Title
Genetic networks controlling structural outcome of glucosinolate activation across development.

Permalink
https://escholarship.org/uc/item/5w15b6rv

Journal
PLoS genetics, 4(10)

ISSN
1553-7390

Authors
Wentzell, Adam M
Boeye, Ian
Zhang, Zhiyong
et al.

Publication Date
2008-10-24

DOI
10.1371/journal.pgen.1000234

Peer reviewed
Genetic Networks Controlling Structural Outcome of Glucosinolate Activation across Development

Adam M. Wentzell1,2, Ian Boeye2, Zhiyong Zhang2, Daniel J. Kliebenstein1,2*

1 Genetics Graduate Group, University of California Davis, Davis, California, United States of America, 2 Department of Plant Sciences, University of California Davis, Davis, California, United States of America

Abstract

Most phenotypic variation present in natural populations is under polygenic control, largely determined by genetic variation at quantitative trait loci (QTLs). These genetic loci frequently interact with the environment, development, and each other, yet the importance of these interactions on the underlying genetic architecture of quantitative traits is not well characterized. To better study how epistasis and development may influence quantitative traits, we studied genetic variation in Arabidopsis glucosinolate activation using the moderately sized Bayreuth × Shahdara recombinant inbred population, in terms of number of lines. We identified QTLs for glucosinolate activation at three different developmental stages. Numerous QTLs showed developmental dependency, as well as a large epistatic network, centered on the previously cloned large-effect glucosinolate activation QTL ESP. Analysis of Heterogeneous Inbred Families validated seven loci and all of the QTL × DPG (days post-germination) interactions tested, but was complicated by the extensive epistasis. A comparison of transcript accumulation data within 211 of these RILs showed an extensive overlap of gene expression QTLs for structural specifiers and their homologs with the identified glucosinolate activation loci. Finally, we were able to show that two of the QTLs are the result of whole-genome duplications of a glucosinolate activation gene cluster. These data reveal complex age-dependent regulation of structural outcomes and suggest that transcriptional regulation is associated with a significant portion of the underlying ontogenic variation and epistatic interactions in glucosinolate activation.

Introduction

Most phenotypic variation present in natural populations is under polygenic control, largely determined by genetic variation at multiple quantitative trait loci (QTLs), which has motivated considerable efforts to elucidate the genetic basis of these polygenic traits [1–3]. A complete understanding of quantitative traits necessitates identification of the underlying genes and their associated additive, dominance, and epistatic effects [2]. In addition, the underlying genetic architecture of many quantitative traits may vary across development and different environments. As such, a comprehensive description of a quantitative traits genetic architecture requires analysis in several developmental or environmental contexts to assess stability of the genetic architecture [2,4,5].

QTL mapping, which measure the association of genetic markers with phenotypic variation, is one of the most common approaches for identifying loci and epistatic interactions controlling polygenic inheritance [1]. Improved statistical models, marker technology, and genomic resources have facilitated QTL mapping experiments for a wide array of quantitative traits, ranging from development and morphology to metabolism and disease resistance [2,4,5]; However, QTL mapping experiments are often limited to a single stage in development and one or few environments. As a consequence, there is little information available to answer the question of how the underlying genetic architecture varies across developmental and environmental contexts.

Accurate characterization of a quantitative trait’s underlying genetic architecture is often limited by practical considerations that limit the number of progeny included in a mapping analysis. Small populations are especially problematic in the presence of epistasis between QTLs, as the pair wise comparisons required to detect these interactions rapidly exhausts the available genotypic variation, leading to an underestimation of numbers of loci and interactions, resulting in an incomplete picture of the genetic architecture [6,7]. One common type of epistasis occurs when a trait is controlled by one or few large effect loci and numerous modifier QTLs of smaller effect, a situation frequently observed in plant disease resistance [8–16]. In such systems, the effects of any modifiers are most detectable in those lines containing the appropriate allele at the large effect locus; however this reduces by half the population in which to detect these smaller effect loci, significantly reducing statistical power [5–7]. Thus, resolution of the underlying genetic basis of complex traits requires the analysis of large populations across different environments or developmental stages [17,18].

To investigate how development and epistasis can interact to control the variation in an adaptive trait, we studied the outcome
Author Summary

A principal interest in biology is to understand how natural genetic variation translates into phenotypic variation. A key component of this connection is how the genetic variation interacts with other sources of variation, such as environment (G × E), development (G × D), or other genetic loci (G × G or epistasis). To analyze the molecular underpinnings of these quantitative genetics interaction terms, we investigated the genetic architecture of an adaptive trait, glucosinolate activation, in Arabidopsis thaliana during the development of what is considered a static mature rosette. Variation in glucosinolate activation was principally controlled by epistatic and G × D interactions. Epistatic interactions identified both Mendelian epistasis, where regulatory loci controlled enzymatic loci, and quantitative interactions between regulatory loci. G × D appeared to involve master regulatory loci as determined by trans-eQTL hotspot analysis. Finally, two common glucosinolate activation QTLs appear to have evolved via gene loss and sub-functionalization following quadruplication of an ancestral genomic fragment, potentially by two whole-genome duplications. Thus, genomic duplication events may facilitate the formation of quantitative genetic variation. This study provides insights into the molecular basis of the link between genetic and phenotypic variation in a potentially adaptive trait.

of glucosinolate activation within Arabidopsis thaliana using a moderately sized recombinant inbred population. Glucosinolates are the inert storage form of a two part phytochemical defense system found throughout the Brassicaceae, where biologically active structures are catabolically produced by the enzyme myrosinase [19–21](Figure 1). The particular structural outcome, as defined by the chemical structure of the end product, of glucosinolate activation plays an important role in plant defense against insect herbivory [22–25], as well as the nutritional and flavor characteristics of brassicaceous crops [26]. Further, the structural outcome shows significant intraspecific diversity such that natural accessions activate a glucosinolate to either a nitrile or isothiocyanate depending upon their genotype. Thus an improved understanding of the genetic basis of variation in structural outcomes has important potential implications in evolution and ecology as well as nutrition and agriculture.

Glucosinolate activation in Arabidopsis provides an excellent model for studying how development and epistasis influence quantitative traits, with a molecularly characterized biochemical pathway comprising demonstrated epistatic interactions and developmental variation. During glucosinolate activation, the myrosinase enzyme catabolically generates the unstable intermediate. The final structural outcome of subsequent rearrangement of this unstable intermediate is influenced by the presence or absence of various structural specifier proteins (Figure 1). The Epithiospecifier Protein (ESP) and an as yet unidentified simple nitrile structural specifier (AtNSP), promote the formation of simple and epithionitrides at the expense of the default isothiocyanate rearrangement via two biochemically related yet separate rearrangements (Figure 1) [22,23,27–31]. The Epithiospecifier Modifier (ESM1) epistatically modulates ESP-mediated epithionitride and simple nitrile rearrangements (Figure 1)[22,23]. The observation of quantitative variation influencing the developmental regulation of glucosinolate activation enabled us to explore the stability of QTLs and epistatic interactions across development [30,32].

