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# Genomic islands of differentiation between house mouse subspecies

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Understanding the genes that contribute to reproductive isolation is essential to understanding speciation, but isolating such genes has proven very difficult. In this study I apply a multilocus test statistic to >10,000 SNP markers assayed in wild-derived inbred strains of house mice to identify genomic regions of elevated differentiation between two subspecies of house mice, *Mus musculus musculus* and *M. m. domesticus*. Differentiation was high through ~90% of the X chromosome. In addition, eight regions of high differentiation were identified on the autosomes, totaling 7.5% of the autosomal genome. Regions of high differentiation were confirmed by direct sequencing of samples collected from the wild. Some regions of elevated differentiation have an overrepresentation of genes with host–pathogen interactions and olfaction. The most strongly differentiated region on the X has previously been shown to fail to introgress across a hybrid zone between the two subspecies. This survey indicates autosomal regions that should also be examined for differential introgression across the hybrid zone, as containing potential genes causing hybrid unfitness.

[Supplemental material is available online at [www.genome.org](http://www.genome.org).]

Speciation in sexual organisms occurs as barriers to gene flow evolve between previously interbreeding populations. Part of a research program to elucidate the origin of species therefore requires an understanding of how and why new traits arise that reproductively isolate populations and to identify the genes controlling these traits (Coyne 1992). The genetic basis underlying the origin of new species has been studied mainly in *Drosophila*. Results of these studies suggest that the genes generally evolved under the influence of positive selection (Barbash et al. 2003; Presgraves et al. 2003; Ting et al. 2004; Greenberg et al. 2006) and involve various normal functions in the organism. While not studied in detail on the molecular level, one gene has also been isolated that is involved in reproductive isolation between two species of mouse, *Mus musculus* and *Mus spretus* (Fossella et al. 2000).

A prerequisite to study the process of speciation and eventually identify the genes involved in it is a close relationship among the taxa. Close relationships among taxa imply that the identified genetic differences are more likely to be involved in the speciation process rather than accumulating after speciation is completed. The house mouse provides an ideal study system. At least three different subspecies are distinguished, which diverged from each other between 0.8 and 1 million years ago (Mya) (Boursot et al. 1993; Guenet and Bonhomme 2003). *M. m. domesticus* and *M. m. musculus* are the best studied subspecies. *M. m. domesticus* ranges across western Europe, northern Africa, and the middle East, whereas *musculus* extends throughout eastern Europe and northern Asia. The two subspecies meet in a narrow hybrid zone, which most likely is maintained by a balance between selection against hybrids and dispersal (Hunt and Selander 1973; Payseur et al. 2004).

One general approach toward the identification of genes involved in speciation relies on identifying regions of the genome that are strongly differentiated between populations or species (Cavalli-Sforza 1966). Regions of elevated genetic differ-

entiation can then be used to search for candidate genes (Akey et al. 2002; Payseur et al. 2002; Kayser et al. 2003; Storz et al. 2004; Storz 2005).

Previously, this approach has been applied in the hybrid zone between the two subspecies of house mice along a transect in eastern Germany (Tucker et al. 1992; Payseur et al. 2004; Payseur and Nachman 2005) and Denmark (Dod et al. 1993). The expectation is that genes causing reproductive isolation introgress at lower rates than genes not involved in reproductive isolation (Barton and Hewitt 1981). However, the number of markers was very limited in these studies (Tucker et al. 1992; Dod et al. 1993) or restricted to a single chromosome (Payseur et al. 2004; Payseur and Nachman 2005). Thus, a more systematic overview over patterns of differentiation over the whole genome is desirable.

In this report I used genomic resources publicly available as thousands of SNP markers typed by The Wellcome Trust Center for Human Genetics in wild-derived inbred strains of the house mouse to systematically identify genomic regions that are highly differentiated between subspecies. I confirm that the X chromosome is strongly differentiated, as expected from previous hybrid zone work (Tucker et al. 1992; Dod et al. 1993), but also identify several regions on the autosomes that are similarly highly differentiated. I consider how these regions might have originated and the genes they contain.

