the intervals with extreme distortion (Fig. 1 and Supplementary Fig. 1). These intervals are not SNP-poor and, therefore, are more prone to statistical fluctuations. Furthermore, intervals with an excess of diagnostic CAST SNPs or intervals with an excess of diagnostic PWD SNPs cluster in different megabase-long regions on particular chromosomes (Fig. 3). These features indicate that the distorted patterns are due to introgression of haplotypes from a different subspecies in one, or more, of the reference strains, which in turn indicates that the three wild-derived strains may not be pure representatives of each subspecies. Intersubspecific introgression has been reported in wild mice and in wild-derived strains^{16,18–20}. Furthermore, MOLF, a wild-derived strain that is considered to be a representative of *M. m. molossinus*, carries an *M. m. domesticus* Y chromosome (Supplementary Fig. 3 online). We conclude that

MOLF also has introgressed haplotypes from a subspecies that is inconsistent with its phylogenetic history.

To delineate regions of the genome in which the reference strains accurately represent the three main subspecies, we first identified intervals with extreme distortion in favor of a single type of diagnostic SNP (Fig. 1c,d and Supplementary Fig. 4 online) and applied a hidden Markov model (HMM) to consolidate larger regions that have a high concentration of unbalanced intervals of the same type. The HMM eliminates unbalanced intervals that lack local support. We left the status of these intervals undetermined. Balanced intervals within unbalanced regions are assigned by the HMM to an introgression class (Fig. 3). The remaining intervals with balanced frequencies of diagnostic SNPs and good local support span 72% of the mouse genome (Supplementary Fig. 4). These intervals probably represent regions in which the three reference strains are true representatives

of the *domesticus*, *musculus* and *castaneus* mouse lineages. In summary, we partitioned the mouse genome into three classes: regions of potential introgression (13% of the genome; Fig. 3), regions with undetermined status (15%) and regions in which the three reference strains provide a balanced representation of the three main subspecies.

Approximately 5.7% of the genome has an excess of diagnostic CAST SNPs and a deficit of diagnostic SNPs for PWD and WSB (shown in purple in Fig. 3). In 5.9% of the genome there is an excess of diagnostic PWD SNPs and a deficit of CAST and WSB diagnostic SNPs (shown in blue in Fig. 3). The third pattern represented by an excess of WSB diagnostic SNPs (shown in orange in Fig. 3) is found in small regions spanning 1.3% of the genome, including the Y chromosome, and is consistent with the hypothesis that *M. m. musculus* and *M. m. castaneus* are sister subspecies²⁹.

We confirmed that, in the regions of potential introgression (Fig. 3), one of the three reference strains carries a haplotype from a different subspecies by sequencing short intervals in the three reference strains and in six additional wild-derived strains (Supplementary Note). These experiments confirm that most regions with extreme distortion in the frequency of diagnostic SNPs (Fig. 1) are due to introgression of *M. m. domesticus* haplotypes into PWD and CAST. Remarkably, some of the introgressed haplotypes span dozens of megabases and are unequally distributed along the genome. For example, there is an excess of *M. m. domesticus* introgression on chromosomes 14 and 9 in PWD and on the X chromosome in CAST (Fig. 3). Other wild-derived strains (CASA and PWK) may also have introgressed haplotypes from *M. m. domesticus*.

In regions of introgression we cannot directly determine the subspecific origin of classical strains. This shortcoming can be addressed by analyzing additional wild-derived strains. Previous conclusions in mouse on effective population sizes and on the rate of variants that are inconsistent with phylogeny owing to incomplete lineage sorting and homoplasy need to be re-evaluated^{18,31}. Wild-derived inbred strains have previously been used to study the genetics of speciation^{32–35}. Although our findings should not affect the hybrid sterility genes mapped using this approach, they may compromise the general conclusions that have been made about the genetic architecture of this critical process^{14,32}.

Ancestry of classical strains

To assign subspecific origins to genomic intervals of the 11 NIEHS classical strains, MOLF and C57BL/6, we examined the bias-corrected phylogenetic trees. In regions where diagnostic SNP frequencies are balanced, we assumed that the root of each tree was located at the node of common ancestry between PWD, CAST and WSB. Splitting the tree at this node partitions each of the remaining strains into one of three groups according to its local subspecific origin.

