α-amatin resistance in Drosophila melanogaster: A genome-wide association approach

Chelsea L. Mitchell¹, Catrina E. Latuszek¹, Kara R. Vogel², Ian M. Greenlund¹, Rebecca E. Hobmeier³, Olivia K. Ingram¹, Shannon R. Dufek¹, Jared L. Pecore¹, Felicia R. Nip³, Zachary J. Johnson⁴, Xiaohui Ji⁵, Hairong Wei⁵, Oliver Gailing⁵, Thomas Werner¹*

¹ Department of Biological Sciences, Michigan Technological University, 1400 Townsend Dr., Houghton, MI, United States of America, ² Department of Neurology, University of Wisconsin School of Medicine and Public Health, 1300 University Ave., Madison, WI, United States of America, ³ College of Human Medicine, Michigan State University, Clinical Center, East Lansing, MI, United States of America, ⁴ U.S. Forest Service, Salt Lake Ranger District 6944 S, 3000 E, Salt Lake City, UT, United States of America, ⁵ School of Forest Resources and Environmental Sciences, Michigan Technological University, 1400 Townsend Dr., Houghton, MI, United States of America

* twerner@mtu.edu

Abstract

We investigated the mechanisms of mushroom toxin resistance in the Drosophila Genetic Reference Panel (DGRP) fly lines, using genome-wide association studies (GWAS). While Drosophila melanogaster avoids mushrooms in nature, some lines are surprisingly resistant to α-amatin—a toxin found solely in mushrooms. This resistance may represent a pre-adaptation, which might enable this species to invade the mushroom niche in the future.

Although our previous microarray study had strongly suggested that pesticide-metabolizing detoxification genes confer α-amatin resistance in a Taiwanese D. melanogaster line Ama-KTT, none of the traditional detoxification genes were among the top candidate genes resulting from the GWAS in the current study. Instead, we identified Megalin, Tequila, and widerborst as candidate genes underlying the α-amatin resistance phenotype in the North American DGRP lines, all three of which are connected to the Target of Rapamycin (TOR) pathway. Both widerborst and Tequila are upstream regulators of TOR, and TOR is a key regulator of autophagy and Megalin-mediated endocytosis. We suggest that endocytosis and autophagy of α-amatin, followed by lysosomal degradation of the toxin, is one of the mechanisms that confer α-amatin resistance in the DGRP lines.

Introduction

The mechanisms that allow some drosophilid species to breed in toxic mushrooms have remained largely elusive for the past half century, despite a series of studies attempting to reveal the genetic basis of this phenomenon [1–8]. α-Amanitin is the principal toxin of several deadly poisonous mushroom species, including the death cap (Amanita phalloides) and destroying angel (Amanita bisporigera, A. ocreata, and A. virosa) [9, 10]. This toxin inhibits the function of RNA polymerase II, halting all eukaryotic messenger RNA production [11]. Among eukaryotes, only a few species,
such as *Drosophila guttifera*, *D. falleni* and *D. recens* [1, 3, 5, 8, 12], are adapted to breeding in and feeding on these very poisonous mushrooms. Thus, there is virtually no competition for this food source [3, 5, 8, 12], and breeding in these mushroom species provides protection against parasitic nematodes, which are susceptible to mushroom toxins [13, 14].

Typically, *D. melanogaster* oviposits in and feeds on fermenting fruit and strictly avoids mushrooms. Although α-amanitin is solely produced by mushrooms, a few lines of *D. melanogaster* were shown to be up to two orders of magnitude more resistant to α-amanitin than susceptible lines of this species. Resistant lines can tolerate up to ~ 10 μg of α-amanitin per gram of food, as compared to ~ 0.1 μg/g in very susceptible lines [2, 7, 15, 16]. In contrast, mycophagous species develop in mushrooms containing up to 1200 μg of total α-amanitin [17] and are deleteriously affected by α-amanitin in the range of 750 to 1000 μg/g [5]. For comparison, the human LC50 is 0.1 μg/g body weight, which can make the consumption of a single mushroom lethal [17]. Because *D. melanogaster* has no known history of mushroom-feeding in the wild, and the resistance is insufficient to safely consume highly toxic mushrooms, α-amanitin resistance may be a pre-adaptation for a possible mushroom niche invasion in the future.

In our previous microarray study [15], we identified four possible molecular mechanisms for α-amanitin resistance in a *D. melanogaster* line from Taiwan (Ama-KTT, collected in 1968 in Kenting): 1) blockage by cuticular proteins, 2) detoxification by phase I and II detoxification enzymes: Cytochrome P450s (P450s), Glutathione-S-transferases (GSTs), and UDP glucuronosyl transferases (UGTs), 3) cytoplasmic sequestration in lipid particles, and 4) cleavage by peptidases. The top three candidate genes resulting from that study, *Cyp6a2*, *Cyp12d1-d*, and *Cyp12d1-p*, have all been shown to detoxify pesticides, such as dichlorodiphenyltrichloroethane (DDT), imidacloprid, dicyclanil, and atrazine [18–22]. Thus, pesticides may have originally triggered the evolution of α-amanitin resistance in this Taiwanese fly line.

Here, lines from the *Drosophila* Genetic Reference Panel (DGRP) [23] were tested to determine whether α-amanitin resistance is a variable trait in a North American population of *D. melanogaster* and to identify the candidate genes/mechanisms that protect these lines from α-amanitin. We measured the phenotypes as the fraction of larvae emerging as adults on media supplemented with different α-amanitin concentrations. We performed four tests: 1) a GWAS for 180 DGRP lines using a low toxin concentration (0.2 μg/g α-amanitin/food), 2) a GWAS for 180 DGRP lines using a relatively high toxin concentration (2.0 μg/g α-amanitin/food), 3) a GWAS using lethal concentration 50 (LC50) values obtained for 37 DGRP lines, for which RNA-seq data exist, and 4) a correlation analysis of larval resistance with adult transcriptome data among the 37 DGRP lines. We show that α-amanitin resistance is a common and variable trait among all tested lines. We identified three genes, *Megalim* (*mgl*), *Tequila* (*teq*), and *widebhorst* (*wdb*) that may interact with the Target of Rapamycin (TOR) pathway to mediate α-amanitin resistance through endocytic and autophagic sequestration and degradation.

**Results and discussion**

**α-amanitin resistance variation and trait definition**

We first tested if the *D. melanogaster* DGRP lines varied in their resistance to α-amanitin, using two α-amanitin concentrations (0.2 μg/g and 2.0 μg/g) that we predicted to allow for an evaluation of the resistance variation among all DGRP lines. We placed ten freshly hatched first-instar larvae per fly line and toxin concentration into vials and performed three replicates for a total of 30 larvae per line and concentration. We defined the phenotypic value for this experiment as the average number of adult flies emerging from the three replicate vials, ranging from 0–10 flies. The data are shown in Fig 1A and 1B. Out of the 189 DGRP lines initially tested, twelve lines displayed relatively high resistance to α-amanitin on 2.0 μg/g, 83 showed...
intermediate resistance, 91 showed virtually no resistance on 2.0 μg/g, while three lines did not produce enough eggs for the experiments. Based on these data (S1 Table), we concluded that α-amanitin resistance in the DGRP lines is a quantitative trait amenable to GWAS.

