

THE VARIABLE GENOMIC ARCHITECTURE OF ISOLATION BETWEEN HYBRIDIZING SPECIES OF HOUSE MICE

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Studies of the genetics of hybrid zones can provide insight into the genomic architecture of species boundaries. By examining patterns of introgression of multiple loci across a hybrid zone, it may be possible to identify regions of the genome that have experienced selection. Here, we present a comparison of introgression in two replicate transects through the house mouse hybrid zone through central Europe, using data from 41 single nucleotide markers. Using both genomic and geographic clines, we found many differences in patterns of introgression between the two transects, as well as some similarities. We found that many loci may have experienced the effects of selection at linked sites, including selection against hybrid genotypes, as well as positive selection in the form of genotypes introgressed into a foreign genetic background. We also found many positive associations of conspecific alleles among unlinked markers, which could be caused by epistatic interactions. Different patterns of introgression in the two transects highlight the challenge of using hybrid zones to identify genes underlying isolation and raise the possibility that the genetic basis of isolation between these species may be dependent on the local population genetic make-up or the local ecological setting.

KEY WORDS: Evolutionary genomics, hybridization, introgression, reproductive isolation, speciation.

In hybrid organisms, the products of meiotic recombination and segregation provide an opportunity to measure the contribution of different genomic regions to reproductive isolation. The fitness effects of individual chromosomal regions define their fate in a population of hybrids, and comparative analysis across the genome allows mapping of the genetic components of isolation between species (Rieseberg et al. 1999; Buerkle and Lexer 2008; Gompert and Buerkle 2009a). By parsing the effects of differ-

ent genomic regions on isolation between taxa, the evolutionary processes and histories that lead to speciation can be revealed.

In many cases, hybrid zones contain organisms that are the result of multiple generations of recombination, and the genetic architecture of isolation may be complex. Fitness variation associated with particular chromosomal blocks in hybrids can promote introgression, can enhance barriers to gene flow, or can be negligible and not associated with isolation between species. Furthermore, the genetic architecture of isolation between two taxa may vary spatially. Across sites of contact and hybridization between species, there may be environmental and ecological

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variation or genetic variation for factors that contribute to isolation. A study that compares the genetic architecture of isolation at multiple points of geographic contact between hybridizing species can identify variable and consistent aspects of the architecture and, in so doing, will point the way toward a more complete characterization of the individual genetic components and their historical contribution to speciation.

Previous hybrid zone studies have considered the similarity of isolating barriers among geographic locales and among individuals. Researchers have compared clines along replicate spatial transects (Szymura and Barton 1991; Barton and Gale 1993; Morgan-Richards and Wallis 2003; Bozikova et al. 2005; Yanchukov et al. 2006; Nolte et al. 2009) and compared the composition of multiple hybrid populations (Buerkle and Rieseberg 2001; Aldridge 2005; Borge et al. 2005). Among these studies, there is a range of concordance in clines and hybrid composition between different samples from the same hybrid zone, indicating that it is difficult to make generalizations about hybrid zone dynamics. Moreover, laboratory studies of hybridization have provided evidence for polymorphism for reproductive isolation (e.g., *Chorthippus*, Shuker et al. 2005; *Drosophila*, Reed and Markow 2004; Kopp and Frank 2005; *Helianthus*, Rieseberg 2000; *Mimulus*, Sweigart et al. 2007; *Mus*, Vyskocilova et al. 2005; Good et al. 2008b; *Tribolium*, Wade et al. 1997).

A recently developed method for characterizing introgression between species' genomes provides a statistical framework to compare the architecture of isolation between multiple sampling locations in hybrid zones (Gompert and Buerkle 2009a). This genomic clines method examines introgression between genomes, rather than the more traditional approach of fitting geographic clines in population allele frequencies. The estimated genomic clines are multinomial regression functions for the genotypes of individuals as a function of their ancestry at all loci. The functions for multiple hybrid populations are compared on the basis of their likelihoods given a focal set of data. In this article, we use the genomic clines approach, in addition to geographic clines (Barton and Hewitt 1985), to make a formal comparison of introgression in two transects across the house mouse hybrid zone in Central Europe.

The house mouse species *Mus domesticus* and *M. musculus* hybridize in a narrow hybrid zone that runs roughly north-south through Europe, from Denmark to Bulgaria. This zone represents secondary contact between these species, and the zone may be as young as one or two thousand years, and possibly less (Cucchi et al. 2005). *Mus domesticus* and *M. musculus* in central Europe can be easily distinguished morphologically based on coat color, tail length, and craniofacial shape (Macholan 1996). There is some evidence for weak conspecific mate preference (Laukaitis et al. 1997; Smadja and Ganem 2002; Smadja et al. 2004; Bimova et al.

2005), and some crosses between *M. musculus* and *M. domesticus* yield sterile male offspring, although the extent of sterility depends on which individuals are used for the crosses (Forejt and Ivanyi 1975; Storchova et al. 2004; Britton-Davidian et al. 2005; Vyskocilova et al. 2005; Good et al. 2008b). Hybrid mice have much higher loads of intestinal parasites than either of the parental species (Sage et al. 1986a; Moulia et al. 1993, 1995; Derothe et al. 2001). This, in addition to the hybrid sterility found in some crosses between *M. musculus* and *M. domesticus*, indicates that fitness of some hybrids is reduced relative to parental *M. domesticus* and *M. musculus*.

Although many studies of this hybrid zone have been conducted (Hunt and Selander 1973; Sage et al. 1986b; Vanlerberghe et al. 1986, 1988a,b; Tucker et al. 1992; Fel-Clair et al. 1996; Orth et al. 1996; Boissinot and Boursot 1997; Prager et al. 1997; Munclinger et al. 2002; Payseur et al. 2004; Bozikova et al. 2005; Raufaste et al. 2005; Macholan et al. 2007; Teeter et al. 2008), none have compared different transects for the same set of nuclear markers. Here, we compare a previously established transect across Bavaria (Tucker et al. 1992; Payseur et al. 2004; Teeter et al. 2008) and a newly established transect 300 km to the north using the same 41 (38 autosomal and three X-linked) single nucleotide polymorphism (SNP) markers. We use the genomic clines method to detect marker-specific patterns of introgression that deviate from neutral expectations, and to compare these patterns of introgression between transects through the hybrid zone. We also compare the genomic clines with the traditional approach of fitting "geographic" clines to population allele frequencies (Barton and Hewitt 1985; Barton and Gale 1993). Finally, we examine pairwise associations between loci. Our results reveal remarkable genomic and geographic complexity in patterns of introgression between species of house mice.