We first determined the matrilineal (mitochondria) and patrilineal (Y chromosome) inheritance patterns. We confirmed that the classical strains share an almost identical mitochondrial haplotype of *M. m. domesticus* origin (Supplementary Fig. 3), which supports the contention that laboratory strains descend from a very small pool of founders^{4,36}. As expected from studies on mitochondrial variation in *M. m. molossinus*²³, MOLF has an *M. m. musculus* haplotype that is significantly different from the one carried by PWD.

Our analysis confirms the prevalence of the *M. m. musculus* (*molossinus*) Y chromosome among classical strains^{6,37}, and also indicates that many strains (FVB, NOD, BTBR and AKR) carry an *M. m. domesticus* Y chromosome. Interestingly, the *M. m. molossinus* strain MOLF carries an *M. m. domesticus* Y chromosome. Flow of mitochondrial DNA across subspecies boundaries³⁸ and discordant phylogenetic patterns between mitochondria and the Y chromosome have been reported in wild populations¹⁹, which indicates that secondary introgression after radiation of the subspecies²⁰ might

have contributed to the pattern of intersubspecific introgression observed in the wild-derived inbred strains, in addition to accidental 'contamination' in the laboratory.

We extended our assignment of the subspecific origin to the 72% of the autosomal and X-chromosomal genomic intervals for which the ancestry of the reference strains is unambiguous. Figure 4 and Supplementary Figure 5 online show the results for four representative chromosomes in which black denotes *M. m. domesticus* intervals, red denotes *M. m. musculus* intervals and green denotes *M. m. castaneus* intervals. In most regions, the subspecific assignments remain stable along a substantial length of the chromosome. This is particularly notable, given that the assignment was carried out automatically without any further attempt to smooth local fluctuations. Small, isolated segments of distinct subspecific origin do occur (Fig. 4a) and in some cases cluster in specific regions of the genome (Supplementary Fig. 5). The 100-kb interval size selected for our analysis may result in intervals that span transition zones between regions that have different subspecific origins, some intervals may contain smaller segments from a different subspecific origin embedded within them, and segmental duplications and copy number polymorphisms^{39–41} may lead to unusual patterns in the assignment of ancestry. Although the subspecific assignment for each classical strain and MOLF is well supported by bootstrap replicates (85 out of 100 replicates in 94.5% of intervals and 99 out of 100 replicates in 86.3% of intervals), the ancestral origin may be

