

A study of allelic series using transcriptomic phenotypes

David Angeles-Albores^{1,2} and Paul W. Sternberg^{1,2,*}

¹*Division of Biology and Biological Engineering, Caltech, Pasadena, CA, 91125, USA*

²*Howard Hughes Medical Institute, