Genome-wide association study of behavioral, physiological and gene expression traits in outbred CFW mice

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Although mice are the most widely used mammalian model organism, genetic studies have suffered from limited mapping resolution due to extensive linkage disequilibrium (LD) that is characteristic of crosses among inbred strains. Carworth Farms White (CFW) mice are a commercially available outbred mouse population that exhibit rapid LD decay in comparison to other available mouse populations. We performed a genome-wide association study (GWAS) of behavioral, physiological and gene expression phenotypes using 1,200 male CFW mice. We used genotyping by sequencing (GBS) to obtain genotypes at 92,734 SNPs. We also measured gene expression using RNA sequencing in three brain regions. Our study identified numerous behavioral, physiological and expression quantitative trait loci (QTLs). We integrated the behavioral QTL and eQTL results to implicate specific genes, including Azi2 in sensitivity to methamphetamine and Zmynd11 in anxiety-like behavior. The combination of CFW mice, GBS and RNA sequencing constitutes a powerful approach to GWAS in mice.

In the last decade, GWAS have demonstrated that common alleles influence susceptibility to virtually all common diseases1–3. The success of GWAS in elucidating the genetic determinants of disease in humans is due in part to the large number of recombinations among unrelated individuals, which permits high-resolution mapping across the genome. One important conclusion from these studies is that most causal loci seem to be due to regulatory rather than coding polymorphisms.

Mice offer a powerful tool for elucidating the genetic architecture of complex traits: environmental factors can be held constant or systematically varied; genome editing permits experimental testing of identified genotype–phenotype relationships; most mouse genes have a human homolog, allowing rapid translation to humans; and relevant tissues can be obtained under tightly controlled conditions and used to identify gene eQTLs. However, the mouse populations used in most previous studies lacked sufficient recombination to narrow the implicated loci to a tractable size and thus generally failed to identify specific genes5,6.

In this study, we mapped QTLs and eQTLs using CFW mice, which are a commercially available outbred population7. Although CFW mice were not developed for genetic research, they have several attractive properties. CFW mice were derived from a small number of founders and have subsequently been maintained as an outbred population for more than 100 generations, thus degrading LD between nearby alleles8–10. Although CFW mice have longer-range LD than most human populations, they have less LD than other commercially available laboratory mice11 and therefore should provide fine-scale mapping resolution. In comparison to humans, the more extensive LD in CFW mice means that fewer markers are needed to perform GWAS and correspondingly lower levels of significance are required because fewer independent hypotheses are tested. We used GBS to overcome another barrier to GWAS in mice, which is the high cost and limited coverage of extant SNP genotyping arrays. Finally, on the basis of the importance of regulatory variation suggested by human GWAS11, we identified eQTLs that co-mapped with behavioral QTLs in an effort to identify the most likely causal genes.

RESULTS

We phenotyped 1,200 male CFW mice for conditioned fear, anxiety-like behavior, methamphetamine sensitivity, prepulse inhibition, fast-twitch skeletal muscle, hindlimb muscle strength, bone mineral density, hippocampal and striatal gene expression, and physiological and behavioral traits in five brain regions. We performed a genome-wide association study (GWAS) of these phenotypes in CFW mice using genotyping by sequencing (GBS). We identified numerous behavioral, physiological and expression quantitative trait loci (QTLs) in an effort to identify the most likely causal genes.
Genotyping
Existing mouse SNP genotyping technologies, such as the Mouse Universal Genotyping Array (MUGA), MegaMUGA\(^{12}\), the more recent GigaMUGA\(^{13}\) and the Mouse Diversity Array (MDA)\(^{14}\), were not designed to capture common genetic variation in the CFW population. Furthermore, we sought to reduce the cost of genotyping, which has been a barrier to GWAS in mice. Therefore, we adapted GBS, which was originally developed in maize\(^{15}\), for use in mice. We used GBS to genotype 1,024 CFW mice and identified 92,734 SNPs, representing a larger number of polymorphic SNPs than were found using MegaMUGA. The remaining 13,450 SNPs (14%) represent autosomal biallelic SNPs after filtering, 79,284 (86%) of which were present in dbSNP (v137). The remaining 13,450 SNPs (14%) represent new SNPs that had not previously been reported. The distribution of GBS SNPs on autosomes is shown in Figure 2a. The non-uniform distribution of SNPs is likely due to differences in the numbers of polymorphic markers among all laboratory mice (Fig. 2a and Supplementary Fig. 5) and regions that are identical by descent among CFW mice. The non-uniform distribution of polymorphic SNPs seems to be a characteristic of CFW mice, as polymorphic SNPs identified by the MegaMUGA array showed a similar pattern ($\chi^2 = 0.43$ on a log scale; Fig. 2a).