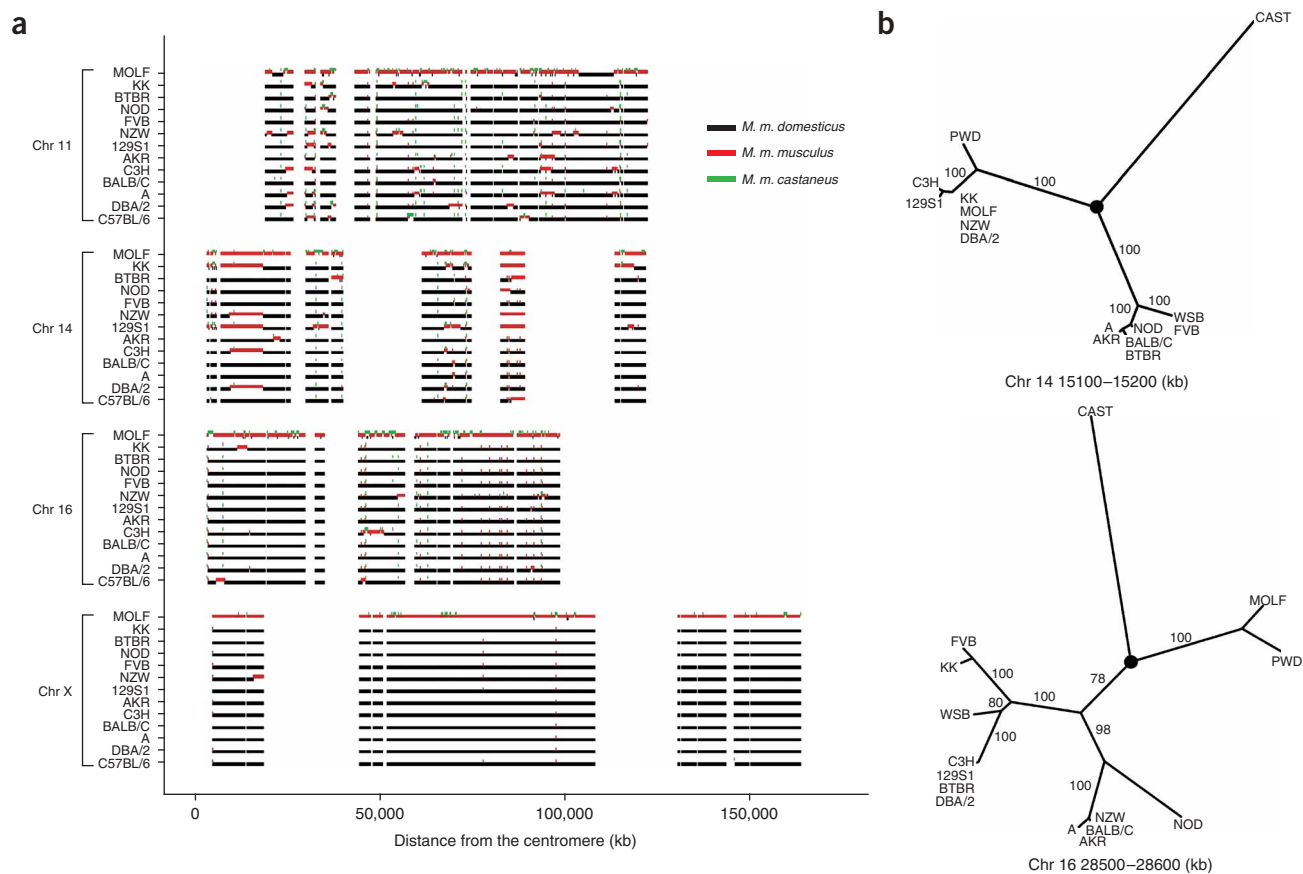


Figure 4 Subspecific origin of classical and hybrid strains. (a) Subspecific assignments for the 12 classical strains (including C57BL/6) for chromosomes 11, 14, 16 and X. Each 100-kb interval is shown as a vertical bar of a color that reflects its subspecific origin. Intervals without color are regions of intersubspecific introgression or of undetermined status. (b) Corrected phylogenetic trees for two 100-kb intervals in chromosomes 14 and 16. The circle denotes the assumed location of the root. Numbers represent bootstrap replicates that support each node.

Table 1 Contribution of the three main subspecific lineages to the genome of the laboratory strains

	B6	DBA/2	A	BALB/C	C3H	AKR	129S1	NZW	FVB	NOD	BTBR	KK	MOLF
<i>M. m. domesticus</i>	0.92	0.91	0.94	0.95	0.92	0.94	0.91	0.87	0.96	0.93	0.92	0.86	0.11
<i>M. m. musculus</i>	0.07	0.07	0.05	0.04	0.07	0.05	0.08	0.11	0.03	0.06	0.06	0.12	0.74
<i>M. m. castaneus</i>	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.02	0.01	0.02	0.02	0.02	0.15

Fractions of balanced 100-kb intervals assigned to each subspecies are shown.

incorrectly inferred in some intervals owing to limitations of the data and methodology.

The genomes of classical strains are overwhelmingly of *M. m. domesticus* origin (Table 1). Although a predominant contribution of that subspecies was predicted⁸, the exceptionally high levels (ranging from 86% to 96%) observed in all strains were unexpected. The *M. m. musculus* subspecies has the second largest contribution to the genome of classical strains, whereas only 1–2% of their genome derives from *M. m. castaneus*. The spatial distribution of subspecific origins and the contribution of subspecies other than *M. m. domesticus* to the genomes of classical strains are not random. On chromosomes 16 and X, the classical strains are largely of *M. m. domesticus* origin, whereas MOLF is of *M. m. musculus* origin (Fig. 4a). By contrast, the classical strains have a more equilibrated contribution of *M. m. domesticus* and *M. m. musculus* on chromosome 14. Finally, chromosome 11 shows an intermediate situation. Many regions of *M. m. musculus* origin are shared among multiple classical strains, which suggests that the history and close relationships between strains has played a part in shaping the distribution of subspecific diversity.