Methods

SAMPLING

Mice used in this study were collected from two transects through the *M. musculus* × *M. domesticus* hybrid zone in central Europe. The southern transect is located in the German state of Bavaria and western Austria, referred to here as the Bavarian transect, and has been reported previously (Payseur et al. 2004; Bozikova et al. 2005; Teeter et al. 2008). The northern transect is located in the German states of Thuringia, Saxony-Anhalt and Saxony (referred to here as the Saxon transect). Collection for this transect was performed by K.C. Teeter in 2001–2003, and a total of 322 *Mus* were collected from this transect (Table 1). In both transects, sampling was performed in a roughly linear, east–west manner. Transect distances (in kilometers) were calculated from the western end of the transect. The location of the hybrid zone, the two transects, and collecting localities for the Saxon mice are shown

Table 1. Collecting localities, transect distances, and number of mice per locality for the Saxon hybrid zone transect. Transect distances are calculated from the western end of the transect. Data for the Bavarian transect can be found in Teeter et al. (2008).

Locality	Name	Distance	No. of mice
1	Remderoda	0	35
2	Benkendorf	21	1
3	Doellnitz-Halle	33.6	5
4	Borau bei Weissenfels	34.2	3
5	Burgliebenau	36	1
6	Muschwitz bei Weissenfels	42.45	1
7	Zeitz	42.9	1
8	Grosspoerthen bei Zeitz	44.7	4
9	Nissma bei Kayna	52.95	3
10	Borna	69.6	6
11	Floessberg	73.5	4
12	Trebishain bei Floessberg	76.05	4
13	Thallwitz	81.3	10
14	Nischwitz	81.9	1
15	Dehnitz bei Wurzen, Family Lehne	83.85	36
16	Dehnitz/NSI	83.85	6
17	Lueptitz	85.95	43
18	Gniebitz bei Trossin	86.7	70
19	Trebelshain	90	14
20	Zschirla	91.8	1
21	Mehderitzsch/Losswig	103.5	1
22	Kreischau	104.4	2
23	Hohenlauft, by Rosswein	112.8	8
24	Troischau, by Rosswein	113.7	16
25	Wilsdruff	138.9	24
26	Lohmen	172.5	1
27	Pulsnitz	172.8	1
28	Kamenz, Museum	178.4	2
29	Kamenz OT Wiesa	179.1	6
30	Deutschbaselitz	181.5	1
31	Piskowitz	184.8	3
32	Skerbersdorf	227.1	3
33	Friedersdorf bei Goerlitz	230.6	4
34	Goerlitz, Tierpark	239.3	1
Total number of mice			322

in Figure 1. All mice were commensal (collected in or near human dwellings).

DNA EXTRACTION

DNA extraction was performed on frozen spleen or kidney tissues. All extractions for the Bavarian transect, and a subset of those for the Saxon transect, were done using standard Proteinase K/phenol-chloroform extractions. For some of the samples from the Saxon transect, a MasterPure™ DNA Purification Kit, manufactured by Epicentre Biotechnologies (Madison, WI), was used.

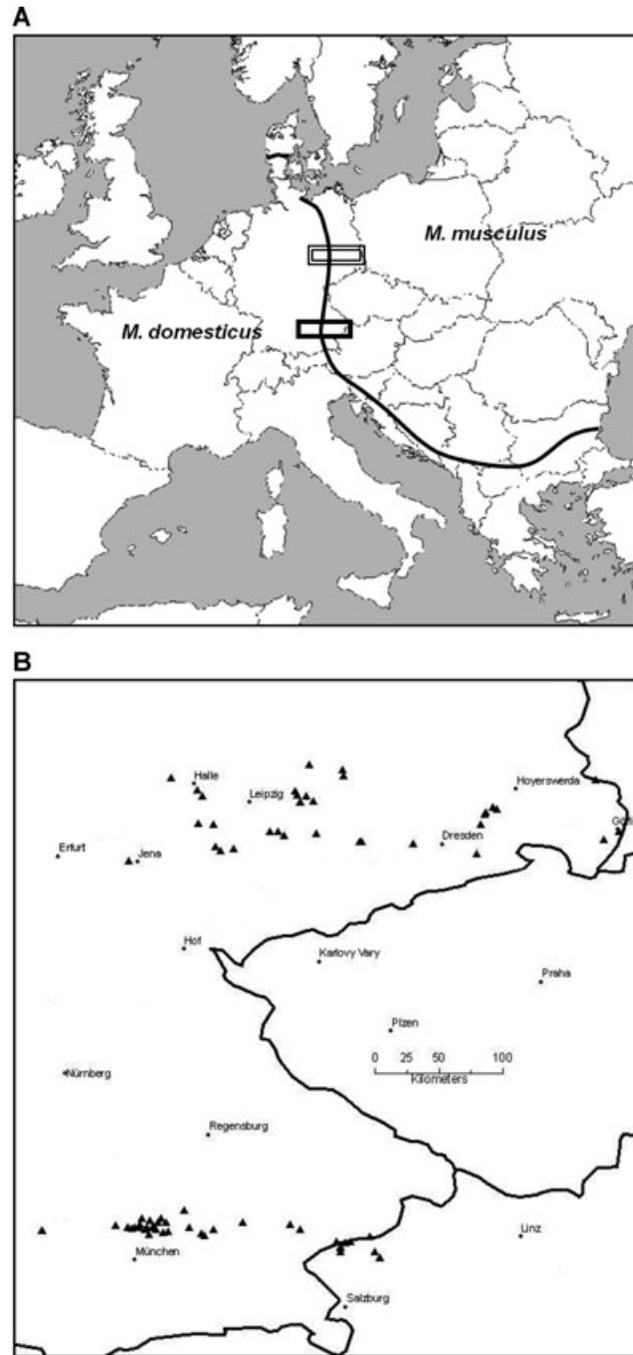


Figure 1. Location of *Mus* hybrid zone and sampling transect in Europe. (A) The solid black line shows the location of the hybrid zone. *Mus domesticus* is located to the west and south of the hybrid zone; *M. musculus* to the east and north. The outlined box shows the location of the Saxon transect, and the solid box shows the location of the Bavarian transect. (B) Detailed view of the locations of sampling localities, marked with triangles.

DEVELOPMENT AND SCORING OF MOLECULAR MARKERS

Thirty-eight autosomal markers identified from the mouse SNP database were previously determined to be diagnostic for

M. domesticus and *M. musculus* using a panel of 10 allopatric *M. domesticus* and *M. musculus* (Teeter et al. 2008). Markers were named so that an integer gives the chromosome number, and the decimal gives the approximate physical location of the marker in megabases (Mb) along the chromosome, e.g., marker 1.014 is on chromosome 1, at 14 Mb (Table 2). Exact marker locations are as in Teeter et al. (2008). All genotyping for samples from the Saxon transect was performed at the University of Michigan. One marker on chromosome 10 (10.055) scored in the Bavarian transect, failed to amplify in mice from the Saxon transect, and was not included in this study. For the Bavarian transect, odd-numbered markers were scored at the University of Michigan, and even-numbered markers at the University of Arizona (Teeter et al. 2008). Genotyping for autosomal markers was completed using TaqMan probes and chemistry from Applied Biosystems (Foster City, CA). Genotyping for three X-linked markers, *Emd*, *Pola1*, and *Xist* was completed with PCR-RFLP methods, following Payseur et al. (2004).

