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Genome Sequencing Reveals Loci under Artificial Selection that Underlie Disease Phenotypes in the Laboratory Rat

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SUMMARY

Large numbers of inbred laboratory rat strains have been developed for a range of complex disease phenotypes. To gain insights into the evolutionary pressures underlying selection for these phenotypes, we sequenced the genomes of 27 rat strains, including 11 models of hypertension, diabetes, and insulin resistance, along with their respective control strains. Altogether, we identified more than 13 million single-nucleotide variants, indels, and structural variants across these rat strains. Analysis of strain-specific selective sweeps and gene clusters implicated genes and pathways involved in cation transport, angiotensin production, and regulators of oxidative stress in the development of cardiovascular disease phenotypes in rats. Many of the rat loci that we identified overlap with previously mapped loci for related traits in humans, indicating the presence of shared pathways underlying these phenotypes in rats and humans. These data represent a step change in resources available for evolutionary analysis of complex traits in disease models.

INTRODUCTION

In the past 100 years, more than 500 inbred rat strains have been derived for a range of physiological and pathophysiological phenotypes (Aitman et al., 2008; Lindsey, 1979) but have been predominantly used to study cardiovascular and metabolic phenotypes, which are complex traits governed by the interaction between multiple genetic factors and the environment. Inbred rat models of cardiovascular and metabolic phenotypes have been derived from various founder colonies or stocks at
various geographic locations by crossing relatively small numbers of rats within the colony and selecting for the desired disease phenotypes over several generations, with simultaneous or subsequent brother-sister mating to develop genetically homogeneous inbred strains (Jong, 1984; Rapp, 2000).

Although the majority of inbred rat models of hypertension and diabetes were generated from outbred Wistar colonies, efforts have been made to derive disease models on various other genetic backgrounds (Jong, 1984). Each strain so derived is therefore expected to show major founder effects and should be genetically and phenotypically distinct from its founder colony as well as from other strains derived from different founder colonies (Doggrell and Brown, 1998). Significant genotypic and phenotypic heterogeneity in the genetic models of hypertension and diabetes therefore provides a unique resource to study molecular mechanisms behind different etiological forms of hypertension. In addition, metabolic phenotypes such as insulin resistance and dyslipidemia, which were frequently co-inherited with hypertension and may form part of the hypertension phenotype, may have inadvertently been coselected with hypertension, as well as compensatory alleles that protect against target organ damage mediated by phenotypes such as hypertension (St Lezin et al., 1999).

We hypothesize that, in these rat strains, phenotype-driven selection may have resulted in “artificial selective sweeps” with fixation of sequence variants that underlie disease phenotypes, as has been observed in the artificial selection of a number of other disparate but benign traits in different species (Rubin et al., 2010; Wright et al., 2005; Xia et al., 2009). Artificial selective sweeps in inbred rat strains may be unique to a particular strain and disease phenotype in comparison with other strains and may contribute to the molecular basis of hypertension and other related phenotypes in that strain.

Genes in a genome evolve at different evolutionary rates due to varying evolutionary constraints on each gene, and interactions between genes underlying complex phenotypic traits are maintained by coevolution (Pagliarini et al., 2008; Tillier and Charlebois, 2009). Because of the polygenic nature of hypertension and other complex phenotypes, genes containing disease-inducing variants might have evolved together at an evolutionary rate that will be different from the rest of the genome. The identification of genes that have coevolved and of artificial selective sweeps in rat disease models may therefore be informative for identifying loci underlying disease phenotypes and for understanding the polygenic architecture of complex disease phenotypes.

Present sequencing technology now permits rapid and accurate sequencing of whole genomes through which near-complete catalogs of genomic variants can be obtained. It is therefore possible to perform, in model organisms, genomic screening for identification of coevolved gene clusters and artificial selective sweeps that harbor potentially pathogenic genes and mutations.

In this study, we sequenced the genomes of 25 new rat strains on high-throughput sequencing platforms, with a major focus on strains with well-characterized cardiovascular and metabolic phenotypes. These included 11 widely used rat models of hypertension, diabetes, and insulin resistance, along with their respective control strains. We identified a comprehensive catalog of genomic variants in 27 rat strains (25 strains sequenced for this study and two strains sequenced previously; Atanur et al., 2010; Simonis et al., 2012). We also identified clusters of genes and genomic loci that coevolved during selective breeding of these laboratory rat strains and established their relationship to genes and loci known to underlie disease phenotypes in these strains. The study therefore provides insights into the genetics and biology of cardiovascular and metabolic phenotypes shown by the sequenced rat strains.

RESULTS

Sequencing and Variant Calling

We sequenced the genomes of 25 rat strains and analyzed them together with the genomes of two rat strains that we sequenced previously (Atanur et al., 2010; Simonis et al., 2012). We also resequenced the genome of BN/Mcwi to correct for errors in the reference genome, using DNA from the same animal that was used for the reference BN genome sequence (Gibbs et al., 2004). Quality filtered sequence reads were mapped to the reference BN/Mcwi genome assembly (referred to hereafter as the BN reference), version RGSC3.4. After removing clonal reads, we achieved at least or close to 20× coverage for all strains except for strains BBDP/Wor and WKY/NHsd, for which we achieved ~10× coverage (Table 1).