We used the Bayreuth (Bay-0) × Shahnara (Sha) recombinant inbred lines (RILs) [33] to map QTLs controlling the structural outcome of glucosinolate activation in Arabidopsis. These parental accessions contain genetic variation for ESP and ESM1 and differ in developmental regulation of structural outcomes [32]. We measured structural outcomes at 30, 35 and 42 days post germination (DPG), and compared the resulting maps to assess the stability of the underlying genetic architecture across development. These DPG were chosen because day 30 represents the end of logarithmic growth in all of the lines, or Stage 1.10, while day 42 is one week away from the earliest RIL flowering (Stage 5.10) in our environmental conditions [32,34,35]. Thus, we can focus on developmental changes in what is typically considered a static rosette and is also the tissue and stages where lepidopteran insects predominate on Arabidopsis in the wild [36]. This analysis identified eleven loci and twelve pair wise epistatic interactions influencing structural outcomes, as well as eight different QTL × DPG interactions. Heterogeneous Inbred Families (HIFs) differing only for their genotypes at each QTL locus validated seven loci and all of the QTL × DPG interactions tested.

![Figure 1. Glucosinolate Activation and the Subsequent Structural Rearrangement.](https://example.com/figure1.png)

Myrosinase enzymes initiate glucosinolate activation by hydrolyzing the thioglucose bond, generating an unstable aglycone intermediate (bracketed). This intermediate can spontaneously rearrange into the isothiocyanate structure. Epithionitride structural specifiers such as ESP can promote the formation of epithionitride structures from glucosinolates with a terminal double bond, and simple nitriles from all other glucosinolates. As yet unidentified Arabidopsis simple nitrile structural specifiers (AtNSPs) promote the formation of simple nitrile structures from all glucosinolates. In parenthesis are the Bayreuth and Shahnara accession showing their predominant glucosinolate activation form.

doi:10.1371/journal.pgen.1000234.g001
The availability of transcript accumulation data within 211 of these RILs [37] enabled comparison of expression QTLs (eQTLs) for structural specifier genes and their homologs, which demonstrated collocation of eQTL clusters with eleven of the identified loci. These data reveal complex age dependent regulation of structural outcomes and suggest that transcriptional regulation is associated with a significant portion of the underlying variation, and may explain the epistatic interactions described here.

**Results**

**Structural Outcomes of Glucosinolate Activation in Bay-0 and Sha**

Bay-0 and Sha contain different glucosinolates due to variation at the *GSL-AOP* and *GSL-Elong* loci, such that Bay-0 has 3-hydroxypropyl glucosinolate as its main short chain aliphatic glucosinolate and Sha has but-3-enyl glucosinolate [38]. These two accessions also differ in the structures they produce following activation of these glucosinolates. Bay-0 lacks functional ESP and produces mixtures of simple nitriles and isothiocyanates, depending on the age of the plant [32]. In contrast Sha possesses a functional allele of ESP and produces mixtures of epithionitriles, simple nitriles and isothiocyanates (Figure 1). In agreement with previously published analysis, interplanted Sha parental controls had an increasing epithionitrile proportion during development for both the exogenous (Figure 2A and C) and endogenous glucosinolate substrates (Table S4) [32]. In contrast to Sha, the Bay-0 parent showed little variation in the structural outcome of exogenous allyl glucosinolate activation between 30, 35 and 42 DPG (Figure 2A and C). The activation products for the endogenous 3-hydroxypropyl glucosinolate produced in Bay-0 could not be detected in this experiment. Thus, there is developmental variation in glucosinolate activation between the Bay-0 and Sha parental accession which allows us to investigate how the genetics of glucosinolate activation interact with plant development.

**Distribution of Structural Outcomes in the RIL Population**

We measured the structural outcome of glucosinolate activation using exogenous allyl glucosinolate in the Bay-0 × Sha RILs and compared the trait distribution to the interplanted parental

![Figure 2. Ontogenic variation of Structural Outcome in Glucosinolate Activation.](http://www.plosbiology.org/content/4/10/e1000234.Fig2.jpg)

**Figure 2. Ontogenic variation of Structural Outcome in Glucosinolate Activation.** GC-FID was used to measure the structural outcome from exogenous allyl glucosinolate activation in the RILs and interplanted Bay-0 (dashed line) and Sha (solid line) parental accessions at 30, 35, and 42 DPG. For the parental analysis, averages with standard errors are shown. For the RILs, the percent simple nitriles were binned into 1 percent intervals from 0 to 20 percent, and percent epithionitriles were binned in 5 percent intervals from 0 to 100 percent and the number of RILs per bin are shown. For the RILs, black bars show the distribution as measured at 30 DPG, grey is 35 DPG and white at 42 DPG. A) Simple nitriles in Bay-0 (dashed) and Sha (solid) parental accessions at 30, 35, and 42 DPG. B) Distribution of simple nitrile formation in the RILs. C) Epithionitrile formation in parental accessions. D) Distribution of epithionitrile formation in the RILs. doi:10.1371/journal.pgen.1000234.g002
controls. The use of exogenous allyl glucosinolate allowed us to mask effects of variation in glucosinolate biosynthesis and accumulation. Considerable transgressive segregation was observed for simple nitrile where there were numerous RILs with values higher and lower than the Bay-0 or Sha parents (Figure 2B). Further, there was transgressive segregation for epithionitrile as evidenced by the RILs with a higher value than the Sha parent (Figure 2D). This suggests that alleles promoting the formation of each structure exist in both parents. This is particularly surprising in the case of epithionitrile proportions, as the Bay-0 parent cannot produce epithionitriles due to a lack of functional ESP and as such might not be expected to contain genes enhancing epithionitrile formation.

This population includes 212 lines producing no epithionitrile structures following activation of allyl glucosinolate (Figure 2D). This is consistent with the previously observed requirement of a functional ESP allele for epithionitrile production [27,30,32]. For the subpopulation of RILs with a functional allele of ESP, the distribution gradually shifted towards increased epithionitrile production from 30 DPG to 42 DPG, in a manner consistent with the epithionitrile increase observed in the Sha parental controls (Figure 2C). However, individual RILs showed possible differences in both the direction and magnitude of change in epithionitrile production from 30 to 42 DPG, suggesting that there is genetic variation in age dependent control of epithionitrile proportion from 30 to 42 DPG (Table S4). These possible differences will however require identification of the QTL and HIF validation to ensure that this is not random variation around the mean.

The distribution of nitrile formation within the RILs did not show a similar ontogenic shift, but there were numerous lines that had no simple nitrile formation at 42 DPG (Figure 2B). This low or undetectable amount of simple nitriles is connected to epithionitrile substrate in these RILs at this DPG. This difference in trile proportions approaching complete utilization of the glucosinolate substrate in these RILs at this DPG. This undetectable amount of simple nitriles is connected to epithionitrile formation.

