Identifying Loci Contributing to Natural Variation in Xenobiotic Resistance in Drosophila

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Abstract

Natural populations exhibit a great deal of interindividual genetic variation in the response to toxins, exemplified by the variable clinical efficacy of pharmaceutical drugs in humans, and the evolution of pesticide resistant insects. Such variation can result from several phenomena, including variable metabolic detoxification of the xenobiotic, and differential sensitivity of the molecular target of the toxin. Our goal is to genetically dissect variation in the response to xenobiotics, and characterize naturally-segregating polymorphisms that modulate toxicity. Here, we use the Drosophila Synthetic Population Resource (DSPR), a multi-parent advanced intercross panel of recombinant inbred lines, to identify QTL (Quantitative Trait Loci) underlying xenobiotic resistance, and employ caffeine as a model toxic compound. Phenotyping over 1,700 genotypes led to the identification of ten QTL, each explaining 4.5–14.4% of the broad-sense heritability for caffeine resistance. Four QTL harbor members of the cytochrome P450 family of detoxification enzymes, which represent strong a priori candidate genes. The case is especially strong for Cyp12d1, with multiple lines of evidence indicating the gene causally impacts caffeine resistance. Cyp12d1 is implicated by QTL mapped in both panels of DSPR RILs, is significantly upregulated in the presence of caffeine, and RNAi knockdown robustly decreases caffeine tolerance. Furthermore, copy number variation at Cyp12d1 is strongly associated with phenotype in the DSPR, with a trend in the same direction observed in the DGRP (Drosophila Genetic Reference Panel). No additional plausible causative polymorphisms were observed in a full genomewide association study in the DGRP, or in analyses restricted to QTL regions mapped in the DSPR. Just as in human populations, replicating modest-effect, naturally-segregating causative variants in an association study framework in flies will likely require very large sample sizes.
Author Summary

A striking feature of biology is that within populations there is substantial interindividual phenotypic variation for traits of biomedical significance. Some fraction of this variation is due to environmental factors, but for many traits segregating genetic differences contribute significantly to phenotypic variation. Elucidating those causative sequence changes that result in complex trait variation is of central importance to biology, and requires the coordinated use of multiple approaches. Here we employ a multi-level strategy to dissect genetic variation in caffeine resistance in Drosophila melanogaster, leveraging powerful genetic screening in a multiparental mapping panel, a genomewide association study, high-throughput RNA sequencing, and gene knockdowns using RNA interference. We identify several short genomic regions that collectively explain a substantial portion of the heritable variation for caffeine resistance, and find that several of these regions harbor members of known detoxification enzyme families. One such gene—Cyp12d1—shows increased expression on exposure to caffeine, and experimentally reducing gene expression leads to a reduction in caffeine resistance. We additionally show that variation in the number of copies of Cyp12d1 is positively associated with resistance. These compelling lines of evidence imply that structural variation at this gene causally contributes to xenobiotic resistance in Drosophila.

Introduction

Living organisms are subjected to a barrage of toxic compounds, or xenobiotics, in their environment and their diet. Animals are frequently exposed to toxins produced by potential prey and/or plant hosts as chemical defenses [1, 2], and are increasingly subject to pressures from human activity, such as pollution and the application of pesticides. Humans themselves are additionally exposed to an array of xenobiotics throughout their lives in the form of pharmaceuticals [3]. Given the evolutionary, agricultural, and medical relevance of the response and resistance to toxins, dissecting the genetic factors responsible for xenobiotic metabolism is essential.

Some of the best understood cases of xenobiotic response mechanisms come from insect populations or species that are able to withstand pesticides or toxic host chemical defense compounds. In some cases, certain individuals are simply insensitive to the xenobiotic due to a change in the structure, and therefore function, of the target of the toxin. For example, the Monarch butterfly (Danaus plexippus) is resistant to cardenolides, a class of secondary metabolites toxic to most animals, produced by their milkweed host plant. This resistance is due to at least one amino acid change in the Monarch Na⁺,K⁺-ATPase gene that prevents dietary cardenolides binding to the protein, making the protein insensitive to cardenolide inhibition [4, 5]. In many other cases, prior to reaching its target, the toxin is metabolized into less harmful substances by the action of a sophisticated three step detoxification system [6, 7]. In the first step, cytochrome P450 monooxygenases (P450s) act on the toxic compounds to decrease their toxicity. The products of these reactions subsequently become substrates for phase two enzymes, such as glutathione-S-transferases (GSTs) and UDP-glucuronosyltransferases (UGTs), which add large, charged side groups onto substrate molecules making them easier to excrete. Finally, membrane-bound ATP-binding cassette (ABC) transporters remove conjugated products from the cell. The Tobacco hornworm (Manduca sexta) is a facultative tobacco specialist, and detoxifies ingested nicotine by inducing P450 enzymes [8, 9]. In addition, as a result of a series of naturally-occurring gene duplication events and transposable element (TE) insertions,
overexpression of the P450 Cyp6g1 is primarily responsible for resistance to the insecticide DDT (dichlorodiphenyltrichloroethane) in Drosophila [10–12].

There are tens to hundreds of P450s, GSTs, UGTs, and ABC-transporters in eukaryotic genomes, and for most xenobiotics the precise series of enzymes responsible for their in vivo metabolism is unknown. We know there is substantial interindividual variation in the response to toxic compounds in animals such as Drosophila (e.g., [13–15]) and in the response to drugs in humans (e.g., [16–18]), but we rarely know which genes in the detoxification cascade segregate for functional variation, or understand the mechanisms by which genetic variation at these loci influences phenotype.

Here we characterize naturally-occurring alleles that contribute to resistance to caffeine in Drosophila. Caffeine is one of the most widely used psychoactive drugs in humans, acting as a stimulant, and has been employed as a convenient model xenobiotic in Drosophila research, where it is known to induce the expression of a number of P450 and GST genes in Drosophila S2 cells, larvae and adults [19–22]. However, simply because a gene responds to caffeine challenge by increasing its expression does not necessarily imply the gene segregates for alleles that give rise to variable drug response.

A powerful method to identify natural alleles that contribute to variable xenobiotic response is to employ a large, stable set of highly-recombinant genotypes derived from a multiparental mapping population [23–29]. The Drosophila Synthetic Population Resource (DSPR [30, 31]) consists of a set of >1,700 genotyped RILs (Recombinant Inbred Lines) derived from a pair of highly-recombinant synthetic populations each founded by eight strains. Our previous work has demonstrated the high statistical power and excellent mapping resolution of this community resource [30], and we have succeeded in resolving strong candidate genes contributing to nicotine resistance [15], and the response to chemotherapeutic drugs [14, 32].

In this study we identify several QTL contributing to variation in Drosophila adult female lifespan during continuous exposure to 1% caffeine, a measure of caffeine resistance. Of the ten mapped QTL, five harbor strong a priori candidate detoxification genes (e.g., P450s). One of these genes—Cyp12d1—is implicated by our largest-effect QTL, and segregates for copy number variation (CNV) that strongly correlates with phenotype in the DSPR. Statistically accounting for the effect of this variant in a genomewide scan eliminates the QTL harboring the locus.

In addition, Cyp12d1 shows expression induction in response to caffeine, and RNAi knockdown of the gene significantly reduces resistance. A follow-up association scan using the Drosophila Genetic Reference Panel (DGRP [33, 34]) reveals no significant associations after genomewide or QTL-specific multiple testing correction. Although not significant, the intermediate-frequency CNV at Cyp12d1 shows a trend in the expected direction in the DGRP; lines harboring the duplication have marginally higher resistance on average (p = 0.065). If the effect of the CNV is modest, and particularly if additional functional alleles contribute to the QTL implicating Cyp12d1, the functional loci may be difficult to individually identify by association in a small sample of natural chromosomes.