After applying rigorous filtering criteria (see Extended Experimental Procedures available online) and excluding genomic variants that arose due to potential errors in the reference genome (in which the reference allele was different from the BN/Mcwi allele detected by resequencing), 9,665,340 single-nucleotide variants (SNVs) and 3,502,117 short indels were identified across the 27 rat strains (Table 1). This includes 839,691 SNVs and 479,974 indels, in which we could not determine the BN/Mcwi allele by resequencing because of either low coverage or poor read mapping and/or base quality. The variant set therefore may still contain a small proportion (0.001%) of false positives due to base call errors in the reference BN genome. We found that 98.3% of SNV calls were homozygous and 1.7% were heterozygous, the latter most likely arising from a combination of true heterozygote loci due to incomplete fixation of the inbred strain and false-positive calls due to regions of copy number variation, sequencing errors, and mapping errors in highly repetitive genomic regions, as previously described (Atanur et al., 2010; Keane et al., 2011). Along with SNVs and indels, we also identified a total of 719,929 structural variants in the 26 rat strains sequenced on the Illumina platform.

To assess the sensitivity and specificity of the variant calls, we compared the LE/Stm Illumina sequence with the publicly available sequence of 13 bacterial artificial chromosomes (BACs) derived from the LE/Stm genome generated by conventional capillary sequencing, which provided a control sequence region of ~2 Mb for LE/Stm. There were 3,382 SNVs and 1,211 indels identified in the LE/Stm BAC sequences compared to the BN reference genome. We estimated 0.38% false-positive and 8.1% false-negative SNV calls in the Illumina LE/Stm sequence, with an estimated false positive and false negative rate for indels of 3.75% and 10.98%, respectively. Thus the overall accuracy of genotype calls, including variant and reference calls, was...
99.99% for the LE/Stm genome, which can be scaled to the remaining 25 rat strains sequenced on the Illumina platform, as the variants were called in these strains simultaneously with the same methods.

Within the total set of 9,665,340 identified SNVs, 67,616 were in protein-coding sequence, among which 29,131 were nonsynonymous coding (NSC) variants and 38,485 were synonymous coding (SC) variants. Of the 29,131 NSC variants, 409 predicted premature truncation of the protein due to gain of stop codons, whereas 27 predicted the loss of stop codons (Table S1 and Figures S1A and S1B). We also identified 2,366 short indels in protein-coding sequence, which predicted disruption of the open reading frame of the encoded protein (Table S2 and Figures S1C and S1D). Of the observed 37,510 large deletions, 1,497 overlapped with the predicted protein-coding sequence.

### Phylogenetic History of Laboratory Rat Strains

The evolutionary history of laboratory rat strains is complex, as they were derived by multiple rounds of interbreeding and inbreeding in different locations at different points in time (Jong, 1984; Saar et al., 2008). To inform understanding of the evolutionary history of laboratory rat strains, we constructed a phylogenetic tree of strains, including the BN reference strain, using our catalog of 9.6 million high-quality SNVs. Laboratory rat strains clustered essentially according to founder stocks and colonies from which they were derived (Figure 1). Moreover, Wistar-derived strains showed two prominent subclusters, consistent with the known breeding history of Wistar-derived strains in Japan (SHR, SHRSP, GK, and WKY) or elsewhere in Europe and the USA (MHS, MNS, LEW, WAG, and BBDP). Sprague-Dawley-derived (SD) strains (SS, SR, LH, LL, and LN) also clustered together, whereas other strains formed more isolated clusters, including BN strains from which the reference genome strain (BN/Mcwi) was selected. The WKY/Gla strain was closer to SHR substrains than other WKY substrains, possibly reflecting the known diversity of WKY strains arising from their distribution to different geographical locations before complete inbreeding (Kurtz et al., 1989) or from the close relationship of SHR and WKY sublines during their establishment and maintenance.