To assess the quality of the GBS genotypes, we estimated the genotyping error rate in two ways. First, we compared genotypes for typing error rate in two ways. First, we compared genotypes for genotypes using GBS. All genome-wide genotyping using GBS.

Figure 2 Components of the study. Our study consisted of four phases, including behavioral testing and measurement of physiological traits (phase A), GBS (phase B), measurement of gene expression in brain tissues using RNA-seq (phase C), and QTL mapping for physiological and behavioral traits and for gene expression (phase D). The red dotted line for phase D corresponds to $P = 2 \times 10^{-7}$.

Genetic architecture of the CFW population
Comparing LD in different populations is useful for gauging mapping resolution\(^{16}\). LD ($r^2$) decays rapidly in CFW mice in comparison to other populations, consistent with previous findings based on a much smaller number of SNPs\(^{9,17}\) (Fig. 2b), supporting the suitability of the CFW population for high-resolution mapping. Notably, the majority of the SNPs we identified in CFW mice segregate among the subspecies\(^{18–20}\) (Supplementary Fig. 11).

SNP heritability
Supplementary Table 1 shows SNP heritability\(^{21,22}\), which is the proportion of variance in a trait explained by available SNP genotypes. SNP heritability estimates ranged from 9–60%, with a mean of 28%. The mean SNP heritability for physiological traits was slightly higher (32%) than for behavioral traits (27%).
Genome-wide association studies
We mapped QTLs for 66 behavioral and physiological phenotypes (Fig. 3 and Supplementary Tables 1 and 2). We used GEMMA to fit a linear mixed model (LMM) and quantify support for an association at each SNP. We also used a simpler linear model that did not correct for population structure and observed that this model produced broadly similar results (Supplementary Fig. 12). However, we have presented the results from the LMM-based analysis because this approach may reduce subtle inflation of the test statistics due to close relationships or fine-scale population structure. We calculated a significance threshold via permutation, which is a standard approach for QTL mapping in mice that controls for the type I error rate\(^3\).\(^{24}\) (Supplementary Fig. 13). This approach identified numerous QTLs for physiological and behavioral traits (Fig. 3 and Supplementary Figs. 14–18) that exceeded \(P = 2 \times 10^{-6}\) (\(P < 0.1\)).

Supplementary Table 2 contains more detailed information about all the most significant physiological and behavioral QTLs.

For testis weight, we found a strong association with rs6279141 on chromosome 13 (\(P = 4.51 \times 10^{-18}\), Fig. 3b) that accounted for 7.5% of variation in this trait. The implicated region contained few genes (Fig. 3d), one of which was Inhba, a gene that has been shown to affect testis morphogenesis, testicular cell proliferation and testis weight in mice.\(^25\)–\(^27\) Thus, both Inhba and Slc4a2 have a critical role in osteoclasts in mice\(^34\), and homozygous deletion of Slc4a2 is associated with the osteopetrosis-like phenotype ‘marble bone disease’ in Red Angus cattle.\(^35\) Thus, both Abcf2 and Slc4a2 are viable candidates for this region. The association on chromosome 11 contains the gene Col1a1. In humans, osteogenesis imperfecta type I can be caused by a null allele of COL1A1 and results in gracile bones with decreased strength.\(^36\)–\(^37\) Col1a1 is also associated with other bone size phenotypes, making Col1a1 a likely causal gene for this locus.

Finally, we identified several associations for behavioral traits, including for methamphetamine sensitivity on chromosome 6 at rs22397909 (\(P = 9.03 \times 10^{-7}\)) and on chromosome 9 at rs46497021 (\(P = 1.58 \times 10^{-4}\)); these associations account for 2.6% and 2.1% of phenotype variance, respectively (Fig. 4). We also identified an association for anxiety-like behavior with rs2384652 on chromosome 13 (\(P = 7.31 \times 10^{-8}\)) that explains 3% of the trait variance. For prepulse inhibition (PPI; 12 dB), we identified associations with rs264716939 on chromosome 7 (Fig. 3c) and rs230308064 on chromosome 13 (\(P = 1.18 \times 10^{-6}\) and 2.17 \(\times 10^{-6}\), respectively). There were many genes in the ~3-Mb region on chromosome 7 that were associated
with PPI (Fig. 3e), making it difficult to identify the causal gene(s). Candidate genes for the associations with behavioral traits are discussed below.

Expression quantitative trait loci

In an effort to identify causal genes in our behavioral QTLs, we mapped eQTLs for three brain regions that are critical for the behaviors that we studied. We performed RNA-seq on mRNA from three brain regions: hippocampus (n = 79), striatum (n = 55) and prefrontal cortex (n = 54). In a cis-eQTL scan that was limited to the region flanking the gene being interrogated (Supplementary Figs. 19 and 20), we identified a total of 6,045 associations for 4,174 genes (Fig. 4a, Supplementary Fig. 21 and Supplementary Table 3) at a permutation-derived significance threshold of P < 0.05 (this threshold reflects a per-gene, per-brain region). For 534 of these genes, we identified a cis-eQTL in all three tissues. For an additional 803 genes, we identified a cis-eQTL in two of the three tissues (Fig. 4b). The RNA-seq data were generated from a set of partially overlapping individuals; therefore, we did not perform a joint analysis of the three brain tissues.

