Selection on cis-Regulatory Variation at B4galnt2 and Its Influence on von Willebrand Factor in House Mice

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The RIIRS/J inbred mouse strain is a model for type 1 von Willebrand disease (VWD), a common human bleeding disorder. Low von Willebrand factor (VWF) levels in RIIRS/J are due to a regulatory mutation, Mvwf1, which directs a tissue-specific switch in expression of a glycosyltransferase, B4GALNT2, from intestine to blood vessel. We recently found that Mvwf1 lies on a founder allele common among laboratory mouse strains. To investigate the evolutionary forces operating at B4galnt2, we conducted a survey of DNA sequence polymorphism and microsatellite variation spanning the B4galnt2 gene region in natural Mus musculus domesticus populations. Two divergent haplotypes segregate in these natural populations, one of which corresponds to the RIIRS/J sequence. Different local populations display dramatic differences in the frequency of these haplotypes, and reduced microsatellite variability near B4galnt2 within the RIIRS/J haplotype is consistent with the recent action of natural selection. The level and pattern of DNA sequence polymorphism in the 5´ flanking region of the gene significantly deviates from the neutral expectation and suggests that variation in B4galnt2 expression may be under balancing selection and/or arose from a recently introgressed allele that subsequently increased in frequency due to natural selection. However, coalescent simulations indicate that the heterogeneity in divergence between haplotypes is greater than expected under an introgression model. Analysis of a population where the RIIRS/J haplotype is in high frequency reveals an association between this haplotype, the B4galnt2 tissue-specific switch, and a significant decrease in plasma VWF levels. Given these observations, we propose that low VWF levels may represent a fitness cost that is offset by a yet unknown benefit of the B4galnt2 tissue-specific switch. Similar mechanisms may account for the variability in VWF levels and high prevalence of VWD in other mammals, including humans.

Introduction

von Willebrand factor (VWF) is a central component of hemostasis, and defects in VWF result in von Willebrand disease (VWD), the most common inherited bleeding disorder in humans (Johnsen and Ginsburg 2006). Type 1 VWD accounts for 70% of all of VWD cases and is characterized by a quantitative reduction in VWF. The RIIRS/J inbred mouse strain has a phenotype similar to Type 1 VWD in humans (Sweeney et al. 1990). The low VWF levels in RIIRS/J are due to a mutation previously termed Mvwf1, for “Modifier of” VWF 1 (Mohlke et al. 1996). Mvwf1 is a naturally occurring, cis-regulatory mutation that results in a unique, tissue-specific switch in the expression of a glycosyltransferase, B4GALNT2, from the intestinal epithelial pattern seen in most mice (and humans; Montiel et al. 2003) to a vascular endothelial cell-specific pattern. Endothelial cell expression of B4GALNT2 leads to aberrant posttranslational modification of VWF, resulting in accelerated clearance from circulation and a reduced plasma VWF phenotype (Mohlke et al. 1999).

Further characterization of the B4galnt2 locus in RIIRS/J identified a ~30-kb genomic region of high sequence divergence (>2% relative to C57BL6/J) flanking B4galnt2 exon 1 (Johnsen et al. 2008). Divergence within this region is heterogeneous, with a peak of nucleotide divergence (up to 8%) in the immediate 5´ flanking region of the gene that is several fold higher than the expected difference between inbred mouse strains (Wade et al. 2002). Transgenic experiments confirmed that the mutation causing the tissue-specific switch in B4galnt2 gene expression lies within this region (Johnsen et al. 2008). A survey of 65 laboratory and wild-derived inbred mouse strains found a haplotype block spanning ~97 kb of nearly identical sequence to RIIRS/J in 12 other inbred strains that also exhibited the Mvwf1 tissue-specific switch in B4galnt2 gene expression. Five of these strains were independent, wild-derived strains of Mus musculus domesticus origin.

These observations raise important evolutionary questions regarding the origin and maintenance of these divergent haplotypes. However, until now, only inbred strains have been analyzed. In this study, we survey both DNA sequence polymorphism and microsatellite variability surrounding the B4galnt2 gene region in natural populations of M. m. domesticus and determine the B4GALNT2 expression pattern and VWF levels in wild-caught mice. We find that the pattern of haplotypic variation previously observed in laboratory and wild-derived inbred strains is also present in natural populations, that there is evidence of a recent shift in frequency of one of these haplotypes due to natural selection, and that the level and pattern of polymorphism in the 5´ flanking region significantly deviates from the neutral expectation. Coalescent simulations performed to evaluate the role of introgression indicate that the heterogeneity in divergence between haplotypes is highly unlikely under an introgression event alone. Furthermore, we demonstrate an association between genotype at B4galnt2, B4GALNT2 expression pattern, and VWF levels in a natural population.

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Key words: balancing selection, introgression cis-regulatory variation, house mouse, Mus musculus, von Willebrand disease (VWD), von Willebrand factor (VWF).