### Table 1. Sequencing Details and Variant Calls in 27 Inbred Rat Strains

<table>
<thead>
<tr>
<th>Rat Strain</th>
<th>Gb of Bases Mapped</th>
<th>Average Coverage</th>
<th>Number of SNVs</th>
<th>Number of Indels</th>
<th>Number of Structural Variants</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACI/EurMcwi</td>
<td>86.64</td>
<td>33.69</td>
<td>3,607,275</td>
<td>1,168,780</td>
<td>17,859</td>
</tr>
<tr>
<td>BBDP/Wor</td>
<td>25.68</td>
<td>9.99</td>
<td>3,322,410</td>
<td>1,131,697</td>
<td>35,387</td>
</tr>
<tr>
<td>BN.Lx</td>
<td>58.40</td>
<td>22.71</td>
<td>51,938</td>
<td>45,404</td>
<td>NA</td>
</tr>
<tr>
<td>F344/NCrI</td>
<td>65.13</td>
<td>25.33</td>
<td>3,433,241</td>
<td>1,132,993</td>
<td>42,138</td>
</tr>
<tr>
<td>FHH/EurMcwi</td>
<td>57.98</td>
<td>22.55</td>
<td>3,471,696</td>
<td>1,170,337</td>
<td>22,628</td>
</tr>
<tr>
<td>FHL/EurMcwi</td>
<td>53.25</td>
<td>20.71</td>
<td>3,422,550</td>
<td>1,117,039</td>
<td>14,217</td>
</tr>
<tr>
<td>GK/Ox</td>
<td>75.25</td>
<td>29.26</td>
<td>3,584,504</td>
<td>1,147,996</td>
<td>58,877</td>
</tr>
<tr>
<td>LE/Stm</td>
<td>56.11</td>
<td>21.82</td>
<td>3,485,480</td>
<td>1,152,163</td>
<td>19,285</td>
</tr>
<tr>
<td>LEW/NCrIBR</td>
<td>51.06</td>
<td>19.86</td>
<td>2,966,945</td>
<td>1,014,796</td>
<td>39,031</td>
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<tr>
<td>LEW/Crl</td>
<td>66.30</td>
<td>25.78</td>
<td>2,941,368</td>
<td>1,002,364</td>
<td>39,146</td>
</tr>
<tr>
<td>LH/MavRrrc</td>
<td>55.84</td>
<td>21.71</td>
<td>3,459,239</td>
<td>1,169,791</td>
<td>21,901</td>
</tr>
<tr>
<td>LL/MavRrrc</td>
<td>58.04</td>
<td>22.57</td>
<td>3,419,697</td>
<td>1,177,989</td>
<td>20,623</td>
</tr>
<tr>
<td>LN/MavRrrc</td>
<td>56.54</td>
<td>21.99</td>
<td>3,406,103</td>
<td>1,171,795</td>
<td>20,561</td>
</tr>
<tr>
<td>MHS/Gib</td>
<td>64.68</td>
<td>25.15</td>
<td>3,270,047</td>
<td>1,112,526</td>
<td>23,551</td>
</tr>
<tr>
<td>MNS/Gib</td>
<td>57.26</td>
<td>22.27</td>
<td>3,278,667</td>
<td>1,123,980</td>
<td>22,769</td>
</tr>
<tr>
<td>SBH/Ygl</td>
<td>65.76</td>
<td>25.57</td>
<td>3,461,088</td>
<td>1,134,320</td>
<td>13,789</td>
</tr>
<tr>
<td>SBN/Ygl</td>
<td>47.17</td>
<td>18.34</td>
<td>3,334,857</td>
<td>1,078,851</td>
<td>9,586</td>
</tr>
<tr>
<td>SHR/NHsd</td>
<td>61.25</td>
<td>23.82</td>
<td>3,795,348</td>
<td>1,222,619</td>
<td>45,284</td>
</tr>
<tr>
<td>SHR/Olapcv</td>
<td>52.72</td>
<td>20.5</td>
<td>3,832,318</td>
<td>1,276,189</td>
<td>11,781</td>
</tr>
<tr>
<td>SHRS/Gla</td>
<td>70.03</td>
<td>27.23</td>
<td>3,735,521</td>
<td>1,174,933</td>
<td>14,665</td>
</tr>
<tr>
<td>SR/Jr</td>
<td>50.28</td>
<td>19.55</td>
<td>3,421,364</td>
<td>1,140,725</td>
<td>36,798</td>
</tr>
<tr>
<td>SS/JrHsdMcwi</td>
<td>55.82</td>
<td>21.71</td>
<td>3,383,380</td>
<td>1,136,032</td>
<td>11,124</td>
</tr>
<tr>
<td>SS/Jr</td>
<td>51.42</td>
<td>19.99</td>
<td>3,377,708</td>
<td>1,110,653</td>
<td>38,535</td>
</tr>
<tr>
<td>WAG/Rij</td>
<td>53.58</td>
<td>20.84</td>
<td>3,167,781</td>
<td>1,084,318</td>
<td>51,423</td>
</tr>
<tr>
<td>WKY/Gla</td>
<td>68.17</td>
<td>26.51</td>
<td>3,819,860</td>
<td>1,234,622</td>
<td>11,310</td>
</tr>
<tr>
<td>WKY/NHsd</td>
<td>30.02</td>
<td>11.67</td>
<td>3,877,157</td>
<td>1,313,104</td>
<td>31,875</td>
</tr>
<tr>
<td>WKY/NCrI</td>
<td>63.77</td>
<td>24.8</td>
<td>3,718,449</td>
<td>1,250,041</td>
<td>45,788</td>
</tr>
</tbody>
</table>

See also Figure S1 and Tables S1 and S2.

*Gb of bases mapped and average coverage were calculated after removing clonal reads.

*Numbers of sequence variants relative to the BN reference genome.*
Coevolutionary Gene Clusters

Because rat models of hypertension, insulin resistance, and diabetes were selectively bred for these genetically complex disease phenotypes, the genes underlying these phenotypes may have been coselected and coevolved because of the selective pressure exerted on them during the derivation of these strains. Functionally related genes with a similar evolutionary rate tend to have a similar phylogenetic history (Pagliarini et al., 2008). To identify clusters of genes that coevolved during selection for complex traits, we used a mirror tree approach (Pazos and Valencia, 2001), as illustrated in Figure 2, to test the hypothesis that genetic effects on disease phenotypes may be detectable as coevolved variation in protein-coding sequence between rat strains. Coevolutionary gene clusters were identified using transcripts that show NSC variants, including SNVs causing a stop gain or stop loss in at least one strain compared to the reference BN genome. We calculated evolutionary rate by dividing the number of NSC variants in a transcript by the number of SC variants in the same transcript, both normalized by the length of the transcript. This defined a phylogenetic vector for each transcript. Similarly phylogenetic vectors were calculated for each transcript showing frameshift due to indels.