In addition, we searched for cis-eQTLs by examining allele-specific expression (ASE), which measures the relative expression of the two possible RNA alleles derived from a heterozygous SNP40,41. We identified 655 genes with ASE in at least one of the three tissues. Of these, 380 (58%) were found only using ASE analysis and 275 (42%) were also identified in the conventional cis-eQTL scan, suggesting that there was more overlap than would be expected by chance. Overlap was likely limited by several factors, including type I errors in the ASE analysis and type II errors in both the ASE analysis and conventional cis-eQTL mapping.
We also mapped eQTLs across the genome for each gene in an effort to detect trans-eQTLs. We identified 2,278 trans-eQTLs that were significant (P < 0.05, permutation-based threshold) after testing 43,414 transcripts across the three brain regions. We expected almost that many tests to be positive under the null hypothesis. Consistent with this expectation, a quantile–quantile plot of these results suggested that only a small number of the results were true positives (Supplementary Fig. 22). As expected, most true positive results seem to be from the hippocampus, which had the largest sample size (n = 79).

Integration of behavioral QTLs with eQTLs

On the basis of evidence from human GWAS, we anticipated that heritable gene expression polymorphisms (eQTLs) would be responsible for most of the observed behavioral associations. Therefore, we tried to identify eQTLs that co-mapped with behavioral QTLs, under the assumption that the eQTL might be the molecular cause of the behavioral QTL. For example, we observed an association between methamphetamine sensitivity and rs238465220 on chromosome 13 (P = 7.3 × 10−6; Supplementary Fig. 18). The implicated region is small (<1 Mb) and contains only two genes: Cmc1 and Azi2 (Fig. 4d). We identified cis-eQTLs for both genes in the striatum, which is the tissue that is most relevant for methamphetamine sensitivity. However, rs46497021 was most strongly correlated with Azi2 expression (P = 1.2 × 10−6; Fig. 4e). In addition, the patterns of SNPs associated with methamphetamine sensitivity and Azi2 expression showed obvious overlap. Therefore, although both Cmc1 and Azi2 are credible positional candidates, the eQTL data suggest that Azi2 is most likely to be the causative gene. To our knowledge, neither gene has previously been implicated in dopaminergic/striatal processes, suggesting that this observation may offer new insights into the biology of this drug-abuse-relevant trait.

Similarly, we identified an association between anxiety-like behavior and rs238465220 on chromosome 13 (P = 7.3 × 10−6; Supplementary Fig. 17). The implicated region spans ~1.5 Mb and contains four genes: Chrm3, Larphb, Dip2c and Zmynd11. Among these genes, rs238465220 was also associated with expression of Zmynd11 in the hippocampus, suggesting that this locus may influence anxiety-like behavior through regulation of Zmynd11 expression (Supplementary Fig. 23). Zmynd11 has not previously been implicated in anxiety; however, copy number variants in ZMYND11 were recently shown to be associated with autistic tendencies and aggressive behaviors in humans42. These examples illustrate the usefulness of combining GWAS with eQTL data to identify the molecular mechanism by which a chromosomal region influences a complex trait.

DISCUSSION

We performed a GWAS in a commercially available outbred mouse population, which identified numerous physiological, behavioral and expression QTLs. In several cases, the implicated loci were smaller than 1 Mb and contained just a handful of genes, including an obvious candidate. In addition, we used the eQTL results to further parse between the genes in the intervals that were implicated for the behavioral traits.

The goal of using CFW mice was to enhance our mapping resolution. CFW mice have shorter-range LD than other commercially available populations9. Using GBS genotypes, we estimated LD in CFW mice and compared it to that of other mapping populations (Fig. 2b). The F14 generation of the LG/SM×AIL advanced intercross line (LG×SM-AIL) that we have used in previous studies43–46 showed more extensive LD than CFW mice. Various outbred heterogeneous stocks, typically made up of eight inbred strains, have also been used in previous mapping efforts47–49. We examined one heterogeneous stock49 and found that it also had longer-range LD than CFW mice. The Hybrid Mouse Diversity Panel (HMDP)50,51, which is a collection of 100 inbred strains, is currently being genotyped using GBS, and we expect that it will also exhibit extensive LD.
of approximately 100 inbred mouse strains that has been used for QTL mapping, also showed greater LD than CFW mice, as did a smaller panel of 30 inbred strains\textsuperscript{52}. DO mice\textsuperscript{12,18,19,53} exhibited LD decay that was almost as degraded as that of CFW mice. Populations like the AIL and heterogeneous stocks (including the DO population) are expected to show decreased LD in the future because of the accumulation of additional recombination (for example, the LG × SM-AIL is now at generation 62). The MF-1 population is another commercially available outbred population that has been used to map QTLs\textsuperscript{50,54}, but we were unable to obtain the data needed to estimate LD decay in this population. Comparing LD patterns in different populations is a common method for estimating mapping resolution\textsuperscript{16}; however, additional factors including the allele frequency distribution\textsuperscript{55}, population structure\textsuperscript{56}, error rates and the number, effect size and frequency of causal variants all influence power and mapping resolution. Despite these limitations, our comparison of LD (Fig. 2b) and our mapping results (Figs. 3 and 4, Supplementary Figs. 14–18 and 21–23, and Supplementary Table 2) suggest that CFW mice are an attractive option for fine-mapping studies.