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Materials and Methods

Animal Material

Population samples of *M. m. domesticus* from France, Germany, and Cameroon are comprised of wild-caught individuals as previously described (Ihle et al. 2006; Baines and Harr 2007). The population sample of *M. m. domesticus* from Chicago was kindly provided by Bettina Harr. DNA from a single *M. famulus* individual (kindly provided by F. Bonhomme) was sequenced as an outgroup. Individuals screened for their VWF levels and B4galnt2 expression pattern were caught in the summer of 2005 at the same sampling localities in the Massif Central of France as described in Ihle et al. (2006). Mice were acclimated to laboratory conditions for 2 months prior to analysis. Animal maintenance and handling was according to the required legal standards and institutionally approved by the Bezirksregierung Köln.

Preparation of Biological Samples

DNA was prepared from either ethanol-preserved or frozen tissues as previously described (Nichols et al. 1994). Platelet-poor plasma was collected as previously described (Nichols et al. 1994) and stored at −80 °C until analysis. Tissues were formalin-fixed and paraffin embedded by the Tissue Core of the University of Michigan Comprehensive Cancer Center.

Analysis of B4galnt2 Expression Patterns and Plasma VWF Levels

*Dolichos biflorus* agglutinin (DBA) lectin staining was performed on formalin-fixed, paraffin-embedded tissues as previously described (Mohlke et al. 1999) using horseradish peroxidase conjugated DBA lectin (H-Y Labs, Inc.). Small intestine from C57BL/6J and RIIIS/J were simultaneously stained as controls.

Plasma VWF levels were determined by an enzyme linked immunosorbent assay in duplicate, as previously described (Mohlke et al. 1996) using aliquots of pooled C57BL/6J plasma to generate a standard curve. Statistical significance in VWF levels between the two groups (with or without RIIIS/J nucleotide polymorphisms at fragment 6) was determined using Student’s unpaired two-tailed t-test.

Polymerase Chain Reaction (PCR) and DNA Sequencing

Genomic sequencing was performed at the University of Cologne. Briefly, nine primer pairs that span the RIIIS/J haplotype block were designed for PCR using RIIIS/J genomic sequence (GenBank EF372924). A list of the PCR primers, cycling conditions, and sequencing primers for each amplified product is provided in supplemental table S1, Supplementary Material online.

Genotyping at B4galnt2

A PCR fragment diagnostic of the RIIIS/J and C57BL/6J haplotypes was sequenced to screen natural pop-
we reconstructed the phase of diploid individuals using the program PHASE version 2.1 (Stephens et al. 2001; Stephens and Donnelly 2003). The algorithm was run five times automatically using the “-x” option and the output providing the best “goodness of fit” was used. Although the phase could be reconstructed with a certainty of $\geq 90\%$ for a majority of individuals and heterozygous positions, we chose the composite genotypic disequilibrium, $A$, for analysis of linkage disequilibrium (LD), as it does not require a known phase of alleles (Weir 1996). In addition, this measure of LD does not require an assumption of random mating, which may be more appropriate for wild mouse populations where deviations from Hardy–Weinberg proportions are common (Laurie et al. 2007). The composite genotypic disequilibrium was computed using the “composite_LD” function submitted to the Bioperl project (Stajich et al. 2002), and LD is reported as the composite genotypic $r^2$, the squared correlation of genotypic indicators at two loci in diploid individuals (Weir et al. 2004; Laurie et al. 2007).

Microsatellite gene diversity estimates (expected heterozygosity, corrected for sample size) were calculated using the program MS Analyzer 3.15 (Dieringer and Schlötterer 2003). The $\ln RH$ statistic (Kauer et al. 2003) was applied to compare variability levels between populations. Significance was estimated by comparing the $\ln RH$ values of the typed microsatellites against a reference data set of microsatellite loci distributed throughout the genome (Ihle et al. 2006). To obtain estimates of within-haplotype class heterozygosity at microsatellites, the phase of the 12 microsatellite loci and two haplotype classes (as determined by diagnostic PCR fragment 6) was reconstructed using PHASE (Stephens et al. 2001; Stephens and Donnelly 2003) as described above. Only the genotypes for which an estimate of the phase with respect to the alternative $B4galnt2$ haplotypes had a certainty $\geq 90\%$ were included in within-haplotype heterozygosity estimates and analyses.

Allele Age Estimates

Two separate methods were used to estimate the age of the RIIIS/J haplotype, one based on the accumulation of new mutations since the common ancestor of RIIIS/J-bearing chromosomes (Rozas et al. 2001) and the other based on allele frequency (Kimura and Ota 1973). The age based on the accumulation of mutations among RIIIS/J chromosomes was estimated assuming a star phylogeny, with the number of mutations following a Poisson distribution. The presence of at most two nucleotide mutations among a sample of eight RIIIS/J chromosomes sequenced at the nucleotide level (one singleton within an RIIIS/J homozygote and one singleton of uncertain phase occurring in a heterozygote within sequence fragments 2 through 8) and two microsatellite loci monomorphic among 23 chromosomes (the minimum number for which reliable phase estimates were obtained) were used for this calculation. The nucleotide mutation rate was assumed to be $2 \times 10^{-9}$ (range $1.67 \times 10^{-9}$ to $2.98 \times 10^{-9}$; Eyre-Walker et al. 2002), whereas microsatellite mutation rates were estimated by the method of Xu and Fu (2004) in the German population, which is less likely to be influenced by the effects of selection, and assuming an effective population size of 58,000 (Salcedo et al. 2007). The 95% confidence interval (CI), $[t_{\text{lower}} - t_{\text{upper}}]$, was estimated as follows: $t_{\text{lower}}$ is the value of $t$ for which the probability of two nucleotide and zero microsatellite mutations plus the probabilities of all outcomes with equally many or more mutations (i.e., two, three, four, etc., mutations at microsatellite and nucleotide loci combined) is equal to 0.025. $t_{\text{upper}}$ is the value of $t$ for which the probability of two nucleotide and zero microsatellite mutations plus the probabilities of all equally or more extreme outcomes (e.g., no mutations at microsatellite or nucleotide loci) is equal to 0.025. The 95% CI for the expected age based on allele frequency (Kimura and Ota 1973) was obtained using the approximation given by Slatkin and Rannala (2000). Estimates are given in units of years, assuming two generations per year.