Results
Phenotypic Variation and Heritability

We measured the lifespan of female flies on 1% caffeine-supplemented media for 1,714 RILs from the DSPR, and 165 inbred lines from the DGRP, testing 16.5 females per genotype on average. There is substantial variation in phenotype in both mapping populations (S1 and S2 Figs). The average RIL mean phenotype in the pA DSPR panel is 37.3 hours (SD = 17.82), and in the pB panel is 35.5 hours (SD = 16.87). The average line mean phenotype in the DGRP is higher at 59.8 hours (SD = 20.80), although the range of the genotype means in the two
mapping populations is similar (DSPR range = 6.8–119.9 hours, DGRP range = 10.9–112.7 hours). The broad-sense heritability of our measure of caffeine resistance is just over 50% in all populations (DSPR pA and pB = 0.53, DGRP = 0.51). Because we employed line means for mapping, thus reducing environmental variance, the broad-sense heritability of the mean measure of caffeine resistance is around 94% (DSPR pA and pB = 0.95, DGRP = 0.94).

We note that the contribution of variation in lifespan under control, drug-free conditions likely plays a minimal role in our measure of caffeine resistance. Ivanov et al. [35] measured the lifespan of virgin females from the DGRP, and for the 155 lines common to both studies there is no statistically-significant correlation between lifespan and caffeine resistance line means (Pearson’s $r = 0.09, p = 0.27$).

**Mapping Caffeine Resistance QTL in the DSPR**

Using the same approach we have used previously [15, 31], we mapped QTL for caffeine resistance separately in both pA and pB DSPR mapping populations, identifying ten QTL contributing to resistance (Fig 1 and Tables 1 and S1). Three QTL are common to both populations (i.e., the 2-LOD support intervals overlap), one is unique to pA, and six are unique to pB. Given that the seven unique QTL generally explain smaller fractions of the heritability than the three common QTL (Table 1), it is possible we simply had insufficient power to detect them in the other panel of RILs. However, we cannot discount the possibility that alleles underlying these panel-specific QTL are private to a given panel. Under the assumption the QTL mapped are independent, and act additively, the total variance explained by all mapped QTL is simply the sum of the individual estimated heritability values. On this basis, in the pA population the four mapped QTL explain 31.9% of the broad-sense heritability for the trait, while in pB the nine mapped QTL explain 50.2% of the heritability.

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Fig 1. Genome scan for caffeine resistance QTL. Scans for population pA and pB are shown in blue and red curves, respectively, and the horizontal dotted lines represent population-specific genomewide 5% permutation thresholds (pA, LOD = 8.1; pB, LOD = 7.4). Genetic distances along the chromosomes are indicated along the x-axis. The centromeres are at positions 54 and 47 on chromosomes 2 and 3, respectively. The positions of the ten QTL we describe in the text are indicated on the plot for ease of reference. Intervals implicated by these QTL are highlighted as vertical bars, with pA-specific QTL in light blue, pB-specific QTL in pink, and QTL identified in both panels in yellow.

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Eight of the ten QTL map away from centromeres (Fig 1), and are mapped to relatively short intervals of 260–750 Kb that include modest numbers of protein-coding genes (mean = 64.8, range = 26–122; S2 Table). Q1 and Q8 are close to the chromosome 2 and chromosome 3 centromeres, respectively, and map to much broader genomic intervals, suggesting it may be more difficult to resolve the underlying causative loci. In terms of a priori candidate genes—genes one might anticipate being involved in the differential metabolism of xenobiotics—four QTL intervals contain one or more P450 genes (Q1, Q2, Q3, and Q9), one contains a carboxylesterase (Q8), and one contains two ABC transporters (Q1, although these two genes are only within the Q1 interval implicated in pA, not also the interval implicated in pB). None contain GSTs or UGTs.

If the locus responsible for each of the three shared QTL (Q1, Q2, and Q3) is the same in both panels, the causative gene should be present at a position implicated by both the pA and pB QTL LOD support intervals. For Q1 the area implicated by both mapped intervals is 500 kb, about half the width of the panel-specific intervals (Table 1), and harbors just 48 protein-coding genes. For Q2 the area implicated by both QTL is the same as that implicated by pA alone, containing 36 genes (Table 1), and for Q3 the area implicated by both QTL is just 60 Kb and contains ten genes. Each of these three small sets of genes suggested by the intersection of the pA and pB QTL support intervals contains a P450 gene (Q1 harbors Cyp310a1, Q2 harbors Cyp12d1, and Q3 harbors Cyp301a1). These are strong a priori candidates to contribute to variation in resistance to xenobiotics.

Table 1. Details of mapped caffeine resistance QTL.

| Name (Chr) | Panel | LOD score | Peak cM (2-LOD CI)a | Peak Mb (2-LOD CI)a | Number of genesb | Percent of H²c |
|------------|-------|------------|---------------------|---------------------|-----------------|-----------------|
| Q1 (2L)    | pA    | 9.9        | 53.3 (53.1–53.6)    | 18.74 (18.22–19.26) | 123             | 5.5             |
|            | pB    | 14.0       | 53.2 (52.8–53.3)    | 18.52 (17.65–18.72) | 81              | 7.7             |
| Q2 (2R)    | pA    | 27.2       | 63.6 (63.3–63.8)    | 7.07 (6.90–7.16)    | 36              | 14.4            |
|            | pB    | 10.3       | 63.6 (62.9–64.3)    | 7.06 (6.68–7.43)    | 122             | 5.7             |
| Q3 (2R)    | pA    | 11.3       | 66.1 (65.8–66.3)    | 8.36 (8.21–8.49)    | 65              | 6.3             |
|            | pB    | 10.6       | 66.3 (66.2–66.8)    | 8.49 (8.43–8.74)    | 48              | 5.9             |
| Q4 (2R)    | pB    | 8.1        | 68.8 (68.5–69.2)    | 9.65 (9.54–9.81)    | 26              | 4.5             |
| Q5 (2R)    | pB    | 9.8        | 75.2 (73.6–76.3)    | 11.49 (11.11–11.76) | 68              | 5.4             |
| Q6 (2R)    | pB    | 9.1        | 83.9 (82.9–84.1)    | 13.77 (13.46–13.83) | 69              | 5.1             |
| Q7 (2R)    | pB    | 9.1        | 88.1 (87.5–89.0)    | 15.23 (15.02–15.58) | 74              | 5.0             |
| Q8 (3L)    | pB    | 10.4       | 46.4 (46.0–47.1)    | 20.56 (19.89–24.36) | 398             | 5.8             |
| Q9 (3R)    | pA    | 10.2       | 54.4 (54.1–55.2)    | 9.87 (9.73–10.41)   | 79              | 5.7             |
| Q10 (3R)   | pB    | 9.1        | 62.5 (62.4–63.4)    | 14.01 (13.98–14.36) | 61              | 5.1             |

a 2-LOD CI indicates the 2-LOD support interval of the QTL. Physical positions are given based on release 5 of the Drosophila reference genome.

b The number of protein-coding genes in the 2-LOD support interval.

c The proportion of the phenotypic variance due to each QTL comes directly from the linear model used for mapping (page 77 of [76]). The percentage of the broad-sense heritability (H²) is simply this estimate divided by the broad-sense heritability of the mean measure of caffeine resistance.

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would also be expected to yield non-biallelic patterns of founder effects, we cannot be confident that QTL are truly multi-allelic.