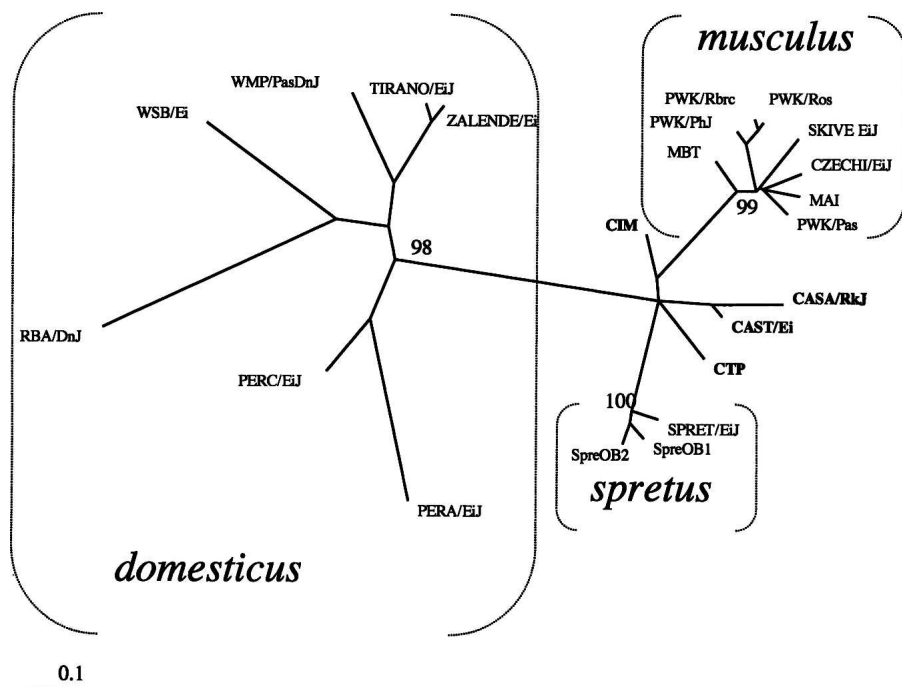
## Results

### Taxa

The genetic makeup of laboratory strains of the house mouse, from which the SNPs available from The Wellcome Trust Center for Human Genetics are derived, is a mixture with the largest contribution from *Mus musculus domesticus* (~60%), an intermediate contribution from *Mus musculus musculus* (~30%), and a small contribution from *Mus musculus castaneus* (~10%) (Wade et al. 2002; Wade and Daly 2005). The tree based on all 13,000 SNPs showed that most wild-derived inbred strains cluster into three well supported clades (Fig. 1). Combined with the information in

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**Figure 1.** Maximum likelihood phylogenetic tree showing the relationships among wild-derived inbred strains of *Mus musculus* and *Mus spretus*. The tree was graphically represented in TreeView (Page 1996).

Table 1, all three clades were unambiguously assigned to one of the three subspecies of *Mus musculus* or to *Mus spretus*. For three strains (PWK/Rbrc, PWK/Ros, PWK/Pas), no information was available from The Jackson Laboratory Web site (<http://jaxmice>).

ties that are closer to the assumed ancestral origin of all house mouse subspecies (i.e., India and Thailand [Boursot et al. 1993]), which could mean that they represent a larger and more ancestral diversity range.

jax.org/query). However, Gregorova and Forejt (2000) state their origin is in the Czech Republic. These strains are substrains derived from PWK/Ph and they all clearly cluster within other strains of the subspecies *musculus*. Four of the wild-derived inbred strains were assigned to the subspecies *castaneus* (Table 1). These strains did not cluster together but are located dispersed and more ancestrally between the sister species *Mus spretus* and the strains from the *Mus musculus* subspecies *musculus*. One explanation for this pattern is ascertainment bias stemming from the selection process of the SNP markers. Since most mutations are polymorphic in *domesticus*, because this is the subspecies with the highest genetic contribution to laboratory strains, *castaneus* is expected to harbor the ancestral state. Thus, these strains are placed more toward the common ancestor of all subspecies, *Mus spretus*. The few SNPs that are polymorphic in *castaneus* provide too little resolution to cluster all four strains together on one branch. An alternative explanation for the position of *castaneus* strains on the tree could be that they stem from localities that are closer to the assumed ancestral origin of all house mouse subspecies (i.e., India and Thailand [Boursot et al. 1993]), which could mean that they represent a larger and more ancestral diversity range.

**Table 1.** Geographic origin and other strain characteristics as obtained from The Jackson Laboratory Web site for the inbred strains included in the SNP survey

Strain name	Geographic origin (The Jackson Laboratory)	Other sources	Robertsonian fusion
CASA/RkJ	Thailand		
CAST/Ei	Thailand		
CIM	India		
CTP	Thailand		
CZECHI/EiJ	Studeneč, Czech Republic		
MAI	Austria		
MBT	Bulgaria		
PWK/PhJ	Lhotka, Czech Republic		
PWK/Rbrc		Prague, Czech Republic <sup>a</sup>	
PWK/Ros		Prague, Czech Republic <sup>a</sup>	
PWK/Pas		Prague, Czech Republic <sup>a</sup>	
SKIVE/EiJ	Skive, Denmark		
PERA/EiJ	Nana Village, Rimac Valley, Peru		Rb(4.12)9Bnr
PERC/EiJ	Nana Village, Rimac Valley, Peru		Rb(1.3)1Bnr; Rb(10.12)5Lub; Rb(11.13)4Bnr; Rb(16.17)7Bnr;
RBA/DnJ	Grisons, Switzerland		Rb(2.8)2Lub; Rb(4.6)2Bnr; Rb(5.15)3Bnr; Rb(9.14)6Bnr;
TIRANO/EiJ	Tirano, Italy		Rb(1.11)2Mpl; Rb(10.17)9Mpl; Rb(13.15)10Mpl; Rb(2.16)3Mpl;
WMP/PasDnJ	Monastir, Tunisia		Rb(3.12)4Mpl; Rb(4.6)5Mpl; Rb(5.14)6Mpl; Rb(7.18)7Mpl;
WSB/Ei	Centreville, Maryland, USA		Rb(8.9)8Mpl;
ZALENDE/EiJ	Zalende, Switzerland		Rb(1.3)1Bnr; Rb(11.13)4Bnr; Rb(16.17)7Bnr; Rb(4.6)2Bnr;
			Rb(5.15)3Bnr; Rb(8.12)5Bnr; Rb(9.14)6Bnr

<sup>a</sup>Origin of strains given in Gregorova and Forejt (2000).