Coalescent Simulations

Coalescent simulations were performed in order to carry out an extension of the haplotype test of Hudson et al. (1994) and to simulate the effect of an introgression event on the pattern of nucleotide variability at $B4galnt2$. The test of Hudson et al. (1994) gives the probability of observing a subset of $i$ alleles with $j$ or fewer segregating sites given an overall sample of $n$ alleles with $S$ segregating sites. To generate genealogies containing mutation events both at nucleotide and microsatellite loci, the program simcoal2 (Laval and Excoffier 2004) was used. Given the larger sample size of individuals typed for microsatellite loci (31–46) than for nucleotide polymorphism (six to eight), the number of segregating sites expected within the RIIIS/J haplotype in the total larger sample was estimated using Watterson’s (1975) estimator of $\theta$. To evaluate an introgression model, coalescent simulations similar to those described by Evans et al. (2006) were performed using the software ms (Hudson 2002). Random genealogies for a sample of 16 chromosomes were simulated for a $\sim 60$-kb chromosomal region. The number of segregating sites was fixed based on the number observed over the entire $\sim 60$ kb given the number observed in fragments 2–8. We used the following scenario to reproduce the effects of introgression: Eight chromosomes were obtained from population A (recipient population) and eight from population B (donor population), representing the same frequency of RIIIS/J haplotypes present in the sequenced sample. Two demographic models were considered for population A, a standard constant size model assuming an effective population size of 58,000 (Salcedo et al. 2007), and the most likely demographic scenario (including a population size bottleneck and migration within and between species) estimated for the French population by Baines and Harr (2007). In both cases, population A was assumed to have split from population B at 500,000 years (1,000,000 generations) in the past. Given the low level of variation observed within RIIIS/J haplotypes, population B was subjected to a severe population bottleneck in the recent past ($N_0/N_0 = 0.000001$, $t_b = 0.000001$, time measured in $4N_e$ generations). This simulates the event that only a single haplotype introgressed into population A and
subsequently increased in frequency. For each model, simulations were performed with both zero recombination and using the rate estimated for this chromosomal region (Jensen-Seaman et al. 2004). For each simulated genealogy, seven representative regions of the same length and relative position of fragments 2–8 were taken to compare the variance in both nucleotide diversity, $\pi$, and the average pairwise divergence between haplotype classes, $\pi_{\text{between}}$, with the observed values. $P$ values are given as the number of simulations out of 20,000 for which the observed variance in either $\pi$ or $\pi_{\text{between}}$ were equal to or greater than the observed value.

**Results**

**Screen of the RIIIS/J Haplotype in Natural Populations**

Our previous survey of laboratory and wild-derived inbred strains (Johnsen et al. 2008) suggests that the haplotype associated with the RIIIS/J strain segregates in natural populations. This was based on extensive sequence characterization of six common laboratory strains and 23 wild-derived strains, including representatives of the *M. musculus* subspecies *musculus*, *domesticus*, *castaneus*, and *molossinus*, as well as the closely related *Mus spreitus* and *Mus spicilegus*. The RIIIS/J haplotype appeared to be confined to laboratory strains and wild-derived strains of *M. m. domesticus* origin.

To screen for the distribution and frequency of the RIIIS/J haplotype in natural populations of *M. m. domesticus*, a diagnostic PCR fragment (fragment 6; see Materials and Methods and fig. 1) was typed in 126 individuals from four separate populations. There are 11 single nucleotide polymorphisms in this fragment between the RIIIS/J and C57BL6/J haplotype classes, and assignment of haplotype class was confirmed by the generation of Neighbor-Joining (Saitou and Nei 1987) trees for each population (supplementary figure S1, Supplementary Material online). The frequency of the RIIIS/J haplotype was highly variable (table 1), with the two *M. m. domesticus* populations in Europe displaying the most striking difference in haplotype frequency (0% vs. 39%). No significant deviation from Hardy–Weinberg equilibrium was observed in Cameroon ($P = 0.56$) or Chicago ($P = 0.53$), whereas an excess of homozygotes was observed in France ($P = 0.015$). Previous studies of both this French population and other natural populations of *M. m. domesticus* indicate that an excess of homozygotes is a common observation at loci throughout the genome, however, likely due to inbreeding effects associated with communal nesting in wild mouse populations (Ihle et al. 2006; Laurie et al. 2007).