Copy number variants (CNVs) are strong candidates to contribute to phenotypic variation [38–40]. One of the best described examples of a functional CNV is at the P450 gene Cyp6g1 in D. melanogaster, where a series of structural alleles confer varying levels of resistance to the pesticide DDT [10–12]. Previous work has observed variation in copy number at the Cyp12d1 locus—under our QTL Q2—in natural populations [41–43], with the D. melanogaster reference
strain possessing two copies with nearly identical gene sequences, Cyp12d1-d and Cyp12d1-p [44, 45]. Using PCR assays we found that DSPR founders A7, B1, B5, and B7 also possess these two gene copies, while the other 11 founders are single copy. Founder haplotypes at Q2 that harbor two copies of Cyp12d1 often have higher caffeine resistance than single-copy founders (Fig 2), although the founder with the highest mean resistance (B4) only harbors one copy of Cyp12d1. Considering individual RIL phenotypes, in both populations RILs with two copies of Cyp12d1 survive significantly longer on caffeine than RILs with only one copy of this gene (Welch’s t-test, p < 0.0001 in each population; S4 Fig). To explore whether the CNV can account for the mapped QTL Q2 we rescanned the genome after statistically controlling for the CNV status of each RIL. While most other QTL are still present, after adjusting for the effect of the Cyp12d1 CNV the signal at Q2 is entirely removed from both DSPR panels (S1 Table and S5 Fig). Thus, it is likely that the CNV at Cyp12d1, and/or a nearby variant or variants in strong LD with this event, plays a role in variable caffeine resistance in the DSPR.

Expression Candidate Genes

Previous work on the genetic basis of xenobiotic resistance has made extensive use of genome-wide expression studies to investigate both the innate differences between strains susceptible and resistant to a given compound, and any changes in gene expression that occur in an organism following exposure to the toxic substance. We took pA RILs with either very high or very low caffeine resistance, generated adult females heterozygous between lines of similar phenotype, and exposed groups of these flies both caffeine-supplemented and control media for short periods. Using heterogeneous animals minimizes inbreeding effects that may contribute to gene expression, but does make the assumption that heterozygous progeny are phenotypically similar to their parents. Following RNA extraction we pooled samples within phenotypic class (high or low resistance) and treatment, and carried out RNAseq, generating genomewide expression measures for four conditions, and leading to four contrasts of interest (Low-control versus High-control, Low-caffeine versus High-caffeine, Low-control versus Low-caffeine, High-control versus High-caffeine). A total of 242 genes were differentially-expressed, i.e., survived a genomewide per-contrast FDR threshold of 5% in at least one contrast (S3 Table). At least 10% of these genes are named members of recognized detoxification families, including P450s (16 genes), GSTs (5 genes), UGT genes (3 genes), and ABC transporters (5 genes).

The first two contrasts allow identification of genes having expression differences between genotypes with different phenotypes; 34 and 37 genes significantly change in expression between Low and High resistance genotypes on control food and caffeine-supplemented food, respectively. Eleven of these genes are shared between contrasts, and for all eleven the direction of the expression change is preserved over contrasts. The second two contrasts allow us to identify genes that change in expression on exposure to caffeine; in the Low (High) resistance genotype 178 (125) genes are differentially-expressed between control and caffeine food. One-hundred and one of these genes are shared between contrasts, and again the direction of the expression changes are consistent between genotypes; caffeine either induces expression in both genotypes or represses expression in both. Willoughby et al. [22] previously used a detoxification gene-specific microarray to identify 16 genes (11 P450 genes, 5 GSTs) upregulated in D. melanogaster third-instar larvae on exposure to caffeine. Ten of these 16 are induced in response to caffeine in both High and Low DSPR resistance genotypes, validating our RNAseq, and suggesting adults and larvae may respond to caffeine challenge via similar mechanisms.

Since our RNAseq study employed pA-derived genotypes, expression changes at genes within QTL intervals mapped in the pA population (Q1, Q2, Q3, and Q9) are of particular interest. Collectively, seven genes under these QTL show a change in expression that survives a
genomewide multiple-testing threshold (5% FDR) in at least one experimental contrast (Table 2). Four of these genes are annotated, but un-named genes with limited functional information in FlyBase [45]. One additional gene, \textit{Cpr49Ae}, under Q3 produces a constituent of chitin, and is expressed at higher levels in genotypes that are more resistant to caffeine (Table 2).

As a class, cuticle genes were found to be enriched among the set of genes overexpressed in a \textit{D. melanogaster} strain resistant to \textalpha-amanitin, a toxin produced by a number of species of mushroom [46]. Higher expression of \textit{Cpr48Ae} in caffeine resistant genotypes could plausibly be responsible for providing additional surface protection against xenobiotic exposure. Nonetheless, it is worth noting that \textit{Cpr48Ae} is implicated only by Q3 mapped in pA, and is not present within the interval mapped in pB.

The strongest expression candidates from this RNAseq screen are P450 genes, \textit{Cyp12d1} under Q2, and \textit{Cyp6d5}, one of the two P450s under Q9 (Table 2). As mentioned previously \textit{Cyp12d1} exists in the \textit{D. melanogaster} reference strain as a tandemly-duplicated pair of genes (\textit{Cyp12d1-d} and \textit{Cyp12d1-p}), differing by just 3/1,563 coding sequence nucleotides. Recognizing the difficulty distinguishing reads from each gene, we eliminated the annotation for \textit{Cyp12d1-d} during RNAseq analysis, allowing reads from both gene copies to pile up on \textit{Cyp12d1-p}, and report these values in Table 2. Results are essentially identical regardless of which gene copy is masked for analysis. \textit{Cyp12d1} shows strong, significant expression induction on caffeine exposure in both High and Low resistance genotypes, as has been shown previously [22, 47]. In addition, \textit{Cyp12d1} gene expression is significantly higher for High resistance genotypes under both control and caffeine treatments. We note that at least some of the difference between Low and High genotypes is plausibly explained by the known CNV at this gene; Of the 18 RILs comprising each genotype pool, 17/18 Low genotypes are single copy for \textit{Cyp12d1}, while 14/18 High genotypes have two copies. Array-based female head-specific expression data derived from the progeny of crosses between DSPR RILs [36] confirms a
significant effect of the Cyp12d1 CNV on expression of this gene (S6 Fig). Cyp12d1 is a clear candidate for the causal locus responsible for Q2.

Cyp6d5 (Q9) shows highly-significant expression induction following caffeine exposure, but there is no significant difference in expression between Low and High genotypes. The second P450 under Q9 (Cyp313a1) shows a reduction in expression in response to caffeine in the Low resistance genotypes at a nominal level ($p = 0.005$), which is not the pattern expected under the assumption that higher expression leads to greater resistance to the drug. Thus, Cyp6d5 is the stronger candidate to causally underlie Q9. Neither of the P450s under Q1 or Q3 mapped in pA (Cyp310a1 and Cyp301a1, respectively) show significant expression variation across samples even at a nominal, $p < 0.05$ level (S1 Dataset). This result doesn’t rule out the possibility the genes contribute to variable resistance to caffeine, but suggests any such role may not involve regulatory variation.