Following this experiment, we calculated the LC₅₀ values of a 37-line subset of these 180 lines. Because α-amanitin is very expensive and LC₅₀ measurements are extremely labor-intensive and time-consuming, we limited the determination of the LC₅₀ values to those 37 DGRP lines, for which adult transcriptome data are available [24] to conduct later LC₅₀/gene expression correlation tests. To calculate the LC₅₀ values, we tested each of the 37 lines on seven α-amanitin concentrations, including controls lacking α-amanitin. For each line and concentration, ten internal replicates of ten first-instar larvae per feeding tube were performed, and each experiment was performed three times. Thus, we tested a total of 300 larvae per concentration and fly line, as compared to only 30 larvae per concentration in the 180-line experiment on 0.2 μg/g.

**Fig 1. Larval viability variation in the DGRP lines in response to α-amanitin.** The y-axis shows individual viability values, while the x-axis represents the individual DGRP lines. The lines are sorted from lowest α-amanitin resistance (left) to highest α-amanitin resistance (right). The error bars represent the standard error of the mean (SEM). A) 180 lines tested on 0.2 μg/g α-amanitin. (Individual line numbers are not shown but can be found in S1 Table). The y-axis represents the average number of flies hatched from 10 larvae placed on toxic food. B) 180 lines tested on 2.0 μg/g α-amanitin. (Individual line numbers are not shown but can be found in S1 Table). The y-axis represents the average hatch counts out of 10 larvae placed on toxic food. C). The y-axis represents the LC₅₀ values of the 37-line subset. The line numbers are shown on the x-axis.
and 2.0 μg/g. The resulting LC$_{50}$ values were defined as the phenotypic trait for this experiment and are shown in Fig 1C and S2 Table.

We note that among the 37-line subset, the resistance data from the 2 μg/g α-amanitin treatment and the LC$_{50}$ experiment were only weakly correlated ($r = 0.24$, $p = 0.075$, linear regression analysis) (S3 Table). This low correlation was caused by at least two factors. First, the 2 μg/g α-amanitin experiment data had a 10X lower sample size than the LC$_{50}$ calculation, as mentioned earlier. Second, the two lines with the highest LC$_{50}$ values (lines DGRP-307 and 315) did not show a very high viability in the 2 μg/g α-amanitin experiment. Although the initial association between both data sets was weak, it increased when these two lines with very high LC$_{50}$ values were excluded from the comparison ($r = 0.39$, $p = 0.02$) (S4 Table). Among the 37 lines, the resistance data at 0.2 μg/g α-amanitin were not associated with the LC$_{50}$ values ($r = 0.17$, $p = 0.31$), and the association remained non-significant after exclusion of the two lines with very high LC$_{50}$ values ($r = 0.32$, $p = 0.06$). Furthermore, the resistance data among all 180 lines at 2 μg/g α-amanitin and 0.2 μg/g α-amanitin revealed a weak and barely significant correlation ($r = 0.13$, $p = 0.037$) (S3 Table). In order to test to what degree viability associated with the α-amanitin resistance phenotype is genetically determined, we calculated the broad-sense heritability values. They were 0.95 for the 37-line LC$_{50}$ data, 0.46 for the 180-line 2.0 μg/g data, and 0.32 for the 180-line 0.2 μg/g data. Thus, the 180-line 0.2 μg/g data contain the greatest level of random mortality that is not associated with the α-amanitin-resistance phenotype.

The 180-line GWAS on 2.0 μg/g versus the 37-line GWAS

After establishing that α-amanitin resistance varies among the DGRP lines, we next performed three separate GWAS to identify candidate genes underlying α-amanitin resistance in the DGRP lines: 1) a 37-line GWAS using the LC$_{50}$ values, 2) a 180-line GWAS on 2.0 μg/g α-amanitin, and 3) a 180-line GWAS on 0.2 μg/g α-amanitin. The phenotypic values associated with each line were submitted to the DGRP website at dgrp2.gnets.ncsu.edu for analysis. The Manhattan plots for all three GWAS are shown in Fig 2. The results for the 37-line GWAS are provided in S5 Table, the results for the 180-line GWAS on 2.0 μg/g α-amanitin are listed in S6 Table, and the 180-line GWAS results on 0.2 μg/g α-amanitin can be found in S7 Table.

Because the 180-line data on 0.2 μg/g showed the lowest heritability value, we will first discuss the 37-line GWAS results using LC$_{50}$ values and the 180-line GWAS on 2.0 μg/g. Table 1 summarizes the top 11 candidate genes from S5 and S6 Tables, which fulfilled at least one of three requirements: 1) the single mixed p-value was below the Bonferroni p-value cutoff of $< 2.60E-08$ ("strict"), 2) the single mixed p-value was above but still close to the Bonferroni cutoff value ("relaxed", $< 1.00E-05$), and/or 3) the candidate genes were identified in both GWAS datasets, for one of which the smallest p-value was at least "relaxed" ($< 1.00E-05$).

The 180-line GWAS on 0.2 μg/g resulted in one gene with a very low p-value. This gene, pHCl, showed a single-mixed p-value of 5.60E-08 (FDR (BH) $p = 0.05423$)). It encodes a neurotransmitter-gated ion channel transmembrane protein [25]. However, none of the genes identified in the 180-line GWAS on 0.2 μg/g had known functions that link them to possible toxin resistance mechanisms (S7 Table). As discussed earlier, the reason for this lack of suitable candidate genes may be random mortality at the larval stage that is not associated with the α-amanitin resistance phenotype. We note that some DGRP lines, including resistant ones, showed poor viability on the low 0.2 μg/g toxin concentration, while they survived better on higher α-amanitin concentrations. Although the reason for this paradox is unknown, it has been observed in two unrelated α-amanitin-resistant lines of D. melanogaster [2, 3], as well as in several mycophagous species [3, 8].
Discussion of the top eleven candidate genes

Megalin (*mgl*) was the most significant candidate gene. We identified this gene in both the 180-line GWAS on 2.0 μg/g α-amanitin and the 37-line GWAS. In the 180-line GWAS on 2.0 μg/g, *mgl* was represented by one intronic single nucleotide polymorphism (SNP) (X\_9293649\_SNP) with a "strict" p-value of p = 4.26E-09. Additionally, we found two "suggestive" intronic *mgl* SNPs in the 37-line GWAS (X\_9355099\_SNP and X\_9355132\_SNP, p = 9.02E-04 each). We speculate that intronic SNPs may alter cis-regulatory modules, and that higher *mgl* gene expression may confer higher resistance to α-amanitin. *Megalin* encodes a conserved endocytosis and transcytosis receptor found in the apical region of many epithelial cell types in both

---

Fig 2. Manhattan plots for the three GWAS. A) 37-line GWAS using LC\(_{50}\) values, B) 180-line GWAS on 2.0 μg/g α-amanitin, C) 180-line GWAS on 0.2 μg/g α-amanitin. Selected significant gene names are printed on the top right of the corresponding dots in the graphs.

doi:10.1371/journal.pone.0173162.g002
vertebrates and invertebrates [26–31]. *Megalin* is a member of the low-density lipoprotein receptor (LDLR) family and has more than 40 different protein ligands (including toxic substances [32]), which are degraded in the lysosome upon receptor binding and endocytosis [26, 27, 33]. Endocytosis is a process known to be regulated in the epithelium and fat body of *Drosophila* by TOR [34]. Furthermore, *Megalin* endocytosis is induced by the Peroxisome Proliferator-Activated Receptor (PPAR), a protein with the capacity to increase autophagy (also see *wdb*, below) and a repressor of TOR [35]. If *Megalin* confers α-amanitin resistance, as suggested by our GWAS results, a possible scenario would be the uptake/sequestration of α-amanitin by the *Megalin* protein and the subsequent degradation by endocytic processes up-regulated in the midgut epithelium, where the food is absorbed in *Drosophila* larvae.