To take into account the effect of population structure on evolutionary rate, we used SNV data across the strains to estimate principal components contributing to variability in the strains (Sato et al., 2005). The first two principal components were found to explain ~70% of the sequence variability across strains (Figure S2A) and clearly separate Japanese Wistar-derived strains, SD-derived strains, and the remaining rat strains (Figure S2B). We adjusted the evolutionary rate for the first two principal components and then calculated correlation coefficients between the residuals for all possible pairwise combinations of transcripts containing NSC SNVs. Coevolutionary clusters were then identified using 310,690 pairs of transcripts with significant correlation (\( r > 0.96; p \text{~value} < 9.33 \times 10^{-15}; \text{FDR} < 0.1\% \)). A total of 1,955 clusters were identified (Table S3A), the majority of them containing two \((n = 1063)\) or three \((n = 383)\) transcripts, whereas 109 clusters contain 10 or more transcripts (Table S3B). Confirming the cluster analysis, for 1,133 clusters, more than 40% of the NSC mutations represented within the cluster showed linkage disequilibrium \((\text{LD}; r^2 > 0.8)\) with each other, and the majority of them showed perfect LD \((r^2 = 1)\) even though SNVs were located on different chromosomes. After random shuffling of the SNVs, this LD structure was completely lost (Figure S3), suggesting that mutations in these genes may have coevolved under the same selection pressures. Relatively smaller numbers \((n = 145)\) of clusters were identified using frameshift coding mutations (Table S3B) because only 1,634 of the transcripts show frameshift coding mutations in at least one strain compared to the BN reference genome.

To corroborate the data generated from principal component analysis (PCA), we also used the R package EMMA (Kang et al., 2008), which is based on a linear mixed model to correct for population structure and genetic relatedness in model organism association mapping, to identify coevolutionary gene clusters. The networks obtained from EMMA showed high concordance with the networks obtained from PCA-based population structure correction, with ~75% of edges shared between the two networks \((p \text{~value} \text{ for significant overlap} < 10^{-16})\). The number of shared edges was raised to an average of 94% (min 72%, max 100%) when considering edges in clusters identified from PCA-derived networks that were unique to a single strain (Table S4).

The majority of the large clusters contain transcripts mutated either uniquely in a single strain or in strains that were derived from the same founder colony and show close genetic proximity in the phylogenetic tree. For example, the two fawn-hooded rat strains FHH and FHL shared NSC variants in 311 transcripts that were unique just to these strains, and the Wistar-derived first two principal components and then calculated correlation coefficients between the residuals for all possible pairwise combinations of transcripts containing NSC SNVs. Coevolutionary clusters were then identified using 310,690 pairs of transcripts with significant correlation \((r > 0.96; p \text{~value} < 9.33 \times 10^{-15}; \text{FDR} < 0.1\% \)). A total of 1,955 clusters were identified (Table S3A), the majority of them containing two \((n = 1063)\) or three \((n = 383)\) transcripts, whereas 109 clusters contain 10 or more transcripts (Table S3B). Confirming the cluster analysis, for 1,133 clusters, more than 40% of the NSC mutations represented within the cluster showed linkage disequilibrium \((\text{LD}; r^2 > 0.8)\) with each other, and the majority of them showed perfect LD \((r^2 = 1)\) even though SNVs were located on different chromosomes. After random shuffling of the SNVs, this LD structure was completely lost (Figure S3), suggesting that mutations in these genes may have coevolved under the same selection pressures. Relatively smaller numbers \((n = 145)\) of clusters were identified using frameshift coding mutations (Table S3B) because only 1,634 of the transcripts show frameshift coding mutations in at least one strain compared to the BN reference genome.

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The majority of the large clusters contain transcripts mutated either uniquely in a single strain or in strains that were derived from the same founder colony and show close genetic proximity in the phylogenetic tree. For example, the two fawn-hooded rat strains FHH and FHL shared NSC variants in 311 transcripts that were unique just to these strains, and the Wistar-derived
SHR, SHRSP, and WKY strains uniquely shared NSCs in 140 transcripts (Table S3A). However, the finding of NSC or frame-shift variants shared between strains that originated from the same founder population irrespective of their disease status, for example, between a hypertensive strain and its respective normotensive control, indicates that they are unlikely to be responsible for disease phenotypes and are more likely a reflection of shared ancestry. We therefore sought clusters of coevolved transcripts that were unique to strains selected for disease phenotypes. Such clusters would be unique to an individual disease strain and would not be found even in the respective control strains with which they share immediate common ancestors, minimizing the likelihood that the clusters have arisen because of shared ancestry.

A unique coevolutionary cluster was identified in each disease model, the largest of which was found in the GK strain (Tables 2 and S5A). We also identified clusters of transcripts that were uniquely mutated in all disease models derived from the same founder, but not in control strains with which they share immediate common ancestors, minimizing the likelihood that the clusters have arisen because of shared ancestry.

The Milan hypertensive rat strain (MHS) was derived from an outbred Wistar colony along with the normotensive MNS control strain. Hypertension in MHS rats has been shown to be due to excess renal sodium reabsorption (Bianchi et al., 1986). We identified a cluster of 65 transcripts (47 genes) showing NSC sequence variants and a cluster of 5 transcripts (4 genes) showing frameshift coding variants in MHS (Tables S5D and S5E). Importantly, the NSC cluster contains the \textit{Add1} gene encoding Alpha-adducin, in which the amino acid substitution F316Y has been identified as a cause of hypertension in MHS (Bianchi et al., 2005; Ferrandi et al., 2010). Polymorphisms in
the human ADD1 gene have also been associated with hypertension and responsiveness to antihypertensive medica-
tions in several populations (Cusi et al., 1997; Li, 2012). We
also identified, along with Add1, genes Slc4a2, Slc12a8, and
Atp13a4, which show NSC mutations in MHS. Slc4a2 encodes
the anion exchange protein 2, which has shown evidence for
association with human hypertension (Söber et al., 2009).
Slc12a8 belongs to the cation chloride cotransporter family
(Hebert et al., 2004), and Atp13a4 encodes a cation-transporting
P-type ATPase (Kwasnicka-Crawford et al., 2005). In light of
longstanding observations on the relationship between ion
transport and hypertension in rats and humans (Bianchi et al.,
1990; Lifton et al., 2001), the physiological consequences of
mutations in these ion transporters merit further investigation.