GENOMIC CLINE ANALYSES

We summarized the ancestry of individual mice with a hybrid index, which is simply the fraction of alleles at the 38 autosomal loci that were inherited from *M. musculus*. This summary of genome-wide admixture in individuals was used to predict the probabilities of observing each of the three possible genotypes at focal loci, which are referred to as genomic clines (Gompert and Buerkle 2009a). The clines were estimated using multinomial regression of the observed genotypes on hybrid index.

To identify loci that do not conform to expectations of neutral introgression, the likelihoods of the regression model and of a neutral model (both given the observed data) can be compared. We used a permutation procedure to simulate neutral introgression (Gompert and Buerkle 2009a), which is based on the logic that all loci should be exchangeable under neutrality and appropriately retains the overall structure of the sampled population, as well as accounting for stochastic variation among loci that would result from genetic drift. We summarized deviations from neutrality on the basis of: (1) whether homozygotes (*M. musculus* or *M. domesticus*) were more or less common than expected under neutrality, which corresponds to expectations under positive or negative selection, and (2) whether heterozygotes were more or less common than expected under neutrality, which corresponds to expectations of over- and under-dominance (as in Nolte et al. 2009). Additionally, we searched for evidence of pairwise associations between alleles at different loci, which might be caused by epistasis, by adding the genotype at a potentially interacting locus to a regression model and determining whether this information improved the fit of the model (using AIC; in the basic regression model, the other predictors of genotype at the focal locus were hybrid index, and genome-wide heterozygosity).

To quantify differences in the genomic clines from the two transects through the mouse hybrid zone, we used a ratio of the likelihoods of the genomic clines (models), given the data from one of the transects ($\ln L(M_{\text{Sax}} | D_{\text{Sax}}) / L(M_{\text{Bav}} | D_{\text{Sax}})$). The null distribution of the likelihood ratios was determined by 1000 replicate simulations in which individuals were permuted between transects.

All analyses associated with genomic clines were performed using the R package INTROGRESS (Gompert and Buerkle 2009b) and additional functions written by the authors. Significance thresholds for genomic clines analyses, including tests for genotype-specific deviations, were adjusted using the false discovery rate procedure (Benjamini and Hochberg 1995).

GEOGRAPHIC CLINE ANALYSES

The shape of geographic clines was estimated individually for each marker using a two-parameter model and the software Clin-eFit (Porter et al. 1997). The simple two-parameter model of cline shape uses cline center and width to describe the cline shape along the length of the transect and was used as in Teeter et al. (2008). The two-parameter model was chosen rather than the more complex six-parameter models for clines (Barton and Hewitt 1985; Barton and Bengtsson 1986; Barton and Gale 1993), because the likelihood surface for the more complex model can be very flat and uninformative, and optima can be difficult to find (results not shown; Raufaste et al. 2005; Macholan et al. 2007).

The data for the X chromosome markers from the Bavarian transect are from Payseur et al. (2004). In the Payseur paper, these markers were analyzed using six-parameter models, whereas here we have used two-parameter cline models to compare data from the two transects. These models return wider cline widths compared to the six-parameter models.

Spearman nonparametric rank correlation tests were used to detect correlations between cline widths and centers for each marker. Correlation tests were also used to evaluate similarity in cline shape between the Saxon transect and the Bavarian transect (Teeter et al. 2008), by comparing cline widths and centers in each transect. These tests were performed in SPSS 11.0 for Macintosh OS X.

Results

GENOTYPING

Thirty-eight autosomal markers and three X-linked markers were scored in mice from both hybrid zone transects (Teeter et al. 2008; Table S1). Hybrid indices for all samples plotted against interspecific heterozygosity (Fig. 2A) indicated that the sampling from the two transects does not result in the same distributions of genomic admixture, with a nearly continuous distribution of hybrids in the Bavaria transect and few intermediate hybrids in

Table 2. Genomic and geographic cline analyses for genetic markers typed in mice collected from the Saxony and Bavaria transects. Markers were named so that an integer gives the chromosome number, and the decimal gives the approximate physical location of the marker in Mb along the chromosome. For the X chromosome markers, a gene name is given. LnL ratios, *P* values, and deviation categories are derived from the genomic clines analyses. The cline centers (CC) and widths (CW) are derived from the two-parameter geographic cline analyses. n, nonsignificant; *, significant; NE, not estimated; +, excess; -, deficit of genotype; respectively following false discovery rate correction (Benjamini and Hochberg 1995). Significant deviations may exist for the probabilities of individual genotypes without an overall significant deviation for all three genotypes at a locus. Genomic clines from the two transects were compared using a ratio of the likelihoods of the genomic clines given the data from one of the transects ($\ln L(M_{Sax}|D_{Sax})/L(M_{Bay}|D_{Sax})$).

Marker name	Saxony					Bavaria					Comparison	
	LnL ratio	<i>P</i>	Deviation category	CW-S (km)	CC-S (km)	LnL ratio	<i>P</i>	Deviation category	CW-B (km)	CC-B (km)	LnL ratio	Comparison <i>P</i>
1.014	15.861	0	*(DD:+ DM:- MM:-)	60	127.4	7.45	0.003	*(DD:+ DM: MM:)	18.5	60.9	18.83	*0.009
1.046	2.638	0.226	n (DD: DM: MM:)	44.3	113.8	17.91	0	*(DD:+ DM: MM:-)	81.9	75.7	3.41	n 0.436
1.159	7.384	0.012	*(DD: DM:+ MM:-)	79.4	123	96.96	0	*(DD:- DM: MM:+)	36.1	52.1	81.93	*0
2.03	33.531	0	*(DD:+ DM:- MM:+)	13.8	102.5	12.4	0	*(DD: DM:- MM:+)	20.7	58.4	33.34	*0
2.078	5.218	0.052	n (DD: DM: MM:)	29.2	105.7	80.98	0	*(DD:- DM:+ MM:+)	74.9	50.5	70.26	*0
2.165	9.273	0.004	*(DD: DM: MM:)	38.3	111.1	12.98	0	*(DD:+ DM: MM:)	21.1	61.7	2.24	n 0.668
3.007	6.042	0.019	*(DD:- DM: MM:)	44.3	107.6	235.54	0	*(DD:- DM:+ MM:+)	101.9	30.9	176.62	*0
3.14	17.09	0	*(DD: DM:+ MM:)	41.1	111.2	12.93	0	*(DD: DM: MM:-)	65.1	69.9	6.37	n 0.19
4.057	45.59	0	*(DD:+ DM: MM:-)	120.4	163.7	45.56	0	*(DD:+ DM:+ MM:-)	140.4	99.9	13.8	*0.01
4.129	30.204	0	*(DD: DM:- MM:+)	17.7	102.1	40.78	0	*(DD: DM:- MM:+)	13	57.5	7.24	n 0.141
5.007	52.262	0	*(DD:- DM: MM:+)	29.6	96.1	79.95	0	*(DD:- DM:+ MM:+)	45.8	53.9	69.52	*0
5.097	45.238	0	*(DD:+ DM: MM:-)	61.4	139.3	66.15	0	*(DD:- DM:- MM:+)	6.5	57.7	143.56	*0
6.088	3.589	0.13	n (DD: DM:- MM:)	36.1	108.4	25.73	0	*(DD: DM:- MM:+)	15.4	57.9	14.97	*0.021
6.113	4.394	0.077	n (DD: DM: MM:)	44	109.4	16.75	0	*(DD: DM: MM:)	26.3	62	13.07	*0.012
7.083	25.902	0	*(DD:- DM: MM:+)	80.7	112.6	100.92	0	*(DD:+ DM:+ MM:-)	172.4	118.2	123.8	*0
7.126	16.519	0	*(DD:- DM:+ MM:)	64.3	111.5	3.37	0.124	n (DD: DM: MM:)	62	68.7	8.76	n 0.048
8.078	3.903	0.104	n (DD: DM: MM:+)	35	104.4	21.49	0	*(DD:+ DM: MM:-)	173.4	95.1	18.72	*0.001
8.101	43.721	0	*(DD:- DM: MM:+)	23.4	97.8	17.6	0	*(DD: DM: MM:)	12.9	58.1	40.58	*0
9.052	19.79	0	*(DD: DM:- MM:)	28.7	107.5	32.49	0	*(DD:- DM: MM:+)	14.8	56.3	19.47	*0.002
9.075	316.091	0	*(DD:- DM:+ MM:+)	66.5	73.3	61.26	0	*(DD: DM:- MM:+)	6.4	56.3	784.18	*0