The *Tequila* (*teq*) gene was represented by five intronic SNPs and indels with "relaxed" p-values in the 180-line GWAS on 2.0 μg/g: 3L_9069423_SNP (p = 6.91E-08), 3L_9069400_SNP (p = 7.15E-08), 3L_9069410_INS (p = 7.15E-08), 3L_9069403_SNP (p = 7.38E-08), and 3L_9069415_SNP (p = 7.71E-08). Interestingly, *teq* activation by insulin signaling has been shown to increase Akt activity, a key modulator of TOR signaling that can activate TOR complex 1 (TORC1) [36]. We hypothesize that the intronic SNPs in the *teq* locus cause a decrease in *teq* gene expression, which

### Table 1. Candidate genes resulting from the 180-line GWAS on 2.0 μg/g and the 37-line GWAS.

| Candidate gene symbol | 180-line GWAS on 2.0 μg/g: lowest single mixed p-value | 37-line GWAS with LC50 values: lowest single mixed p-value | 180-line GWAS on 2.0 μg/g: lowest FDR (BH) p-value | 37-line GWAS with LC50 values: lowest FDR (BH) p-value | Cyto. map location/ chromosome arm | Selected known or predicted functions, as per FlyBase [25] |
|-----------------------|------------------------------------------------------|-----------------------------------------------------|---------------------------------------------------|---------------------------------------------------|-----------------------------------|------------------------------------------------------------|
| mgl                   | 4.26E-09 ***                                        | 9.02E-04 *                                         | 0.00734                                           | 0.41622                                           | 8D10 / X                          | LDL receptor class A, chitin-based cuticle development, endocytic receptor in epithelial cells |
| CG13611               | 7.66E-09 ***                                        | N/A                                                | 0.00734                                           | N/A                                              | 95F8-95F9 / 3R                    | Oxidoreductase activity                                               |
| NnaD                  | 3.45E-08 **                                         | N/A                                                | 0.01478                                           | N/A                                              | 12B4 / X                          | Peptidase M14, carboxypeptidase A                                      |
| teq                   | 6.91E-08 **                                         | N/A                                                | 0.01478                                           | N/A                                              | 66F4-66F5 / 3L                    | LDL receptor class A, scavenger receptor activity; chitin binding; serine-type endopeptidase activity |
| Scr                   | 2.30E-06 **                                         | 3.54E-06 **                                         | 0.06678                                           | 0.24164                                           | 84A5 / 3R                         | Midgut development, transcription factor activity                     |
| Pli                   | 1.99E-05 *                                          | 3.98E-07 **                                         | 0.13086                                           | 0.09639                                           | 95C5-95CB / 3R                    | Immune response                                                           |
| Rbp6                  | 1.69E-04 *                                          | 3.82E-06 **                                         | 0.28190                                           | 0.13828                                           | 73E5-74A1 / 3L                    | mRNA-binding, stem cell development                                   |
| Hs6st                 | 5.39E-05 *                                          | 4.71E-06 **                                         | 0.18521                                           | 0.13828                                           | 92B8-92C1 / 3R                    | Sulfotransferase activity                                                |
| Ets65A                | 7.47E-04 *                                          | 6.65E-06 **                                         | 0.45526                                           | 0.13828                                           | 65A6 / 3L                         | Regulation of transcription                                             |
| wdb                   | 4.59E-04 *                                          | 7.20E-06 **                                         | 0.39087                                           | 0.13828                                           | 98A6-98A8 / 3R                    | Protein phosphatase 2A                                                  |
| siti                  | 5.40E-04 *                                          | 7.27E-06 **                                         | 0.40926                                           | 0.13828                                           | 69C4 / 3L                         | Threonine kinase activity                                               |

The single mixed p-values are.

*** = strict (below the Bonferroni cutoff value of 2.60E-08)

** = relaxed (between the Bonferroni cutoff value and 1.00E-05), or

* = suggestive (between 1.00E-04 and 9.99E-04).

N/A = these SNPs do not segregate in the 37-line GWAS. We also provide a multiple test correction from the entire set of 1.9 million GWAS tests for each phenotype and calculated the False Discovery Rate from the single mixed p-values: FDR (BH) = Benjamini Hochberg False Discovery Rate (bold values are significant).