Functional Validation of P450 Gene Effects on Caffeine Resistance

To confirm effects of a subset of the P450 genes uncovered by our mapping and expression studies (Cyp12d1-d and Cyp12d1-p, Cyp301a1, and Cyp6d5 under Q2, Q3, and Q9, respectively) we carried out knock-down experiments using the Gal4-UAS RNAi system. When knocked down ubiquitously (RNAi in all cells at all timepoints), all but one of the Gal4-UAS showed a significant reduction in caffeine resistance compared to controls (Fig 3A and 3B), with the remaining genotype showing a minor reduction in resistance (Fig 3B; transformant ID 21235, Cyp12d1-p, Welch’s $t$-test, $p = 0.066$). In two cases the RNAi appears to interfere with some essential developmental process, since we struggled to generate individuals of the appropriate genotypes for assay (Fig 3A; genes Cyp12d1-d, transformant ID 109248, and Cyp301a1). Since Cyp301a1 appears to be involved in cuticle development, and reduced expression of the gene results in cuticle malformation [48], this may have played a role in our failure to generate large numbers of progeny. We also cannot rule out any minor developmental abnormalities in the other RNAi genotypes that may have led to reduced adult fitness, and a nonspecific reduction in caffeine resistance. To control for any such pleiotropic effects of the test genes during development, we subsequently used RU486-inducible Gal4 to drive RNAi in all cells in young adults (Fig 3C and 3D). Over two trials (differing only in the amount of time the test animals were fed RU486 prior to caffeine+RU486 exposure), the only RNAi knockdown to show a consistent, highly-significant reduction in caffeine resistance was for Cyp12d1-d (Fig 3C and 3D). Since the effect of RU486 on expression knockdown is dose-dependent [49], it is possible higher concentrations of RU486 could have replicated our findings using the ubiquitous Gal4 driver (Fig 3A), and Cyp301a1 and Cyp6d5 remain excellent candidates to underlie QTL Q3 and Q9.

Critical to interpreting RNAi results with Cyp12d1-d and Cyp12d1-p is the fact that these genes appear to be the result of recent duplication [42], and have nearly identical sequences [44, 45]; The genes differ by 3/1,563 coding sequence nucleotides, leading to 3/521 amino acid differences. The hairpin sequences of the UAS-RNAi constructs targeting one member of the gene pair exactly, exhibit zero (transformant ID 50507), one (transformant IDs 21235, 109248, 109256), or two (transformant ID 49269) nucleotide differences from the other member of the gene pair (S2 Dataset). So it is highly likely that in each Cyp12d1 knockdown experiment we are reducing the expression of both genes, albeit perhaps to different degrees.

Association Mapping Caffeine Resistance Loci in the DGRP

The DSPR has excellent power to identify QTL contributing to trait variation [30], but given the haplotype structure of the DSPR RILs the approach suffers from a lack of resolution in
comparison with population-based association mapping. It is possible to carry out local, QTL-centric association scans in the DSPR by imputing SNP genotypes using the estimated mosaic founder haplotype structure of each RIL, but again LD (linkage disequilibrium) is such that large numbers of SNPs tag the exact same haplotype combinations, precluding localization of causative variants [15, 32].

To attempt to identify candidate causative polymorphisms we used the DGRP [33, 34], a sample of fully-resequenced inbred lines derived from a single collection, to carry out association mapping. Following a GWAS (Genomewide Association Study), testing 1.9 million SNPs, we identified several loci that showed significant association with resistance.
polymorphic events for association with caffeine resistance, the site with the highest association statistic \( (p = 1.7 \times 10^{-6}) \) was far from a genomewide Bonferroni significance threshold of \( p \sim 10^{-8} \) (S4 Table). The lack of a strong association is unsurprising given the likelihood that causative variants explain relatively small fractions of the variance (Table 1), and the low power in a GWAS with \( < 200 \) lines to find sites with such effect sizes [50, 51]. To avoid a harsh, genomewide significance threshold we instead restricted our focus to those variants present within QTL intervals mapped in the DSPR. This procedure is more appropriate given we are attempting to validate mapped QTL, rather than attempting de novo discovery in the DGRP. Nevertheless, no site reached a QTL-specific Bonferroni threshold, and only one site beneath a mapped QTL reached \( p < 10^{-5} \) (one of the 24,782 tested variants within the large Q8 interval). Tens to hundreds of variants within QTL intervals have \( p \)-values that reach a nominal 5% threshold, but in each case the fraction of such variants is essentially what would be expected by chance.

We additionally examined associations within 5-kb of the start and end of each of the six annotated P450 genes we have discussed previously (Cyp12d1-d and Cyp12d1-p, Cyp301a1, Cyp310a1, Cyp313a1, Cyp6d5), but again no site in these regions survived a per-gene Bonferroni correction for multiple tests.

In the DSPR we found evidence supporting the role of copy number variation at Cyp12d1 in conferring resistance to caffeine. Just as for the DSPR we genotyped lines from DGRP using PCR based assays [12, 42], and confirmed our data with bioinformatic calls based sequencing read-depth [41]. Using only those 151 DGRP lines for which we had caffeine phenotypes, where we were confident the lines were homozygous at the Cyp12d1 locus, and where PCR-based calls and bioinformatics calls were identical, the mean resistance of DGRP lines carrying one Cyp12d1 copy was 57.6 hours (SD = 20.88, \( N = 119 \)), and the mean resistance of lines with two copies was 65.4 hours (SD = 20.67, \( N = 32 \)). While there was a trend in the anticipated direction, the effect of the Cyp12d1 duplication on caffeine resistance is not significant at the 5% level in the DGRP (Welch’s \( t \)-test, \( p = 0.065 \); S4 Fig). Given power constraints in the DGRP [50] we may be unable to detect the CNV in this population, particularly if we have overestimated the heritability explained by Q2 (Beavis effects [52, 53]), or if the effect of Q2 in the DSPR is not entirely due to the CNV, and other variants at Cyp12d1 are additionally causative. However, we cannot rule out a scenario in which the Cyp12d1 CNV has no effect on caffeine resistance in the DGRP.

**Discussion**

In this study we employed a large mapping population, the DSPR, to identify loci segregating for naturally-occurring alleles with variable effects on xenobiotic toxicity in D. melanogaster. Heritability of resistance to caffeine is around 50% in all mapping panels queried in this study, and we succeeded in mapping a number of QTL that collectively explain between a third and half of the broad-sense heritability. Aside from those QTL mapping to centromeric regions, we map QTL to genomic intervals containing fairly modest numbers of protein-coding genes (26–122), and in those cases where a QTL is mapped to the same location in the pA and pB RIL panels, we can further refine the list of the most likely candidates to those genes implicated by both intervals (i.e., 48, 36 and 10 genes for Q1, Q2, and Q3, respectively). Dissection with this level of resolution is difficult with standard, 2-parental QTL mapping using a single generation of meiotic recombination (i.e., using an F2 or backcross mapping design), and can only reasonably be achieved for linkage mapping designs employing populations subjected to multiple rounds of meiotic recombination (e.g., [54]).

Short QTL mapping windows facilitate the identification of true causative gene(s), and in this study our ability to suggest plausible functional loci was additionally enhanced by the...
nature of our target phenotype. Genes involved in metabolic detoxification pathways represent strong a priori candidates to harbor segregating variation contributing to caffeine resistance. Indeed, four caffeine resistance QTL harbor P450 genes (Q1—Cyp310a1, Q2—Cyp12d1, Q3—Cyp301a1, Q9—Cyp6d5 and Cyp313a1), many of which appear to be involved in detoxification [55], and these present excellent targets for future research to identify the precise sequence variants impacting resistance to caffeine. Our mapping results, in combination with RNAseq and RNAi data, imply that a significant fraction of natural variation in resistance to caffeine is due to metabolic detoxification driven by changes in P450 genes (see also [47]).

The strongest candidates from our study are Cyp12d1 and Cyp6d5. These loci are present within QTL intervals (Cyp12d1 is additionally present within QTL mapped in both pA and pB mapping panels), show expression induction in the presence of caffeine (Table 2), and a reduction in resistance on RNAi gene expression knockdown (Fig 3). In addition, a number of previous studies have shown effects of these loci consistent with a role in xenobiotic metabolism: Both genes are expressed in the larval midgut [56], a likely site of xenobiotic metabolism, both have been shown to be induced by caffeine and phenobarbital [22], and silencing both genes alters downstream metabolism of caffeine [47].