The BBDP rat strain is an extensively studied model of
type 1 diabetes, occurring in association with profound T cell
lymphopenia with severe reduction or absence of CD4 and
CD8 T cells (Jackson et al., 1983). We identified a cluster of
63 transcripts (47 genes) showing NSC variants and also 16
transcripts (12 genes) showing frameshift variants that are
unique to BBDP (Tables S5F and S5G). The frameshifts include
a 1 base pair (bp) deletion in Gimap5, a member of the GTPase
of the immunity-associated protein family. This frameshift
mutation was previously identified as the cause of T cell lym-
phopenia in BBDP (Dalberg et al., 2007; Hornum et al., 2002;
MacMurray et al., 2002). The finding that two genes identified
previously as disease susceptibility genes in MHS and
BBDP were among the small gene sets that we defined as
uniquely variant in these strains using coevolutionary cluster
analysis suggests that genes underlying other QTLs are
likely to be found within the coevolved gene sets of these
and other disease strains and that these gene sets should be
prioritized in future investigations of disease phenotypes in
these strains.

Although we identified clusters that were shared between dis-
ease models derived from different founder colonies such as
BBDP and MHS (n = 17 transcripts) or GK and SBH (n = 17 tran-
scripts), no clusters were found that revealed potentially func-
tional mutations across all models of hypertension, diabetes,
To assess whether the PASS regions identified by the RSD analysis were due to the varying demographic history of the rat strains or, alternatively, were due to the effects of selection pressures exerted during derivation of the strains, we performed coalescent simulations that accounted for the known demographic history of the 22 laboratory rat strains, but not for the selection pressure exerted on them. In our null model, obtained by simulation on demographic history without phenotypic selection, no genomic region unique to a strain with length greater than 40 kb was observed, with a statistically unlikely probability of observing unique haplotype blocks of length 40 kb (p = 0.0003) or 30 kb (p = 0.002). All of the PASS regions that we identified were longer than or equal to 30 kb, and some were more than 1M b (Table S6). The identification of PASS regions that were considerably longer than those observed under the null model without selection provides evidence that PASS regions are enriched for positively selected haplotype blocks.

As further evidence that the identified PASS regions were positively selected during the phenotype-driven derivation of these strains, we performed coalescent simulations that accounted for the known demographic history of the 22 laboratory rat strains, but not for the selection pressure exerted on them. In our null model, obtained by simulation on demographic history without phenotypic selection, no genomic region unique to a strain with length greater than 40 kb was observed, with a statistically unlikely probability of observing unique haplotype blocks of length 40 kb (p = 0.0003) or 30 kb (p = 0.002). All of the PASS regions that we identified were longer than or equal to 30 kb, and some were more than 1M b (Table S6). The identification of PASS regions that were considerably longer than those observed under the null model without selection provides evidence that PASS regions are enriched for positively selected haplotype blocks.

As further evidence that the identified PASS regions were positively selected during the phenotype-driven derivation of these strains, we estimated the ratio of NSC variants to SC variants in PASS regions. PASS regions were highly enriched for NSC mutations as compared to the rest of the genome, with an NSC to SC ratio of 1.65 and 0.76, respectively, in PASS regions and across the whole genome (Fisher’s exact test p value = 2.19 × 10⁻⁵; Figure 3D).

We next assessed what proportion of PASS regions colocalized with known physiological quantitative trait loci (pQTLs) for the metabolic and cardiovascular phenotypes manifested in these strains. Of the 96 PASS regions that were unique to 1 of the 11 cardio-metabolic disease strains, 25 were colocalized with pQTLs reported in RGD (http://rgd.mcw.edu/) in the same strain in which each PASS region was identified. This overlap was marginally significant compared to the overlap expected by chance (permutation test p value = 0.04). Interestingly, the majority (78%) of the PASS regions were localized either directly under or in close proximity (<10 Mb) to a peak of at least one QTL linkage (Table S7).

To identify genes and pathways that may have contributed to phenotype-driven selection during generation of disease strains, we carried out gene ontology (GO) and KEGG pathway analyses of genes in the PASS regions in these strains. Only two strains, SS and FHH, showed significant enrichment (p < 0.05) for specific biological processes or pathways. In FHH, 44 genes reside within 14 FHH-specific PASS regions (Figure 3A), and these were enriched for genes involved in the renin-angiotensin system (RAS) and in proteolysis (p = 3.4 × 10⁻⁴ and 1.6 × 10⁻⁸, respectively).
Three genes within FHH PASS regions were components of RAS, including Cma1 and CtsG, both of which are reported to mediate non-angiotensin conversion enzyme (ACE)-dependent conversion of angiotensin I to angiotensin II (Uehara et al., 2013). The RAS system is central to the regulation of blood pressure in all mammals, and FHH rats show reduced sensitivity to angiotensin II, which has been proposed as a cause of FHH susceptibility to renal disease (van Rodijnen et al., 2002). In humans, a haplotype at CMA1 has been weakly associated (p = 2 × 10^{-4}) with hypertension in Han Chinese (Wu et al., 2012), and in mice, chymase 1 has been proposed to modulate regulation of blood pressure by angiotensin II (Li et al., 2004). In addition, ACE inhibitors and angiotensin receptor blockers are among the most widely used antihypertensive agents (Kobori et al., 2007). CMA1 and CTSG have NSC sequence variants that are unique to FHH among rat strains (His → Arg in CMA1; Ala → Thr in CTSG). The histidine at codon 94 in CMA1 is conserved in all mammals. CMA1 and CTSG are therefore compelling candidates as regulators of blood pressure in FHH and, because of their enzymatic function in non-ACE-dependent conversion of angiotensin I to angiotensin II, are potential novel antihypertensive drug targets.