doi:10.1371/journal.pone.0173162.t001
would have a repressing effect on TOR. Previous studies in Drosophila have associated teq silencing with increased lifespan [36] and long-term memory formation [37]. Our previous work also correlated α-amanitin resistance to increased lifespan and reduction of body size; both could potentially be explained by reduced TOR activity [16] (Fig 3). We note that both Megalin and Tequila had highly significant FDR (BH) values of 0.00734 and 0.01478, respectively. The widerborst (wdb) gene was represented by four intronic SNPs with "relaxed" p-values in our 37-line GWAS results: 3R_23395889_SNP, 3R_23395891_SNP, 3R_23395908_SNP (p = 7.20E-06 each), and 3R_23395900_SNP (p = 8.24E-06). Also, several SNPs of mostly intronic nature were identified in this GWAS with "suggestive" p-values. Additionally, we identified one intronic SNP with a "suggestive" p-value in 180-line GWAS on 2.0 μg/g. Widerborst is a key regulator of the catabolic process of autophagy and has also been shown to have a lifespan-increasing effect in Drosophila [38]. Current knowledge indicates that common machinery contributes to the regulation and mechanisms involved in degradation by the endosomal and autophagic pathways [28]. For example, TORC1 regulation and the early and recycling endosomes of the endocytic pathway contribute to phagophore formation and maturation of the phagolysosome [28] (Fig 3). The Ser/Thr Phosphatase Protein Phosphatase 2A (PP2A) regulatory subunit Widerborst inhibits signaling upstream of TOR via the heterotrimeric holoenzyme PP2A-A/Wdb/PP2A-C complex (Fig 3). The Widerborst protein acts upstream to inhibit TOR via a genetic interaction with PtdIns3K/PTEN/Akt [39]. Thus, our results and previous studies cumulatively suggest that a TOR-inhibiting mechanism is involved in α-amanitin resistance. Ets at 65A (Ets65A) is a transcription factor gene of largely unknown function. Ets65A is situated at the same chromosomal position to which α-amanitin resistance has been mapped in two previous studies [2, 7], making this factor particularly interesting. Begun and Whitley (2000) [7] suggested that Mdr65 may be the α-amanitin resistance-conferring gene at that locus. In the 37-line GWAS, we identified two Ets65A SNPs with "relaxed" p-values: 3L_6097046 SNP (p = 6.65E-06, intronic) and 3L_6096774 SNP (p = 7.89E-06, synonymous coding). In the 180-line GWAS on 2.0 μg/g, we found one intronic SNP of Ets65A with a "suggestive" p-value. The Ets65A gene is expressed in adipose tissue [40] and associated with cold adaptation [41] and olfactory behavior [42] in D. melanogaster. It is worth noting that the TOR pathway not only responds to cellular energy cues, but has also been shown to play a critical role in heat shock and cold-induced stress responses [43]. However, Ets65A has not been shown to mediate any detoxification responses yet. Sex combs reduced (Scr) encodes a transcription factor involved in anterior midgut development in Drosophila embryos [44, 45]. We identified four intronic Scr SNPs with "relaxed" p-values in our 180-line GWAS on 2.0 μg/g: 3R_2658997 SNP (p = 2.30E-06), 3R_2668803 SNP (p = 3.54E-06), 3R_2672660 SNP (p = 7.12E-06), and 3R_2665283 SNP (p = 9.41E-06). The 37-line GWAS uncovered seven additional SNPs for Scr with "suggestive" p-values, five of which were present in the 3’ UTR, and the other two were located in introns. Because food absorption happens in the midgut, this organ is a strategic place for mushroom toxin resistance mechanisms to be deployed. CG13611 is a gene with only predicted functions. We identified it in the 180-line GWAS on 2.0 μg/g as the second of two genes that show at least one SNP with a "strict" p-value. CG13611 was represented by three SNPs: 3R_20106448 SNP (p = 7.66E-09, non-synonymous coding), 3R_20106547 SNP (p = 1.08E-06, non-synonymous coding), and 3R_20105165 SNP (p = 6.11E-05, upstream). The CG13611 protein is predicted to have oxidoreductase activity [25]. We have previously shown that oxidoreductase activity may be an important process to help detoxify α-amanitin [15]. It may thus be possible that CG13611 exerts a function similar or identical to phase I detoxification. The observed non-synonymous coding changes might alter
the enzyme activity, possibly leading to differences in the enzyme’s ability to chemically modify α-amanitin. We note that CG13611 had a highly significant FDR (BH) p-value of 0.00734. Pellino (Pli) was identified as the top candidate gene in the 37-line GWAS, represented by a single intronic SNP (3R_19707888_SNP, p = 3.98E-07). We further found five additional intronic SNPs in the 180-line GWAS on 2.0 μg/g with “suggestive” p-values. The Pli gene encodes an intracellular, positive regulator of the innate immune response in animals ranging from insects to man [46]. The Pellino protein regulates the Toll/Toll-like pathways, which respond to microbial antigens [47–49]. Interestingly, human Pellino-1 has been shown to confer resistance to two chemotherapeutic drugs in lung cancer cells [50]. Although it is not entirely clear how an innate immune pathway might lead to α-amanitin resistance in fruit flies, the innate immune system is capable of identifying and removing foreign substances from the body, and links between the innate immunity and autophagy have been described earlier [51, 52].

NnaD was represented by one SNP in the 180-line GWAS on 2.0 μg/g: X_13585381_SNP (p = 3.45E-08), which is located upstream of the NnaD gene. Its protein function is described as peptidase M14 and carboxypeptidase A [25]. The mouse orthologue protein Nna1 is known to hydrolyze polyglutamate-containing substrates in neurons [53]. Because α-amanitin is a

---

**Fig 3. The TOR pathway may be central to the evolution of α-amanitin resistance in the DGRP lines.** The schematic represents a suggested model for how α-amanitin resistance interplays with the TOR pathway via three of the top candidate genes suggested by our GWAS: Widerborst, Tequila, and Megalin. The proteins Widerborst and Tequila are known upstream regulators of TOR, influencing autophagy. TOR is a critical repressor of autophagy and Megalin-mediated endocytosis. Both the endocytic and autophagic catabolic processes end with the degradation and recycling of macromolecules in lysosomes. We hypothesize that in α-amanitin-resistant flies, Widerborst protein levels are up-regulated, while Tequila protein levels are down-regulated to collectively repress TOR, through heterotrimeric PP2A-A-Wbd-PP2A-C/TOR and Akt/TOR interactions, respectively. Megalin protein is hypothesized to be up-regulated and to sequester α-amanitin to the endosome. TOR inactivation may play a role in the elimination of cytoplasmic α-amanitin by de-repression of the autophagic process. As a result, the toxin would become entrapped in an elongating phagophore, and the autophagosome would then undergo lysosomal fusion, followed by degradation of α-amanitin. Down-regulation of TOR in α-amanitin-resistant flies is also consistent with the reduced thorax size (in the presence of α-amanitin) and increased longevity observed in α-amanitin resistant flies [16].
peptide, we speculate that NnaD may be involved in the proteolytic degradation of α-amanitin.

We do not currently offer an explanation for the possible roles of RNA-binding protein 6 (Rbp6), Heparan sulfate 6-O-sulfotransferase (Hs6st), and sticky (sti) in the detoxification of α-amanitin. All three genes showed SNPs with "relaxed" p-values in the 37-line GWAS and SNPs with "suggestive" p-values in the 180-line GWAS on 2.0 μg/g.

The Pearson correlation analysis: Comparing larval resistance to the adult transcriptome

We decided to focus on the 37-line subset because their adult transcriptomes were sequenced along with their genomes. We performed a Pearson correlation analysis, using our larval LC_{50} values and the publicly available RNA-seq data of untreated male and female adults. As a result, we identified the Translocator protein (Tspo) gene (S8 Table) with negative correlation coefficients in females of -0.72 (p = 6.35E-07) and males -0.67 (p = 2.00E-05). The female p-value was below the Bonferroni cut-off for this analysis (p = 4.95E-06), while the male correlation p-value was above the cutoff value. We also calculated the FDR-corrected p-values for the genes in this analysis and found 187 genes with corrected p-values < 0.05. We were unable to link the identified genes to toxin resistance. Tspo was the only gene resulting from the Pearson analysis that was also identified in one of our two GWAS. This gene was highly significant (corrected p = 0.00160 in females and = 0.01686 in males). In the 180-line GWAS on 2.0 μg/g, two SNPs with "suggestive" p-values were identified: one in the 3’ UTR and one forming a synonymous stop codon (AGG to AGA). Tspo encodes an outer mitochondrial membrane protein. Studies in Tspo mutants revealed important roles for this gene in lifespan determination, oxidative phosphorylation, positive regulation of apoptosis, control of ethanol-mediated behavior, as well as neurodegeneration [54, 55].

Different genes and mechanisms may underlie α-amanitin resistance in different populations and species

The rationale of using D. melanogaster to study mushroom toxin resistance is that D. melanogaster is currently our best genetic Drosophila model organism, and that there are no sequenced lines available for mycophagous species to allow for GWAS analyses. Studies in D. melanogaster have thus far suggested six possible mechanisms to evolve resistance to α-amanitin: 1) In a lab mutagenesis screen, an RNA-polymerase II mutant was identified that made the line 250-fold more resistant to α-amanitin than wild type [56]. 2) Our own laboratory has previously shown that the Taiwanese D. melanogaster line Ama-KTT [15] may have evolved resistance through 2) phase I detoxification (mediated by Cyp genes), 3) phase II detoxification (mediated by Gst and Ugt genes), 4) sequestration of α-amanitin in lipid particles, and 5) peptidase cleavage. Interestingly, the CYP450 inhibitor piperonyl butoxide (PBO) caused a dramatic reduction in α-amanitin resistance in the mycophagous species D. phalerata [8], suggesting that Cyp genes may play a role even in some mycophagous species. 6) In the current study, we show evidence that the TOR pathway may play a role in mediating the degradation of α-amanitin through endocytosis and autophagy. It is possible that mycophagous Drosophila species use some or a combination of the mechanisms that we have found in D. melanogaster to gain resistance to α-amanitin.