We also show that the copy number variation segregating at Cyp12d1 [12, 41, 42] associates with our caffeine resistance phenotype in the DSPR RILs. Gene duplications are known to be important contributors to complex variation (e.g., [57]), and evidence from other studies demonstrates that structural variation at P450 genes can be associated with toxin resistance. For instance, an allelic series of CNV events and TE insertions at Cyp6g1 impacts DDT resistance in D. melanogaster [12], although in this same study no association was seen between Cyp12d1 copy number and DDT resistance. In the DGRP, the Cyp12d1 CNV shows an effect on caffeine resistance in the same direction as in the DSPR, though this trend is not significant at the 5% level (p = 0.065). Heterogeneity in the genetic architecture of caffeine resistance across mapping panels could explain the disconnect among studies, with copy number at Cyp12d1 positively, and causally, associating with resistance in the DSPR but not in the DGRP. Alternatively, the CNV could be in strong LD with the true causative variant in the DSPR, whereas these events may not be in LD in the DGRP. Notably however we did not identify any other associations around Cyp12d1 in the DGRP that could represent this indirect association. Given the multiple lines of evidence supporting an effect of the Cyp12d1 locus, it remains highly likely that this gene is involved in caffeine resistance. Future targeted fine-mapping and/or functional genomics work will be required to localize the causative variant(s) in the Cyp12d1 region that lead to natural variation in caffeine resistance.

Other P450s implicated (Cyp301a1, Cyp310a1, and Cyp313a1) in our mapping did not show changes in expression, either between susceptible or resistant genotypes, or between control and caffeine-supplemented media (S1 Dataset). This may imply that the effect of these genes on variable caffeine resistance is not regulatory in origin, and thus cannot be captured by RNAseq. Alternatively, overall expression levels may be too low—which is likely true for Cyp301a1 (FPKM = 1.88–2.24) and Cyp310a1 (FPKM = 0.52–0.67)—and any expression differences may be subtle, precluding the identification of significant expression changes without high levels of biological replication, or by carrying out RNAseq on the likely tissues in which the detoxification process takes place (e.g., [56]). However, without further validation it is not possible to exclude the possibility that these P450 genes are not the causative loci, and instead are merely closely linked to the true caffeine resistance genes.

For the six QTL not containing P450s, all of which have modest effects and were mapped only in the pB panel of DSPR RILs, it is difficult to suggest likely causative genes solely from the lists of implicated loci. QTL Q8 does harbor a carboxylesterase, Est-Q, that is known to show expression in the larval midgut, a site of xenobiotic metabolism [56]. However, while this...
is a plausible candidate gene, because Q8 implicates a total of 398 genes, other evidence for a
role in caffeine resistance would be desirable prior to any attempt to functionally validate this
gene. Similarly, the Smc5 gene, shown to be required for resistance to caffeine exposure during
development [58] is also present within Q8, and remains a plausible candidate. The genes cnc, Hr96, and Keap1 are known to regulate the transcriptional response to xenobiotic challenge in
Drosophila [20, 59], and ERR is associated with the regulation of a number of P450 genes [60].
However, none of these genes are located under the caffeine resistance QTL we map here, sug-
gestig that natural variation in the regulation of the overall transcriptional response to xenobi-
otics is not strongly involved in caffeine resistance, at least in our assay. Gustatory receptors
have been shown to be involved in the avoidance of noxious substances, and Gr66a and Gr93a
are required to avoid caffeine ingestion [61, 62]. Neither of these genes are implicated by our
mapped QTL, although other members of the family are present in mapped intervals (Gr47b is
present within the Q2 interval mapped in pB, and Gr77a is present under Q8).

The absence of clear candidate genes underlying mapped QTL is emblematic of the chal-
lenge facing most linkage mapping studies. Here, we used a combination of RNAseq and RNAi
to validate the effects of certain P450 genes on caffeine resistance, and to refine our list of
potential genes for those QTL without obvious a priori candidates. Moving from causative
genomic regions to the true causative sites is an ongoing challenge for mapping studies. Com-
bining multiple sources of data to provide several lines of evidence supporting candidate loci
may prove generally valuable to resolve functional loci within mapped genomic windows, and
data on the positions of QTL from this study represent the starting point for such exploration
(Tables 1 and S2).

A goal of our research is not only to identify the causative genes, but also to identify the
causative sequence variants, and we employed the DGRP association mapping panel to this
end [33, 34]. Using a similar phenotyping strategy to that employed in the DSPR, and a
straightforward GWAS analysis [33], we found no genomewide significant associations after
controlling for multiple tests. This result is likely explained by the low power of the DGRP on a
genomewide scale: With 158 lines and a genomewide significance threshold of \( p < 10^{-8} \), the
power to find an intermediate-frequency causative polymorphism (minor allele frequency,
MAF = 0.4) that explains 10% of the broad-sense heritability is just 3% (see [50] for description
of power calculation).

Due to the lack of power for de novo identification of associations in a small GWAS panel,
we instead attempted to validate associations within intervals implicated by DSPR QTL, and
find associations directly at the six P450 genes suggested by our study. However, even with a
lower significance threshold (approaching \( p < 2 \times 10^{-4} \) for the gene-centric association scans)
and a concomitant increase in power (62% for a site at a frequency of 0.4 that explains 10% of
the variation among lines), we found no QTL-specific variants, and no variants at the set of
P450 genes that survive multiple test correction. Thus, power deficits in the DGRP may not
entirely explain the lack of overlap between findings in the DSPR and DGRP.

A lack of overlap between the DSPR and DGRP might be expected under a situation in
which we have overestimated the heritability explained by QTL mapped in the DSPR. If true
QTL effects are much smaller than we estimate (Table 1), power to validate such effects is in
turn reduced in the DGRP. Beavis [52] showed that the effects of mapped QTL tended to be
overestimated, with the greatest overestimation associated with experiments employing small
numbers of samples. Given that the sample size for each DSPR panel was >800 genotypes, and
our power to map QTL contributing 5% to variation is over 80% [30], we anticipate that Beavis
effects will not be large (see [52] and Figure 15.8 in [63]). Nonetheless, work on the Beavis
effect [52, 53] has focused on two-way crosses involving a single-generation of meiotic
recombination, and there has been no exploration of the expected magnitude of the Beavis effect in highly-recombinant, multiparental mapping populations.

One possible reason for the failure to find associations in the DGRP within mapped QTL intervals could be that causative variants present in the DSPR are absent in the DGRP. Variants at intermediate frequency will typically be shared by both panels, and given sufficient power, should replicate. Rare variants will be shared less often, and will only replicate some of the time. Those SNPs present in two or more DSPR founders are found in the DGRP >91% of the time, while those SNPs private to a single DSPR founder are seen in the DGRP ~70% of the time (see DSPR founder SNP data from FlyRILs.org and DGRP variant data from dgrp2.gnets.ncsu.edu). In addition, founder means at QTL do not show a pattern consistent with a single founder having a private allele (Figs 2 and S2). Thus, there is limited evidence that the variants responsible for our QTL are unique to the DSPR.

A related possibility is that causative variants have very different frequencies in the DSPR and DGRP. Population-based association mapping approaches struggle to identify low frequency causative variants [50, 64, 65]. Whereas provided the minor allele is captured among one of the founders, a multiparental mapping population has good power to find a low frequency causative variant [30]. Thus, DSPR QTL generated by rare causative loci may not replicate in the DGRP. One piece of evidence against this possibility is the observation that Q1, Q2, and Q3 were all mapped in both DSPR panels. This implies these QTL are not due to rare causative variants present solely in one panel of the DSPR, and suggests the causative variants underlying these QTL are likely present at intermediate-frequency in the DGRP. For instance, the frequency of the duplication allele at the plausible causative CNV at the Cyp12d1 gene within Q2 is 12.5% among DSPR pA founders, 37.5% among pB founders, and 21.2% among the DGRP lines. Since we do not implicate a causative polymorphism for all other QTL we map in the DSPR it is not possible to determine whether the (unknown) polymorphisms responsible have similar frequencies in the DGRP.