Structural Proportion QTLs

To identify loci controlling the diversity of structural outcomes in Bay-0 and Sha, we independently mapped QTLs for all traits at 30, 35, and 42 DPG. Analysis of epithionitrile proportions for exogenous allyl glucosinolate revealed nine loci (Figure 3A, Table S3), including seven novel QTLs and the previously identified ESP and ESM1 loci [22,23]. Three loci (ESP, GSL.Activ.II.13, and ESM1) were detected in the RILs at all three DPG, although the additive effect of ESM1 switched direction from promoting nitrile formation at 42 DPG relative to promoting isothiocyanate formation at 30 and 35 DPG. In spite of the fact that Bay-0 lacks functional ESP, four QTLs showed a positive impact of the Bay-0 allele on epithionitrile production, which is consistent with the observed transgressive segregation in the RILs (Figures 2D and 3A).

We identified ten loci affecting the proportion of simple nitrile structures produced from exogenous allyl glucosinolate, including ESP, ESM1 and eight novel loci (Figure 3B, Table S3). The average allelic substitution effect of these QTLs was 27% and the median was 20%. ESP, ESM1 and six of the novel loci overlapped with epithionitrile production QTLs with all but ESP showing the same direction of allelic effect upon epithionitrile and nitrile production (Figure 3). Six of the simple nitrile proportion QTLs were significant at a single DPG, three were detected at two consecutive DPG, and one locus (GSL.Activ.II.13) was detected in all three QTL maps. Increased simple nitrile proportions were fairly evenly distributed between the Bay-0 and Sha alleles (Figure 3B). One locus (GSL.Activ.IV.16) exhibited significant additive effects in opposite directions at 30 and 35 DPG (Figure 3B). Isothiocyanates identified a combination of the nitrile and epithionitrile QTLs (Figure S1). This suggests that the genetic architecture underlying glucosinolate activation is much more complex than the two locus model previously assumed [22,41].

Endogenous Glucosinolate Activation

We proceeded to compare QTLs identified using the exogenous allyl glucosinolate to those identified with the endogenous but-3-epin and 4-methylsulfinylbutyl glucosinolate, the two glucosinolates with the highest level of accumulation in this population. The

| Table 1. Heritability of Structural Outcomes of Glucosinolate Activation. |
|-----------------------------|----------------|----------------|
| Trait                       | P Value | Type III Sums of Squares | Heritability |
| % Simple Nitriles           | Geno    | DPG | Model | Geno | DPG | |
| % Epithionitriles           | <0.001  | 0.397 | 13083.4 | 6603.8 | 17.1 | 50.5 |
| Butenyl % Simple Nitriles   | <0.001  | <0.001 | 874209.6 | 427015.0 | 166069.7 | 48.8 |
| Butenyl % Epithionitriles   | <0.001  | <0.001 | 54639.5 | 28550.7 | 19106.5 | 52.8 |
| 4MSO % Simple Nitrile       | <0.001  | <0.001 | 124753.5 | 82196.3 | 15517.3 | 65.9 |

Geno represents the genotype term in the model. DPG is the days-post-germination factor in the model. doi:10.1371/journal.pgen.1000234.t001
analysis of structural outcomes of endogenous glucosinolate activation is complicated by independent assortment at the GSL.Elong and GSL.AOP biosynthetic loci, limiting each measurable endogenous glucosinolate to one quarter of these RILs [38,42,43]. QTL analysis of endogenous but-3-enyl glucosinolate activation detected three loci and nine epistatic interactions affecting simple nitrile formation, and only ESP and three epistatic interactions for epithionitrile proportions. All QTLs were consistent with those observed using the exogenous allyl glucosinolate (Figure 3A and Table S3).

QTL analysis of 4-methylsulfinylbutyl glucosinolate activation identified three loci affecting simple nitrile formation, which were
all identified using exogenous allyl glucosinolate (Table S3). The Sha allele of ESP increased simple nitrile formation from this glucosinolate, which lacks a terminal double bond and cannot form epithionitrile structures. In contrast, the non-functional Bay-0 allele increases simple nitrile formation from the exogenous allyl glucosinolate, possibly because this eliminates substrate competition between \( \text{AtNSP} \) and ESP. As such, the use of exogenous allyl glucosinolate provides the greatest power to independently map both structural outcomes utilizing the entire RIL population.

**QTL \( \times DPG \) Interactions**

To test how plant age in DPG altered QTL identification, we conducted an ANOVA analysis of the significant genetic loci using the data from all three assays in a single model. Each genotype is replicated within each DPG allowing for a test of marker \( \times DPG \) interactions. Analysis of epithionitrile proportions using the full data set identified five marker \( \times DPG \) interactions, suggesting that these loci may be involved in controlling the increase in epithionitrile formation observed from 30 to 42 DPG (Table S3). Analysis of simple nitrile proportions in the full data set also detected five significant QTL \( \times DPG \) effects, suggesting age dependent regulation of simple nitrile rearrangements (Table S3). Three of the marker \( \times DPG \) interactions significantly affected both simple nitrile and epithionitrile proportions, possibly as a consequence of these two rearrangements competing for the same pool of substrate or co-regulation of the two glucosinolate activation outcomes. These loci with DPG interactions provide the potential to begin understanding how ontogenic variation and genetic variation interact at the molecular level.

**Numerous Epistatic Interactions Influence Glucosinolate Activation**

Given the numerous QTLs controlling glucosinolate activation and the requirement of a functional allele at the ESP locus for epithionitrile production, we hypothesized that there would be significant epistasis affecting structural outcomes in this mapping population. We utilized an ANOVA to test all possible pair wise QTL interactions for significant epistasis. We identified a total of eleven different pair wise epistatic interactions for epithionitrile proportion, including the previously described ESP \( \times \) ESM1 interaction \([22,23]\) (Figure 3A, Table S3). Consistent with the requirement of functional ESP for epithionitrile production, all other QTLs for epithionitrile formation showed a significant epistatic interaction with ESP for at least one of these data sets (Table S3). Epistatic interactions involving ESP represent classical epistasis, where genotypes with the nonfunctional Bay-0 ESP allele produce no detectable epithionitriles, hiding the function of the interacting loci (Figure 4A/C). This suggests that the QTLs epistatic to ESP may function to modulate the activity of the functional Sha allele of ESP. Interestingly, the highest levels of epithionitriles were not exclusively observed in RILs with Sha genotypes at the interacting locus. For example, the ESP \( \times \) GSL.Activ.V.18 interaction produced the highest epithionitrile proportions in lines with the Bay-0 allele at GSL.Activ.V.18 (Figure 4C).
While all epithionitrile QTLs were epistatic to ESP, four epistatic interactions were detected for epithionitrile proportion that did not involve ESP (Figure 3B and 4D and Table S3). These interactions were examples of quantitative epistasis, where the effects of one locus was modified by the other locus in a quantitative manner, rather than the absolute dependence of one locus on the other locus as exhibited by interactions involving ESP (Figure 4D versus 4A/C). Additionally, these epistatic interactions formed networks such that GSL.Activ.II.13, GSL.Activ.II.61 and ESM1 showed all possible pair wise epistatic interactions with each other as well as with ESP (Figure 3B and Figure 4B). Likewise, a similar network involves GSL.Activ.I.16xGSL.Activ.IV.16 and ESP (Figure 3). This suggests that complex epistasis may begin to identify underlying regulatory or protein interaction networks controlling the structural outcome of glucosinolate activation.