Continued.

Table 2. Continued.

Marker name	Saxony						Bavaria						¹ Comparison	
	LnL ratio	P	Deviation category	CW-S (km)	CC-S (km)	LnL ratio	P	Deviation category	CW-B (km)	CC-B (km)	LnL ratio	Comparison	P	
10.045	8.291	0.004	*(DD:+ DM: MM:-)	68.9	126	28.91	0	*(DD:- DM: MM:+)	131.9	63.6	65.87	*0		
11.053	8.558	0.002	*(DD: DM:+ MM:)	87.7	122.8	15.9	0	*(DD: DM: MM:)	28.6	60.7	31.94	*0		
11.089	4.604	0.049	n (DD: DM: MM:)	32.1	108.8	32.75	0	*(DD:+ DM:- MM:+)	10.1	58.2	7.72	n 0.091		
12.031	5.75	0.027	*(DD: DM: MM:)	49.3	111.7	118.25	0	*(DD:+ DM:- MM:-)	125.2	112.5	86.46	*0		
12.099	6.94	0.01	*(DD: DM: MM:)	45.1	112.1	23.92	0	*(DD:- DM: MM:+)	23.6	57.1	14.41	*0.007		
13.029	72.252	0	*(DD:+ DM: MM:-)	72.2	156	150.4	0	*(DD:+ DM:+ MM:-)	341.8	176.4	15.04	*0.002		
13.056	7.713	0.011	*(DD:- DM: MM:+)	52.5	108.3	56.71	0	*(DD:- DM: MM:+)	26.5	54.6	29.45	*0		
14.031	7.862	0.01	*(DD: DM: MM:)	55.3	119.2	6.44	0.01	*(DD:- DM:+ MM:)	95.6	69.9	27.48	*0		
14.074	12.32	0.002	*(DD:+ DM: MM:)	37.4	113.4	18.64	0	*(DD:+ DM: MM:-)	84.4	78.5	2.44	n 0.649		
15.065	12.308	0	*(DD:+ DM: MM:-)	76.3	130.1	65.99	0	*(DD:+ DM: MM:-)	91.2	89.1	27.2	*0.004		
15.099	23.235	0	*(DD: DM:- MM:)	20.7	101.9	24.72	0	*(DD: DM:+ MM:)	25.7	61	46.3	*0		
16.014	5.948	0.023	*(DD: DM: MM:)	18.4	107.2	38.12	0	*(DD: DM:- MM:+)	7	56.5	9.5	n 0.092		
17.046	6.187	0.02	*(DD: DM: MM:-)	93.1	130.9	68.09	0	*(DD:- DM: MM:+)	14.7	54.6	54.52	*0.001		
17.091	11.458	0.001	*(DD:+ DM: MM:-)	74.7	132.2	118.26	0	*(DD:+ DM:- MM:-)	54.9	87.2	27.52	*0.033		
18.028	15.983	0	*(DD:+ DM: MM:-)	84.5	135.7	69.43	0	*(DD:+ DM: MM:-)	135.4	105	6.22	n 0.141		
18.064	5.911	0.023	*(DD: DM: MM:)	37	110	28.68	0	*(DD:+ DM:- MM:)	20.6	61.7	9.07	n 0.064		
19.044	15.806	0	*(DD: DM: MM:+)	25.6	100.4	11.55	0	*(DD: DM: MM:)	16.2	58.4	13.58	n 0.042		
19.052	13.795	0	*(DD:+ DM: MM:-)	86.9	136.5	18.26	0	*(DD:+ DM:+ MM:-)	72.2	75	10.52	*0.028		
<i>Emd</i>	10.104	0.007	*(DD: DM:- MM:)	18.4	101.4	50.31	0	*(DD: DM:- MM:+)	5.14	55.39	3.44	n 0.513		
<i>Potal</i>	20.66	0	*(DD:+ DM:- MM:)	NE	NE	48.24	0	*(DD: DM:- MM:+)	4.41	55.02	2.35	n 0.622		
<i>Xist</i>	21.316	0	*(DD:+ DM:- MM:)	15.2	102.5	25.93	0	*(DD:+ DM: MM:-)	295.19	108.75	39.56	*0		
Mean	24.420512	0.0198293		50.2	114.9	48.9926829	0.0033415		66.3	70.8	53.2014634	0.0933415		
Median	12.308	0.001		44.3	111.1	32.49	0		28.6	60.9	18.83	0.004		