Why D. melanogaster has evolved α-amanitin resistance is not clear. At least in the case of the Taiwanese line Ama-KTT, pesticide exposure may have triggered the up-regulation of detoxification enzymes, which may display cross-resistance to α-amanitin [15]. We speculate that the evolution of α-amanitin resistance is a pre-adaptation, preparing the species to invade the mushroom niche in the future. Considering that D. melanogaster can complete its
development on non-toxic fungi in the laboratory, such as Baker’s yeast, a niche expansion to include other fungi seems plausible. Such a niche invasion may be aided by our previous observation that females of resistant strains show a higher fecundity when they grew up on food containing sublethal α-amanitin concentrations [16]. This increased fecundity might help the species to adapt to a diet of increasingly toxic mushrooms.

Limitations

Although it would be ideal to establish highly accurate LC$_{50}$ values for all 180 DGRP lines, it would be very time-consuming and take many resources to set up such an experiment. Working with α-amanitin requires a tradeoff between the number of fly lines used and the accuracy of the phenotypic values to measure. For our 180-line GWAS on 2.0 μg/g and on 0.2 μg/g, we had to accept a relatively low accuracy of the resistance measurement (30 larvae per line). In the 37-line GWAS, we were able to use highly accurate phenotypic data (2100 larvae per line to establish the LC$_{50}$ values). We note that the average fly hatch numbers of 3 X 10 larvae used as a trait for the 180-line GWAS on 2.0 and on 0.2 μg/g were not a great predictor of the LC$_{50}$ values among the 37 lines.

Although the main reason for using the 37-line subset of DGRP lines was to discover links between larval α-amanitin resistance and constitutive gene expression differences in adults, we did not identify many meaningful candidate genes by correlating our larval LC$_{50}$ values with the publicly available RNA-seq data of untreated male and female adults, which may have several explanations: 1) larval and adult resistance mechanisms differ from one another, 2) adults of lines that show larval resistance are not α-amanitin-resistant, 3) the resistance mechanism (s) cannot be detected at the transcriptome level in untreated adults, and/or 4) the low number of 37 lines may have limited the chance to identify highly significant associations.

Future research

Based on the known interplay of three of our eleven top candidate genes (Fig 3), the next line of experiments should focus on the mgl, teq, and wdb genes to test if changes in endocytosis, autophagy, and TOR signaling cause the observed variation in α-amanitin resistance. In order to obtain functional data, gene overexpression experiments, CRISPR-Cas9 knockout, and/or RNAi should be employed to manipulate susceptible lines to become more resistant and resistant lines to become less resistant than they were before the genetic manipulation.

Conclusions

Our data generated an interesting hypothesis that endocytosis and autophagy may mediate α-amanitin resistance in D. melanogaster, possibly through the involvement of the TOR pathway. We found that genes identified in our previous microarray study did not show genome-wide significance in the current study. Thus, α-amanitin resistance may have evolved independently in the Taiwanese and the North American populations, utilizing different genes and molecular mechanisms.

Materials and methods

Fly lines

We obtained all 189 D. melanogaster DGRP lines that were publicly available in April of 2012 from the Bloomington Drosophila Stock Center at Indiana University. The fly lines were maintained at room temperature at ~ 50% humidity on standard food containing cornmeal,
granulated sugar, Brewer’s yeast, agar, and methylparaben as an antifungal agent. Baker’s yeast sprinkles were added to new bottles and vials just before the flies were transferred to them.

Quantification of larval α-amanitin resistance for the two 180-line GWAS

Four- to six-day-old adult flies were allowed to lay eggs on molasses agar caps that contained fresh Baker’s yeast paste, while being housed in incubators at 25˚C, 70% humidity, and a 12:12 hour light/dark cycle. Freshly hatched first-instar larvae were then transferred in groups of ten into 2-mL plastic tubes (USA Scientific), each containing 125 mg of dry instant *Drosophila* food (Carolina) mixed with 375 µL of water that contained either 0.1 or 1.0 µg of dissolved α-amanitin, resulting in 500 mg of hydrated food of the concentrations 0.2 or 2.0 µg/g of α-amanitin, respectively. For each DGRP line (including the 37-line subset lines), three such replicate vials were seeded with 10 larvae, and the average number of adult flies hatching from the three replicates was used as the phenotypic value for the two 180-line GWAS.

Dose-response studies to quantify larval α-amanitin resistance for the 37-line GWAS

Before calculating the LC$_{50}$ values of the 37 lines, we pre-evaluated the α-amanitin resistance of each fly line based on the data obtained for 2.0 µg/g and a second α-amanitin concentration, 0.2 µg/g. This estimate of resistance allowed us to roughly sort the 37 lines into three categories: low, medium, and high α-amanitin resistance. For the exact LC$_{50}$ value calculation, adult flies were allowed to lay eggs on molasses agar caps that contained fresh Baker’s yeast paste, while being housed in incubators at 25˚C, 70% humidity, and a 12:12 hour light/dark cycle. Freshly hatched first-instar larvae were transferred in groups of ten into 2-mL plastic tubes (USA Scientific), each containing 125 mg of dry instant *Drosophila* food (Carolina) mixed with 375 µL of liquid (either sterile water or dissolved α-amanitin in water), thus resulting in 500 mg of hydrated food. All lines were then tested on seven α-amanitin concentrations, including the control without α-amanitin: low-resistance lines were tested on 0, 0.025, 0.05, 0.075, 0.1, 0.25, and 0.375 µg/g; medium-resistance lines on 0, 0.33, 0.66, 1.0, 1.33, 1.66, and 2.0 µg/g; and high-resistance lines were tested on 0, 0.66, 1.33, 2.0, 2.66, 3.33, and 4.0 µg/g food. Ten tubes were prepared for each toxin concentration and experiment, and each experiment was performed in three replicates. Only flies that completely hatched from their pupal cases were scored, and experiments in which at least 80% of the control flies hatched were used for the analysis. The experiments were then normalized by setting the control fly hatch numbers to 100%, and the LC$_{50}$ values for each fly line were calculated using scatter plots and the logarithmic trendline function in Microsoft Excel.