Another important parameter for GWAS is allele frequency, as power to identify associations increases with greater MAF. Laboratory mouse populations have higher average MAF than humans or wild mouse populations\textsuperscript{57}. We found that 73% of SNPs genotyped in this study had MAF \textgtr 0.05, although our SNP filtering steps may have underestimated the number of rare SNPs. Populations produced by crossing inbred strains, such as F\textsubscript{2} crosses, recombinant inbred lines, AILs and heterogeneous stocks, typically have even more desirable MAF distributions\textsuperscript{43}. Because the ascertainment of SNPs included on genotyping platforms directly influences the estimated MAF distribution, we did not attempt to use publicly available data to compare MAFs in commonly used mapping populations.

We found that CFW mice lacked genetic variability in certain regions; for example, chromosome 16 had a low density of polymorphic markers, as measured using both GBS and MegaMUGA (Fig. 2a), and no significant QTLs (Fig. 3a). This region is an example of a previously described tendency for laboratory mouse populations to harbor regions that are identical by descent\textsuperscript{43,57}.

Several other advantages of CFW mice include their commercial availability, their low cost and the ability to acquire non-siblings upon request. We also found that the CFW mice were easy to handle, and their uniform coat color simplified automated scoring of certain behavioral traits.

One barrier to more widespread adoption of GWAS in mice has been the lack of universal and economical SNP genotyping platforms. In this report, we demonstrate the use of GBS to overcome this obstacle. GBS is a reduced-representation sequencing approach in which a small fraction of the genome is sequenced at moderate depth to obtain genotypes at a subset of markers. Although GBS shares some characteristics with low-coverage whole-genome sequencing\textsuperscript{58–60}, GBS yields high coverage for a subset of the genome, thus acquiring information about fewer SNPs but with greater confidence. Our GBS methods included a custom-designed library preparation protocol (which reduced per-sample costs) and used the standard software toolkits GATK\textsuperscript{61} and IMPUTE2 (ref. 62). An advantage of GBS was that it did not require preselection of polymorphic SNPs. We chose conservative criteria for SNP calling, which yielded 92,734 SNPs, of which 14% were newly discovered and possibly unique to the CFW population. These 92,734 SNPs provided extensive coverage of the genome (Fig. 2a) and allowed for fine-mapping (Figs. 3c,d and 4d,e, and Supplementary Figs. 17 and 23). The number of markers obtained using GBS can be titrated by varying the restriction enzymes used, the fragment sizes selected and the degree of sample multiplexing. GBS involves imputation to correct errors and to populate missing genotypes, requiring more expertise than analysis of SNP genotyping arrays. In comparison to conventional array-based SNP genotyping, GBS had a higher error rate, which is expected to modestly decrease power but should not produce false positive QTLs, as errors will not be correlated with the traits. We are currently improving genotype imputation methods for populations in which the founder haplotypes are known, such as the AIL and heterozygous stock populations\textsuperscript{12,24,46,63,64}. Because the monetary advantage of GBS over array-based genotyping will continue to increase as sequencing prices decrease, we anticipate that GBS and other sequencing-based approaches will supplant array-based methods in the coming years.

The majority of human GWAS findings implicate regulatory rather than coding differences\textsuperscript{4,11}. The identified haplotypes frequently contain several genes. It is now widely appreciated that, even when an association can be localized to a single gene, that gene may not be the cause of the association\textsuperscript{65}, meaning that proximity to the peak SNP is not sufficient to identify the causal gene. eQTLs can provide the crucial link between a region implicated by GWAS and the biological processes that underlie the association. Therefore, a major goal of our study was to integrate behavioral QTL and eQTL data. We used RNA-seq to examine gene expression in three brain regions that are known to be important for the behavioral traits that we studied. Although Azi2 was not an obvious candidate for the behavioral QTL for methamphetamine sensitivity, our data showing the co-mapping of an eQTL for Azi2 expression in the striatum provide an additional layer of evidence. Similarly, Zmynd11 has not previously been implicated in anxiety-like behavior, but the eQTL for Zmynd11 expression in the hippocampus suggests that it is the most promising of the four genes within the behavioral QTL. These examples demonstrate the power of integrating fine-mapping of behavioral QTLs and eQTLs, and extend on multiple previous mouse studies that have used similar approaches in conjunction with F\textsubscript{2} crosses\textsuperscript{66}, recombinant inbred lines\textsuperscript{67–69}, selected lines\textsuperscript{70}, heterogeneous stocks\textsuperscript{71}, outbred MF-1 mice\textsuperscript{50} and the HMDP\textsuperscript{51,72,73}.