Our analysis also confirms the presence of extensive introgression of *M. m. domesticus* haplotypes in MOLF (for example, distal chromosome 11; Fig. 4 and Supplementary Fig. 5). The genome of this strain is a mosaic of three subspecies with 74% of *M. m. musculus* origin, 15% of *M. m. castaneus* origin and 11% of *M. m. domesticus* origin (Table 1).

Once the subspecific assignment of classical and hybrid strains was completed, we estimated the specificity and sensitivity of diagnostic SNPs. Remarkably, 88.7% of the 3,220,959 diagnostic SNPs that can be tested are completely specific; in other words, the diagnostic allele is not present among any of the NIEHS strains that have a different subspecific assignment. Conversely, the allele present at diagnostic SNPs is shared by all strains assigned to that subspecies in 59% of the 2,354,446 diagnostic SNPs in which this test can be carried out. Thus, despite the limited number of reference strains, diagnostic SNPs identified under our definition can be used collectively to assign subspecific origin.

Detailed images of regions with intersubspecific introgression, their subspecific ancestry and the supporting phylogenetic trees are available at the following website: <http://www.genomedynamics.org>.

Genetic variation in classical strains

In contrast with previous analyses^{8–11,17}, we have determined that, on average, 9% of the genome has a different subspecific origin between any given pair of classical strains, whereas 91% of the genome shares the same subspecific origin (Fig. 5). This indicates that many of the regions with high variation identified in previous studies might have the same subspecific origin. The deep branching

of the *M. m. domesticus* lineage in many of the phylogenetic trees supports the model of high intrasubspecific variation (Fig. 4b).

To investigate the extent of genetic variation within versus between subspecies, we measured the pairwise distances along the bias-corrected trees in each 100-kb interval. These distances were then normalized to the average distance between pairs of strains from different subspecies in that interval. This normalized measure of variation allows direct comparisons between genomic regions that have different coverage, gene density and mutation rates, and makes it possible to calculate the normalized variation for all 25,400 100-kb intervals, regardless of the distribution of diagnostic SNPs. The distribution of within versus between subspecies variation (Fig. 5) demonstrates that our subspecific assignments that are based on tree topology are correct for the vast majority of intervals because pairs of classical strains thought to inherit segments from different species that are based on tree topology (Fig. 4) show a unimodal frequency

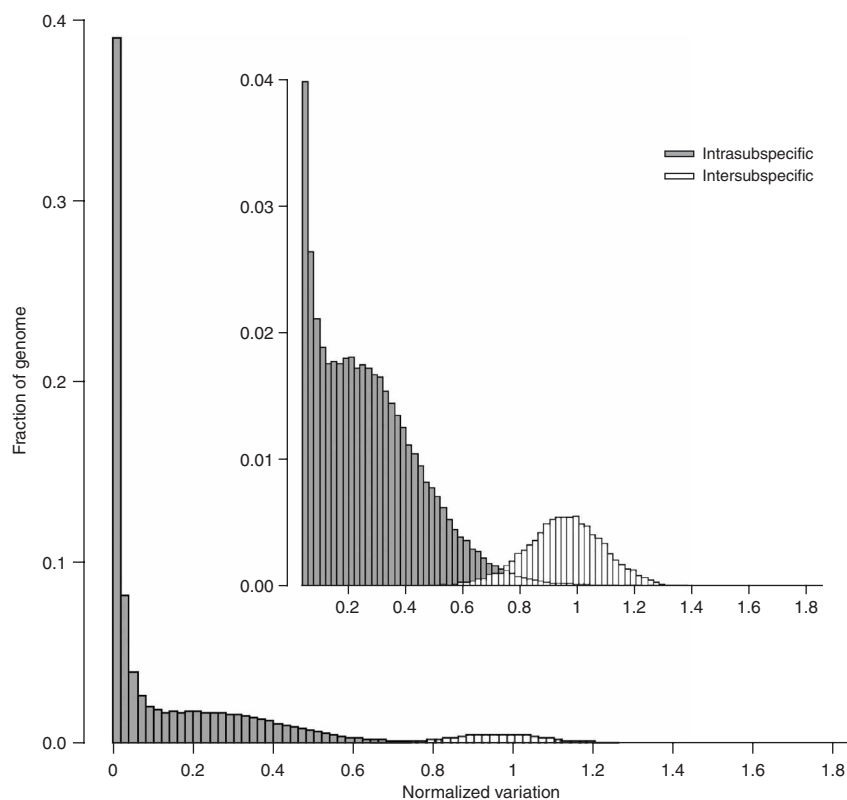


Figure 5 Frequency distribution of the normalized variation in pairwise comparisons between classical strains. The horizontal axis shows the normalized variation over 100-kb intervals for the 55 pairwise comparisons between 11 classical strains. The average variation in intersubspecific comparisons is set at one. White bars correspond to comparisons in which the two classical strains have haplotypes that are derived from different subspecies. Gray bars denote intrasubspecific comparisons. The inset expands the frequency distribution to emphasize the component of variation that is >0.4%.

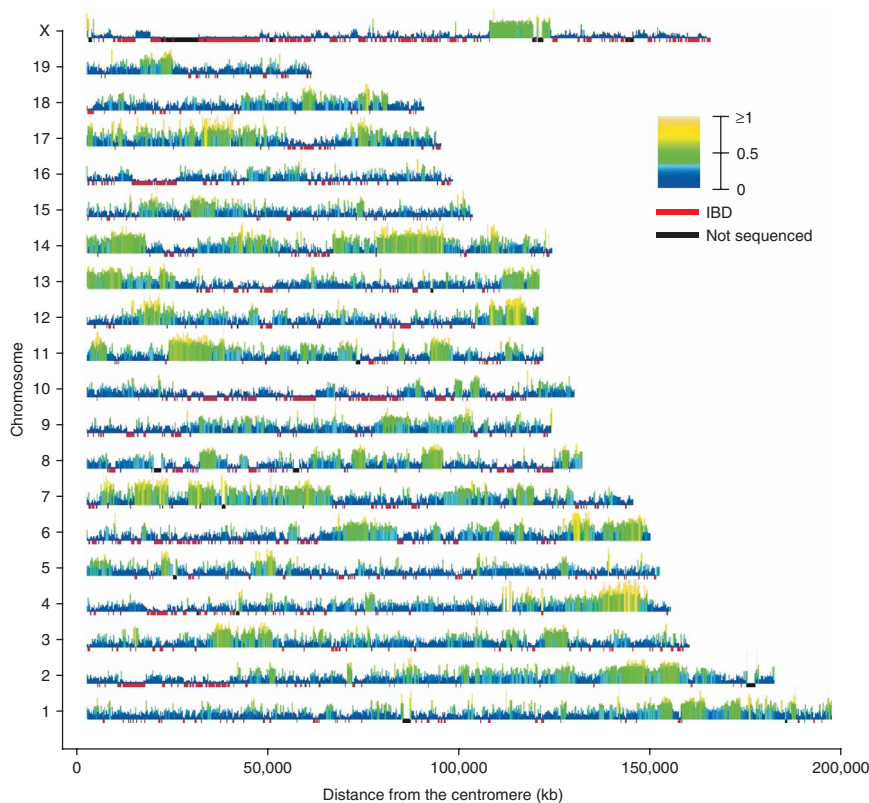


Figure 6 Frequency and spatial distributions of the mean normalized genetic variation observed among 11 resequenced strains. Spatial distribution of the mean normalized variation in the 11 resequenced classical strains is shown as vertical bars of different color and height for each 100-kb interval. IBD, identical by descent.

distribution that is centered on the average level of variation between subspecies (white bars in Fig. 5). On the other hand, more than 99.5% of intervals in which both strains were thought to have the same subspecific origin based on tree topology have normalized variation that is less than one, which would be expected if the subspecific assignments were correct.

The distribution of variation between pairs of inbred strains seems to be a composite of three distinct but overlapping distributions (Fig. 5). Two of these distributions contain the intrasubspecific variation, and the third encompasses the intersubspecific variation. This third distribution, as expected, has the highest level of variation. Direct resequencing indicates that, on average, there is one SNP every 151 bp in intersubspecific comparisons between wild-derived strains, which is similar to previous reports¹⁸. The more prominent of the two intrasubspecific components has variation that is less than 2% of the intersubspecific variation. These regions are expected to have, on average, one SNP every 20 kb, and they probably represent inherited regions that were identical by descent (IBD) within the recent derivation of the classical strains. The fraction of the genome in IBD regions ranges between 36% and 62% in 55 pairwise comparisons among the 11 NIEHS classical strains. These IBD regions represent blind spots in genetic studies that use crosses between classical strains. The remaining intrasubspecific variation encompasses 50% of the genome and has a broad distribution (Fig. 5). The distribution peaks at one-third of the typical level of intersubspecific variation. We propose that this variation is representative of the natural intrasubspecific variation found in every *M. musculus* subspecies.