Simple nitrile proportion from exogenous allyl glucosinolate identified fewer epistatic interactions than epithionitrile formation with most interactions also involving the ESP locus. The ESPxESM1 epistatic interaction affected both simple nitrile and epithionitrile proportion, but with different effects on each structural proportion (Figure 4 A and B). The lower number of epistatic interactions for simple nitrile formation is partly explained by the lower variation present for simple nitrile formation within these RILs (Figure 2A/B).

**QTL Validation using HIFs**

To provide additional support for the identified GSL.Activ QTLs, we obtained HIFs that contain appropriate variation for seven of the loci detected in this study (Table S1). For the GSL.Activ.II.13 and GSL.Activ.III.64 loci, the available HIFs only contained a non-functional ESP, thus we were unable to test the effects of these loci on epithionitrile production. We assayed glucosinolate activation in each HIF line at 24 and 38 DPG, to confirm the QTLs and any interaction between the QTLs and DPG. The HIFs confirmed seven of the GSL.Activ QTLs. This included four loci for epithionitrile formation, two for simple nitrile production and several for the total production of nitriles or isothiocyanates (Figure 5, Table 2, Table S5). HIF-241 and HIF-425 vary for ESP and confirm that a functional Sha allele is necessary for epithionitrile production (Figure 5A, Table S5). Interestingly, the efficiency of epithionitrile formation significantly differs between these two HIFs (\(P = 0.047\) for HIFxESP genotype and \(P = 0.048\) for HIFxESP genotypexDPG), confirming the presence of background ESP modifiers. The level of validation observed in the HIF analysis is strongly supportive of the QTL mapping results, as each HIF genotype was only analyzed in six-fold replication per DPG whereas each marker genotype in the RIL study was analyzed in roughly 200-fold replication, lending more power to the RIL analysis.

The original RIL QTL analysis did not replicate each line within each DPG, and as such, we designed the HIF analysis to confirm that these QTLs do interact with the plant age in DPG. All four confirmed epithionitrile proportion QTLs exhibited significant genotypexDPG effects (Figure 5, Table 2 and Table S5). For example, HIF-157 which varies for GSL.Activ.V.18, showed a significant difference in epithionitrile proportions between alleles at 24 but not 38 DPG (Figure 5B and Table S5). In contrast, the locus GSL.Activ.V.48 in HIF-350 showed no
difference in epithionitrile proportions between alleles at 24 DPG, but a significant effect at 38 DPG in agreement with the QTL prediction from the RIL analysis (Figures 3A, 5C and Table S5). These results confirm that we have identified ontogenic dependent QTLs in our study.

### HIF Analysis Reveals an Additional QTL Tightly Linked to ESP

QTL mapping analysis of epithionitrile proportion consistently produced a large peak in the LOD plot at the ESP locus but there was also frequently a small shoulder (Figure 6A). This could be explained by residual significance from the large-effect ESP locus or suggest the presence of a tightly linked QTL in this region. Due to tight genetic linkage with ESP, we did not include this putative locus in our statistical models. We did however identify two HIFs in this region, HIF149 with functional ESP and HIF107 with non-functional ESP. These HIFs allow us to test for the existence of this additional locus as well as its potential dependency upon ESP (Table S1). Analysis of structural outcomes in HIF149 confirmed the existence of an additional QTL telomeric of ESP on Arabidopsis chromosome 1. This QTL, *GSL.Activ.I.69*, affects both simple nitrile and epithionitrile proportions (Table S5 and Figure 6B/C). *GSL.Activ.I.69* also displays an age dependent effect on simple nitrile proportion, but not epithionitrile proportion (Table S5 and Figure 6B/C). Interestingly, HIF107 did not identify a significant QTL, suggesting that *GSL.Activ.I.69* is epistatic to ESP (Table S5). This HIF analysis of a QTL shoulder suggests there may be even further additional QTLs for glucosinolate activation in this RIL population.

### eQTLs for Glucosinolate Activation Genes

The *TGG1* and *TGG2* myrosinases, ESP, and ESM1 are the primary genes with a demonstrated role in controlling glucosinolate activation and associated structural outcomes within the Arabidopsis rosette [22–25,27,30,31]. However, these genes and the MBPs form gene families in Arabidopsis and some of these uncharacterized homologs may determine the genetic variation observed in the structural outcome of glucosinolate activation. To identify any potential expression level polymorphisms in these four gene families that may control the glucosinolate activation QTLs, we identified eQTLs for the full list of potential glucosinolate activation homologs (Table S2) [37].

We first obtained the estimated heritability for transcripts encoding these potential glucosinolate activation genes [37]. The average transcript heritability for the 30 measurable probe sets was 63.0%, which is significantly higher than the genome wide average transcript heritability (t-test, $P = 0.045$). Further, the glucosinolate activation genes had more eQTLs, both cis and trans than the average Arabidopsis transcript. This excess was most dramatic with trans-acting eQTLs, with an average of 3.1 trans-eQTLs detected for each glucosinolate activation gene in comparison to the average Arabidopsis transcript, with 1.5 trans-eQTLs in this population [37]. These results agree with previous studies showing the transcripts for glucosinolate biosynthetic genes had higher heritability and variance than the average Arabidopsis transcript [38,44].

The previous analysis of eQTLs within the Bay-0×Sha RIL population was conducted at 35 DPG in the same growth chamber allowing their direct comparison [37]. The eQTLs controlling transcript accumulation of the glucosinolate activation genes revealed eQTL clusters collocating with ten of the eleven structural proportion QTLs (Figure 7B). The eQTL clusters partitioned into both cis and trans-eQTL clusters. The two cis-eQTL clusters are associated with the genomic regions around ESP and ESM1, which contain several ESP, ESM1, and MBP homologues but do not overlap with known trans-eQTL hotspots (Figure 7A and B [see the black arrows]). These regions appear to be the result of two separate whole-genome duplications that copied an original region containing the ancestral genes for ESP, ESM1 and the MBPs (Figure 7A). These consecutive
duplications generated four genomic regions that contain additional non-glucosinolate genes whose paralogs also have a conserved order (Figure 7A). In these two cis-eQTL clusters, we found six and eight cis-eQTLs for putative glucosinolate activation genes in the ESP and ESM1 regions respectively (Figure 7). While variation in ESP and ESM1 have been shown to control glucosinolate activation phenotypes, these local cis-eQTL clusters suggest that additional genes at each locus could contribute to the effects on the structural outcome of glucosinolate activation.