RNA-seq offers a number of advantages relative to array-based gene expression measurements\textsuperscript{74–80}. In particular, we were able to map cis- and trans-eQTLs using a traditional mapping approach and simultaneously map cis-eQTLs by quantifying ASE. Because only a fraction of genes can be studied using ASE analysis, we did not anticipate complete overlap between genes identified using these two approaches. Using ASE, we identified 655 cis-eQTLs, of which 42% were also identified as cis-eQTLs using conventional mapping.

We found that physiological traits typically had slightly higher heritabilities than behavioral traits (Supplementary Table 1). We also found that the effect sizes of individual associations tended to be higher for physiological traits (Supplementary Table 2), consistent with findings from another recent study in rats\textsuperscript{81}. However, it was not always true that traits with the highest heritabilities also showed the largest effect sizes for individual associations. Because the effect size of individual QTL alleles is of paramount importance for assessing power at a given sample size and because this parameter is never known in advance, it is not possible to provide general guidelines about the sample size needed for future studies. On the basis of our results, we suggest that a sample size of 1,000 or more CFW mice should be used for most traits, although traits like testis weight and abnormal BMD would have yielded significant results with just a few hundred mice. Although our use of CFW mice was intended to increase mapping precision, there is a direct tradeoff between mapping precision and statistical power\textsuperscript{6}; therefore, sample sizes required
for studies using CFW mice will necessarily be larger than for those using F₂ crosses, recombinant inbred lines or other traditional mapping populations that offer less precision.

Our data do not directly address the reasons that the effect sizes we observed are so much larger than the effect sizes observed in most human GWAS. We can speculate that the unique population history of laboratory mice (involving domestication, selection and repeated population bottlenecks) has increased the frequency of alleles that may have been rare in ancestral wild mouse populations. It is also true that, unlike many traits studied in human GWAS, the traits we are examining are not disease traits (and thus may not influence fitness); the underlying alleles may therefore not have been influenced by natural selection, even among ancestral wild mouse populations from which laboratory populations were originally derived. Furthermore, laboratory mice are drawn from a much more uniform environment, potentially diminishing gene-by-environment interactions that may reduce effect sizes in human GWAS. Finally, because LD in the CFW population is more extensive than in humans, we are effectively testing fewer hypotheses and therefore applied a lower (permutation-derived) significance threshold.

We have shown that use of CFW mice in conjunction with GBS and RNA-seq provides a powerful and efficient means for identifying genetic associations and for nominating candidate genes within the associated regions. In comparison to other outbred mouse populations, CFW mice showed rapid decay of LD (Fig. 2b), were less expensive and primarily allowed examination of M. m. domesticus-derived alleles (Fig. 2c). In comparison to human GWAS, this approach provided dramatically reduced costs, the ability to examine phenotypes associated with different alleles, a broader spectrum of mouse populations in behavioral genetics research. More broadly, our results demonstrate methods and principles that apply to a variety of other model systems.

URLs. Phenotype, genotype and RNA-seq gene expression data, http://dx.doi.org/10.5061/dryad.2rs41; R code implementing data analyses, http://github.com/pcarbo/cfw; Quanto program, http://biostats.usc.edu/software.

METHODS

Methods and any associated references are available in the online version of the paper.

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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AUTHOR CONTRIBUTIONS

A.A.P. conceived the study. C.C.P. and A.A.P. supervised the project. S.G. and P.C. designed and implemented the statistical and bioinformatics analyses with contributions from C.C.P., J.K.P. and A.A.P. N.M.G. designed and executed the RNA-seq and GBS protocols with assistance from E.A. and J.D. C.C.P. performed the behavioral phenotyping with assistance from E.L. and Y.J.P. A.A.P. performed the muscle and bone phenotyping with input from D.A.B. C.L.A.-B. performed the TMD phenotyping. C.C.P., S.G., P.C. and A.A.P. wrote the manuscript, with input from all co-authors.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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ONLINE METHODS