We also determined the distribution of variation in comparisons between the three reference strains and MOLF and between the reference strains and each one of the classical strains independently (Supplementary Fig. 6 online). The intrasubspecific variation in comparisons involving MOLF has a similar range and mode as the distribution between classical strains, which indicates that this feature is neither restricted to these strains nor restricted to one subspecies. The absence of a peak that corresponds to variation near zero in comparisons involving MOLF indicates that this strain shares little or no IBD regions with the three reference strains. On the other hand, the classical strains do share regions of IBD with WSB, although to a lesser extent than observed within classical strains. These results demonstrate that there is substantial intrasubspecific variation in the *M. m. domesticus* and *M. m. musculus* subspecies, and that the classical strains have captured a fraction of that variation.

In addition to pairwise analyses, we have determined the subspecific origin and variation level in comparisons that include all 11 classical strains. Our analyses indicate that, for approximately two-thirds of the fraction of the genome in which subspecific origin was assigned, all classical strains are derived from a single subspecies, primarily *M. m. domesticus* (Supplementary Fig. 6). Most of the remaining one-third has two subspecific origins, with a predominant con-

tribution from the *M. m. domesticus* lineage. In those genomic regions with two subspecific origins, the minor subspecies is represented on average in 2 out of the 12 classical strains. These conclusions also hold true for the 28% of the genome for which we were not able to assign subspecific origins in the classical strains (Supplementary Fig. 6).

We determined the mean of the normalized variation among the 11 classical strains in every 100-kb interval (Supplementary Fig. 6). This analysis revealed that 11% of the genome is IBD when all strains are considered together. The level of identity is remarkable given that these strains include Castle, Swiss and Asian-derived strains³. This finding reinforces the conclusion that there is a very limited pool of founders and raises questions about whether, in addition to drift, selection for desirable traits in ‘fancy’ mice was involved in establishing the IBD regions. The addition of the C57BL/6 strain does not substantially reduce the fraction of the genome within IBD regions.

The spatial distribution of variation (Fig. 6) reveals substantial heterogeneity at the chromosomal, regional and local levels. For example, chromosomes X, 10 and 16 have low variation in most intervals, whereas chromosomes 14, 17 and 11 have high variation in most intervals. The most striking cases of regional variation are the island of high variation found on chromosome X and the distal regions of chromosomes 4 and 12. IBD regions are also clustered, spanning megabase-long regions in some chromosomes.

Assigning a subspecific origin in any given strain remains constant for extended regions, and consecutive trees with similar topologies are the norm. However, there are frequent minor changes in the topology of trees across consecutive 100-kb intervals that are due to historical

recombination events between haplotypes from within the same subspecies. High historical recombination rates should be beneficial for mapping complex traits. However, we also found that the most common strain distribution patterns in classical strains, representing 99.4% of all complete SNPs, are found on average in almost half of mouse chromosomes. This indicates that false positives will be a formidable obstacle in association mapping studies.

DISCUSSION

In summary, our analyses of the NIEHS data show that the genomes of classical inbred strains are largely derived from the *M. m. domesticus* subspecies. The distribution of genetic variation within the classical strains is nonrandomly distributed. Large regions of the genome are essentially IBD, whereas in other regions the level of diversity approaches that found in intersubspecific comparisons. More than half of the genome of the classical strains shows intermediate variation that is consistent with an intrasubspecific origin. We also found unexpected and frequent intersubspecific introgressions in the wild-derived strains. These features, and the limited amount of diversity that segregates among the classical strains (26% of the estimated total variation in the NIEHS data set), argue for the development of new mouse inbred lines that harbor greater allelic diversity and more complete randomization of ancestry. In particular, our results support the use of larger, heterogeneous populations⁴² and the Collaborative Cross⁴³, a large panel of recombinant lines that randomizes the natural variation in inbred strains from the three main mouse subspecies.