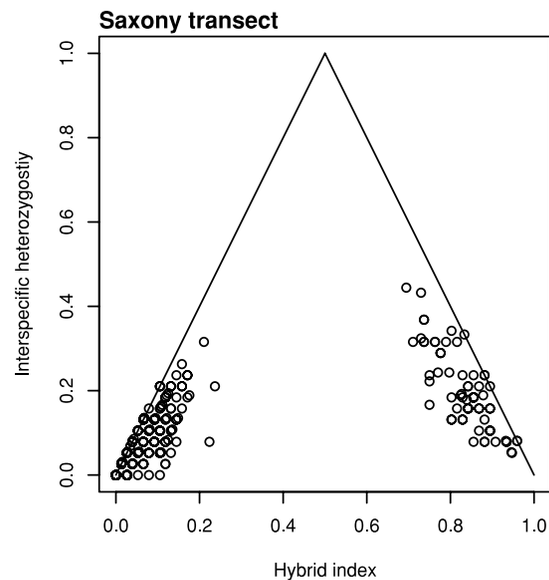
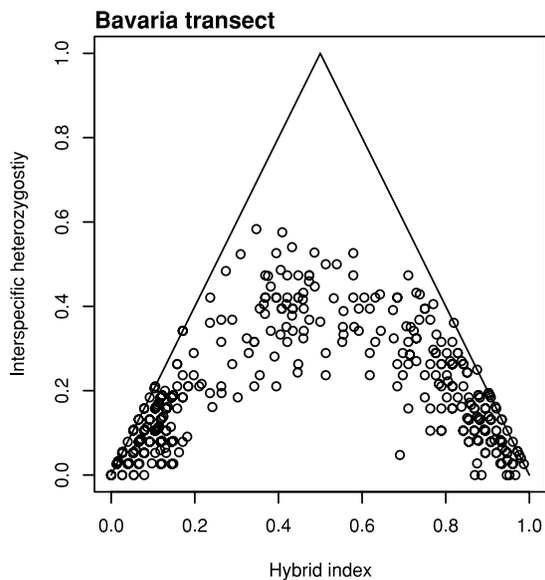
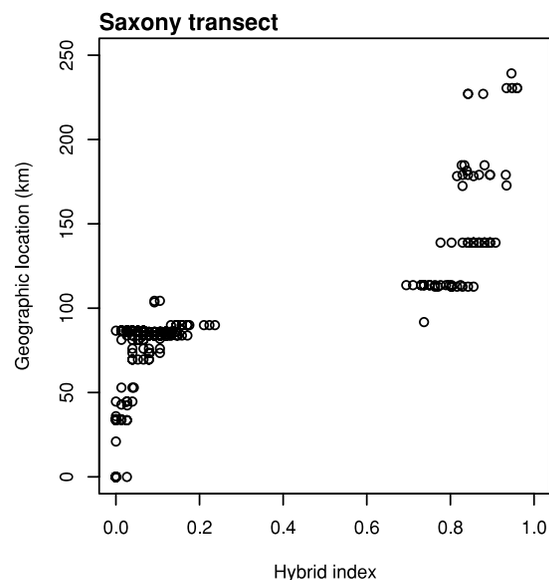
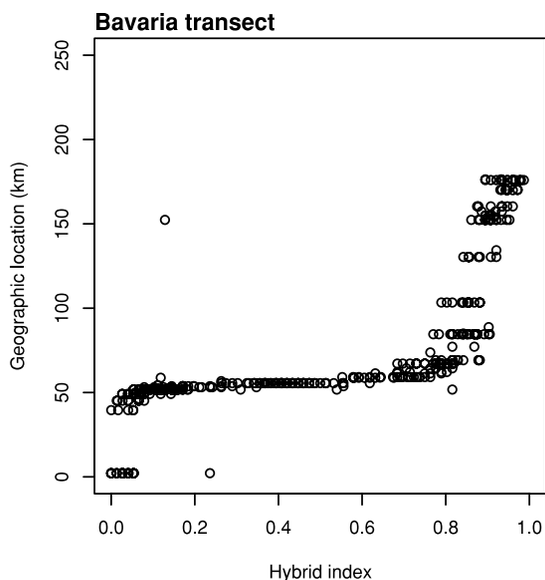
A Hybrid index vs. heterozygosity**B** Hybrid index vs. geographic location

Figure 2. Heterozygosity of individual mice versus (A) hybrid index, measured as the proportion of alleles with *M. musculus* ancestry, and (B) geographic location of mouse collection sites measured in kilometers from western-most locality versus hybrid index, of individuals at those sites.

the Saxony transect. Hybrid indices for all samples plotted against distance from the western-most locality showed roughly similar patterns between transects, with populations on the *M. musculus* side of the hybrid zone having a greater variance of hybrid indexes among individuals (Fig. 2B; Fig. S1).

GENOMIC CLINE ANALYSES

There was extensive heterogeneity among loci in the patterns of introgression between species, which was visually evident in the raw data (Fig. 3) and in the fitted genomic clines (Fig. 4). There

was also statistical evidence for heterogeneity among loci, in the form of significant deviations from neutrality (based on exchangeability of loci) for the majority of markers (Table 2). Deviations from neutrality included loci with excess introgression into the genomic background of each species (e.g., excess *M. musculus* in *M. domesticus* background: 1.159 and 3.007 in Bavaria, and excess *M. domesticus* in *M. musculus* background: 12.031 (Bavaria) and 17.091 (Bavaria and Saxony); Fig. S2) and a few loci that exhibited patterns of introgression that were consistent with under-dominance (*Emd* and *Pola1*).