Animal models. We phenotyped 1,200 male CFW mice (Mus musculus) that were obtained from the Charles River Laboratories facility in Portage, Michigan, USA (strain code, CRL.CFW(SW); facility code, P08). We performed a power analysis using the program Quanto (see URLs). This analysis indicated that 1,200 mice would provide 80% power to detect QTLs that accounted for ~3% of total trait variance with $P < 5 \times 10^{-7}$. Since our study was completed, the Portage colony has been relocated to Kingston, New York, USA (new code K92). It has been reported that the ancestors of the CFW mice were obtained from a large colony of Swiss mice in 1926 and maintained by Leslie Webster at the Rockefeller Institute. A single pair of highly inbred albino mice was later acquired by Carworth Farms and used to initiate an outbred mouse stock. Several mice from this colony were later acquired in 1974 by Charles River Laboratories and were subsequently maintained as an outbred population8–10.

Every 2 weeks, 48 male CFW mice were shipped from Charles River Laboratories in Portage to our laboratory in Chicago. We requested that Charles River Laboratories send only one mouse from each litter to avoid obtaining siblings, as the use of close relatives reduces power to map QTLs and complicates analysis. The average age of the mice upon arrival in our laboratories was 35 d (ranging from 34 to 46 d), and the average weight of the mice was 25.5 g (ranging from 13.4 to 38.7 g). Mice were housed four per cage and given ~15 d to adapt to their new environment (Supplementary Fig. 1).

Standard laboratory chow and water were available ad libitum, except during the behavioral procedures and before testing for fasting glucose levels. Mice were maintained on a standard 12 h light/12 h dark cycle (lights on at 6:30 a.m.). All phenotyping occurred during the light phase between 8:00 a.m. and 4:00 p.m., over the period of August 2011 to December 2012. All procedures were approved by the University of Chicago Institutional Animal Care and Use Committee (IACUC) in accordance with National Institutes of Health guidelines for the care and use of laboratory animals.

Phenotyping. The order of phenotyping was identical for each mouse and is shown schematically in Supplementary Figure 1. One day after arrival, mice were fasted for 4 h before measurement of blood glucose levels. Fourteen days later, we assessed the response to a new environment and to administration of 1.5 mg/kg methamphetamine in a 3-d paradigm48,49. Twelve days later, we tested mice for conditioned fear46. Nine days after that, we tested mice for PPI44. At 14 d after PPI testing, we collected one leg for measurement of muscle and another leg for conditioned fear46. Nine days after that, we tested mice for PPI44. Finally, after 15 d, we weighed and sacrificed the mice. Immediately after sacrifice, we weighed testes and collected one leg for measurement of muscle and another leg for conditioned fear46. Nine days after that, we tested mice for PPI44. Finally, after 15 d, we weighed and sacrificed the mice. Immediately after sacrifice, we weighed testes and collected one leg for measurement of muscle and another leg for conditioned fear46. Nine days after that, we tested mice for PPI44. Finally, after 15 d, we weighed and sacrificed the mice. Immediately after sacrifice, we weighed testes and collected one leg for measurement of muscle and another leg for conditioned fear46. Nine days after that, we tested mice for PPI44. Finally, after 15 d, we weighed and sacrificed the mice. Immediately after sacrifice, we weighed testes and collected one leg for measurement of muscle phenotypes and the other leg for measurement of bone phenotypes. We also measured tail length at this time (Supplementary Note).

RNA sequencing. We collected brain tissue from a subset of the mice as a source of mRNA from the hippocampus ($n = 79$), striatum ($n = 55$) and frontal cortex ($n = 54$). We used RNA-seq48,49 to quantify gene transcript abundance in these brain tissues. Library preparation was performed with the TruSeq RNA Sample kit (Illumina). Samples were multiplexed five per lane and sequenced on an Illumina HiSeq 2000 sequencer, using single-end 100-bp reads. We processed the RNA-seq short reads using the Tuxedo software suite50; (i) first, we aligned the short reads to the reference genome assembly (NCBI release 38, mm10) with Bowtie 2 (ref. 86); (ii) next, we used TopHat2 (ref. 79) to align the short reads to known splice junctions; and (iii) finally, we used Cufflinks61 to calculate, for each gene, a gene-level measure of expression based on the mapped reads. This measure is reported in RPKM. This measure does not depend on the length of the coding sequences or the sequencing depth of each sample (so mapping eQTLs will not be biased by these factors). We focused on this gene-level measurement for subsequent investigation, including eQTL mapping and assessment of ASE. See the Supplementary Note for further details.