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As expected from the ascertainment bias during the selection process of SNP markers, the branch lengths for members of subspecies *domesticus* were longer than branches for other subspecies. However, within *musculus* strains there was still a substantial amount of variation evident, which was roughly proportional to the contribution of this subspecies to the genetic makeup of laboratory strains.

Only strains clustering within *musculus* and within *domesticus* were used for further analysis (seven *domesticus* and eight *musculus*). Among the 13,000 SNPs, 10,265 were polymorphic in either *domesticus* (6281 SNPs) or *musculus* (1146 SNPs) or both subspecies (1650 SNPs) or showed a fixed difference between the subspecies (1188 SNPs). These 10,265 “variable” SNPs were used for further analysis. The number of SNPs per chromosome was correlated with chromosome size (Pearson  $r = 0.767$ ,  $N = 20$  chromosomes,  $P < 0.0001$ ). The largest chromosome (chromosome 1) contained 804 SNPs, and the smallest (chromosome 19) contained 235 SNPs. The total number of variable SNPs was used to generate the expected values for the  $\chi^2$  test. The ratio of SNPs showing a fixed difference between strains to those that were polymorphic in at least one strain was  $1188/9077 = 0.13$ , which yields an expected number of fixed SNPs of 7.9 and polymorphic SNPs of 52.1 in a 60-SNP sliding window.

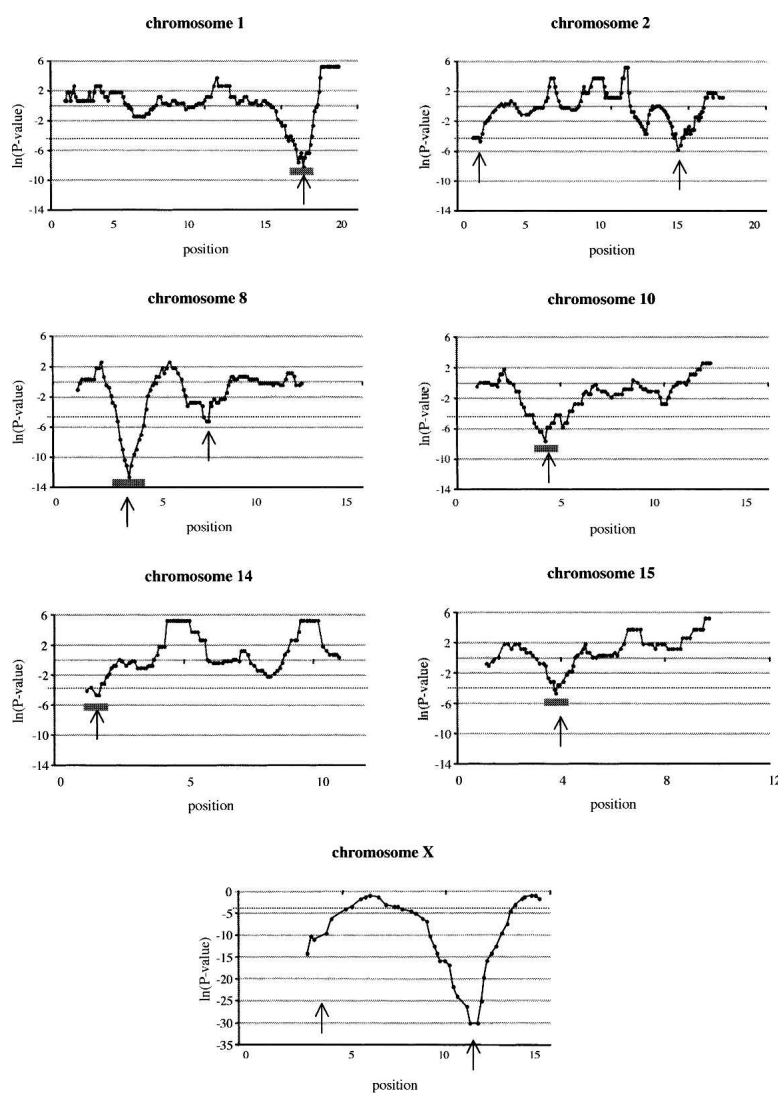
### Differentiation based on SNP analysis from lab strains

I searched for significant regions of differentiation using a  $\chi^2$  test and a permutation test, as explained in the Methods. In the Supplemental material I describe analyses using SNP windows from 20–100 SNPs in size. In the main text I confine the discussion to the results obtained from a sliding window size of 60 SNPs, which provides a resolution similar to that obtained from smaller window sizes, whereas windows of  $>60$  SNPs give results that are inconsistent with the permutation test (see Supplemental material).