A final possibility for our failure to replicate our QTL mapping results in the DGRP is suggested by the observation from a number of different multiparental mapping studies that QTL frequently segregate for multiple alleles [26, 36, 37]. Similarly, the strain effects for some of the QTL we map here do not show a clear pattern indicative of biallelic QTL simply segregating for a single "high" and "low" allele (Figs 2 and S3). A multiparental panel tests the effect of a local haplotype on phenotype, integrating over any causative variants in a short genomic window. So if QTL mapped in the DSPR are routinely generated by the combined action of multiple variants, each with very small effects, the power to identify any of these individually in the DGRP will be extremely limited. Detection of very subtle effect, intermediate-frequency causative variants, contributing <5% to the natural variation will require massive sample sizes as is common in the human genetics community (e.g., [66]). If instead the multiple causative variants present at genes contributing to polygenic trait variation are all rare, as is the case for genes leading to single-gene, Mendelian disorders, typical GWAS analytical approaches are very poorly powered to detect such genes [65].

In summary, we have used the DSPR to map QTL that collectively explain a large fraction of the natural variation for caffeine resistance in flies. Four of these QTL map to genomic regions containing members of the large family of cytochrome P450 detoxification enzymes. These loci present excellent candidates to harbor variant(s) contributing to phenotypic variation, and for three of these loci—Cyp6d5, Cyp12d1, and Cyp301a1—we validate their effects on caffeine resistance via RNAseq and/or RNAi, suggesting an important role for variation in metabolic detoxification in the control of caffeine resistance. In addition, our data suggests that the number of copies of Cyp12d1 is a strong determinant of caffeine resistance. We were unable to resolve any additional candidate causative sequence changes by association in a small panel of
natural chromosomes, likely due in part to concerns related to power, and potentially as a result of our QTL being generated by the combined action of multiple, small-effect causative alleles. Nonetheless, within each interval implicated by a QTL, tens to hundreds of variants show associations at nominally-significant levels (e.g., $p < 0.05$). If some of these modest-effect associations could be validated in an independent, large association mapping panel this would provide strong evidence for their causative role in affecting phenotypic variation. Ultimately, for strong candidate regions, in the future it will be possible to employ CRISPR-Cas9 genome editing to directly compare alleles at candidate loci in an otherwise identical genetic background to localize regions, and ultimately nucleotide variants directly contributing to phenotypic variance.

**Materials and Methods**

**Mapping Populations**

To map QTL for caffeine resistance we used RILs from the DSPR (FlyRILs.org). Full details of the mapping panel, its development, the analytical methods employed to map QTL, and simulations supporting the power and resolution of the approach have been described previously [30, 31]. Briefly, the DSPR consists of two panels of >800 RILs (pA and pB). Each set is descended from an advanced generation intercross among eight highly-inbred founder lines, seven of which are specific to a single panel (A1-A7 or B1-B7), with one is common to both panels (AB8). Each intercross population was maintained as a pair of independent replicate subpopulations (pA1, pA2, pB1, pB2) at large population size for 50 generations, after which RILs were generated via 25 generations of full sibling mating. The 15 founder lines have been resequenced, and the RILs genotyped via Restriction site Associated DNA (RAD) tags. These data, along with a hidden Markov model (HMM), allowed the underlying mosaic haplotype structure of each RIL to be determined at a fine scale.

To carry out the GWAS we used strains from the DGRP, a set of inbred lines descended from gravid females caught at a single collection location (Raleigh, NC, USA), and inbred via 20 generations of full sibling mating. DGRP lines were purchased from the Bloomington *Drosophila* Stock Center (flystocks.bio.indiana.edu/). The lines have been fully resequenced, and molecular variation in the panel has been described [33, 34], along with the computational pipeline used to make variant calls [33, 67].

**Caffeine Resistance Assay**

Flies from each genotype were allowed to lay eggs for up to 48 hours on standard cornmeal-yeast-molasses food in regular narrow, polystyrene fly vials, and adults were cleared to maintain a roughly equal density of eggs across experimental vials. Eight days after the start of egg laying, any newly emerged adults were cleared from the experimental vials, and two days later 20–25, likely mated female flies were harvested from each vial under CO2 anesthesia. These flies were kept as groups on standard media and allowed to recover for 24 hours prior to caffeine challenge.

Our measure of caffeine resistance for a genotype is taken as mean lifespan on media supplemented with 1% caffeine (C0750, Sigma), the same concentration as employed in a previous study of caffeine resistance variation in *D. melanogaster* [13]. To increase throughput, and provide for relatively automated data collection, we made use of the *Drosophila* Activity Monitoring System (DAM2, TriKinetics, Inc.). Twenty-four hours before the start of the assay we made cornmeal-yeast-dextrose media, adding caffeine at ~50°C to minimize any heat-induced degradation. We replaced the molasses from our typical culture medium with dextrose to achieve better batch-to-batch reproducibility since small volumes of molasses proved difficult to
dispense accurately. The molten media was poured into 100mm diameter petri dishes and allowed to set for 2–3 hours. We pushed the ends of large bundles of polycarbonate activity monitor tubes (5mm diameter × 65mm length) into the media, filling each tube to a height of ~10mm, cleaned the tubes to remove excess media, and sealed the food plug in each tube with molten paraffin wax. On the day of the assay we aspirated individual, 1–3 day old experimental females into monitor tubes without anesthesia, capped the tubes with small foam plugs cut from Droso-Plugs (Genesee Scientific Corporation), and inserted tubes into monitors. All experimental individuals were reared and assayed at 25°C and 50% relative humidity on a 12 hour light/12 hour dark cycle.

The time when experimental flies were introduced to caffeine food in monitor tubes was recorded using a custom Excel macro (Microsoft Corporation), facilitated by arbitrary bar-codes on experimental vials and monitors. Activity data was recorded every minute for six days for each experimental block using the DAMSystem3 data collection software (TriKinetics, Inc.), and any permanent cessation of activity was interpreted to be a result of fly death. The data were filtered to remove animals with very low activity levels from the start of the assay, presumably due to damage during aspiration into monitor tubes, and the time until death on caffeine food for each fly was extracted using a series of custom scripts written in R (www.R-project.org).

The above assay was carried out on 1,714 DSPR RILs (853 pA and 861 pB lines) and 165 DGRP lines across 37 experimental blocks. For 1,623 DSPR and 159 DGRP lines we generated a single experimental vial of flies, and tested 16 females from that vial in a single experimental assay block. For a small minority of lines we independently tested 16 flies in each of two blocks, and the correlation between means calculated for each block is high (Pearson’s $r = 0.89$, $p < 10^{-15}$), giving us confidence in our phenotypic measurements. Over all mapping panel genotypes we scored an average of 16.5 females per genotype (SD = 3.62, range = 9–32). Raw phenotype data is presented in S5 Table.

Heritability Estimates

Broad-sense heritability of our measure of caffeine resistance (the time until death of an adult female fly on caffeine-supplemented media) was estimated by calculating the genetic and phenotypic variance components from a linear mixed model using the lme and VarCorr functions in the nlme package in R [15, 68]. The estimated genetic variance component divided by the total variance of line means yielded the heritability of the line means that were used for mapping. Heritabilities were separately calculated for both panels of DSPR RILs and the DGRP.