Genotyping by sequencing. GBS is a reduced-representation genotyping method for obtaining genotyping information by sequencing only regions that are proximal to a restriction enzyme cut site15. Our protocol was adapted from previously described procedures62. GBS libraries were prepared by digesting genomic DNA with a restriction enzyme, PstI, and annealing oligonucleotide adaptors to the resulting overlaps. Samples were multiplexed 12 per lane and sequenced on an Illumina HiSeq 2000 sequencer, using single-end 100-bp reads. We obtained an average of 4.8 million reads per sample. By focusing the genotyping effort on PstI restriction sites, we obtained high coverage (~15×; Supplementary Fig. 24) at a subset of genomic loci, although the reads were very non-uniformly distributed. We aligned the 100-bp single-end reads to Mouse Reference Assembly 38 from the NCBI database (mm10) using BWA86. We used GATK94,95 to discover variants and to obtain genotype probabilities.

For the VQSR step, we calibrated variant discovery against (i) whole-genome sequencing data that we ascertained from a small set of CFW mice and (ii) SNPs and indels from the Wellcome Trust Sanger Institute Mouse Genomes Project104 and SNPs available in dbSNP release 137. We used IMPUTE2 (ref. 62) to improve low-confidence genotypes or genotypes that were not called in individual mice. Supplementary Table 4 and the Supplementary Note detail our efforts to estimate the error rate of GBS in this study. The Supplementary Note also contains a description of a small number of SNPs that were discarded because a large proportion of the genotypes were imputed with low certainty. Finally, the Supplementary Note details our identification of 110 DNA samples that seemed to be mislabeled and were therefore excluded from our study (Supplementary Figs. 25–28).

Treemix analysis. We estimated phylogenetic relationships between CFW mice and different laboratory strains sequenced as part of the Wellcome Trust Sanger Institute Mouse Genomes Project using Treemix28. We used the genotypes for the laboratory strains sequenced by the Wellcome Trust to obtain the locations of SNPs that were identified in the CFW mice using our GBS pipeline. We excluded the Mus spretus strain from the Wellcome Trust data, as this strain was included as an outgroup. Because the laboratory strains were all inbred, we assumed that allele frequency was 1 or 0. We represented each strain by only a single individual. We used a subset of 100 CFW mice to compute allele frequencies from the genotype likelihoods of GBS SNPs in our sample. Treemix was used to fit a maximum-likelihood tree to all the laboratory strains and CFW samples.

QTL mapping for behavioral and physiological traits. We performed a GWAS for the behavioral and physiological phenotypes using all SNPs with MAF >2% and good imputation quality (defined as 95% of the samples having a maximum-probability genotype greater than 0.5). Although our analyses did not suggest the presence of close relatives or population structure, we used the LMM implemented in the program GEMMA92,93. GEMMA is similar to a standard linear regression, in which a quantitative trait (y) is modeled as a linear combination of genotype (x) and covariates (z), except that it includes an additional ‘random’ or ‘polygenic’ effect capturing the covariance structure in the phenotype that is attributed to genome-wide genetic sharing.

$$y_i = \mu + \mathbf{z}_i \mathbf{\alpha} + \mathbf{z}_i \mathbf{\beta} + \mathbf{z}_i \mathbf{\epsilon} + u_i + \epsilon_i$$

The notation in this expression is defined as follows: $y_i$ is the ith phenotype sample; $\mathbf{z}_i$ is the sample of covariate k, in which k ranges from 1 to the number of covariates included in the regression (m); $\mathbf{\alpha}$ is the coefficient corresponding to covariate k; $\mathbf{z}_i$ is the genotype of sample i at SNP j; $\mathbf{\beta}$ is the coefficient corresponding to SNP j; $u_i$ is the polygenic effect for the ith sample; $\epsilon_i$ is the residual error; and $\mu$ is the intercept. The genotype, $\mathbf{z}_i$, is represented as the expected allele count, in which 0 represents homozygosity for the major allele and 2 represents homozygosity for the minor allele, and $\mathbf{\beta}$ is the additive effect of the expected allele count on the phenotype. The residuals $\epsilon_i$ are assumed to be i.i.d. normal with mean zero and covariance $\mathbf{\Sigma}$, whereas the polygenic effect $u_i$ is a random vector drawn from the multivariate normal distribution with mean zero and $n \times n$ covariance matrix $\mathbf{\Sigma}_K$, where n is the number of samples.

We estimated the relatedness matrix $K$ from the genotype data. We specified the covariance matrix using the realized relationship matrix $K = XX^T/p$, where $p$ is the number of SNPs and $X$ is the $n \times p$ genotype matrix with entries $x_{ij}$. This formulation was derived from a polygenic model of the phenotype in which all SNP effects helped explain variance in the phenotype and the contributions of individual SNPs were i.i.d. normal$^{94,96}$.

The inclusion of a genetic marker in both the fixed and random terms can deflate the test statistic for this marker, leading to a loss of power to detect
a QTL; this problem has been termed 'proximal contamination' (ref. 95). To avoid proximal contamination, we computed 19 different \( K \) matrices, each one excluding 1 of the 19 autosomes. To scan markers on a given chromosome, we used the version of \( K \) that did not include that chromosome. We have previously proposed this leave-one-chromosome-out (LOCO) approach as a simple solution for avoiding the problem of proximal contamination\(^\text{95}\).