Figure 2 shows heterogeneity in differentiation along those chromosomes where both the  $\chi^2$  test and the permutation test identified at least one region to be significantly highly differentiated between the two subspecies. The permutation test identifies three additional chromosomes (3, 7, 9) over the  $\chi^2$  test with a window of larger differentiation than expected (see Supplemental Table 2). In the following, however, I consider only regions for which both tests were significant.

Eight regions of elevated differentiation are present on the autosomes (Fig. 2). Their exact genomic position

based on the most recent assembly of the mouse genome (build 34) is given in Table 2. Two chromosomes contained more than one region where sliding windows of high differentiation ( $P > 0.015$ ) did not overlap (chromosome 2 and 8). Most chromosomes showed no region of elevated differentiation (chromosomes 3–7, 9, 11–13, 16–19). Regions of significant differentiation covered  $\sim 7.5\%$  of the autosomes in total, with an average length of 20 Mb (from  $\sim 11$  Mb [chromosome 15] to 31 Mb [chromosome 10]). In contrast to the autosomes, elevated differentiation spanned almost across the entire X chromosome (88%, Fig. 2), with two large regions of elevated differentiation separated by a region of lower differentiation. However, the sliding windows of both high differentiation regions overlap on the X chromo-



**Figure 2.** Signed  $\ln$ -transformed  $P$ -values from a  $\chi^2$  analysis of a sliding window plotted against chromosomal position (midpoint of a sliding window). Positive values on the  $y$ -axis indicate a deficiency in the number of fixed differences between the subspecies while negative values indicate more fixed differences than expected. Sliding windows consist of 60 SNPs moved for five SNP each step. (Dashed line) Significance level for a surplus of fixed differences among subspecies, (arrows) regions of significantly elevated differentiation, (short horizontal gray bars) regions that were chosen for direct sequencing of independent samples of *M. m. musculus* and *M. m. domesticus*. Only chromosomes where significant differentiation was found are shown. Note that the  $y$ -axis of the X chromosome differs from those of the autosomes because of the overall much higher differentiation of the X. Base positions on the  $x$ -axis need to be multiplied by  $10^7$ .

**Table 2.** Positions of significantly elevated differentiation regions on each chromosome (based on build 34 of the mouse genome) identified using a sliding window technique and a window size of 60 SNPs

Chromosome	Begin	End	Size
1	148,046,005	174,537,672	26,491,667
2	6,377,329	23,423,751	17,046,422
2	136,738,822	151,795,810	15,056,988
8	25,166,987	51,402,145	26,235,158
8	65,731,752	81,716,391	15,984,639
10	28,936,290	59,816,232	30,879,942
14	9,366,031	23,841,110	14,475,079
15	32,698,808	43,617,391	10,918,583
X	5,371,526	147,372,655	142,001,129

some. Thus, a contiguous region of elevated differentiation on the X chromosome is given in Table 2.

### Natural populations

The results from direct sequencing of natural populations of *musculus* (from the Czech Republic, 14 chromosomes sequenced) and *domesticus* (from Germany, 16 chromosomes sequenced) are shown in Figure 3 and Table 3. The average  $G_{st}$  value within the sequenced fragments, which is based on polymorphic sites only, was significantly higher for fragments located in the candidate regions of high differentiation (as identified from the SNP survey of wild-derived inbred strains) than for fragments that are located outside of these regions ( $P = 0.0003$ ,  $t$ -test,  $df = 12$ ). Thus, regions identified by the SNP analysis were confirmed in an independent sample. Nei's statistic (Nei 1987) is based on the average pairwise divergence across all sites, including those that are identical in each species, and appears less powerful. Although not significant, Nei's average divergence averages 22% higher for high differentiation regions than for low differentiation regions (average divergence in high differentiation region = 0.011, average divergence in reference region = 0.008, Table 3). Also, sequences from each individual more often coalesce within the assigned subspecies for the loci located in high differentiation regions than for loci located in low differentiation regions (Supplemental Fig. 1).

I found no measurable differences in divergence to *M. famulus* between regions of high and low differentiation ( $P = 0.551$ ,  $t$ -test,  $df = 11$ ): Of the 14 genomic fragments sequenced, I was unable to amplify one fragment in *famulus*, hence the number of degrees of freedom is 11).

### Functional annotation of candidate regions

The full list of functional themes that were significantly ( $P \leq 0.05$ , binomial test) overrepresented among the genes located in the high differentiation regions is available as Supplemental Table 3. The most significant entry in this list was found for chromosome 1, where genes related to immunity and defense were strongly ( $P < 0.000001$ ) overrepresented. Next most overrepresented functional categories in the highly differentiated regions are genes related to proteolysis on chromosome 2 ( $P = 0.00004$ ) and olfactory genes on chromosome 1 ( $P = 0.00011$ ). These three identified categories of genes are also significant when corrected for testing of multiple categories (Bonferroni  $P < 0.0002$ , 244 functional categories tested).