METHODS

PCR-directed resequencing. DNA was obtained from The Jackson Laboratory, with the exception of CIM/Pas, which was a gift. To determine the FPR and FNR, we resequenced 70 fragments located in 14 chromosomes and spanning 28 kb (Supplementary Table 1 online). The position of each base pair in these fragments was tiled in the Perlegen arrays and was completely sequenced in this study in the NIEHS strains and C57BL/6J. In the intersubspecific introgression studies, we resequenced 15 fragments located in ten chromosomes and spanning almost 12.5 kb (Supplementary Table 1) in the following strains: *M. m. castaneus*: CAST/Eij, CASA/RkJ and CIM/Pas; *M. m. musculus*: CZECHII/Eij, PWK/PhJ and PWD/PhJ; and *M. m. domesticus*: WSB/Eij, PERC/Eij and TIRANO/Eij. Amplification and purification of PCR products were carried out as previously described¹⁸. Sequencing was carried out at the Automated DNA Sequencing Facility, University of North Carolina at Chapel Hill, on an ABI Prism 3730 (Applied Biosystems). All sequences were initially aligned using the Sequencher (GeneCodes) software. Aligned sequences were trimmed to retain only high-quality sequences. We determined the genomic positions of each SNP in Build 36 of Ensembl, and the region of complete overlap between our sequences and the NIEHS sequences. Overlapping regions were then compared, and shared and non-shared SNPs were identified. We considered the SNPs to be shared if both data sets had a SNP at the same position, with the same alternative alleles and the same strain distribution pattern.

Frequency of diagnostic SNPs using the raw data. We defined a SNP to be diagnostic if the genotypes in the three reference strains (CAST/Eij, PWD/PhJ and WSB/Eij) were complete and the SNP was polymorphic among these strains. There are three types of diagnostic SNP that correspond to the three strain distribution patterns among the reference strains. The number of diagnostic SNPs was determined in every 100-kb interval and their proportions were mapped to a simplex.

False-negative rate. Based on a comparison of the NIEHS SNPs with our resequencing data, we determined the FNR for the different classes of MAFs (Fig. 2b). Regression of a log-transformed MAF on the FNR provides a robust smoothed estimate of the MAF-specific FNR. The estimated FNRs that correspond to SNPs with the minor allele shared by 1 to 7 strains were 0.76, 0.64, 0.57, 0.52, 0.48, 0.45 and 0.42, respectively. The average genome-wide

FNR was computed as a weighted average of the FNRs across the MAF classes. To estimate the proportion of SNPs that are variable within the classical strains, we calculated the proportion that are variable among classical strains within each MAF class, applied bias correction (see Branch-length correction) to each MAF class and calculated the weighted average.

Branch-length correction. The FNRs of SNPs that have different MAFs were used to correct the estimated branch lengths in the phylogenetic trees. The corrected length is proportional to the expected total number of SNPs (observed plus unobserved). To obtain this correction, we multiplied the estimated length of each branch by the factor $1/(1 - \text{FNR})$, corresponding to the MAF of that branch. Terminal branches of the tree, with lower MAFs and correspondingly higher FNRs, expand more than the inner branches, which have higher MAFs and lower FNRs.

Phylogenetic analyses. Phylogenetic analyses were carried out using the PHYLIP version 3.6 phylogeny inference software package (Felsenstein, J., Department of Genome Sciences, University of Washington, Seattle, 2005). A tree was generated for each 100-kb interval using the SNP genotypes for the 15 NIEHS strains. Branch lengths were corrected as described above. We used Dnapars (DNA parsimony algorithm version 3.6) with default options as described online (see URLs section below), although using the search option to be 'Rearrange on one best tree'. Although we were aware that this search option is less thorough, our pilot study showed that, in most cases, trees found from searches that rearrange from all the possible most parsimonious trees were similar except some discordant results for the terminal branches. We determined the robustness of the tree by bootstrap analysis (Seqboot software (100 replicates)) using the Consense (majority rule) software program. The mitochondrial and Y-chromosomal analyses were carried out with the genotypes at 286 and 4935 SNPs, respectively. Similar results were obtained using both distance (neighbor joining) and maximum likelihood (Dnaml) approaches.

Frequency of diagnostic SNPs using corrected data. Using the corrected trees, we determined the distance from the common node to each of the three reference strains, WSB/Eij, PWD/PhJ and CAST/Eij, in each 100-kb interval. These distances were transformed into fractions representing the local contribution of diagnostic SNPs and were represented in the simplex as described above.