**DNA Sequence Variation in European *M. m. domesticus***

For a more detailed analysis of DNA sequence variation at *B4galnt2*, we chose to focus on the *M. m. domesticus* populations from Germany and France for several reasons. Although populations from Chicago and Cameroon both display frequencies of the RIIIS/J haplotype of −20% to 25%, these represent much more recent colonizations and are thus more likely to be confounded by demography.

**Table 1**

<table>
<thead>
<tr>
<th>Population</th>
<th>C57 Haplotype</th>
<th>RIIIS/J Haplotype</th>
</tr>
</thead>
<tbody>
<tr>
<td>France ($n = 51$)</td>
<td>$60.8% (62/102)$</td>
<td>$39.2% (40/102)$</td>
</tr>
<tr>
<td>Germany ($n = 36$)</td>
<td>$100% (72/72)$</td>
<td>$0% (0/72)$</td>
</tr>
<tr>
<td>Cameroon ($n = 29$)</td>
<td>$77.6% (45/58)$</td>
<td>$22.4% (13/58)$</td>
</tr>
<tr>
<td>Chicago ($n = 10$)</td>
<td>$75% (15/20)$</td>
<td>$25% (5/20)$</td>
</tr>
</tbody>
</table>

Additionally, the German and French populations have been extensively characterized at both the microsatellite (Ihle et al. 2006) and DNA sequence level (Baines and Harr 2007), which provide powerful points of comparison, and phenotypic screening of wild-caught mice from the same locations sampled by Ihle et al. (2006) was possible.

Nine sequence fragments spanning approximately 100 kb of the *B4galnt2* structural gene and 5′ flanking region were sequenced in six to eight individuals each in the German and French populations (fig. 1; table 2), for which data from seven unlinked autosomal reference loci also exist (Baines and Harr 2007). These fragments were chosen to encompass the region previously identified as highly divergent between the C57BL6/J and RIIIS/J strains (Johnsen et al. 2008), in addition to sequence flanking this region. An outgroup sequence from *Mus famulus* was obtainable for six of these nine segments. Figure 2 displays a plot of nucleotide diversity across the *B4galnt2* region relative to the genome average as determined from seven unlinked autosomal reference loci (Baines and Harr 2007). A clear peak of polymorphism in the immediate 5′ flanking region of the gene is present in the French population, which decreases on either side. Polymorphism levels at this peak are nearly 10-fold higher than the genome average ($\pi = 2.43\%$ vs. $\pi_{\text{average}} = 0.26\%$) for this population. In contrast, the German population displays consistently low polymorphism across the region comparable to levels seen at the reference loci.

**Haplotype Structure and LD**

To analyze the pattern of haplotypic variation at *B4galnt2*, the phase of the diploid sequence data was reconstructed using the program PHASE (Stephens et al. 2001; Stephens and Donnelly 2003). This could be estimated with a certainty of >90% for a majority of polymorphic sites and individuals (supplementary figure S2, Supplementary Material online). Inspection of these results reveals the presence of two major haplotype classes spanning sequence fragments 2–8, which are very similar to those present in the C57BL6/J and RIIIS/J inbred strains. Although the C57BL6/J haplotype class appears fixed in the German population, both haplotype classes segregate at intermediate frequency in the French population.

Although the haplotype reconstruction was reliable over most of the sampled region, there was less certainty at the 5′ and 3′ extremes (i.e., fragments 1 and 9). Thus, to quantify LD across the entire region, we report the composite genotypic $r^2$, as it does not require a known phase of alleles (see Materials and Methods). LD between sites in the
French population is high, with an average \( r^2 \) over the entire sequenced region of 0.686. In particular, the association between sites spanning the ~60-kb interval between fragments 2–8 reaches the maximum (\( r^2 = 1.00; P = 0.0047 \)). This value far exceeds the mean LD between sites at similar distance in a large-scale study of a *M. m. domesticus* population from Arizona (\( r^2 / C2 = 0.15; Laurie et al. 2007 \)). However, due to the large number of pairwise comparisons (1,771) associated with the many divergent sites between haplotypes, no pairwise comparison remains significant after Bonferroni correction. LD in the German population, where only the C57BL6/J haplotype class is present, is comparatively lower (average \( r^2 = 0.201 \)) and also displays no significant association between sites.

The region spanning fragments 4–7 is particularly striking. These fragments total \( 2 \text{ kb} \) and span a \( 23 \text{-kb} \) region (of which \( 19 \text{ kb} \) lies upstream of the start codon). In this region, only three polymorphic sites were identified “within” haplotype classes (one within RIIIS/J, one within C57BL6/J in the German population, and one singleton of uncertain haplotypic phase in a heterozygous individual), whereas 36 polymorphic sites are found “between” them.