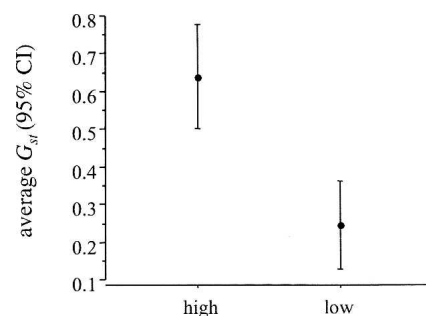
### Recombination rates

Previously, it has been shown that elevated differentiation was confined to regions of reduced recombination (Turner et al. 2005). Recombination rates are available only for a cross between a laboratory strain (B6-Ob) and *M. m. castaneus* (Dietrich et al. 1996), which limits the comparison between *musculus* and *domesticus*. Using these rates, the association between recombination rate and differentiation in my study is very low ( $r = 0.047$ ; Fig. 4).

### Discussion

Genetic changes that lead to the split of one species into two are of major interest in evolutionary biology. The house mouse is an ideal model system because it consists of three "incipient" species, i.e., subspecies. Initial differentiation of these subspecies probably took place on the Indian subcontinent ~0.5–1 Mya (Boursot et al. 1993; Guenet and Bonhomme 2003). It is thought that house mice started to colonize most of the Eurasian continent and Northern Africa ~10,000 yr ago, in association with the origin of human commensalism. Each subspecies followed its own colonization route, and the resulting distribution is nearly non-overlapping. Where the subspecies come into contact they form hybrid zones (Boursot et al. 1993).

My survey of ~10,265 SNP loci on all chromosomes except the Y is the first to address the question of how genetic differences between subspecies of the house mouse are distributed across the genome. I found several "genomic islands" of large differentiation between strains derived from *musculus* and *domesticus*. The largest difference between the two subspecies was along the X chromosome, but some regions of the autosomes were also strongly differentiated. Some potential sources of bias can be eliminated. Within *domesticus* a number of chromosomal races are known, which possess different combinations of two chromosomes joined together in Robertsonian fusions (Nachman and Searle 1995). It is therefore possible that the result is an artifact of considering one or other chromosomal race. However, four of the seven *domesticus* strains contained different Robertsonian fusions (Table 1), while the other three contained the standard karyotype. This heterogeneity should increase the variance within *domesticus*, making it harder to detect fixed differences between the subspecies. In addition, I was able to confirm the pattern of high and low differentiation regions using independent animal samples collected from nature.



**Figure 3.** Mean and 95% confidence interval of average  $G_{st}$  based on direct sequencing of fragments within high ( $N = 8$ ) and low ( $N = 6$ ) differentiation regions in natural population of house mice.

**Table 3.** Summary of polymorphism data from natural populations of *M. m. musculus* and *M. m. domesticus*

Associated gene name	Chrom	Gene start position	Gene stop position	Sub-species	No. alleles	No. bp	Seg sites	$\pi$	$\theta$	Average div. ( <i>famulus</i> )	Average $G_{st}$ ( <i>dom</i> vs. <i>mus</i> )	Average div. ( <i>dom</i> vs. <i>mus</i> )
<b>I. High differentiation</b>												
<i>Dusp2</i>	1	166,019,931	166,049,702	<i>musculus</i>	14	668	2	0.0012	0.0009	0.02	0.63	0.014
				<i>domesticus</i>	16	590	4	0.0028	0.002			
<i>Ppp1r3b</i>	8	34,178,803	34,191,218	<i>musculus</i>	14	699	1	0.0002	0.0004	0.03	0.40	0.009
				<i>domesticus</i>	14	697	9	0.0052	0.0041			
<i>Pcm1</i>	8	40,177,408	40,269,979	<i>musculus</i>	14	634	0	0	0	0.02	0.39	0.002
				<i>domesticus</i>	16	607	2	0.0006	0.001			
<i>Q61875_MOUSE</i>	8	42,253,455	42,255,026	<i>musculus</i>	14	717	2	0.0011	0.0009	0.04	0.82	0.019
				<i>domesticus</i>	16	719	2	0.0006	0.0008			
<i>Sec63</i>	10	42,867,857	42,938,871	<i>musculus</i>	12	672	1	0.0005	0.0005	0.02	0.72	0.005
				<i>domesticus</i>	16	614	0	0	0			
<i>Nr1d2</i>	14	16,724,647	16,757,089	<i>musculus</i>	14	616	1	0.0009	0.0005	0.02	0.67	0.003
				<i>domesticus</i>	16	620	0	0	0			
<i>Rims2</i>	15	39,098,034	39,584,795	<i>musculus</i>	14	423	1	0.0006	0.0007	0.02	0.66	0.012
				<i>domesticus</i>	16	423	2	0.0016	0.0014			
<i>Atp6v1c1</i>	15	38,224,368	38,231,577	<i>musculus</i>	14	546	1	0.0005	0.0006		0.82	0.020
				<i>domesticus</i>	16	633	2	0.0012	0.001			
<b>II. Low differentiation</b>												
<i>Mme</i>	3	62,963,799	63,050,158	<i>musculus</i>	14	683	9	0.0054	0.0041	0.02	0.35	0.017
				<i>domesticus</i>	16	683	13	0.0073	0.0057			
<i>Wdt2</i>	1	72,455,560	72,508,425	<i>musculus</i>	10	278	3	0.0037	0.0038	0.04	0.20	0.005
				<i>domesticus</i>	12	278	3	0.0039	0.0036			
<i>Bnc1</i>	7	75,774,810	75,800,430	<i>musculus</i>	14	637	5	0.0019	0.0025	0.01	0.39	0.01
				<i>domesticus</i>	16	633	7	0.0024	0.0033			
<i>XP_620246</i>	5	111,762,098	111,855,645	<i>musculus</i>	14	448	2	0.002	0.0014	0.02	0.13	0.004
				<i>domesticus</i>	16	425	7	0.0045	0.005			
<i>Ggh</i>	4	20,143,471	20,167,391	<i>musculus</i>	16	499	1	0.0006	0.0006	0.04	0.13	0.002
				<i>domesticus</i>	16	499	2	0.0018	0.0012			
<i>Melk</i>	4	44,216,840	44,280,598	<i>musculus</i>	16	918	11	0.0039	0.0036	0.03	0.25	0.010
				<i>domesticus</i>	14	919	13	0.0033	0.0044			