GWAS

Out of the original 189 DGRP lines, three lines (DGRP-492, 727, and 894) did not produce enough eggs for experiments, while six lines (DGRP-274, 378, 398, 476, and 554) produced a submission error on the DGRP2 website (they were marked as invalid lines). The remaining 180 lines listed in S1 Table were used in the current study. In order to identify SNPs associated with α-amanitin resistance in larvae, we submitted the average numbers of hatching flies from three replicates on 2.0 µg/g and on 0.2 µg/g α-amanitin (the two 180-line GWAS) and the LC$_{50}$ values of the 37-line study, not sexed, to the DGRP for analysis, using their web portal at dgrp2.gnets.ncsu.edu. Genotype-phenotype associations were calculated by the DGRP team, using a linear mixed model, which accounts for any effects of *Wolbachia* infection, common polymorphic inversions, and cryptic relatedness in the DGRP lines, as described in detail in [57]. Because 1.9 million SNP variants were tested in the GWAS, the genome-wide
significant threshold at the 5% significance level was determined after Bonferroni correction for multiple testing [58] as 2.60E-08 (0.05/2,192,980) [59]. However, 1.00E-05 was often used as a “relaxed” threshold in previous DGRP GWAS studies [60, 61]. For comparability with other studies, we report significance at the “strict” (< 2.60E-08) and “relaxed” (<1.00E-05) significance threshold. For candidate genes that were identified in both GWAS, we further report a “suggestive” significance threshold between 1.00E-04 and 9.99E-04. Candidate genes were selected based on the significance levels in the 180-line and the 37-line GWAS. The associations between LC\textsubscript{50} data with the average number of hatching flies at 2.0 and 0.2 \textmu g/g \alpha-amanitin among the 37 lines were calculated using a linear regression analysis (S3 and S4 Tables). Broad-sense heritability values were estimated as the ratio of the genetic variance and phenotypic variance according to [62].

RNA expression correlation analysis

In order to test what candidate genes resulting from the 37-line GWAS were dysregulated in adult flies, we used the publicly available RNA-seq data (http://dgrp2.gnets.ncsu.edu/data.html), which were originally published in [24]. We performed a Pearson correlation analysis, using R software (http://www.R-project.org/). Because the larvae used for the LC\textsubscript{50} experiments were not sexed, we correlated the LC\textsubscript{50} values yielded from the DGRP lines in our 37-line experiment with both adult male and female RNA expression values yielded from the same lines independently.

Supporting information

S1 Table. Viability analysis of all 180 DGRP lines on two \alpha-amanitin concentrations. All DGRP lines were tested on two \alpha-amanitin concentrations (0.2 \textmu g/g and 2.0 \textmu g/g), using 30 larvae per concentration and fly line (three replicates of 10 larvae each). Twelve lines displayed relatively high, 83 intermediate, and 91 low resistance to \alpha-amanitin. The values are also graphically represented in Fig 1A and 1C.

S2 Table. Viability analysis of the 37-line subset expressed as LC\textsubscript{50} values derived from seven \alpha-amanitin concentrations. The LC\textsubscript{50} values of the lines are shown in the first tab, and the raw data used to calculate the LC\textsubscript{50} values are presented in the second tab of the Excel file. The LC\textsubscript{50} values are also graphically represented in Fig 1C.

S3 Table. Correlation analysis among the 37-line GWAS and the two 180-line GWAS raw data sets on both 2.0 and 0.2 \textmu g/g of \alpha-amanitin. The two lines with the highest LC\textsubscript{50} values were not excluded in this file.

S4 Table. Correlation analysis among the 37-line GWAS and the two 180-line GWAS raw data sets on both 2.0 and 0.2 \textmu g/g of \alpha-amanitin. The two lines with the highest LC\textsubscript{50} values were excluded in this file, leading to an improved correlation among the 37-line and the 180-line data on 2.0 \textmu g/g of \alpha-amanitin.

S5 Table. Complete candidate SNP list resulting from the 37-line GWAS. A cutoff single mixed p-value of 1.00E-03 was used.
S6 Table. Complete candidate SNP list resulting from the 180-line GWAS on 2.0 μg/g of α-amanitin. A cutoff single mixed p-value of 1.00E-03 was used.
(XLSX)

S7 Table. Complete candidate SNP list resulting from the 180-line GWAS on 0.2 μg/g of α-amanitin. A cutoff single mixed p-value of 1.00E-03 was used.
(XLSX)

S8 Table. Pearson correlation analysis. Larval LC₅₀ values and the publicly available RNA-seq data of untreated male and female adults were correlated.
(XLSX)

Acknowledgments

We would like to thank our past and current undergraduate students for technical assistance with the experiments: Stephanie D’Annunzio, Lisa Diduch, Valerie Sidock, Andrea Kubicki, Amanda Bybee, Rianne Chittenden, Alyssa Sipes, Alec Steadman, Ty Koenigs, Kyle Glasper, Taylor Luginbill, Brian Borowicz, Stephanie Bonenfant, Ashima Chhabra, Jennifer Hackman, Claire Allison, Jennifer Connors, Andrew Lamoureux, Spencer Olson, Luke Cayemberg, Evan Bachman, Amber Peabody, Ross Michaels, Audri Mills, and Mary Stangis. We would also like to thank Wen Huang, and Trudy Mackay at North Carolina State University for helpful discussions.

Author Contributions

Conceptualization: CLM CEL KRV IMG REH OKI SRD JLP FRN ZJJ XJ HW OG TW.
Data curation: CLM CEL KRV IMG REH OKI SRD JLP FRN ZJJ XJ HW OG TW.
Formal analysis: CLM KRV XJ HW OG TW.
Funding acquisition: TW.
Investigation: CLM CEL KRV IMG REH OKI SRD JLP FRN ZJJ.
Methodology: CLM CEL KRV IMG REH OKI SRD JLP FRN ZJJ XJ HW OG TW.
Project administration: CLM TW.
Resources: HW OG TW.
 Supervision: CLM KRV TW.
Validation: CLM XJ HW OG TW.
Visualization: CLM KRV XJ HW OG TW.
Writing – original draft: TW KRV CLM.
Writing – review & editing: CLM CEL KRV IMG REH OKI SRD JLP FRN ZJJ XJ HW OG TW.

References

1. Jaenike J, Selander RK. Ecological generalism in Drosophila falleni—genetic evidence. Evolution. 1979; 33(2):741–8.
2. Phillips JP, Wills J, Pitt A. Alpha-amanitin resistance in three wild strains of Drosophila melanogaster. Canad J Genet Cytol. 1982; 24(2):151–62. PMID: 6812932
3. Jaenike J, Grimaldi DA, Sluder AE, Greenleaf AL. Alpha-amanitin tolerance in mycophagous Drosophila. Science. 1983; 221(4606):165–7. doi: 10.1126/science.221.4606.165 PMID: 17769215

4. Jaenike J, Grimaldi D. Genetic variation for host preference within and among populations of Drosophila tripunctata. Evolution. 1983; 37(5):1023–33.

5. Jaenike J. Parasite pressure and the evolution of amanitin tolerance in Drosophila. Evolution. 1985; 39(6):1295–301.

6. Jaenike J. Genetic population structure of Drosophila tripunctata—patterns of variation and covariation of traits affecting resource use. Evolution. 1989; 43(7):1467–82.

7. Begun DJ, Whitley P. Genetics of alpha-amanitin resistance in a natural population of Drosophila melanogaster. Heredity. 2000; 85(2):184–90.

8. Stump AD, Jablonski SE, Bouton L, Wilder JA. Distribution and mechanism of alpha-amanitin tolerance in mycophagous Drosophila (Diptera: Drosophilidae). Environ Entomol. 2011; 40(8):1604–12. doi: 10.1603/EN11136 PMID: 22217779

9. Benjamin DA. Mushrooms: poisons and panaceas: a handbook for naturalists, mycologists, and physicians. New York: W.H. Freeman and Company; 1996.