Tests of Neutrality

To test if the pattern of polymorphism at *B4galnt2* is consistent with a neutral model of molecular evolution, we first performed a test based on the distribution of allele frequencies. Tajima’s *D* (Tajima 1989) is consistently positive across the *B4galnt2* region in the French population, indicating an excess of low-frequency variants. We next compared the ratio of polymorphism with divergence across the six fragments for which outgroup data were attainable and seven unlinked autosomal reference loci (Baines and Harr 2007) using a multilocus version of the HKA test (Hudson et al. 1987). The deviation from the neutral expectation for both polymorphism and divergence is plotted in figure 3. The neutral model could be rejected for both populations (*P < 10^-5*

### Table 2

<table>
<thead>
<tr>
<th>Fragment (Position*)</th>
<th>Population</th>
<th>Sample Size</th>
<th>Length (bp)</th>
<th>( S^b )</th>
<th>( \pi^c ) (%)</th>
<th>( \theta^d ) (%)</th>
<th>( K^e ) (%)</th>
<th>Tajima’s <em>D</em></th>
</tr>
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<tbody>
<tr>
<td>1 (−55.3 kb)</td>
<td>France</td>
<td>14</td>
<td>749</td>
<td>1</td>
<td>0.07</td>
<td>0.04</td>
<td>—</td>
<td>1.434</td>
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<td></td>
<td>Germany</td>
<td>12</td>
<td>736</td>
<td>2</td>
<td>0.02</td>
<td>0.05</td>
<td>—</td>
<td>−1.140</td>
</tr>
<tr>
<td>2 (−38.3 kb)</td>
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<td>16</td>
<td>427</td>
<td>3</td>
<td>0.31</td>
<td>0.21</td>
<td>4.85</td>
<td>1.376</td>
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<tr>
<td></td>
<td>Germany</td>
<td>12</td>
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<td>2</td>
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<td>0.08</td>
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</tr>
<tr>
<td></td>
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<td>638</td>
<td>0</td>
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<td>0</td>
<td>3.04</td>
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<td>3.069***</td>
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<td>0</td>
<td>4.81</td>
<td>—</td>
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<tr>
<td></td>
<td>France</td>
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<td>476</td>
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<td>1.30</td>
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<td>0</td>
<td>4.47</td>
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<td></td>
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<td>1</td>
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<td>0.08</td>
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<td>0.15</td>
<td>—</td>
<td>0.672</td>
</tr>
</tbody>
</table>

*P < 0.05; ***P < 0.001.

* Position relative to *B4galnt2* start codon.

\( S \) = the number of segregating sites.

\( \pi \) = nucleotide diversity (Tajima 1983).

\( \theta \) = nucleotide diversity (Watterson 1975).

\( K \) = divergence.

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[Table 2](#)

**Summary of Polymorphism and Divergence across *B4galnt2* Region**

![Fig. 2.—Nucleotide diversity, \( \pi \) (Nei 1987), for *Mus musculus domesticus* populations across the *B4galnt2* gene region. Diversity in France (black line) and Germany (gray line) are strikingly different in the region near *B4galnt2* exon 1. Dashed lines represent the genome average at seven unlinked autosomal reference loci (Baines and Harr 2007).**

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and \( P = 0.002 \) for France and Germany, respectively). However, the loci contributing most to the overall chi-square statistic differ between populations. In particular, significance in the French populations is due to fragments 5 and 6, which display both an excess of polymorphism and a deficiency of divergence. In contrast, sequence fragments spanning \( B4galnt2 \) appear to be evolving similarly to those throughout the genome in the German population, although the test remains significant due to elevated polymorphism at a single unlinked reference locus in this population. It should be noted, however, that this reference locus displays no evidence of a deviation from neutrality by other measures (e.g., Tajima’s \( D \)), nor does it display evidence of selection in other \( M. m. domesticus \) populations, including the French population (Baines and Harr 2007). Furthermore, removal of this locus does not change the significance of the results in the French population due to \( B4galnt2 \) fragments 5 and 6.

Microsatellite Variability

The observation of high divergence between haplotypes, low variation within haplotypes, and large differences in haplotype frequency between populations suggest complex dynamics at \( B4galnt2 \). To further explore the nature of the selective forces operating, we genotyped 12 microsatellites across this region in 31–46 individuals per population (supplemental tables S2 and S3, Supplementary Material online) and compared them with the pattern seen at 118 microsatellites distributed randomly throughout the genome in the same German and French population samples (Ihle et al. 2006). Due to their high polymorphism levels and high mutation rate, these microsatellite markers serve as sensitive indicators of selection at linked sites and have the ability to capture more recent events.