In addition to cis-eQTL clusters, there were several GSL.Activ loci that co-located with clusters of trans-eQTLs suggesting that these may be regulatory loci with a measurable effect on gene expression and structural outcomes. In particular, GSL.Activ.II.13 collocated with a large trans-eQTL cluster but not with the genomic position of any of the identified glucosinolate activation gene homologs (Figure 7B – double black arrows highlight homologe clusters). Interestingly, this locus also collocates with a large trans-eQTL hotspot controlling ~1,200 genes suggesting that it may contain a polymorphism in a pleiotropic developmental regulator [37]. Genetic variation at the GSL.Activ.II.13 locus alters the expression of all four ESM1 homologues, three of four ESP homologues, four MBPs, and two myrosinases. A closely linked locus, GSL.Activ.II.42, also collocates with a large trans-eQTL cluster controlling two myrosinases, eight of twenty potentially interacting genes and ~2,500 other genes (Figure 7B)[37]. Thus, these two loci show that global regulatory loci can have a measurable consequence on glucosinolate activation. Interestingly, the GSL.Activ.I.69 locus identified in the HIF analysis is associated with eQTLs controlling transcript accumulation for TGG2, ESP, ESM1 and an MBP (At3g16470), suggesting that the GSL.Activ.I.69 may also be a regulatory locus. However, it does not co-locate with a previously identified global trans-eQTL hotspot suggesting that it may be more specific to glucosinolate activation [37].

**Discussion**

The structural outcome of glucosinolate activation strongly influences plant defense, and has been thought to be predominantly controlled by genetic variation in ESP and its epistatic modifier locus, ESM1 [22,23,28–30]. Recently this model has been shown to be overly simplistic by the identification of interactions between developmental, environmental and genetic factors influencing the regulation of the structural outcomes in Arabidopsis, Brassica and Nasturtium [30,32,45–48]. These analyses suggested that the mixture of biologically active product structures is generated by two biochemically distinct rearrangements that divert the glucosinolate substrate away from the default isothiocyanate structure, each subject to complex regulatory patterns in rosette leaves. As such, we conducted QTL mapping analysis to simultaneously map loci controlling both simple nitrile and epithionitrile formation using allyl glucosinolate in the Bay-0 x Sha RIL population. The power afforded by this population containing a moderate number of independent lines revealed considerable additional complexity underlying the critical rearrangement step in glucosinolate activation.

Our analysis identified twelve loci and 17 epistatic interactions controlling the mixture of activation structures produced from allyl glucosinolate, including the previously described ESP and ESM1 QTLs (Figures 3 to 6, Table 2, Table S3 and S5). HIF analysis confirmed seven loci as well as their proposed interaction with plant development (Figure 5 and 6, Table 2 and S3). We also found extensive collocation between eQTLs for the genes potentially involved in glucosinolate activation and the structural outcome phenotypic QTLs (Figure 7 and Table S2). These results

---

**Figure 6. HIF Analysis Identifies an Additional Locus, GSL.Activ.I.69.** HIF149 was utilized to test a putative QTL near the ESP locus. Glucosinolate activation was assayed using allyl glucosinolate and the mean and standard error are shown (N=6). Significant differences (P<0.05) are indicated by different letters. A) Illustration of the described shoulder in the LOD profile near the ESP locus on chromosome I generated by CIM based QTLcartographer analysis (35 DPG is illustrated). The black bar shows the region of the shoulder that varies in HIF149. B) Effect of HIF149 upon epithionitrile production. C) Effect of HIF149 upon epithionitrile production.

doi:10.1371/journal.pgen.1000234.g006
Figure 7. Genomic Duplication and eQTLs for Glucosinolate Activation Genes. Phylogenetic analysis was used to identify homologs of the four known families of glucosinolate activation genes (Myrosinase, ESP, ESM1 and MBPs) within Arabidopsis. The genomic position of these genes and their associated eQTLs within this population were then identified and are plotted below. A) The genomic arrangement of the glucosinolate activation genes at the identified genomic duplication underlying the ESM1 (top) and ESP (bottom) QTLs. Genes are colored based on the glucosinolate activation gene with which they share homology; ESP (red), ESM1 (green), and MBP (blue). Genomic relationships for eight additional paralogous sequences are indicated by dashed lines. The AGI locus designation for the genes used as the beginning and end of each illustrated regions are indicated, and the loci corresponding to ESP, ESM1, MBP1 and 2 are labeled, with the scale indicated. B) eQTLs for the identified Myrosinase, ESM1, ESP, and MBP homologs are shown [37]. The five Arabidopsis chromosomes are indicated with roman numerals and represented contiguously from left to right. The position of the identified structural outcome QTLs are indicated at the top of the heat plots. Horizontal lines separate the four gene families with the defining member labeled. Within each family the genes are ordered by physical position from top to bottom. eQTLs shaded blue indicate that increased transcript accumulation for that gene is associated with the Bay-0 allele, and red indicates Sha, with the intensity of the color proportional to the magnitude of the additive effects. The double black arrows show the orientation and position of the tandem gene clusters at the ESP and ESM1 loci. The vertical dashed lines show the likely position for the GSL.Activ.II.13 and 42 loci.

doi:10.1371/journal.pgen.1000234.g007
support a model wherein control of structural outcomes during glucosinolate activation results from multi-locus regulation of both simple nitrile and epithionitrile formation and the interplay between these competing rearrangement processes generates the final mixture of biologically active structures.

Age Dependent Regulation of the Structural Outcome of Glucosinolate Activation

The Bay-0 and Sha parents had previous been shown to have different ontogenic control of glucosinolate activation [32]. Our analysis of epithionitrile proportions revealed several differences between 30, 35 and 42 DPG, and analysis of the full data set identified seven significant QTL×DPG interactions (Figure 3A and Table S3). Further, direct assessment of glucosinolate activation at two DPG in the HIFs confirmed these QTL×DPG interactions (Figure 5, Table 2, S3 and S5). This suggests that these loci are responsible for the age dependent regulation of epithionitrile production, and may act to regulate ESP expression. Similar levels of ontogenic QTL dependency were identified for epithionitrile production, and may act to regulate loci are responsible for the age dependent regulation of epithionitrile production, and may act to regulate activation at two DPG in the HIFs confirmed these QTL×DPG interactions (Figure 5, Table 2, S3 and S5). This suggests that these loci are responsible for the age dependent regulation of epithionitrile production, and may act to regulate ESP expression. Similar levels of ontogenic QTL dependency were identified for simple nitrile production in spite of the absence of ontogenic variation between Bay-0 and Sha for simple nitrile production (Figures 2A/B and 3B and Table S3). Thus, transgressive effects on simple nitrile proportions lacked ESP activity, suggesting that the ability to detect small effects on simple nitrile formation can be negatively impacted as a consequence of reduced flux in the presence of the competing epithionitrile rearrangement.