The 44 genes in FHH-specific PASS regions have a markedly increased NSC-to-SC ratio (Figure 4C) and are highly enriched for GO biological processes related to proteolysis. The 12 genes in this GO category are all mast cell proteases or belong to the family of granzymes, which are also expressed in mast cells and reside within five FHH PASS regions on a 2.13 Mb segment of chromosome (Chr) 15 that shows highly negative Tajima’s D (Figures 4A and 4B). Ten of these genes have NSC variants that are unique to FHH. FHH rats develop glomerulosclerosis and interstitial fibrosis with glomerular hyperfiltration, albuminuria, and end-stage renal failure (de Keijzer et al., 1998; Kreisberg and Kamovsky, 1978; Kirz et al., 1998). A consomic study in which FHH chromosomes were substituted with BN chromosomes showed that substitution of Chr 15 resulted in reduced blood pressure, albuminuria, and glomerular damage, indicating that genetic determinants of these phenotypes reside on FHH Chr 15 (Mattson et al., 2007). Mast cell numbers are increased in human hypertensive nephropathy and are associated with renal disease in the context of nephrectomy-induced hyperfiltration (Weiker et al., 2008). Among the granzymes and mast cell proteases within the Chr 15 PASS regions, GzmF and Mcpt10 contain nonconservative amino acid substitutions (Lys → Gln and Gly → Cys, respectively). Taken together, these data implicate this segment of Chr 15 and specifically the unique FHH variation in granzymes and mast cell proteases in the development of hypertension-induced renal damage in this rat strain.

In SS, 16 protein-coding genes reside within the non-SS-specific PASS regions (Figure 3A and Table S6), and these were enriched for aromatic compound catabolic processes (p = 3.9 × 10^{-5}), owing to the presence of the three paraoxonase genes Pon1, Pon2, and Pon3, in one of the SS PASS regions on Chr 4. Paraoxonase genes protect against oxidative stress and have been implicated in atherosclerosis and diabetes mellitus in humans (Précourt et al., 2011) and blood pressure regulation in mice (Gamlil-Lazarovich et al., 2012). Oxidative stress is associated with endothelial dysfunction and hypertension in humans and in a wide range of experimental models, including the Dahl SS rat (Kushiro et al., 2005; Swei et al., 1997; Vaziri and Rodriguez-Iturbe, 2006). Because the paraoxonase genes overlap with SS QTLs for hypertension (Table S7), the paraoxonase

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**Figure 4. FHH PASS Regions on Chromosome 15**

(A) Blue dots represent relative SNV density (RSD) in 10 kb windows for the FHH rat strain on chromosome 15. The genomic region unique to the FHH strain is highlighted by the gray dotted lines. Heatmap (bottom) showing RSD in the genomic region unique to FHH and flanking regions across all laboratory rat strains sequenced. Only the FHH rat strain showed an RSD value equal to 100. Remaining strains showed an RSD value of zero across almost all of the PASS regions, indicating that only the FHH strain shows SNVs against the BN reference genome in this region. Linkage disequilibrium (LD) structure of SNVs in the genomic region unique to FHH show that ~98% of the SNVs in this region were in significant LD.

(B) Tajima’s D in genomic region unique to FHH rat strain. The PASS region unique to FHH is highlighted, showing highly negative Tajima’s D value. (C) The genomic region unique to FHH shows a markedly increased ratio of NSC to SC variants.
genes are compelling candidates as contributors to the hypertension phenotype in the SS strain.

**DISCUSSION**

Laboratory rat strains have been studied extensively for cardiovascular and metabolic phenotypes for several decades, but the vast majority of the genes and mutations underlying these phenotypes remain to be identified. This is at least in part because of the absence of the genome sequences and the lack of comprehensive catalogs of genomic variants for these strains. Previous catalogs of variants in the rat were restricted to 20,238 SNVs in 167 inbred rat strains, which have been used mostly as markers for QTL mapping (Saar et al., 2008). More recently, the report of 3.6 million SNVs and 0.3 million indels in the SHR strain has facilitated research aimed at understanding the mechanisms underlying complex traits in this strain (Atanur et al., 2010; Heining et al., 2010; McDermott-Roe et al., 2011; Simonis et al., 2012). Another constraint has been that existing data sets of rat variants are biased toward single-nucleotide changes, and apart from SHR, other types of variants such as indels were partially or completely missing. This study reports an extensive catalog of 9.6 million SNVs, 3.5 million short indels, and 719,929 structural variants across 27 laboratory rat strains, which are among the most widely used animal models of hypertension and diabetes.

We used ~2 Mb of high-quality BAC sequence from the non-reference strain LE/Stm to estimate the false-positive and false-negative rates of variant calling. Estimates of false positive and false negatives reported in other model organisms have been based on only the accessible part of the genome (Keane et al., 2011), which may underestimate the false-negative rate at the level of the whole genome. Although we estimate relatively high false-negative rates (8.1% and 10.98% for SNVs and indels, respectively), these false-negative rates are lower than those reported for mouse genome sequences (Keane et al., 2011) and represent upper estimates for missing variants as measured against the entire reference genome, rather than just the accessible genome. The false-positive rates were also lower in our sequences than those reported by Keane et al. for the mouse. The reasons for the lower false-positive and -negative rates may relate to the longer sequence reads and improved algorithms for mapping and variant calling used in the present studies.