The relative amount of heterozygosity between the two populations at individual loci may be described by the ln RH statistic (Kauer et al. 2003). After standardizing the values of the 12 loci using the mean and standard deviation of ln RH from the random data set, we detected no significant outlier after Bonferroni correction. However, when we analyzed heterozygosity with respect to the RIIIS/J and C57BL6/J haplotype classes individually, a striking pattern emerged (fig. 4). At six loci within a \( \sim 160 \)-kb window surrounding \( B4galnt2 \), the average heterozygosity within the RIIIS/J haplotype class is nearly half that of the C57BL6/J class (0.446 vs. 0.807), whereas the average values between the two classes at six more distant flanking loci (1–2.7 Mb from the gene) display nearly identical values (0.732 vs. 0.733). Even more striking, the two microsatellite loci located within the haplotype structure associated with the RIIIS/J and C57BL6/J classes are monomorphic (i.e., contain no variation) within the RIIIS/J class. Specifically, a (GT)\(_{6,23} \) repeat located \( \sim 30 \) kb upstream of the \( B4galnt2 \) start codon is polymorphic within the C57BL6/J haplotype, but harbors only a (GT)\(_{23} \) allele within the RIIIS/J class, and a (GA)\(_{10,37} \) locus \( \sim 1 \) kb downstream of the \( B4galnt2 \) start codon is polymorphic within the C57BL6/J class, but harbors a single (GA)\(_{10} \) allele within the RIIIS/J class.

The observation of a single allele at two microsatellite loci within a haplotype of intermediate frequency is unlikely under a neutral model and suggests a young age of the RIIIS/J haplotype. To assess the probability of observing a single allele at two microsatellite loci within a haplotype of given frequency, we performed an extension of the haplotype test of Hudson et al. (1994) including microsatellites (see Materials and Methods). Coalescent simulations were performed under both a standard constant size model and under the most likely demographic model for the French population (including a population size bottleneck) estimated by Baines and Harr (2007) under the conservative assumption of no recombination. In both cases, the observed level of nucleotide and microsatellite variation within the RIIIS/J haplotype significantly deviated from the neutral expectation (\( P < 0.0002 \) and \( P = 0.0006 \) for constant size and demographic models, respectively).

In addition, we compared the age of the RIIIS/J haplotype estimated from within-haplotype nucleotide and microsatellite polymorphism with that predicted from its frequency. Assuming a star phylogeny and that the mutations among RIIIS/J chromosomes follow a Poisson distribution, an estimate of the age of the RIIIS/J haplotype can
be made given the observation of at most two nucleotide mutations (one singleton within an RIIIS/J homozygote and one singleton of uncertain phase occurring in a heterozygote within fragments 2–8) and assuming that no mutations have occurred at the two monomorphic microsatellite loci, giving a maximum likelihood estimate of 97 years (95% CI 12–350 years). In contrast, the age predicted by the frequency of the RIIIS/J haplotype is 70,040 years (95% CI 12,650–109,246 years) (Kimura and Ota 1973).

Taken together, these observations suggest the RIIIS/J haplotype has risen to intermediate frequency very recently in the French population due to the action of natural selection.

Introgression Simulations

The high sequence divergence, LD, and low variation within the RIIIS/J haplotype suggest introgression from a closely related species as a viable explanation for the origin of this allele in $M$. $m$. domesticus, given that genetic exchange with even more distant, noncommensal species such as $M$. spretus (average nucleotide divergence ~2%; Galtier et al. 2004) is known to occur (Greene-Till et al. 2000; Orth et al. 2002; Bonhomme et al. 2007). However, although the region of LD spans a large distance (~60 kb), as might be expected if the RIIIS/J haplotype recently introgressed, the level of sequence divergence between haplotypes is highly heterogeneous across this region. For example, the average pairwise divergence, $\pi_{\text{between}}$, between haplotypes at fragment 5 (4.6%), near the center of the region of LD, is roughly 10-fold higher than at fragments 2 and 8 (0.47% and 0.54%, respectively). In the absence of comparable heterogeneity in the mutation rate across the region, this pattern would be unexpected under a simple introgression model, and the range of divergence to the more distantly related $M$. famulus across sequence fragments (2.7–4.9%) varies less than 2-fold (table 2).

To test if the heterogeneity in divergence between haplotypes is compatible with a simple introgression model, we performed coalescent simulations allowing for the outcome of sampling alleles from two divergent populations to be evaluated (see Materials and Methods). Genealogies of a ~60-kb chromosomal region (corresponding to the region of LD associated with fragments 2–8) were simulated under a standard constant size model and a demographic model suited to the history of house mice as above (Baines and Harr 2007), both with and without recombination. The observed variances of both $\pi$ and $\pi_{\text{between}}$ across the region are significantly greater than expected under all scenarios ($P < 5 \times 10^{-5}$). Thus, this heterogeneity, largely resulting from the extreme peak of divergence between haplotypes, appears highly unlikely under a simple introgression model.

Mvwf1 Phenotype in a Natural $M$. $m$. domesticus Population

To determine if the presence of the RIIIS/J haplotype in the 5’ flanking region of $B4galnt2$ in wild mice is associated with the same tissue-specific switch in expression conferred by the $Mvwf1$ mutation in inbred strains, the same diagnostic fragment (fragment 6) as above was typed in 24 wild-caught $M$. $m$. domesticus individuals from the Massif Central region of France and small intestine was stained with DBA lectin (which is specific for $B4GALNT2$ glycosylation products; Mohlke et al. 1999). Individuals...
homozygous for the RIIIS/J haplotype uniformly exhibited DBA–lectin staining of blood vessels but demonstrated no intestinal epithelial DBA lectin staining (fig. 5), consistent with the tissue-specific switch in \( \text{B4galnt2} \) expression first characterized in the RIIIS/J inbred mouse strain. All individuals lacking the RIIIS/J haplotype demonstrated intestinal epithelial DBA lectin staining and never exhibited vascular DBA lectin staining (fig. 5). Individuals heterozygous for RIIIS/J polymorphisms exhibited both intestinal epithelial and vascular endothelial DBA lectin staining, consistent with codominant expression of \( \text{B4galnt2} \) alleles in these tissues (fig. 5). In addition, these data indicate that fragment 6 is a reliable diagnostic marker of (i.e., is tightly linked to) the mutation conferring the regulatory switch.