We used a permutation-based approach to calculate the genome-wide significance threshold for \( P \) values calculated in GEMMA. We estimated the distribution of \( P \) values under the null hypothesis by mapping QTLs in 1,000 randomly permutated data sets, then taking the threshold to be the 100(1 – \( \alpha \))th percentile of this distribution, with \( \alpha = 0.1 \). Although this permutation test is technically only valid under the assumption that the samples are exchangeable\(^\text{97}\), we have previously suggested that 'naive' permutations are generally sufficient\(^\text{23}\). Furthermore, given our observation that population structure is subtle, we expect that this simulation provides a good approximation to the null (Supplementary Note).

Heritability estimates. Instead of computing a point estimate for SNP heritability (\( h^2 \)), which is the usual approach (for example, using the REML estimate\(^\text{88}\)), we evaluated likelihood over a regular grid of values for \( h^2 \), which allowed us to directly quantify uncertainty in \( h^2 \) under the reasonable assumption of a uniform prior for the proportion of variance explained\(^\text{96}\).

We estimated \( h^2 \) for our phenotypes\(^\text{21}\). Because the GBS SNPs did not completely tag all causal variants (and because we excluded the sex chromosomes), our estimates of \( h^2 \) underestimate a trait's true narrow-sense heritability. To estimate \( h^2 \), we assumed that all genetic markers made some small contribution to variation in the trait and that these contributions were normally distributed with the same variance\(^\text{96,98,99}\). Under this polygenic model, the covariance of the phenotype measurements was \( \text{Cov}(y_1, \ldots, y_n) = \sigma^2H \), where \( H = (I + \sigma_a^2 K) \), \( I \) is the \( n \times n \) identity matrix, \( K \) is the \( n \times n \) realized relatedness matrix, \( \sigma_a^2 \) is the variance of the additive genetic effects and \( \sigma_e^2 \) is the variance of the residuals. Under this formulation, \( \sigma_a^2 \) represents the relative contribution of the additive genetic variance, and we can use this parameter to provide an estimate for \( h^2 \)

\[
h^2 = \frac{\sigma_a^2}{\sigma_a^2 + \sigma_e^2 + 1}
\]

where \( y_n \) is the mean sample variance of all the available SNPs, or the mean of the diagonal entries of \( K \) assuming that the columns of \( X \) are centered so that each of the columns has a mean of zero. See the Supplementary Note for further details.

Expression QTL mapping. We used the RPKM measurements from RNA-seq and the GBS genotype data to map eQTLs. We performed an eQTL scan separately in each brain tissue (hippocampus, striatum and prefrontal cortex). First, we discarded genes with low levels of expression (RPKM <1) and genes that showed no variability in expression. For the remaining genes, we quantile normalized the expression data. To account for unknown confounders, we removed the linear effects of the first few principal components calculated from the \( k \times n \) gene expression matrix, with \( k \) genes and \( n \) samples (20 principal components for hippocampus, 10 principal components for striatum and 20 principal components for prefrontal cortex\(^\text{41}\)). After removing the linear effects of the principal components, we again quantile normalized the expression data. We then used an LMM as implemented in GEMMA to scan for cis-eQTLs, as described above for the behavioral and physiological phenotypes. To define cis-eQTLs, we only considered SNPs within 1 Mb of the gene's transcribed region (preliminary analyses indicated that 1 Mb captured most of the significant signals; Supplementary Figs. 19 and 20). We used a permutation-based approach to calculate significance thresholds for \( P \) values in cis-eQTL mapping. We used 1,000 permutations of the expression values to compute a separate significance threshold for each gene, using only SNPs that were included in the cis-eQTL scan. In addition to cis-eQTL scans, we also performed genome-wide trans-eQTL scans for all genes. The genome-wide scans were performed using the same LMM that was used for cis-eQTL analyses, except that all SNPs outside a 2-Mb region flanking the gene were included in the trans-eQTL analysis. The significance threshold for trans-eQTLs was computed using permutations of 1,000 randomly selected genes in each tissue; this approach is permissible because all expression traits were quantile normalized (Supplementary Note).

Allele-specific expression. We performed an analysis of ASE to identify genes that had ASE QTLs. This analysis was conducted independently from the mapping of cis-eQTLs described above. We identified variants that had at least ten samples with high-confidence heterozygote genotype calls. For genes that contained at least one such variant, we compared the relative expression of the two alleles across these heterozygote samples. To account for overdispersion, we used a beta binomial model to fit the counts of the two alleles for each sample. We then used a likelihood-ratio test to test for significant deviation of the observed data from the expectation of equal counts for both alleles (Supplementary Fig. 29 and Supplementary Note).

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