10. Vetter J. Toxins of Amanita phalloides. Toxicon. 1998; 36(1):13–24. PMID: 9604278

11. Lindell TJ, Weinberg F, Morris PW, Roeder RG, Rutter WJ. Specific inhibition of nuclear RNA polymerase II by alpha-amanitin. Science. 1970; 170(956):447–9.

12. Spicer GS, Jaenike J. Phylogenetic analysis of breeding site use and alpha-amanitin tolerance within the Drosophila quinaria species group. Evolution. 1996; 50(6):2328–37.

13. Perlman SJ, Jaenike J. Infection success in novel hosts: an experimental and phylogenetic study of Drosophila-parasitic nematodes. Evolution. 2003; 57(3):544–557. PMID: 12703944

14. Perlman SJ, Spicer GS, Shoemaker DD, Jaenike J. Associations between mycophagous Drosophila and their Howardula nematode parasites: a worldwide phylogenetic shuffle. Mol Ecol. 2003; 12(1):237–49. PMID: 12492892

15. Mitchell CL, Saul MC, Lei L, Wei H, Werner T. The mechanisms underlying alpha-amanitin resistance in Drosophila melanogaster: a microarray analysis. PLoS One. 2014; 9(4):e93489. doi: 10.1371/journal.pone.0093489 PMID: 24695618

16. Mitchell CL, Yeager RD, Johnson ZJ, D’Annunzio SE, Vogel KR, Werner T. Long-term resistance of Drosophila melanogaster to the mushroom toxin alpha-amanitin. PLoS One. 2015; 10(5):e0127569. doi: 10.1371/journal.pone.0127569 PMID: 25978397

17. Hallen HE, Luo H, Scott-Craig JS, Walton JD. Gene family encoding the major toxins of lethal Amanita species. Heredity. 2000; 85(2):184–90.

18. Kalaiedzic P, Oehler S, Reczko M, Pavlidi N, Vontas J, Hatzegeorgiou AG, et al. Use of mutagenesis, genetic mapping and next generation transcriptomics to investigate insecticide resistance mechanisms. PLoS One. 2012; 7(6):e42926. doi: 10.1371/journal.pone.0042926 PMID: 22782870

19. Amichot M, Tares S, Brun-Barale A, Arthaud L, Bride JM, Berge JB. Point mutations associated with insecticide resistance in Drosophila melanogaster. Insect Biochem Mol Biol. 2007; 37(5):512–9. doi: 10.1016/j.ibmb.2007.02.008 PMID: 17456446

20. Daborn PJ, Lumb C, Boey A, Wong W, Ffrench-Constant RH, Batterham P. Evaluating the insecticide resistance potential of eight Drosophila melanogaster cytochrome P450 genes by transgenic over-expression. Insect Biochem Mol Biol. 2007; 37(5):512–9. doi: 10.1016/j.ibmb.2007.02.008 PMID: 17456446

21. Festucci-Buselli RA, Carvalho-Dias AS, de Oliveira-Andrade M, Caixeta-Nunes C, Li HM, Stuart JJ, et al. Expression of Cyp6g1 and Cyp12d1 in DDT resistant and susceptible populations of Drosophila melanogaster. Insect Mol Biol. 2005; 14(1):69–77. doi: 10.1111/j.1365-2933.2005.00932.x PMID: 15663776

22. Le Goff G, Hilliou F, Siegfried BD, Boundy S, Wajnberg E, Sofer L, et al. Xenobiotic response in Drosophila melanogaster: sex dependence of P450 and GST gene induction. Insect Biochem Mol Biol. 2006; 36(8):674–82. doi: 10.1016/j.ibmb.2006.05.009 PMID: 16876710

23. Mackay TF, Richards S, Stone EA, Barbadilla A, Ayroles JF, Zhu D, et al. The Drosophila melanogaster Genetic Reference Panel. Nature. 2012; 482(7384):173–8. doi: 10.1038/nature10811 PMID: 22318601

24. Ayroles JF, Carbone MA, Stone EA, Jordan KW, Lyman RF, Magwire MM, et al. Systems genetics of complex traits in Drosophila melanogaster. Nat Genet. 2009; 41(3):299–307. doi: 10.1038/ng.332 PMID: 19234471

25. Attrill H, Falls K, Goodman JL, Millburn GH, Antonazzo G, Rey AJ, et al. FlyBase: establishing a Gene Group resource for Drosophila melanogaster. Nucleic Acids Res. 2016; 44(D1):D786–92. doi: 10.1093/nar/gkv1046 PMID: 26467478
26. Marzolo MP, Farfan P. New insights into the roles of megalin/LRP2 and the regulation of its functional expression. Biol Res. 2011; 44(1):89–105. doi: 10.4067/S0716-976020110000100012 PMID: 21720686

27. Auderset L, Landowski LM, Foà L, Young KM. Low density lipoprotein receptor related proteins as regulators of neural stem and progenitor cell function. Stem Cells Int. 2016; 2016:2108495. doi: 10.1155/2016/2108495 PMID: 26949399

28. Lamb CA, Dooley HC, Tooze SA. Endocytosis and autophagy: Shared machinery for degradation. Bioessays. 2013; 35(1):34–45. doi: 10.1002/bies.201200130 PMID: 23147242

29. Nielsen R, Christensen EI, Birn H. Megalin and cubilin in proximal tubule protein reabsorption: from experimental models to human disease. Kidney Int. 2016; 89(1):58–67. doi: 10.1016/j.kint.2015.11.007 PMID: 26759048

30. Fisher CE, Howie SE. The role of megalin (LRP-2/Gp330) during development. Dev Biol. 2006; 296(2):279–97. doi: 10.1016/j.ydbio.2006.06.007 PMID: 16828734

31. Riedel F, Vorkel D, Eaton S. Megalin-independent yellow endocytosis restricts melanization in the Drosophila cuticle. Development. 2011; 138(1):149–58. doi: 10.1242/dev.056309 PMID: 21138977

32. De S, Kuwahara S, Saito A. The endocytic receptor megalin and its associated proteins in proximal tubule epithelial cells. Membranes (Basel). 2014; 4(3):333–55.

33. Hussain MM, Strickland DK, Bakillah A. The mammalian low-density lipoprotein receptor family. Annu Rev Nutr. 1999; 19:141–72. doi: 10.1146/annurev.nutr.19.1.141 PMID: 10448520

34. Gleixner EM, Canaud G, Hermle T, Guida MC, Kretz O, Helmstadter M, et al. V-ATPase/mTOR signaling regulates megalin-mediated apical endocytosis. Cell Rep. 2014; 8(1):10–9. doi: 10.1016/j.celrep.2014.05.035 PMID: 24953654

35. Cabezas F, Lagos J, Cespedes C, Vio CP, Bronfman M, Marzolo MP. Megalin/LRP2 expression is induced by peroxisome proliferator-activated receptor-alpha and -gamma: implications for PPARs’ roles in renal function. PLoS One. 2011; 6(2):e16794. doi: 10.1371/journal.pone.0016794 PMID: 21311715

36. Huang CW, Wang HD, Bai H, Wu MS, Yen JH, Tatar M, et al. Tequila regulates insulin-like signaling and extends life span in Drosophila melanogaster. J Gerontol A Biol Sci Med Sci. 2015; 70(12):1461–9. doi: 10.1093/gerona/glv094 PMID: 26265729

37. Didelot G, Molinari F, Tchenio P, Comas D, Milhiet E, Munnich A, et al. Tequila, a neurotrypsin ortholog, regulates long-term memory formation in Drosophila. Science. 2006; 313(5788):851–3. doi: 10.1126/science.1127215 PMID: 16902143

38. Funakoshi M, Tsuda M, Muramatsu K, Hatsuda H, Morishita S, Aigaki T. A gain-of-function screen identifies wdb and lkb1 as lifespan-extending genes in Drosophila. Biochem Biophys Res Commun. 2011; 405(4):667–72. doi: 10.1016/j.bbrc.2011.01.090 PMID: 21281604

39. Banreti A, Lukasovich T, Csikos G, Erdelyi M, Sass M. PP2A regulates autophagy in two alternative ways in Drosophila. Autophagy. 2012; 8(4):623–36. doi: 10.4161/auto.19081 PMID: 22330894

40. Baltzer C, Tiefenbock SK, Marti M, Frei C. Nutrition controls mitochondrial biogenesis in the Drosophila adipose tissue through Delg and cyclin D/Cdk4. PLoS One. 2009; 4(9):e6935. Epub 2009/09/11. doi: 10.1371/journal.pone.0006935 PMID: 19742324

41. Bozicevic V, Hutter S, Stephan W, Wollstein A. Population genetic evidence for cold adaptation in European Drosophila melanogaster-populations. Mol Ecol. 2016; 25(5):1175–91. doi: 10.1111/mec.13464 PMID: 26558479

42. Anholt RRH, Mackay TFC. The genetic architecture of odor-guided behavior in Drosophila melanogaster. Behav Genet. 2001; 31(1):17–27. PMID: 11529271

43. Relling JH, Sabatini DM. Stress and mTORtue signaling. Oncogene. 2006; 25(48):6373–83. doi: 10.1038/sj.onc.1209889 PMID: 17041623

44. Bilder D, Scott MP. Genomic regions required for morphogenesis of the Drosophila embryonic midgut. Genetics. 1995; 141(3):1087–100. PMID: 8582615

45. Reuter R, Scott MP. Expression and function of the homeotic genes Antennapedia and Sex combs reduced in the embryonic midgut of Drosophila. Development. 1990; 109(2):289–303. PMID: 1976087

46. Kounatidis I, Ligoxygakis P. Drosophila as a model system to unravel the layers of innate immunity to infection. Open Biol. 2012; 2.

47. Ji S, Sun M, Zheng X, Li L, Sun L, Chen D, et al. Cell-surface localization of Pellino antagonizes Toll-mediated innate immune signalling by controlling MyD88 turnover in Drosophila. Nat Commun. 2014; 5:3458. doi: 10.1038/ncomms4458 PMID: 24632597

48. Jiang Z, Johnson HJ, Nie H, Qin J, Bird TA, Li X. Pellino 1 is required for interleukin-1 (IL-1)-mediated signaling through its interaction with the IL-1 receptor-associated kinase 4 (IRAK4)-IRAK-tumor
necrosis factor receptor-associated factor 6 (TRAF6) complex. J Biol Chem. 2003; 278(13):10952–6. doi: 10.1074/jbc.M212112200 PMID: 12496252

49. Hagheyeghi A, Sarac A, Czernicki S, Grosshans J, Schock F. Pellino enhances innate immunity in Drosophila. Mech Dev. 2010; 127(5–6):301–7. doi: 10.1016/j.mod.2010.01.004 PMID: 20117206

50. Jeon YK, Kim CK, Koh J, Chung DH, Ha GH. Pellino-1 confers chemoresistance in lung cancer cells by upregulating cIAP2 through Lys63-mediated polyubiquitination. Oncotarget. 2016; 7(27):41811–24. doi: 10.18632/oncotarget.9619 PMID: 27248820

51. Delgado M, Singh S, De Haro S, Master S, Pompua M, Dinkins C, et al. Autophagy and pattern recognition receptors in innate immunity. Immunol Rev. 2009; 227(1):189–202. doi: 10.1111/j.1600-065X.2008.00725.x PMID: 19120485

52. Deretic V. Links between Autophagy, Innate Immunity, Inflammation and Crohn's Disease. Digest Dis. 2009; 27(3):246–51.

53. Wu HY, Wang T, Li L, Correia K, Morgan Ji. A structural and functional analysis of Nna1 in Purkinje cell degeneration (pcd) mice. FASEB J. 2012; 26(11):4468–80. doi: 10.1096/fj.12-205047 PMID: 22835831

54. Lin R, Angelin A, Da Settimo F, Martini C, Taliani S, Zhu S, et al. Genetic analysis of dTSPO, an outer mitochondrial membrane protein, reveals its functions in apoptosis, longevity, and Ab42-induced neurodegeneration. Aging Cell. 2014; 13(3):507–18. doi: 10.1111/acel.12200 PMID: 24977274

55. Lin R, Rittenhouse D, Sweeney K, Potluri P, Wallace DC. TSPO, a mitochondrial outer membrane protein, controls ethanol-related behaviors in Drosophila. Plos Genet. 2015; 11(8):e1005366. doi: 10.1371/journal.pgen.1005366 PMID: 26241038

56. Greenleaf AL, Borsett LM, Jiamachello PF, Coulter DE. Alpha-amamin resistance D. melanogaster with an altered RNA polymerase II. Cell. 1979; 18(3):613–22. PMID: 117900

57. Huang W, Massouras A, Inoue Y, Peiffer J, Ramia M, Tarone AM, et al. Natural variation in genome architecture among 205 Drosophila melanogaster Genetic Reference Panel lines. Genome Res. 2014; 24(7):1193–208. doi: 10.1101/gr.171546.113 PMID: 24714809

58. Rice WR. Analyzing Tables of Statistical Tests. Evolution. 1989; 43(1):223–5.

59. Battlay P, Schmidt JM, Fournier-Level A, Robin C. Genomic and transcriptomic associations identify a new insecticide resistance phenotype for the selective sweep at the Cyp6g1 locus of Drosophila melanogaster. G3. 2016; 6(6):2573–81. doi: 10.1534/g3.116.031054 PMID: 27317781

60. Vonesch SC, Lamparter D, Mackay TF, Bergmann S, Hafen E. Genome-wide analysis reveals novel regulators of growth in Drosophila melanogaster. Plos Genet. 2016; 12(1):e1005616. doi: 10.1371/journal.pgen.1005616 PMID: 25751788

61. Durham MF, Magwire MM, Stone EA, Leips J. Genome-wide analysis in Drosophila reveals age-specific effects of SNPs on fitness traits. Nat Commun. 2014; 5:4338. doi: 10.1038/ncomms5336 PMID: 25000897

62. Marriage TN, King EG, Long AD, MacDonald SJ. Fine-mapping nicotine resistance loci in Drosophila using a multiparent advanced generation inter-cross population. Genetics. 2014; 198(1):45–57. doi: 10.1534/genetics.114.162107 PMID: 25236448