Plasma VWF levels were determined in all 24 wild-caught individuals and correlated to genotype at fragment 6. The low VWF phenotype conferred by the RIIIS/J allele has been extensively characterized in laboratory mice and is inherited in an autosomal dominant manner (i.e., mice both heterozygous and homozygous for the RIIIS/J haplotype exhibit similarly low VWF levels) (Nichols et al. 1994; Mohlke et al. 1996, 1999). Individuals either heterozygous or homozygous for the RIIIS/J haplotype class displayed less than half the VWF levels of individuals that did not have the RIIIS/J haplotype (5.5 ± 2.0 vs. 13.3 ± 2.3, \( P < 1 \times 10^{-7} \) using Student’s unpaired two-tailed \( t \)-test; displayed in fig. 6), consistent with vascular endothelial expression of \( \text{B4galnt2} \) alleles leading to altered glycosylation and subsequent accelerated clearance of VWF from circulation. Thus, the phenotypic effect of the RIIIS/J haplotype is confirmed in this natural population of \( M. \text{m. domesticus} \), where it clearly confers the \( \text{Mvwf1} \) tissue-specific switch and causes a significant reduction in plasma VWF.

**Discussion**

The data reported here indicate that the pattern of haplotypic variation at \( \text{B4galnt2} \) in natural populations is similar to that observed in laboratory mouse strains (i.e., two main divergent haplotype classes similar to either C57BL6/J or RIIIS/J). The level and pattern of polymorphism and divergence in populations in which these haplotypes segregate are extreme, suggesting that the RIIIS/J haplotype may have been introduced into \( M. \text{m. domesticus} \) by introgression and/or is maintained by balancing selection, whereas analysis of microsatellite variation within this region suggests the action of very recent natural selection, which may account for the differences in haplotype frequency observed between different local populations. Furthermore, wild-caught individuals display a clear association between haplotype class and variation in the tissue-specific expression pattern of \( \text{B4galnt2} \), and individuals homo- or heterozygous for the RIIIS/J haplotype class display significant reductions in VWF level.

**Origin and Maintenance of Divergent \( \text{B4galnt2} \) Alleles**

Compared with the genomic regions between laboratory mouse strains deemed as “ancestrally divergent” (i.e.,...
originating from different *M. musculus* subspecies), where a nucleotide difference occurs on average every ~200 bp (~0.5% divergence) (Wade et al. 2002), the level of sequence divergence between the RIIIS/J and C57BL6/J inbred strains and the haplotypes that largely correspond to these sequences in natural populations is highly unusual. The overall high sequence divergence, LD, and low variation within a haplotype at intermediate frequency suggest that selection may have operated on an allele derived from a rare interbreeding event between partially isolated populations to give rise to the pattern of variation observed at B4galnt2. However, the sliding window analysis of the RIIIS/J sequence (shown adapted from Johnsen et al. 2008, in fig. 1) demonstrates that this level of divergence is not uniform, displaying a peak of ~8% divergence in the immediate 5′ flanking region of the gene, which decreases on either side. Under a simple neutral introgression model, divergence between haplotypes would be largely governed by the mutation rate, and thus be expected to be more uniform.

Although introgression may be a contributing factor to the history of these two divergent allele classes at B4galnt2, the peak of divergence between them is also consistent with a prediction of long-term balancing selection: linked neutral mutations are expected to accumulate between allelic classes with increasing proximity to the selected site(s), where the opportunity to “migrate” (i.e., to recombine) between allelic classes becomes increasingly smaller (Hudson and Kaplan 1988; Charlesworth et al. 1997; Takahata and Satta 1998; Charlesworth 2006). However, simple long-term maintenance of alleles within a single population would also predict an accumulation of within-allele-class variants, which is not observed in our survey. Together with the observation of large differences in frequency between local populations, this suggests that the selective force(s) maintaining this variation may be heterogeneous over space and/or time. Stahl et al. (1999) proposed a model termed “trench warfare” to explain the heterogeneous distribution of alternative wild-type and disease-resistant alleles segregating at the *Rpm1* locus of *Arabidopsis thaliana* across its geographic range. Under this model, if a resistance allele carries a cost, it will be disfavored in the absence of the pathogen against which it confers resistance, and recurrent advances and retreats of the resistance allele/pathogen will act to maintain variation. It is possible that the German and French populations, which harbor frequencies of the RIIIS/J haplotype of 0% and 39%, represent different stages of such a fluctuating model, with the RIIIS/J haplotype having very recently increased in frequency in the French population.