QTL Mapping in the DSPR

Identification of QTL in the DSPR has been described previously [30, 31]. In essence, for each region in each RIL an HMM is used to assign a probability that the genotype is one of 36 possible homo- or heterozygous founder genotype combinations. Subsequently we assume that the very small number of heterozygous states we find are intermediate between the respective pair of homozygous states, generating eight additive probabilities per position. We then regress the mean line phenotype on these eight probabilities, analyzing the pA and pB panels separately. We did not include a subpopulation covariate since there was no significant difference between the two pA or pB subpopulations in mean RIL caffeine resistance (pA, Welch’s $t = 1.8$, $p = 0.06$; pB, Welch’s $t = 1.9$, $p = 0.06$). Following Churchill and Doerge [69] we use 1,000 permutations of the data to find appropriate genomewide significance thresholds for QTL identification. We used 2-LOD support intervals to estimate the true positions of causative loci, which simulations
suggest correspond to 93–94% confidence intervals for our experimental design, sample size, and mapped QTL effect sizes [30].

We note that there is a clear visual, positive correlation between line mean phenotypes and within-line variance (S1 Fig). This pattern is eliminated following a square root transformation of the line means. However, such a transformation does not change the genomewide likelihood profiles, or alter the QTL we map, and we only report analysis based on the raw, untransformed dataset here.

To account for any effect of the CNV at Cyp12d1 we performed a similar genomewide scan for QTL as described above, but additionally included a covariate describing the CNV status of each RIL. For this analysis we only used those RILs for which we could be confident of the CNV status, i.e., where the founder allele at the gene was estimated with at least 95% certainty. This resulted in a reduction in the sample size for mapping from 853 to 803 RILs in pA, and from 861 to 812 RILs in pB.

**Genomewide Association Mapping in the DGRP**

Phenotype means for 158 DGRP strains assayed were provided to the Mackay lab’s web-based analytical engine (dgrp2.gnets.ncsu.edu) to carry out a GWAS using nearly 2 million common (minor allele frequency > 0.05) SNP and non-SNP variants identified in the panel. Prior to the association tests the line means are adjusted to account for any effect of the Wolbachia bacteria that infects around half of the lines, and to account for any effects of five major chromosomal inversions: In(2L)t, In(2R)NS, In(3R)P, In(3R)K, and In(3R)Mo. After which association tests are carried out for each variant using a linear mixed model accounting for variation in relatedness across the lines. See Huang et al. [33] for full details.

**RNAseq**

We selected two sets of pA RILs that had either very low (in the bottom 2.3% of lines) or very high (in the top 4.2% of lines) caffeine resistance, and crossed RILs in pairs within each phenotypic class. From each cross, we collected 2–3 day old adult female trans-heterozygous progeny under CO₂ anesthesia, allowed the flies to recover for 24 hours, then exposed flies from each genotype either to control food or to 1% caffeine food for 4 hours. No flies died during this short exposure. Flies from each genotype and treatment were then snap-frozen in liquid nitrogen, and total RNA was isolated from each sample using TRIzol reagent (15596–018, Life Technologies). Equal amounts of RNA from each sample were then pooled within each phenotype/treatment combination: susceptible genotypes/control food, susceptible genotypes/caffeine food, resistant genotypes/control food, and resistant genotypes/caffeine food. Ultimately, each of these four pooled samples contained RNA from ~100 experimental females, and haplotypes from 18 pA RILs. The four pooled samples were cleaned through RNeasy Mini columns (74104, Qiagen) following the manufacturer’s protocols, used to generate Illumina TruSeq RNAseq libraries, and sequenced over four lanes of an Illumina HiSeq 2500 instrument to generate single-end 50bp reads (Genome Sequencing Facility, KU Medical Center). Raw sequencing reads were deposited in the Sequence Read Archive (SRA) under project accession number SRP051835.

We trimmed raw sequencing reads with sickle (version 1.200, github.com/najoshi/sickle), and used TopHat (version 2.0.9, tophat.cbcb.umd.edu; [70, 71]) to assemble reads from each sample to the D. melanogaster reference genome (NCBI build 5.3, tophat.cbcb.umd.edu/igenomes.shtml). In order to get an accurate expression measure for the Cyp12d1 gene, a gene that exists as two, nearly identical copies in the reference (Cyp12d1-d and Cyp12d1-p) and is subject to copy number variation in the DSPR founders, we modified the reference annotation...
to eliminate the Cyp12d1-d gene. Following TopHat we had 135–169 million quality-trimmed reads per sample, and 89.1–90.4% of these aligned to the reference genome. Subsequently we used Cuffdiff (version 2.1.1, cufflinks.cbcb.umd.edu; [72, 73]) to identify differentially expressed genes (see S1 Text for details of the code and settings). For each comparison between samples we considered only those tests that were successfully executed ('status' column in "gene_exp.diff" Cuffdiff output file has 'OK' flag), and unless otherwise stated only considered genes to be significantly differentially expressed if they survived a Benjamini-Hochberg correction for multiple-testing ($q < 0.05$).

**RNAi**

To functionally test four P450 genes implicated in our mapping and RNAseq experiments we employed RNAi using the bipartite Gal4-UAS system, and UAS strains from the Vienna *Drosophila* Resource Center (VDRC [74]). We used strains harboring phiC31-based UAS transgenes, or "KK" lines (107641, Cyp6d5; 109248, Cyp12d1-d; 109256, Cyp12d1-p; 109771, Cyp301a1), and strains harboring P-element transgenes, or "GD" lines (50507, Cyp12d1-d; 21235 and 49269, Cyp12d1-p), along with the respective background control lines (60100, KK landing site control strain; 60000, w1118 host strain for GD library).

It has been reported that the 60100 control strain for the KK library has two landing sites [75]. We used the primers described in Green *et al.* [75] to determine which of the landing sites are occupied in the KK-UAS strains we employed: Strain 109248 has the vector integrated into the annotated target site only, strains 107641 and 109256 have vectors in the newly-identified, previously unannotated site, and 109771 has vectors in both landing sites. These differences in transgene position and number make it more difficult to directly compare effects across genotypes.

We tested the phenotypic effects of RNAi-based expression knockdown using two Gal4 drivers. First, we crossed males from a ubiquitous Gal4 driver under the control of the Actin 5C promoter (Bloomington *Drosophila* Stock Center, BDSC number 25374) to females of each UAS or control strain, testing female progeny from these crosses in our standard caffeine resistance assay. Second, we used an RU486-inducible "GeneSwitch" Gal4 driver [49] also under the control of the Act5C promoter (BDSC number 9431). Using this strain we again ubiquitously expressed Gal4 in all cells, but exerted some temporal control by driving Gal4 only in young adults. Males from the RU486-inducible Gal4 strain were crossed to females of the KK-UAS and 60100 control strains, and the resulting female progeny fed on media containing RU486 (M8046, Sigma) at 160μg/ml for 24 hours (trial 1) or 48 hours (trial 2) prior to the assay. The resistance assay was conducted as described, except that in addition to 1% caffeine the test media contained 160μg/ml RU486.

**Cyp12d1 Copy Number PCR**

We used primer sets provided in Schmidt *et al.* [12] and McDonnell *et al.* [42] to call copy number variation at the Cyp12d1-d/Cyp12d1-p locus in lines used in our experiments, and for the DGRP validated our genotype calls against bioinformatic, sequencing read-depth calls from Good *et al.* [41]. CNV genotype calls for the DSPR founders and the DGRP are presented in S3 Dataset, while inferred CNV calls in the DSPR RILs—based on the mosaic haplotype structure of each RIL—are presented in S6 Table.