Discrimination between introgressed and balanced regions. The simplex was divided into five regions (Supplementary Fig. 6). Three regions located at the vertices of the triangles contain the three possible types of unbalanced interval. In these regions, the ratio between the length of the longest and shortest branch for the three reference strains in the corrected tree is $> 4:1$. Geometrically, this corresponds to the inside of three circles centered at the vertices of the simplex, whose radius is $1/\sqrt{12}$ (inside circle in Supplementary Fig. 6). Intervals were classified as potential intersubspecific introgression after running a HMM to fill isolated balanced intervals within large blocks of unbalanced intervals and to remove isolated unbalanced intervals. The HMM has four hidden states that correspond to three types of introgression pattern and a fourth balanced state. Here the 'true' introgression status of each 100-kb interval is considered a hidden state. The 'output' of the HMM is an indicator of which region of the simplex an interval was assigned. The HMM parameters were set to revisit the same state with a probability of 0.99 and to tolerate 1% of intervals that are inconsistent with the 'true' state. The HMM inference algorithm has been described previously⁴⁴. We considered an interval balanced if it was not found to be in an introgression region by the HMM and if the ratio of the longest versus the shortest branch length for the three reference strains in the corrected phylogenetic tree was $< 3:1$ (central circle in Supplementary Fig. 6). Unbalanced intervals excluded from the putative introgression regions by the HMM and intervals located in the periphery between the unbalanced and balanced regions of the simplex were considered undetermined regions.

Normalized variation in pairwise comparisons. In pairwise comparisons, we used the distance between a pair of strains in the corrected phylogenetic tree as an estimate of genetic variation. In each interval, we estimated the variation among the three reference strains. The normalized variation is the ratio between the distance separating a given pair of inbred strains and the average distance among all pairs of strains from different subspecific origins. For balanced

regions of the genome, the intersubspecific average includes seven pairwise comparisons (the variation between the three pairs of reference strains and the variation between the four possible combinations of the three reference strains and the two classical strains from different subspecies). For unbalanced regions of the genome, the intersubspecific average was determined using the two pairs of reference strains that are from distinct subspecies.

Subspecific origin in unbalanced intervals. For unbalanced regions, it was not possible to assign an ancestral subspecific origin to segments of each classical strain. We inferred the number of ancestral subspecies present among all classical strains in each interval, using the observed distribution of intersubspecific and intrasubspecific variation in the balanced regions of the genome (Fig. 5). Specifically, we calculated the ratio between each pair of classical inbred strains and the mean distance between the two pairs of wild-derived strains that have no evidence for introgression. We considered that a pair of classical strains belonged to the same subspecies if the ratio was <0.73 or had haplotypes derived from different subspecies if the ratio was >0.73 . This threshold was derived from the distributions of the mean intrasubspecific versus intersubspecific variation observed in classical strains (Fig. 5).

URLs. NIEHS Mouse Genome Resequencing and SNP Discovery Project: <http://www.niehs.nih.gov/crg/cprc.htm>; NIEHS/Perlegen mouse SNP and genotype data: <http://mouse.perlegen.com/mouse/download.html>; NIEHS/Perlegen strain selection criteria: http://mouse.perlegen.com/mouse/strain_selection.html. Jackson Laboratory: <http://www.jax.org>. Default options for Dnapars: <http://evolution.genetics.washington.edu/phylip/doc/dnapars.html>.

Note: Supplementary information is available on the Nature Genetics website.

ACKNOWLEDGMENTS

We thank S. Ahmed for technical assistance; K. Paigen, K. Broman and B. Payseur for helpful comments during the preparation of the manuscript; J. Felsenstein for advice and L. Wu for assistance with the phylogenetic tree computations and A. Smith for developing a genome browser format for displaying phylogenetic trees. CIM/Pas was provided by F. Bonhomme (University Mont Pellier II). We thank the NIEHS and Perlegen for making the SNP data set freely available prior to publication. This work was supported by the US National Institute of General Medical Sciences as part of the Center of Excellence in Systems Biology (1P50 GM076468).

AUTHOR CONTRIBUTIONS

This study was designed by F.P.-M.V. and G.A.C. The genome-wide analyses were carried out by H.Y. The sequence data used to determine the false-negative and false-positive rates and to confirm the presence and direction of intrasubspecific introgression were generated by T.A.B.

COMPETING INTERESTS STATEMENT

The authors declare no competing financial interests.

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