Structural Outcome QTLs May Result from an Ancient Duplication and Subsequent Neo-Functionalization

Analysis of the genomic regions underlying ESP and ESM1 revealed two distinct and tightly linked clusters of structural specifiers and myrosinase interacting proteins at each locus (Figure 7A). These linked clusters appear to be the product of an ancestral locus, which contained ESP, ESM1, and an MBP and underwent a tandem duplication followed by segmental duplication to generate the ESP and ESM1 loci, with the subsequent loss of some paralogs. These four genomic regions contain the majority of the ESP, ESM1 and MBP homologs but differ in their specific composition. Further, a number of cis-eQTL were detected for these genes, and the associated QTLs have divergent effects on the structural outcome of glucosinolate activation [22,23]. The presence of a large number of duplicated homologs suggests that these QTL may be complex loci with numerous tightly linked polymorphisms contributing to the observed phenotypic effects. Support for this idea comes from the observation that while the ESP and ESM1 proteins were shown to explain most of the effects of their respective QTLs, complementation of both QTLs did not completely recapitulate the phenotypes associated with each QTL [23,42].

The association of genomic duplications with glucosinolate activation QTLs suggests that such duplications may facilitate quantitative genetic variation by creating duplicate genes. The duplicate genes can then undergo genetic sub-functionalization such that the genes have differential functions across natural genotypes [44,49–51]. This role of genome duplications and QTL association has been previously seen in polyploid plants but not characterized in diploids [32,53]. This relationship between genome duplications and QTLs requires the analysis of more traits and cloning of more loci to establish the generality of this connection.

Simple Nitrile and Epithionitrile Rearrangements Involve Distinct Chemistry but Overlapping Genetics

Nine of twelve QTLs affected both simple nitrile and epithionitrile proportions formed from allyl glucosinolate (Figure 3 and 6, and Table S3). Additionally, four epistatic interactions were detected in both the simple nitrile and epithionitrile structural outcome data (Figure 3 and 4 and Table S3). While there was considerable overlap in the QTLs for the two distinct structural outcomes, there was not complete correspondence in the directionality of their effects. The majority of the QTLs altered the accumulation of nitrile and epithionitrile in the same direction, but the ESP locus had opposite effects on simple nitrile and epithionitrile proportions (Figure 3). In those cases where the same pair wise interactions via the ESP locus affected both simple nitrile and epithionitrile formation, they affected each proportion differently, supporting the concept of two independent rearrangements competing for the same substrate. This suggests that simple nitrile and epithionitrile formation share regulatory loci, but that there are likely separate proteins producing nitrile and epithionitrile structures from allyl glucosinolate. This is in agreement with previous observations suggesting the presence of an unidentified simple nitrile forming enzyme in Arabidopsis [23,25,32].

Conclusion

This study shows that there is considerable natural genetic variation controlling the age dependent regulation of structural
outcomes in Arabidopsis [32]. Elucidating the basis of this regulation is necessary to obtain a better understanding of the evolution and ecological significance of developmental trajectories in this important plant defense system. Further, we describe epistatic networks that appear to link regulatory loci with the genes that they regulate. Future analyses will be required to test if quantitative epistasis can be used to generate networks in a fashion similar to Mendelian epistasis but this has potential applications in most species. Finally, the potential for whole genome duplications to be associated with multiple QTLs for the same trait may help to enhance the rate at which additional QTL can be cloned. Once one QTL is cloned for a given trait, it may immediately suggest candidate genes for QTL in genomic regions that share an ancestry through whole genome duplications. The glucosinolate system is a useful model system for quantitative genetics to begin addressing these fundamental issues in quantitative genetics, ecology and evolution.

Materials and Methods

Mapping Population

The population of 411 Bay-0 × Sha RILs [33] were chosen for QTL mapping analysis of the structural outcome of glucosinolate activation. The parents of this population differ in their glucosinolate profile and content, as well as in the structures formed upon glucosinolate activation and the developmental regulation of the structural outcome following glucosinolate activation [32,38]. A subset of 212 lines from this population have also been analyzed for variation in gene expression [37], enabling comparison of gene expression to phenotypic variation. Finally, there are available HIFs, pairs of near isogenic lines fixed for alternate alleles at a single locus in an otherwise identical recombinant inbred background, which offer the opportunity to validate some of the QTLs detected in this study [54].

RIL Growth Conditions and Experimental Design

Seeds were imbibed and cold stratified at 4 degrees for three days to break dormancy. All seeds were sown directly onto Premier ProMix B potting soil (Premier Brands, Inc., Red Hill, Pennsylvania) in 36-cell (approximately 125 cm³ soil per cell) flats, and grown in controlled environment chambers at 20°C with 8 h light at 100–120 μE. Each flat contained one Bay-0 and Sha parents. All plants were free of insect pests by visual inspection. The population was grown three independent times to independently phenotype the parents and all RILs. Further, day 42 is one week away from the attainment of Stage 1.10 or 10 mature leaves per plant for both the parents and all RILs. Further, day 42 is one week away from the earliest RIL flowering (Stage 5.10) in our environmental conditions [32,34,35]. Thus, this range of DPG allows us to focus on rosette age rather than leaf age although the two may be intricately linked in some fashion. Exogenous allyl glucosinolate was added to enable comparisons of structural outcomes using a common substrate for all RILs despite the segregating biosynthetic variation [38]. Further, the allyl glucosinolate allows us to measure all three potential glucosinolate activation endpoints, isothiocyanate, epithionitrile or simple nitrile, whereas half the RILs do not have this capacity due to the lack of alkenyl glucosinolates [32,38]. Upon complete tissue homogenization the reaction vial was capped and incubated for five minutes. The reaction was stopped and glucosinolate activation products extracted with 4 mL of dichloromethane. The organic phase was removed, dried and concentrated to 200 μL for gas chromatography (GC) analysis using an Agilent HP 5890 with a flame ionization detector [22].

Peak identities were confirmed using a GC-mass spectral detector (Agilent HP 6890 with an Agilent 5973N MSD), by comparison with published mass spectra [55]. Quantification was conducted using published response factors that were corrected using propyl isothiocyanate standards as previously described [22]. Structural outcomes are reported as the percent of simple nitrile, epithionitrile, or isothiocyanate products for a particular glucosinolate. For instance, the percent simple nitrile for allyl glucosinolate is defined as [allyl simple nitrile] / [allyl simple nitrile + allyl epithionitrile + allyl isothiocyanate]. Proper chemical names for this equation are allyl simple nitrile is 3-butenyl nitrile and allyl epithionitrile is 2,4-epithiobutyl nitrile. Each structural outcome for each glucosinolate was measured using a similar equation. By dividing the absolute amount of a particular structure by the sum of all possible products, the effects of myrosinase activity and differences in biosynthesis and accumulation of the endogenous substrates are cancelled, since they affect both the numerator and denominator equally. This assay is not a quantitative measure of total myrosinase activity because it reaches saturation for some samples.