We used the new sequence data from the 27 rat strains for two main purposes: to provide a firm foundation for understanding the phylogenetic history of the laboratory rat and to give new insights into the genetic and pathophysiological basis of the disease phenotypes for which the various rat strains were selected. Our phylogenetic analysis indicates that differences in laboratory rat strains are primarily due to genetic differences in the founder colonies from which the rat strains were derived, reflecting differences that were most likely present in the original outbred founder populations (Figure 1).

We used two approaches to give new insights into the genetic and pathophysiological basis of disease phenotypes in rat strains. First, we used the phylogenetic history of the strains to identify clusters of genes with correlated evolutionary rates. Such methods have been widely used to identify coevolving and functionally related proteins and small RNA pathways (de Juan et al., 2013; Pagliarini et al., 2008; Pazos and Valencia, 2001; Tabach et al., 2013). We defined coevolutionary clusters that were unique to each disease strain and among these clusters of genes identified two genes, Add1 and Gimap5, previously shown by positional cloning to underlie susceptibility to hypertension in MHS and lymphopenia and diabetes in BBDP, respectively (Bianchi et al., 2005; Ferrandi et al., 2010; Hornum et al., 2002; MacMurray et al., 2002). Given the small number of genes shown to underlie rat disease phenotypes (Aitman et al., 2008), the finding of Add1 in MHS and Gimap5 in BBDP in coevolutionary clusters for these strains provides proof of concept of the efficacy of this approach in these rat strains. We also found among the coevolutionary clusters for MHS three further ion transport genes (Slc4a2, Slc12a8, Atp13a4), of which one, Slc4a2, shows a predicted nonconservative Gly→Asp amino acid substitution. The longstanding relationship between dysregulated ion transport and hypertension in MHS rats and in humans (Bianchi et al., 1990; Lifton et al., 2001) suggests these ion transporters as compelling candidates as hypertension susceptibility genes. Given that the human orthologs of rat genes within coevolutionary clusters showed significant enrichment for GWAS hits for hypertension and metabolism-related phenotypes, further study of these genes may shed light on the pathogenetic mechanisms for these phenotypes in both species.

Our second approach to gaining new insights into disease phenotypes was to identify putative selective sweeps, which we term PASS regions, that were unique to individual strains and may have been retained during the derivation of these disease models. Precedents for this type of analysis exist in the study of domestication of other species, including maize, silkworm, chicken, and dog (Axelsson et al., 2013; Rubin et al., 2010; Wright et al., 2005; Xia et al., 2009), but the approach has not been applied previously to the study of disease phenotypes. Our analyses took account of the known demographic history of the rat strains and used several methods to distinguish genomic segments that arose due to selection from those that may have arisen from population structure.

First, the PASS regions that we identified as unique to individual rat strains ranged in length from 30 kb to 1.57 Mb, whereas a null model obtained by coalescent simulation that took into account demographic history identified no regions with length > 30 kb. Second, we showed that LD was markedly increased across PASS regions compared to the rest of the genome. Third, we showed highly negative Tajima’s D values within PASS regions, strongly supporting the presence of true selective sweeps among the identified PASS regions. Finally, we found that the ratio of nonsynonymous-to-synonymous coding variants in PASS regions was more than double that across the rest of the genome (1.65 and 0.76, respectively). These various approaches, used as indicators of evolutionary selection in a variety of scenarios (Axelsson et al., 2013; Rubin et al., 2010; Wright et al., 2005; Xia et al., 2009), strongly support the view that the identified PASS regions are significantly enriched for genomic segments retained by phenotype selection during derivation of these strains.
The PASS regions that we identified were significantly enriched for cardiovascular or metabolic QTLs previously mapped in these strains. In addition, where there was overlap between QTLs and PASS regions, the majority of PASS regions were localized at or close to the peak of the QTL linkage, in most cases suggesting candidates for these QTLs and in some cases single genes. These genes may now be tested in targeted gene knockin or knockout experiments that may now be easily performed in the rat on any genetic background (Jacob et al., 2010).

To identify specific pathways that may have contributed to phenotype-driven selection during derivation of disease strains, we carried out GO analyses of genes within PASS regions and identified enrichment for specific biological processes in two strains, FHH and SS. PASS regions in FHH showed enrichment for genes in the RAS and for proteolytic enzymes. The finding of enrichment for genes in the RAS is not unexpected, given the central role of this system in blood pressure regulation. However, canonical RAS genes that are either hypertension susceptibility genes or drug targets are related to ACE, whereas two of the genes in the FHH PASS regions mediate non-ACE dependent conversion of angiotensin I to angiotensin II. These genes, Cma1 and Ctg, are therefore compelling hypertension candidate genes, particularly as the histidine at codon 94 in Cma1 that is uniquely variant in FHH among all rat strains is conserved in all mammals. The finding in FHH PASS regions of enrichment for mast cell proteins, some of which show NSC variants at highly conserved amino acids, suggests a previously unidentified pathway for hypertension-induced nephropathy that is consistent with previous observations in human and rat hypertension, the mechanism for which is at present unknown. Similar considerations apply to the enrichment of genes associated with protection against oxidative stress, found to be enriched in PASS regions in SS.