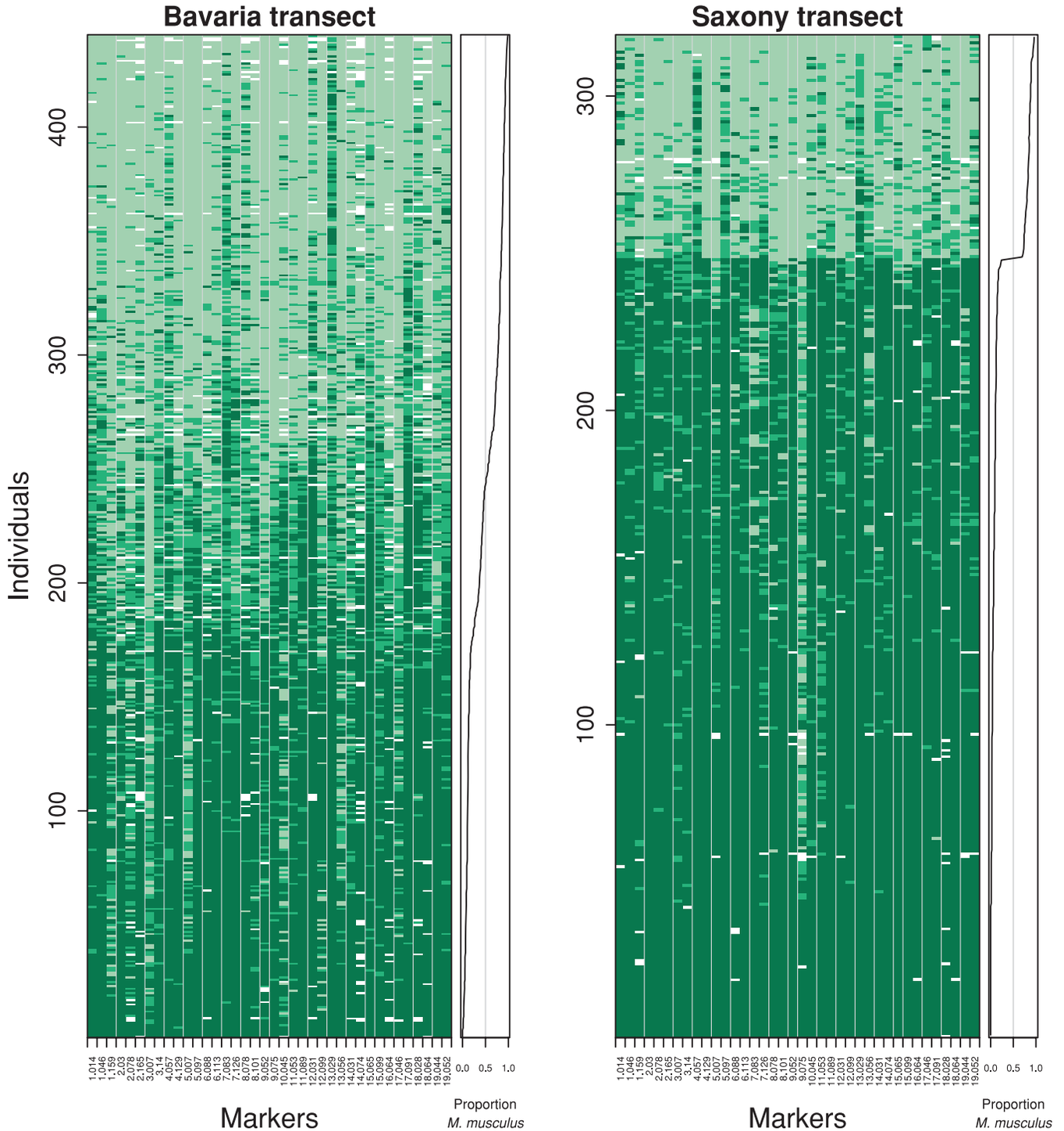


Figure 3. Genotypes of mice in two transects across the European hybrid zone. Markers are on the 19 autosomes and are named according to the chromosome on which they are found and their position on the physical map (as in Teeter et al. 2008). Dark green blocks indicate homozygotes for *M. domesticus* alleles, light green blocks represent homozygotes for *M. musculus* alleles, and intermediate green blocks correspond to heterozygotes. White blocks indicate missing data. The plots to right in each pane indicate the proportion of each individual's genome that has *M. musculus* ancestry, which is equivalent to the hybrid index. Individuals are sorted, with those individuals with genome compositions that resemble *M. domesticus* at the bottom and increasing similarity to *M. musculus* toward the top.

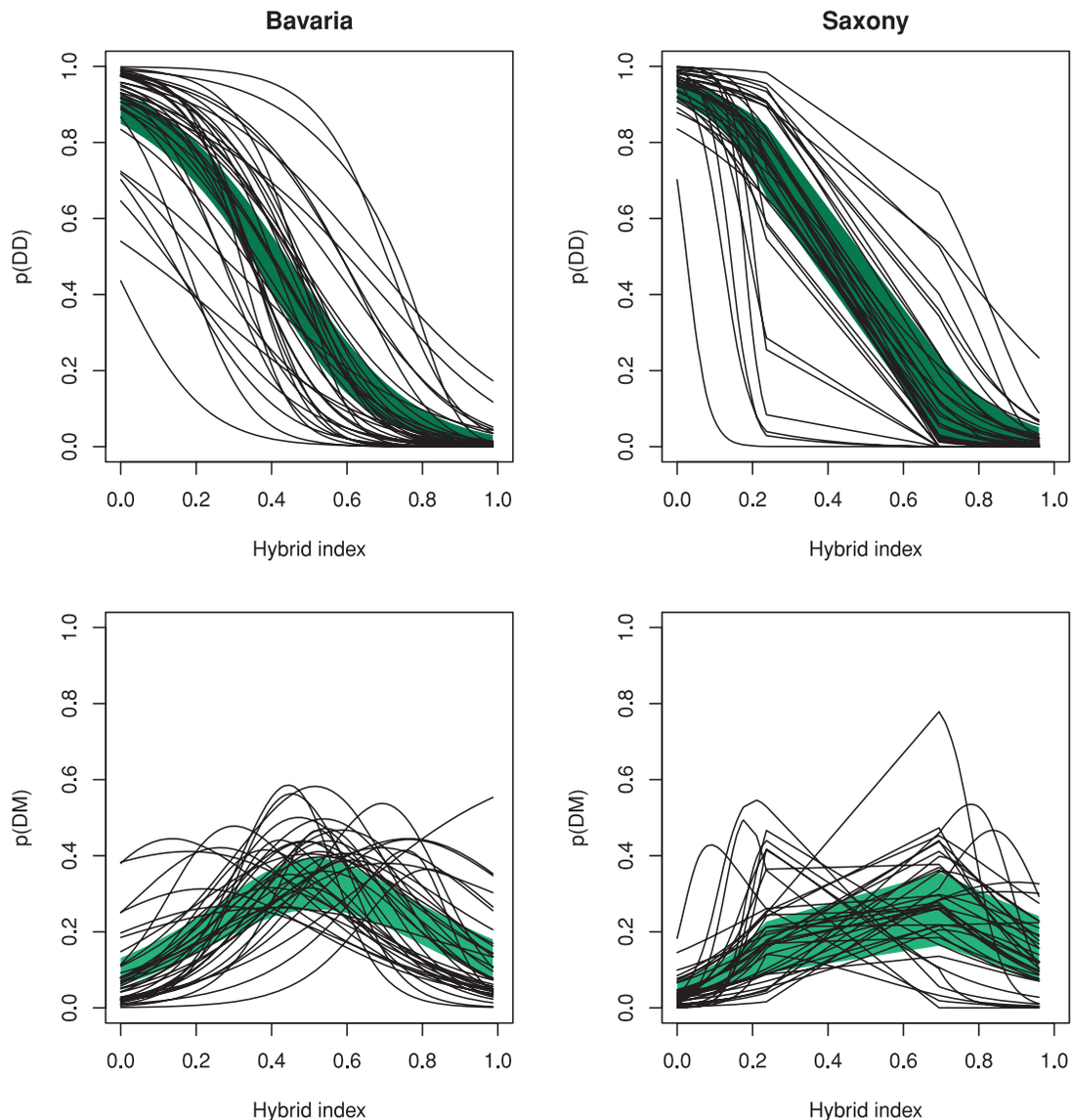


Figure 4. Genomic clines for homozygous *M. domesticus* and heterozygous genotypes for the Bavaria and Saxony hybrid zones. Hybrid index corresponds to the proportion of marker alleles with *M. musculus* ancestry. The dark green and light green shaded regions denote the expected genomic clines (95% CI) for the homozygous *M. domesticus* and heterozygous genotypes, respectively. Solid black lines denote the genomic clines for individual loci based on multinomial regression models for the homozygous (top panels) and heterozygous (bottom panels) genotypes.

In addition to variation among loci, the genomic clines from the two transects were significantly different for 28 of the 41 loci (Table 2). Equivalent differences between transects were observed even if the analysis involved only individuals from the Bavaria transect with hybrid indexes that fell within the distribution of hybrid indexes in the Saxony transect.

ASSOCIATIONS BETWEEN ALLELES AT DIFFERENT LOCI

The Bavaria transect offered stronger evidence for nonrandom associations between loci than did the Saxony transect (Fig. S3). This was likely due to the difference in distribution of hybrid index

values in the two transects. Within the Bavaria transect, there were many nonrandom associations between loci (Fig. S3), and 98.8% of all pairwise associations between loci involved alleles derived from the same species, as evidenced by a consistently positive sign of the regression coefficient for the predictor locus. Importantly, these associations exist after accounting for ancestry through hybrid index and for genome-wide heterozygosity.