div: divergence according to Nei (1987)

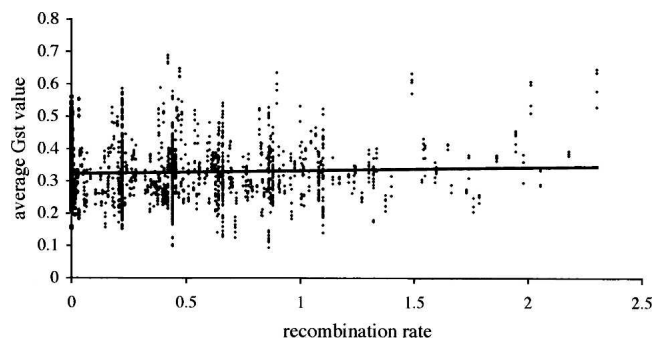
Seg Sites: number of segregating sites

While I found several regions of high differentiation on the autosomes, the overall proportion of the autosomal genome that showed strong differentiation was relatively small (~7.5%). Similarity across most regions of the autosomal genome could reflect (1) significant introgression of genetic material across the subspecies boundaries, (2) ancestral polymorphism, and (3) lower mutation rates in the region without significant differentiation. To examine this I analyzed the 10,265 SNP loci in three individuals of the closely related species *Mus spretus*. 9% (897) of the SNPs that were polymorphic within at least one of the *musculus* subspecies were also polymorphic in *spretus*. Ideraabdullah et al. (2004) sequenced 62 genomic fragments from 14 chromosomes in several wild-derived strains of the species *musculus*, *spretus*, and *spicilegus* and also found a relatively high frequency of ancestral polymorphisms (~13%). Thus, ancestral polymorphism likely contributes to a fraction of the sites that are not fixed between the subspecies.

Under neutral expectations, divergence should be proportional to the mutation rate in a region. Since the mutation rate of the X chromosome is lower than the mutation rate on the autosomes (it is thought that the mutation rate of the X chromosome is ~70% that of the autosomes [McVean and Hurst 1997]), the X chromosome is expected to show lower differentiation. However, I observe the opposite: The level of differentiation between the subspecies is greatly elevated on the X chromosome relative to the autosomes. In addition, mutation rate effects are unlikely to explain differences between regions of high and low differentiation along the autosomes. I tested this by calculating the average

divergence between natural populations of the *musculus* subspecies and the outgroup *Mus famulus* for each of the sequenced fragments. I found divergence from *M. famulus* was similar in the regions of high and low differentiation.

Divergent selection between the subspecies is the most likely cause of regional differentiation. Genes located in the regions of elevated differentiation are candidate targets for selection. Although there are many genes in these regions, the most overrepresented category is genes related to immune response and proteolysis (proteolysis occurs during immune response [Liu 2004]). Genes such as these may be involved in host-pathogen interaction; based on comparisons of non-synonymous/synony-

**Figure 4.** Correlation between recombination rate and average  $G_{st}$  in a sliding window size of 60 SNPs.

mous ( $d_N/d_S$ ) divergence, these genes have been shown to be common targets of directional selection (Tanaka and Nei 1989; Schlenke and Begun 2003). The other genes most strongly over-represented in regions of high differentiation are those related to olfaction. A high  $d_N/d_S$  ratio observed among members of the olfactory gene clusters in mice also suggests positive selection for these genes, possibly favoring taxon-specific pheromone detection (Zhang et al. 2004). Immune defense genes and genes related to olfaction were also found in a recent survey of human and chimpanzee genomes to be evolving particularly fast under positive selection (Bustamante et al. 2005; Nielsen et al. 2005).

Evidence for high regional differentiation within and among chromosomes does not directly address the question of whether these regions are involved in reproductive isolation. In mice, olfaction is known to be important for pre-mating isolation (Talley et al. 2001). In general, only a few genes involved in post-mating isolation have been identified (Orr et al. 2004), and their normal functions within populations are still largely unknown (e.g., Sun et al. 2004). In mice, two methods have recently been used in an attempt to narrow down regions that may contain candidate genes causing hybrid unfitness. First, Payseur and Hoekstra (2005) screened multiple laboratory inbred strains of house mice for physically unlinked SNP pairs that are in linkage disequilibrium. In this way, they identified 14 pairs of gene complexes that could be causing reproductive incompatibilities. While suggesting candidate regions, their approach has the complication that the unknown breeding history of the strains plus selection in the laboratory cannot completely be ruled out as a cause of the linkage disequilibrium (Payseur and Hoekstra 2005).