Another aspect of the data inconsistent with simple long-term maintenance of alleles is the large size of the region of LD, which would not be maintained over long periods unless recombination is suppressed across this region. Although the chromosomal region containing B4galnt2 experiences greater than average levels of recombination for the mouse genome (1.3 cM/Mb; Jensen-Seaman et al. 2004), it is possible that the many nucleotide differences between C57BL6/J and RIIIS/J, in conjunction with several larger insertions/deletions (Johnsen et al. 2008), are acting to suppress recombination between these alleles. The large region of LD would, however, be consistent with the recent increase in frequency of an allele due to natural selection.

**VWF Variability in Mammals**

In addition to identifying signatures of selection at the B4galnt2 locus in these natural populations, we tested wild-caught individuals from a French house mouse population for the *Mvwfl* phenotype initially described in the RIIIS/J inbred mouse strain. RIIIS/J exhibits a low VWF phenotype similar to human type 1 VWD (Sweeney et al. 1990), the most common inherited bleeding disorder in humans. In addition to the mouse, VWD has been reported in other mammals, including pigs, calves, dogs, cats, horses, and rabbits (Mohlke and Ginsburg 1997). VWD in humans exhibits variable expressivity and incomplete penetrance, and this observation, in conjunction with the wide range of VWF levels found in normal subjects (from 50% to 150%), implies the influence of modifier genes outside the VWF gene itself (Levy and Ginsburg 2001). Although one major modifier, “ABO” blood group (Orstavik et al. 1985), and other minor modifiers of VWF have been described (O’Donnell et al. 2002), the cause of the majority of genetic variability in VWF levels in humans remains unexplained. The reason such wide variability in a critical coagulation factor is tolerated within human populations is unknown, and recent work has uncovered similar VWF variability in mice (Lemmerhirt et al. 2006, 2007).

In the current study, we demonstrate that the RIIIS/J haplotype is associated with a remarkably significant decrease in VWF in a wild-caught mouse population despite numerous uncontrolled acquired, environmental, and genetic factors that would be expected to influence VWF levels. Though we cannot exclude an offsetting potential benefit from low VWF levels, we propose another, as yet unknown, benefit of carrying the RIIIS/J haplotype, resulting in balancing selection at the B4galnt2 locus, with low VWF levels representing a pleiotropic byproduct.

**Possible Selective Forces Operating on Variation in B4galnt2 Expression**

The endothelium plays an active role in thrombosis, inflammation, and vascular wall remodeling (Michiels 2003) in response to injury. Consequently, a number of cell adhesion molecules and/or soluble endothelial cell-derived products, such as those critical for the host inflammatory response (Reinhart et al. 2002), are potential candidates for glycosylation modification by endothelial expressed B4GALNT2. The resulting altered function could lead to an advantageous phenotype under specific circumstances by modifying the host response to inflammation, injury, or a pathogen (Laschke et al. 2005; Doulet et al. 2006; Biswas et al. 2007). Such a scenario could favor the RIIIS/J haplotype under certain conditions, such as an infectious outbreak.

Although there is a higher frequency of individuals gaining expression of B4galnt2 in blood vessels than losing expression in intestine in this wild mouse population, the loss of bowel B4galnt2 expression in animals homozygous
for the RIIIS/J haplotype is also a candidate to affect fitness in these populations. Glycosylation of intestinal mucins has been shown to be critical for colonization, adhesion, and infection of enteric pathogens (Karlsson et al. 2000; Petri et al. 2002; Robbe et al. 2003; Kawakubo et al. 2004), and loss of carbohydrate ligands for an invading pathogen could result in a fitness advantage for mice homozygous for RIIIS/J-like sequences. However, expression of B4galnt2 orthologs is conserved in intestine in mice and humans (Capon et al. 2001; Montiel et al. 2003) and likely other mammals (Chae 1997; Karlsson et al. 1997), suggesting that intestinal B4GALNT2 may serve a function that would be lost in mice homozygous for the RIIIS/J haplotype.

Conclusion

Differences in transcriptional regulation appear to account for a significant proportion of variation subject to selection. Previous application of population genetics to the study of variation in cis-regulatory DNA has identified several examples of balancing selection between disease and protective states (reviewed by Hahn 2007). The Mtwf1 regulatory switch in gene expression from one tissue-restricted pattern to another is an unusual molecular mechanism with few reported precedents (Mohlke et al. 1999). The multifaceted phenotype resulting from the tissue-specific switch in B4galnt2 expression pattern likely underlies complex selective forces at work at the B4galnt2 locus. Whether a fitness advantage is conferred by the gain of B4galnt2 vascular expression or the loss of B4galnt2 intestinal expression, a specific advantageous phenotype conferred by the RIIIS/J haplotype class in natural populations may be difficult to identify. The molecular evidence at the B4galnt2 locus in this natural mouse population is consistent with tolerance of the presumably detrimental low VWF phenotype in exchange for an as yet unknown benefit. We speculate that similar selection may account for the tolerance of variability in VWF and high prevalence of VWD in other mammalian species, including humans.

Supplementary Materials

Supplementary tables S1, S2, and S3 and figures S1 and S2 are available at Molecular Biology and Evolution online (http://www.mbe.oxfordjournals.org/).

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