GEOGRAPHIC CLINE ANALYSES

Estimates of cline width for the Saxon transect ranged from 13.8 to 120 km, and estimates of the cline center ranged from 73.3 to 163.7 km along the transect (Table 2, Table S2). The mean

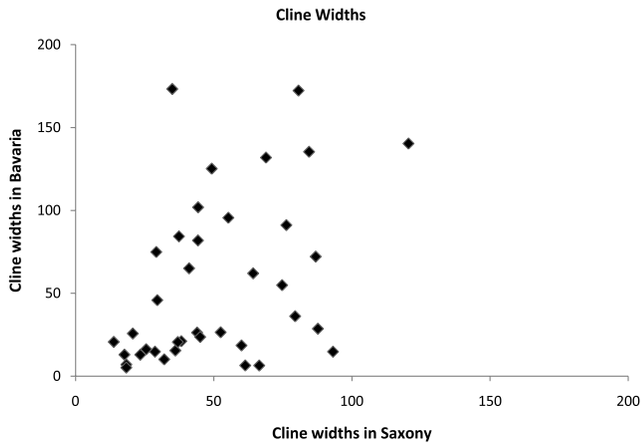


Figure 5. Cline widths (two-parameter) from the Bavarian transect (Teeter et al. 2008), plotted against cline widths from the Saxon transect. A linear regression of the cline widths for the autosomes in the Saxon and Bavarian transects gives an r^2 value of 0.144.

cline center from these models for the Saxon transect was located at 114.9 km along the transect, and the mean cline width was 50.2 km. The positions of the cline width and the cline center from the two-parameter model estimates of autosomal markers were found to have a strong positive correlation in both transects (Bavaria: Spearman's $\rho = 0.618$, $P < 0.001$, Saxony: Spearman's $\rho = 0.812$, $P < 0.001$) (Fig. S4), indicating that the wider clines had centers shifted toward the eastern end of the transects. The 13.029 marker had an estimated cline width of 341.8 km in Bavaria, longer than the actual transect, and therefore it was excluded from these analyses as an outlier. The positions of the cline centers from the two-parameter model estimates of autosomal markers were significantly correlated between the two transects (Spearman's $\rho = 0.458$, $P = 0.003$), as were the cline widths (Spearman's $\rho = 0.371$, $P = 0.018$), indicating that markers show some similarity between geographic clines in both transects. Although there was variation in cline width between the two transects for many markers, there is a set of markers that have narrow cline widths (low introgression) in both transects (Fig. 5).

Discussion

Geographic and genomic cline analyses of markers reveal remarkable differences between the Bavarian and Saxon transects, as well as a few similarities. The differences between transects raise the possibility that there may not be a single genetic architecture of isolation between these species. In addition, it is likely that genetic drift has occurred independently in each of these transects and thereby contributed to these differences. The analyses of clines also identified significant diversity among loci. Clines for some loci were consistent with selection against hybrid genotypes and limited introgression, whereas clines for other

loci offered evidence for positive selection, in the form of genotypes introgressed far into a foreign genetic background. Next we discuss each of the results and conclusions in greater detail.

GENOMIC CLINE ANALYSES

In comparing the two transects through the hybrid zone (Fig. S2), we find that 28/41 markers differ significantly between transects, whereas 13/41 do not (1.046, 2.165, 3.14, 4.129, 7.126, 11.089, 14.074, 16.014, 18.028, 18.064, 19.044, *Emd*, and *Polal*). For the 28/41 markers that differ between transects, it is possible that stochastic variation, differences in sampling between transects, or a combination, could have contributed to these differences. However, given that the majority of these markers were significantly different from the null model of introgression in one or both transects (Table 2), another explanation is that the mouse populations in the transects have experienced different histories of natural selection. Genetic factors contributing to reproductive isolation may be polymorphic in this hybrid zone system. Polymorphism for factors contributing to sterility has previously been documented in other hybridizing taxa (e.g., Reed and Markow 2004) as well as in the house mouse (Vyskocilova et al. 2005; Good et al. 2008b). Additionally, ecological differences (both biotic and abiotic) between these transects may affect which genomic regions contribute to reproductive isolation. Among the 13 markers that have consistent patterns of introgression in the two transects, two markers on the X chromosome (*Emd*, and *Polal*) as well as a few autosomal markers (e.g., 2.030, 4.129) show a deficiency of heterozygotes in each transect, suggesting the presence of nearby genes with consistent, possibly intrinsic, negative effects on fitness in heterozygotes. This could be caused by simple under-dominance or by classic Dobzhansky–Muller incompatibilities.

Genomic cline analyses also reveal a diversity of patterns of introgression among loci. Clines for the majority of markers were inconsistent with neutral introgression in hybrids (40/41 in Bavaria and 35/41 in Saxony; Table 2). Excess and deficits of the three genotypes at each locus (Table 2) are consistent with the action of selection at linked genes. Whereas the permutation approach for testing for deviations from the null model incorporates stochastic variation among loci (including increased variance due to their independent genetic drift), some deviations could result from the action of genetic drift, particularly if drift occurred independently at different sampling localities along the transect (Gompert and Buerkle 2009a). We also note that the sensitivity of the model to complicated forms of population and demic structure, such as are known to exist in house mice, has not been explored.

The most common type of deviation in the Saxon transect (8 out of 41) is DD+, DM, MM–, consistent with positive selection for homozygous *M. domesticus* alleles. However, the

most common type of deviation in the Bavarian transect (7 out of 41) is DD, DM−, MM+, consistent with positive selection for homozygous *M. musculus* alleles. It is not possible to determine whether any specific deviations are false positives with the available data. Functional assays in controlled crosses would be useful for testing the fitness effects of individual loci. Future modeling will also help us understand how population structure within a hybrid zone affects inferences based on genomic clines. More markers per chromosome will allow fine-scale mapping and will determine whether the major factors associated with fitness variation can be identified. Nevertheless, the diversity among loci and among transects highlights the remarkable complexity of genes and geography in this hybrid zone.

GEOGRAPHIC CLINE ANALYSES

Markers that have narrow geographic clines in both transects include *Emd*, 16.014, 8.101, 4.129, and 9.052. It was not possible to estimate a valid two-parameter cline for the *Polal* marker in the Saxon transect using the ClineFit program, but this marker has very limited introgression in Saxony, and a narrow cline in Bavaria (Fig. S1). These markers represent good candidate regions for genes involved in reproductive isolation between *M. domesticus* and *M. musculus*. Linkage disequilibrium analyses in Teeter et al. (2008) found conspecific linkage disequilibrium between 9.052 and several X-linked and autosomal markers, which indicates that this region may be involved in Dobzhansky–Muller incompatibilities (Dobzhansky 1937; Muller 1942; Coyne and Orr 2004). Additionally, *Emd*, 16.014, and 4.129 also participate in significant conspecific associations with other markers, and in the genomic analysis they show similar patterns of introgression in hybrid mice from both transects.