The proximal part of chromosome 17 has been associated with hybrid male sterility in crosses between laboratory strains of house mice and certain *Mus musculus* strains (Forejt and Ivanyi 1974). The responsible locus (*Hst1*) has recently been narrowed down to a region encompassing two candidate genes (Trachtulec et al. 2005). Interestingly, neither the study by Payseur and Hoekstra (2005) nor my analysis of genome-wide differentiation highlighted this region as being significant. While Payseur and Hoekstra (2005) found one SNP on chromosome 17 to be in disequilibrium with a SNP on chromosome 5, this SNP mapped 10 Mb distal to the *Hst1* locus. One explanation for the absence of significant differentiation of the proximal part of chromosome 17 is that the *Hst1* alleles, which are causing hybrid sterility, are still segregating in natural populations of both subspecies (Forejt 1996).

A second method to identify regions involved in post-mating isolation is to study introgression across a natural hybrid zone; portions of the genome that do not introgress may carry genes causing post-mating incompatibilities. Dod et al. (1993) and Tucker et al. (1992) used a small number of markers and found a lower rate of introgression across the hybrid zone for X-linked markers. Among the three X-linked loci that were analyzed by Dod et al. (1993), one locus showed especially low introgression rates across the hybrid zone. The exact physical position of this anonymous marker is not known. However, the genetic map position is available from The Jackson Laboratory Web site. The marker is very closely linked to the *Pgk1* (phosphoglycerate kinase 1) gene. The physical position of the *Pgk1* gene (as determined from the Ensembl Web site) is within 1 Mb to the most strongly differentiated region on the X chromosome discovered in my SNP differentiation screen. Also, Payseur et al. (2004), using 13 microsatellite markers on the X, identified this general region on the X chromosome as being especially retarded

in its movement across the zone, and the same region harbors the *Hstx1* hybrid sterility locus (Storchova et al. 2004). The region of highest differentiation on the X chromosome in the SNP scan described here (from mice largely taken far from the hybrid zone) is thus identical with the region identified previously as showing the lowest introgression and containing a gene involved in reproductive isolation.

The autosomal regions of high differentiation identified in SNP scan should now be examined for differential introgression across the hybrid zone. Those regions showing limited introgression are particularly likely to contain genes involved in reproductive isolation.

## Methods

### SNP database

The Wellcome Trust Center for Human Genetics provides information on 13,370 Single Nucleotide Polymorphisms (SNPs) from 480 *Mus musculus* inbred strains that have been maintained by sibling matings for 20 or more consecutive generations. Most of the SNPs were identified by direct sequence comparisons between five strains (DBA/2J, A/J, C57BL/6J, 129S1/SvImJ, 129X1/SvJ), and some SNPs were included from other studies (see Web site: <http://www.well.ox.ac.uk/mouse/INBREDS/>). The SNPs are more or less equally distributed over all chromosomes, but no SNPs are available that map close to the centromeres and telomeres of each chromosome. The strains from which the SNPs were identified are all laboratory inbred strains, which are mixtures between different subspecies of the house mouse but with a predominant influence of *Mus musculus domesticus* (Wade et al. 2002; Wade and Daly 2005). Most of the strains are rather artificial laboratory lines from multiple sources, but the data set also includes 22 wild-derived strains, among them three strains from the sister species *Mus spretus*. These 22 strains formed the focus of this study. Using the Web interface of the Web site given above, I downloaded all SNPs where at least one of the strains showed a different nucleotide state from the others.

### Phylogenetic clustering of the strains based on SNP data

Information about subspecies affiliation of most of the 22 strains is available on The Jackson Laboratory Web site (Table 1). For some strains, the subspecies affiliation was not available on The Jackson Laboratory Web site. To both confirm the subspecies affiliation of each strain and identify those whose identification was uncertain, I estimated the phylogenetic relationships of the strains. I used the maximum likelihood program tree-puzzle version 5.2 (Schmidt et al. 2002), having first concatenated the nucleotide states of each SNP locus to obtain a single sequence. Based on the phylogenetic grouping and available strain information (Table 1), I assigned the genotyped wild-derived strains to individual subspecies of house mice. A sufficient number of strains was available for further analysis only for the subspecies *domesticus* (seven strains) and *musculus* (eight strains).

### SNP polymorphism analysis

I employ a multilocus test statistic, which searches for a clustering of sites showing a fixed difference between the subspecies when compared with the genome-wide average. To search for a regional clustering of fixed differences, a sliding window technique was used. The window size ranged from 20 SNPs to 100 SNPs in each window, and the window was moved five SNPs at each step along each chromosome. A  $\chi^2$  statistic was used to test whether the number of fixed SNPs in each window was more than expected by chance. The expectation is based on the overall

frequency of fixed SNPs in the whole data set. I selected all windows with  $\chi^2$  test  $P$ -values  $\leq 0.015$  as candidate regions for regional high differentiation.