QTL Analysis

We obtained genotypes and genetic map information for the Bay-0 × Sha RIL population from the Arabidopsis Biological Resource Center [ABRC; www.arabidopsis.org] [33]. To maximize our ability to detect QTLs, we utilized the data from each DPG experiment separately and as a combined data set. For each RIL, the proportion of each activation structure for each glucosinolate were independently used for QTL mapping within Windows QTL Cartographer v2.5 [56–58]. Although the proportions of glucosinolate activation structures obtained from a given substrate are not mathematically independent of one another, the simple nitrile and epithionitrile rearrangements can be separately measured for allyl glucosinolate, allowing simultaneous assessment of these partially independent processes [32]. Composite interval mapping (CIM) was implemented using Zmap (Model 6) with a 10 cM window and an interval mapping increment of 2 cM. The declaration of statistically significant QTL is based on permutation derived empirical thresholds using 1,000 permutations for each trait mapped [59,60]. The Eqtl module of QTL Cartographer was used to automatically identify the location of each significant QTL for each trait [58].

To further test each QTL identified and query for potential epistasis, we conducted an ANOVA for the proportion of each glucosinolate activation structure. The markers most closely linked
to each significant main-effect QTL were used as main effect cofactors. An automated SAS script then tested all main effects and all possible pair wise interactions between main-effect loci. Significance values were corrected for multiple testing within a model using false discovery rate adjustment within the automated script. The script returned all significance values as well as QTL main-effect estimates in terms of allelic substitution values (Table S3). In addition, the combined data were used to estimate the heritability of the different structural outcomes of glucosinolate activation. This was conducted using the general linear model procedure within SAS where broad sense heritability was defined as $\sigma_g^2 / \sigma_p^2$ (Table 1), where $\sigma_g$ is the estimated genetic variance for the structural proportion phenotypes among different genotypes in these RILs, and $\sigma_p$ is the estimated phenotypic variance [3].

### Analysis of HIFs for QTL Validation

To confirm the identified QTLs in this study we obtained sixteen HIFs, corresponding to nine of the loci detected in this experiment, from INRA [http://dbsgap.versailles.inra.fr/portail] (Table S1). There were no HIFs available to test GSL.Activ.IV.16 and GSL.Activ.II.61. Within each HIF, only the genotypes in the region of one QTL differ while the rest of the genome is random.

For most QTLs there was a HIF available with functional ESP, except for GSL.Activ.II.13 and GSL.Activ.II.64. To test simple nitrile proportion QTLs for dependence on ESP genotype, separate HIFs with the functional Sha and non-functional Bay-0 alleles of ESP were chosen when possible.

Each HIF was planted with twelve independent biological replicates per allele per HIF. These were planted and grown under identical conditions as described above for the RIL population. For each HIF, six replicates of each genotype were assayed for structural outcomes at 24 DPG and six were assayed at 38 DPG to validate each QTL and survey for age dependence. These time points were chosen such that there was a sufficient developmental time difference to detect genotype×DPG effects but before epithionitrile formation reached saturation at the later time points. Due to poor germination, HIF191, 244, 338, and 364 were only assayed at 38 DPG.

The data for each class of glucosinolate activation product were analyzed for the effects of genotype, DPG, and DPG×genotype within each HIF using the general linear model procedure in SAS. Given that each HIF is a separate and independent test, we did not correct for multiple testing within these models. We also directly compared HIF241 to HIF425, two independent HIFs differing at the ESP QTL, to assess background dependent effects upon ESP.

### Analysis of Gene Expression QTLs

We used previously published sequence data to identify the known myrosinases and structural specifier genes [22–24,61]. We utilized protein sequence data to identify Arabidopsis homologs of each major glucosinolate activation gene, myrosinase, ESP, ESM1, and Myrosinase Binding Protein 1 (MBP1) and MBP2. For all four gene families we had two criteria to define a gene as potentially associated with glucosinolate activation. First, each included gene had to be similar to known genes based on a BLASTP score of at least e$^{−45}$. Secondly, we further restricted this list to genes that were phylogenetically limited to Arabidopsis when protein sequences from the poplar, grape and rice plant genomes were included [62,63]. This assumes that genes with more similar, non-cruferous homologues are unlikely to be involved in glucosinolate activation as this system is not present in poplar, grape or rice (Table S2). Heritability, eQTL position, eQTL effect and transcript accumulation values were obtained from a previously published analysis of the Bay-0×Sha population [37]. Because these global transcription studies were conducted in the same mapping population grown under the same conditions and in the same growth chambers, it was possible to directly compare the gene expression and structural outcome data.

### Accession Numbers

There are no new accession numbers associated with this dataset. The microarray data set used in this study has been deposited at EBI ArrayExpress [http://www.ebi.ac.uk/arrayexpress/] under numbers E-TABM-126 and E-TABM-224.
Acknowledgments

We also thank Heather Rowe (Genetics Graduate Group, UC Davis) for critical review of the manuscript and plant growth as well as Dr. Joshua Chang Mell (Genetics Graduate Group, UC Davis) for numerous helpful discussions. We are extremely grateful to Dr. Andrew Waterhouse, Mauri Anderson, and Dr. Susan Ebeler (Department of Viticulture and Enology, UC Davis) for the use of their GCs.

Author Contributions

Conceived and designed the experiments: AMW DJK. Performed the experiments: AMW. Analyzed the data: AMW DJK. Contributed reagents/materials/analysis tools: AMW IB ZZ. Wrote the paper: AMW DJK.