Some markers show a pattern along the transects where there is a transition from *M. domesticus* to *M. musculus* genotypes over a short distance near the center of the hybrid zone, but *M. domesticus* alleles then reappear at higher frequencies further from the center of the hybrid zone (e.g., 15.099 and 16.014, Fig. S1). This pattern is suggestive of stronger selection in the center of the hybrid zone and weaker selection further away from it (at least for the *M. domesticus* alleles on a *M. musculus* genetic background). Alternatively, it is possible that some markers are not fixed for different alleles in *M. domesticus* and *M. musculus*, and may instead be shared polymorphisms. In the genomic analysis, some of these markers (e.g., 16.014) show similar patterns of introgression in hybrid mice from both transects whereas others (e.g., 15.099) do not.

Local geography may determine the location of the hybrid zone in some cases, as suggested by Raufaste et al. (2005) for the *Mus* hybrid zone in Denmark. However, there is no clear association of the position of the hybrid zone in either the Saxon or the Bavarian transect with local geographic features. Thus, in the cases in which geographic clines differ between transects,

variation in the local environment or genetic variation could play a role.

ASSOCIATIONS BETWEEN LOCI

The Dobzhansky–Muller model of reproductive isolation is based on epistatic interactions among alleles at different genes. Such epistasis can give rise to nonrandom associations among alleles (i.e., linkage disequilibrium). Positive associations of conspecific genotypes are pervasive in this dataset (Fig. S3), and appear to be particularly strong in the data from the Bavarian transect. Nearly all (98.8%) of the pairwise associations found in the Bavarian transect were between alleles derived from the same species (after accounting for ancestry through hybrid index and for genome-wide heterozygosity). The set of markers with particularly strong associations differs between transects. This observation coupled with the large number of markers involved in significant associations (most combinations did not involve physically-linked markers) raises the possibility of a highly complex basis for reproductive isolation between these taxa, with a web of many interacting loci contributing to isolation. Although these associations could result from divergent selection on the two taxa, as shown in experimental populations of yeast by Dettman et al. (2007), they may also have arisen through other population genetic mechanisms and in the absence of selection, as shown in a grasshopper (*Chorthippus*) hybrid zone by Shuker et al. (2005).

SYMMETRICAL AND ASYMMETRICAL PATTERNS OF INTROGRESSION

Although at individual loci there was significant introgression into both parental genomic backgrounds (Fig. 3), the overall genomic composition of hybrids, as summarized by hybrid index, shows evidence for biased gene flow from populations in which *M. domesticus* alleles predominate into populations dominated by *M. musculus* (Fig. 2B; Fig. S2). In particular, populations at the eastern, *M. musculus*, end of both transects have higher variability in the hybrid indexes of individuals, consistent with a higher rate of gene flow and mixture of populations. These data suggest some decoupling and independence between geography and genetic background and suggest that geographic cline analyses, alone, may not provide an accurate view of gene flow through a hybrid zone. Data from previous studies of the *Mus* hybrid zone also indicate asymmetric patterns of gene flow, biased in the direction from *M. domesticus* into *M. musculus* populations (Vanlerberghe et al. 1988a; Tucker et al. 1992; Fel-Clair et al. 1996; Boissinot and Boursot 1997; Raufaste et al. 2005). However, Munclinger et al. (2002) and Macholan et al. (2007) found opposite patterns for some markers.

A possible explanation for the observed asymmetrical clines is that the hybrid zone has shifted over time, and some loci are “trailing” the majority of the genome. Demographic patterns in

M. domesticus and *M. musculus* could have contributed to this shift. Alternatively, asymmetry in introgression may be due to asymmetric genetic incompatibilities. Studies using experimental crosses between strains of *M. domesticus* and *M. musculus* have identified some genome segments that are associated with hybrid sterility (Vyskocilova et al. 2005; Oka et al. 2007; Good et al. 2008a). In particular, introgression of the *M. musculus* X chromosome onto a *M. domesticus* genetic background causes male sterility in many cases, and there is polymorphism in wild populations for sterility factors (Oka et al. 2004; Britton-Davidian et al. 2005; Good et al. 2008b). It also is possible that behavioral factors have influenced cline shape and patterns of introgression. Mate preference and genetic incompatibilities may interact, as the signals that determine mate selection in these species are at least in part genetically determined. There is behavioral evidence that *M. domesticus* mice tend to dominate in male–male conflicts (Munclinger and Frynta 1997, 2000; Frynta et al. 2005). This evidence suggests that *M. domesticus* could disperse more easily into *M. musculus* territories than vice versa (van Zegeren and van Oortmerssen 1981).

OVERALL PATTERNS AND THE COMPLEX NATURE OF GENOME INTERACTIONS IN HYBRID MICE

We have used two complementary methods of analysis to explore this dataset. The genomic clines method identified features of this hybrid zone that were not apparent from the geographic clines alone. The differences found between the two transects highlight the challenges of using patterns of introgression in hybrid zones to identify a common set of genes underlying reproductive isolation. However, the markers with similar, nonneutral patterns of introgression in both transects are good candidates for further study of invariant components of isolation. The extensive positive associations of conspecific alleles detected in the hybrid zone contributes an additional component to our picture of isolation and speciation.

Although there have been multiple excellent experimental mapping studies performed to identify loci involved in isolation between *M. domesticus* and *M. musculus*, the patterns observed in this natural system provide a more complex scenario than what might be predicted from experimental studies. Our results illustrate the importance of using a combination of studies of natural populations and laboratory studies in constructing a model of speciation. More detailed studies of the hybrid genomes, including denser sampling of the genome in a greater number of hybrid mice, will help illuminate the specific mode of selection acting to create isolation between these taxa and complementary studies on fitness and behavior of mice from the hybrid zone, such as studies of hybrid sterility (Forejt and Ivanyi 1975; Storchova et al. 2004; Britton-Davidian et al. 2005; Vyskocilova et al. 2005; Good et al. 2008a,b), mate choice preference (Laukaitis et al. 1997; Talley

et al. 2001; Smadja and Ganem 2002; Smadja et al. 2004; Bimova et al. 2005), and susceptibility to parasites (Sage et al. 1986a; Mouliia et al. 1993; Derothe et al. 2001; Derothe et al. 2004), may help link specific phenotypes to the patterns of introgression documented here.

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Supporting Information

The following supporting information is available for this article:

Table S1. Genotype data from all 41 markers used for mice collected from the Saxon transect.

Table S2. Two-parameter cline estimates generated in ClineFit for each marker used for the Saxon transect sample.

Figure S1. Scatterplots for data from the Saxon and Bavarian transects for all 41 markers used.

Figure S2. Comparison of genomic clines for the Bavaria and Saxony hybrid zones.

Figure S3. Plot of pairwise epistatic interaction for the two hybrid zone transects.

Figure S4. Plots of two-parameter estimates of cline width versus two-parameter estimates of cline center.

Supporting Information may be found in the online version of this article.

(This link will take you to the article abstract).

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