I also used a second test to detect regional elevated differentiation based on permutation. For this test I set up a null hypothesis based on the distribution of SNPs along each chromosome, rather than the whole genome as was the case for the  $\chi^2$  test. I randomized SNPs within a chromosome and performed the sliding window analysis (as before using window sizes between 20 and 100 SNPs) on the randomized data (1000 random data sets per chromosome). For each chromosome, the window with the highest number of fixed SNPs in each permuted data set was recorded and compared with the window with the highest number of fixed SNPs in the observed (non-randomized) data set. The number of times where I found a window with higher differentiation in the randomized data set compared with the most differentiated window in the observed data divided by the number of permutations corresponds to the  $P$ -value. Chromosomes with a  $P$ -value  $\leq 0.015$  were considered to harbor at least one region of significant elevated differentiation. All statistical analyses were performed using Perl scripts available from the author upon request.

### Sequencing

In order to confirm the results from the SNP analysis, I sequenced wild-caught individuals. I used seven individuals from a natural *M. m. musculus* population collected in Kazakhstan in 2002 and eight individuals from a natural *M. m. domesticus* population (Germany, collected in 2001) for direct sequencing. All individuals were collected >400 km away from the hybrid zone using a sampling strategy that eliminates the chance of collecting related animals (Ihle et al. 2006). As an outgroup I sequenced one animal of *Mus famulus* collected in India and kindly provided by F. Bonhomme. I chose *M. famulus* as the outgroup for sequencing because its more distant relationship to *M. musculus* (estimated 2.8 Myr [Chevret et al. 2003]) means that it is less likely to share ancestral polymorphisms with *Mus musculus* than *Mus spretus* (see Results; Ideraabdullah et al. [2004]).

I sequenced eight genomic fragments located in five of the regions that were identified by the SNP analysis to be highly differentiated and an additional six fragments located outside of highly differentiated regions. Each fragment consisted of 700–800 bp of non-coding sequence (either 5' or 3' flanking regions or from an intron). Information about the sequenced fragments can be found in Supplemental Table 4.

I used a 50- $\mu$ l PCR containing 100 ng of genomic DNA, 1.5 mM MgCl<sub>2</sub>, 200  $\mu$ M dNTPs, 1  $\mu$ M of each primer, and 0.5 U Taq polymerase (Eppendorf Master Taq). A typical cycling profile consisted of 30 cycles with 50 sec at 94°C, 50 sec at 60°C, and 2 min at 72°C. PCR products were purified using 96-well plates (Millipore Montage) according to the manufacturer's protocol. All PCR products were directly sequenced in both directions using the BigDye sequencing chemistry on an ABI3700 automated sequencer. In the case of individuals being heterozygous for an insertion/deletion, I cloned both alleles (TA cloning kit, Invitrogen) and sequenced eight clones per individual. Sequencing reactions were purified using Sephadex columns (Millipore). All sequence data is downloadable in aligned Nexus format from the author's Web site (<http://www.genetik.uni-koeln.de/groups/Tautz/meg/>).

### DNA sequence data analysis

DNA sequences were aligned using MUSCLE (Edgar 2004). As an estimate of differentiation between the subspecies, I calculated

the  $G_{st}$  value (Nei 1973) for each polymorphic site in the sample of both subspecies and took the average over each fragment. These averages were compared between the loci from inside the identified differentiated regions to those from outside these regions by means of a  $t$ -test. I also calculated the nucleotide divergence (average proportion of nucleotide differences) between both subspecies according to Nei (1987) using the program DN-Asp (Rozas et al. 2003).

### Identification of functional categories

I used the Ensembl Web site ([www.ensembl.org](http://www.ensembl.org)) to download all genes located in regions with significantly elevated differentiation. The gene lists were subsequently submitted to PANTHER (<http://www.pantherdb.org>), which identifies overrepresented functional categories among the significant regions (Mi et al. 2005). Significance is established by comparing the gene list with a reference list (in this case the full gene content of the mouse genome) by means of a binomial test (Cho and Campbell 2000).

### Recombination rates

Reduced recombination might be an important cause of large blocks of the genome being restricted to one or other subspecies (Turner et al. 2005). To test this I used recombination rates estimated for each chromosome in the mouse genome in 5-Mb non-overlapping windows (Jensen-Seaman et al. 2004). The midpoint of each window together with the respective recombination rate can be downloaded from [http://www.genome.org/content/vol14/issue4/images/data/528/DC1/Jensen-Seaman-19703\\_SupTable\\_B.txt](http://www.genome.org/content/vol14/issue4/images/data/528/DC1/Jensen-Seaman-19703_SupTable_B.txt). These data were used to extract recombination rates for each sliding window analyzed in the SNP differentiation scan. To test whether genomic regions of elevated differentiation are located in areas of reduced recombination, I calculated the correlation coefficient between the average  $G_{st}$  value (Nei 1973) of all SNPs located within a sliding window and the recombination rate in this window.

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