We note that our analysis of PASS regions and coevolved clusters across disease models did not identify shared clusters of genes that were common to all strains that carry the same disease phenotype, suggesting differing genetic etiologies for complex traits across these disease models, as previously suggested (Stoll et al., 2000). This may reflect the differing pathophysiology of traits such as hypertension across these strains (Jong, 1984) but is also likely due, in part, to the different genetic backgrounds on which these models were developed.

The data from these studies, including the genome sequences, catalogs of sequence variants, sets of coevolved gene clusters, and PASS regions across 27 rat strains, represent a step change in the genome resources available for study of complex phenotypes in the rat model. Identification of genes underlying complex phenotypes in this model has until now rested on laborious and lengthy gene-mapping studies that have mostly localized disease genes to the genome at a resolution of tens of mega base pairs. By taking advantage of whole-genome sequences and population structure in 27 strains, these studies identify coevolved segments and haplotype blocks that are unique to individual disease strains. These genes and pathways, including those previously identified as QTL genes and unidentified genes such as non-ACE-dependent angiotensin converting enzymes, mast cell proteases, and proteins that protect against oxidative stress, were most likely highlighted in these studies because they were selected in the original phenotype-driven derivation of these strains. We believe this to be the first evolutionary analysis of artificial selection for disease phenotypes and suggest that further analysis of these genes will confirm many of them as new genes for these phenotypes that will shed new light on the pathogenesis of these conditions in rats and humans.

**EXPERIMENTAL PROCEDURES**

Also see Extended Experimental Procedures.

**Rat Strains and Genome Sequencing**

We sequenced the genomes of 25 rat strains on the Illumina sequencing platform (Table 1) and analyzed these together with two rat genomes that we sequenced previously.

**Mapping to the Reference Genome**

Quality filtered Illumina paired-end and/or mate pair reads were mapped to the BN reference genome RGSC-3.4 (Gibbs et al., 2004) using BWA-0.5.8c (Li and Durbin, 2009). The genome (BN.Lx) sequenced previously on the SOLID sequencing platform was mapped using BFAST-0.8.4e (Homer et al., 2009).

**Single-Nucleotide Variants and Short Indel Detection**

Genomic variants (single-nucleotide variants and short indels [1–15 bp]) were detected using the Genome Analysis Toolkit (GATK version 1.0.6001) (DePristo et al., 2011; McKenna et al., 2010).

**Structural Variant Prediction**

To identify structural variants, we used two independent methods. Structural variants including insertions, deletions, inversions, and translocations were predicted using the software tool BreakDancer1.2 (Chen et al., 2009). In addition, large deletions were predicted using a custom Perl script as described previously (Atanur et al., 2010), using mapping flags provided by BWA for paired-end reads (Li and Durbin, 2009).

**Functional Consequence Analysis of Predicted Variants**

Functional consequences of predicted variants (SNVs and short indels) were estimated using “variant effect predictor (VEP)-v2.4” (McLaren et al., 2010) on the ENSEMBL 66 genotype set.

**Validation of Genomic Variants**

To estimate sensitivity and specificity of variant calls, the sequence of 13 bacterial artificial chromosome (BAC) clones from various chromosomes (~2 Mb in length) of LE/Stm rat were downloaded from the EMBL web site (accession numbers FO117624-FO117632 and FO181540-FO181543).

**Reconstruction of the Phylogenetic Tree**

A distance matrix, derived from SNVs between all possible pairs of strains, was used to construct the phylogenetic tree using the Fitch-Margoliash method with contemporary tips, version 3.69 from package Phylip (Felsenstein, 1989), with 1,000 bootstraps.

**Identification of Coevolutionary Gene Clusters**

Coevolutionary gene clusters were identified using a mirror tree approach. We used two independent methods to correct for population structure: (1) principal component analysis and (2) a linear mixture model.
Identification of Selective Sweeps between Different Rat Strains
To identify genomic regions shared between different rat strains, relative SNV density (RSD) in nonoverlapping 10 kb windows was calculated in all 26 rat strains (excluding BN.Lx) using the following formula:

$$\frac{\text{Number of SNVs in strain } i}{\text{Total number of SNVs}} \times 100$$

Chi-square statistics were used to determine the goodness of fit to a model of equal distribution of SNVs within each 10 kb window between "n" strains. Chi-square statistics were calculated for all possible combinations of strains according to:

$$SC = \min\left(\sum_{k=1}^{n} \left(\frac{{O}_k - {E}_k}{E_k}\right)^2\right)$$

Adjacent 10 kb windows were merged if they were shared between the same strain combinations.

Calculation of Linkage Distribution
Linkage distribution between the SNVs within the coevalolutionary gene clusters and SNVs within PASS regions was calculated using Haploview 4 (Barrett et al., 2005).

Coalescent Simulations
Coalescent simulations were performed using the program msms (Ewing and Hermisson, 2010).

Calculation of Population Genetics Statistics Tajima’s D
Tajima’s D was calculated in nonoverlapping 10 kb windows using custom Perl script as described in (Tajima, 1989). FDR was calculated using the null distribution of Tajima’s D from simulated data.

ACCESSION NUMBERS
All sequence reads from this study are deposited in the EBI Sequence Read Archive under accession number ERP002160. All sequence variants are deposited in RGD (http://rgd.mcw.edu/).

SUPPLEMENTAL INFORMATION
Supplemental Information includes Extended Experimental Procedures, four figures, and eight tables and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2013.06.040.

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