**Statistical Analysis**

All statistics were carried out using core, or custom written, routines using the R statistical programming language (www.R-project.org).
Supporting Information

S1 Fig. Variation in phenotype among DSPR RILs. Means (filled circles) and 1 SDs (vertical lines) for caffeine resistance.
(PDF)

S2 Fig. Variation in phenotype among DGRP inbred lines. Means (filled circles) and 1 SDs (vertical lines) for caffeine resistance.
(PDF)

S3 Fig. Founder haplotype means and 1-SDs for QTL mapped in just one DSPR population. The number of RILs for which we assign a founder genotype (probability > 0.95) is listed at the bottom of each bar. Only founder means associated with at least 5 observations are presented.
(PDF)

S4 Fig. Effect of copy number variation at Cyp12d1 on caffeine resistance in the DSPR and DGRP mapping panels. The plots show the mean phenotype (± 1-SD) of those lines with one copy of Cyp12d1 (“single”) or two copies (“Dup”), with the number of lines in each class shown at the bottom of each bar. There is a significant effect of the duplication on caffeine resistance in the DSPR pA population (Welch’s t-test, \(p < 10^{-15}\)) and in the DSPR pB population (Welch’s t-test, \(p < 10^{-5}\)). However, the effect on the DGRP is not formally significant ( Welch’s t-test, \(p = 0.065\)).
(PDF)

S5 Fig. Genome scan for caffeine resistance QTL after controlling for Cyp12d1 CNV status. The format of this plot is identical to that of Fig 1 in the main text, but with genomewide 5% permutation thresholds of 8.0 LOD (pA) and 7.2 LOD (pB). In the original genome scan we identified 10 QTL, three in both populations (Q1, Q2, Q3), one in pA only (Q9), and six in pB only (Q4, Q5, Q6, Q7, Q8, Q10). After controlling for the CNV, we continue to identify Q1 in both panels, Q9 in pA only, and Q5, Q6, Q7, Q8, and Q10 in pB only. Q2—the QTL that harbors Cyp12d1—disappears in both panels, suggesting much of the effect of that QTL is due to the CNV and/or factors in LD with this variant. Q3 is now only identified in pB, and is absent in pA. The peak at Q4 is also absent. Finally, a peak towards the end of 3R in pA (3R:21960000..22250000), that was just below the significance threshold in our original analysis, is now significant.
(PDF)

S6 Fig. Effect of CNV on Cyp12d1 expression in female heads. King et al. [36] generated array-based, genomewide expression data from female head tissue for 600 genotypes. Each genotype was the result of a cross between an independent pair of DSPR RILs. Using the normalized expression data for Cyp12d1, which represents a composite measure of expression from any and all gene copies present in the target genotype since the gene copies are not distinguished on the array, and the CNV status provided in the current study, there is a strong effect of CNV status on Cyp12d1 expression (linear regression, \(p < 10^{-15}\)).
(PDF)

S1 Dataset. Expression levels of the five P450 genes under mapped QTL. Data is taken directly from the "gene_exp.diff" Cuffdiff output file.
(PDF)

S2 Dataset. Vienna Drosophila Resource Center (VDRC) UAS-RNAi hairpin sequences for Cyp12d1-d and Cyp12d1-p genes. All data is taken directly from VDRC website (stockcenter.
Hairpin sequences designed to target one member of the gene pair have strong similarity with sequence from the other gene. Bases in hairpin sequences that are different between the two genes are highlighted in green.

S3 Dataset. Copy number variation (CNV) genotype calls for the Cyp12d1 gene region in the DSPR and DGRP mapping panels.

S1 Text. Code used for RNAseq analysis.

S1 Table. LOD scores for genomewide QTL scans in the DSPR. The "Chromosome" column indicates the chromosome arm of the position under test (X, 2L, 2R, 3L, and 3R). The "PhysicalPosition" and "GeneticPosition" columns indicate the physical and genetic positions of the site under test, respectively. The "LODscore" columns present the LOD scores at each position in each of the two mapping populations (pA and pB), and in both populations after accounting for Cyp12d1 CNV status ("CNVadjusted"). The two "VarianceExplained" columns indicate the fraction of the phenotypic variance in the population that is explained by among-line genetic variation at the site under test.

S2 Table. Protein-coding genes implicated by mapped QTL. The "QTL" column indicates the QTL interval the gene is within. The "MappingPopulation" column indicates which population (pA and pB) the QTL was mapped in. The "ProteinCodingGene" column gives the symbol for the gene, and "GenePosition" gives the position of the gene in Release 5 of the Drosophila melanogaster reference genome.

S3 Table. RNAseq-based expression data for all genes surviving a genomewide FDR threshold of 5% in at least one contrast. The data is taken directly from the "gene_exp.diff" Cuffdiff output file, and simply trimmed to remove data for those genes failing to reach genomewide significance (q < 0.05) in at least one contrast. The "Chromosome", "GeneStart", and "Gene-Stop" columns give the position of the gene in the D. melanogaster reference genome (NCBI build 5.3). The "Sample1" and "Sample2" columns give the names of the two samples being compared, with four contrasts possible: Low-control versus High-control, Low-caffeine versus High-caffeine, Low-control versus Low-caffeine, and High-control versus High-caffeine. The "TestStatus" column is derived from Cuffdiff; Only data from tests marked "OK" was used in this study. The columns "Sample1.FPKM" and "Sample2.FPKM" give the FPKM (Fragments Per Kilobase of exon model per Million mapped fragments) values for each sample. The "Log2. FoldChange" column gives the log2 fold change in expression (Sample2 divided by Sample1). The "TestStatistic" column gives the test statistic used by Cuffdiff to compute the significance of the observed change in FPKM between samples, the "Pvalue" column provides the uncorrected p-value of the test statistic, and the "Qvalue" column provides the Benjamini-Hochberg FDR adjusted p-value of the test statistic.

S4 Table. Results of GWAS for caffeine resistance loci using the DGRP. The data is filtered directly from the "gwas.all.assoc" file returned by the Mackay lab DGRP2 analysis pipeline (dgrp2.gnets.ncsu.edu; Accessed June 3, 2014). The "Chromosome" and "Position" columns give information on the variant under test, and the "MAF" column gives the minor allele frequency of the variant in the DGRP. The "Pval" column is the p-value of an association test.
using a mixed-effects model accounting for relatedness among lines. Full details on the analytical pipeline is provided in Huang et al. [33]. Only sites significant at the 1% level ($p < 0.01$) are provided.

**S5 Table.** Raw caffeine resistance phenotypes measured in the DSPR and DGRP mapping panels. The “MappingPanel” column indicates which population the experimental genotype is derived from (the DSPR or DGRP). For DSPR genotypes, columns “Popn” and “PopnRep” indicate which population (pA or pB), and which subpopulation (pA1, pA2, pB1, pB2) the genotype is derived from (for DGRP genotypes all values in these two columns are ‘NA’). The “Genotype” column presents the RIL names for DSPR genotypes and the Bloomington Drosophila Stock Center codes for the DGRP genotypes. There are multiple rows for any given genotype, with each row containing data for a single female. The “RepVial” column indicates which replicate vial each experimental individual is derived from. In most cases all individuals tested for a genotype come from the same replicate vial, listed as ‘1’. The “CaffeineResistance” column is the time, in hours, each experimental female survived during continuous exposure to medium supplemented with 1% caffeine.

**S6 Table.** Inferred Cyp12d1 duplication status for all DSPR RILs. The “RIL” column provides the line code for all 1,715 genotypes phenotyped as part of this study. The “Duplication-Status” column codes the copy number present at the Cyp12d1 gene: 0 = single copy, 1 = duplicated, NA = unknown. Only those RILs where the founder genotype at the Cyp12d1 locus is known with at least 95% certainty are called for copy number variation.

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**Author Contributions**

Conceived and designed the experiments: ADL SJM. Performed the experiments: BRS CAH JLH MAN. Analyzed the data: MAN SJM. Contributed reagents/materials/analysis tools: EGK. Wrote the paper: SJM.

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