References

1. Lynch M, Walsh B (1998) Genetics and analysis of quantitative traits. Sunderland, Massachusetts: Sinauer Associates, Inc.
2. Mackay TFC (2001) The genetic architecture of quantitative traits. Annu Rev Genet 35: 303–339.
3. Falconer DS, Mackay TFC (1996) Introduction to Quantitative Genetics. Essex: Longman, Harlow.
4. Falconer DS, Mackay TFC (1996) Introduction to Quantitative Genetics. Essex: Longman, Harlow.
5. Holland JB (2007) Genetic architecture of complex traits in plants. Curr Opin Plant Biol 10: 156–161.
6. Melchinger AE, Utz HF, Schon CC (1998) Quantitative trait locus (QTL) mapping using different testers and independent population samples in maize reveals low power of QTL detection and large bias in estimates of QTL effects. Genetics 149: 383–403.
7. Stich B, Yu JM, Melchinger AE, Pfeifer HP, Utz HF, et al. (2007) Power to detect higher-order epistatic interactions in a metabolic pathway using a new mapping strategy. Genetics 176: 563–570.
8. Visker M, Keizer LCP, Van Eck HJ, Jacobsen E, Colon LT, et al. (2003) Can the QTL for late blight resistance on potato chromosome 5 be attributed to a fungal mating type? Theor Appl Genet 106: 317–325.
9. Clarke JD, Aarts N, Feys BJ, Dong NX, Parker JE (2001) Constitutive disease resistance requires EDS1 in the Arabidopsis mutants ep and epf and is partially EDS1-dependent in cpr5. Plant J 26: 409–420.
10. Zhao JW, Meng JH (2003) Genetic analysis of loci associated with partial resistance to Sclerotinia sclerotiorum in rapeseed (Brassica napus L.). Theor Appl Genet 106: 759–764.
11. Ahmadi N, Albar L, Pressoir G, Pinel A, Fargette D, et al. (2003) Genetic basis and mapping of the resistance to Rice yellow mottle virus III. Analysis of QTL efficiency in introgressed progenies confirmed the hypothesis of complementary epistasis between two resistance QTLs. Theor Appl Genet 103: 1091–1092.
12. Calenge F, Drouet D, Denance C, Van de Weg WE, Brisset MN, et al. (2005) Identification of a major QTL together with several minor additive or epistatic QTLs for resistance to fire blight in apple in two related progeny. Theor Appl Genet 111: 128–135.
13. Croaker GL, Francis DM (2004) Mapping, genetic effects, and epistatic interaction of two bacterial canker resistance QTLs from Erwinia herbicola. Theor Appl Genet 108: 1047–1055.
14. Thabuis A, Follain A, Servin B, Daubeze AM, Phaly T, et al. (2005) QTL analysis and mapping strategy. Genetics 176: 563–570.
15. Thabuis A, Lelebre V, Bernard G, Daubeze AM, Phaly T, et al. (2004) Phenotypic and molecular evaluation of a recurrent selection program for a polygenic resistance to Phytophthora capsici in pepper. Theor Appl Genet 109: 392–401.
16. Martin GB, Brommonschinkel SH, Chunvongse J, Frary A, Gianul MW, et al. (1993) Map-based cloning of a protein-kinase gene conferring disease resistance in tomato. Science 262: 1432–1436.
17. Melchinger AE, Utz HF, Schon CC (1998) Quantitative trait locus (QTL) mapping using different testers and independent population samples in maize reveals low power of QTL detection and large bias in estimates of QTL effects. Genetics 149: 383–403.
18. Malmberg RL, Held S, Waits A, Mauricio R (2005) Epistasis for fitness-related traits in barley. Plant J 43: 201–210.
19. Malmberg RL, Mauricio R (2005) QTL-based evidence for the role of epistasis in Arabidopsis. Theor Appl Genet 111: 128–135.
20. Malmberg RL, Mauricio R (2005) QTL-based evidence for the role of epistasis in Arabidopsis. Theor Appl Genet 111: 128–135.
21. Malmberg RL, Mauricio R (2005) QTL-based evidence for the role of epistasis in Arabidopsis. Theor Appl Genet 111: 128–135.
22. Lambrix V, Reichelt M, Mitchell-Olds T, Kliebenstein D, Gershenzon J (2001) The Arabidopsis epithiospecifier protein promotes the hydrolysis of glucosinolates to nitriles and influences Thapsigargin sensitivity. Plant Cell 13: 2793–2807.
23. Zhang Z-Y, Ober JA, Kliebenstein DJ (2006) The gene controlling the quantitative trait locus EPITHIOSPECIFIER MODIFIER alters glucosinolate hydrolysis and insect resistance in Arabidopsis. Plant Cell 18: 1524–1536.
24. Barc H, Jander G (2006) Arabidopsis myrosinases TG61 and TG62 have redundant function in glucosinolate breakdown and insect defense. Plant J 46: 549–562.
48. Charron CS, Saxton AM, Sams CE (2005) Relationship of climate and genotype to seasonal variation in the glucosinolate-myrosinase system. II. Myrosinase activity in ten cultivars of Brassica oleracea grown in fall and spring seasons. J Sci Food Agric 85: 682–690.

49. Walsh JB (1995) How Often Do Duplicated Genes Evolve New Functions. Genetics 139: 421–428.

50. Lynch M, Conery JS (2000) The evolutionary fate and consequences of duplicate genes. Science 290: 1151–1155.

51. Lynch M, Force A (2000) The probability of duplicate gene preservation by subfunctionalization. Genetics 154: 459–473.

52. Somorjai IML, Danzmann RG, Ferguson MM (2003) Distribution of temperature tolerance quantitative trait loci in Arctic char (Salvelinus alpinus) and inferred homologies in rainbow trout (Oncorhynchus mykiss). Genetics 165: 1443–1456.

53. Axelson T, Shavorskaya O, Lagercrantz U (2001) Multiple flowering time QTLs within several Brassica species could be the result of duplicated copies of one ancestral gene. Genome 44: 856–864.

54. Loudet O, Gaudon V, Trubuil A, Daniel-Vedele F (2005) Quantitative trait loci controlling root growth and architecture in Arabidopsis thaliana confirmed by heterogeneous inbred family. Theor Appl Genet 110: 742–753.

55. Spencer GF, Daxenbichler ME (1988) Gas chromatography–mass spectrometry of nitriles, isothiocyanates and oxazolidinethiones derived from cruciferous glucosinolates. J Sci Food Agric 31: 359–367.

56. Zeng Z-B, Kao C-H, Baten CJ (1999) Estimating the genetic architecture of quantitative traits. Genet Res 73: 345–355.

57. Baten CJ, Weir BS, Zeng Z-B (1999) QTL Cartographer, Version 1.13 Department of Statistics, North Carolina State University: Raleigh, N.C.

58. Wang S, Baten CJ, Zeng Z-B (2006) Windows QTL Cartographer 2.5 Department of Statistics, North Carolina State University: Raleigh, NC.

59. Churchill GA, Doerge RW (1994) Empirical Threshold Values For Quantitative Trait Mapping. Genetics 138: 906–917.

60. Doerge RW, Churchill GA (1996) Permutation tests for multiple loci affecting a quantitative character. Genetics 142: 285–294.

61. Takechi K, Sakamoto W, Usagi S, Murata M, Motoyoshi F (1999) Characterization of a flower-specific gene encoding a putative myrosinase binding protein in Arabidopsis thaliana. Plant Cell Physiol 40: 1287–1296.

62. Sonderby IE, Hansen BG, Bjarnholt N, Ticoni C, Halkier BA, et al. (2007) A systems biology approach identifies a R2R3 MYB gene subfamily with distinct and overlapping functions in regulation of aliphatic glucosinolates. PLoS ONE 2: e1322.

63. Hansen BG, Klebentnert DJ, Halkier BA (2007) Identification of a flavin-dependent oxygenase as the S-oxygenating enzyme in aliphatic glucosinolate biosynthesis in Arabidopsis. Plant J 50: 902–910.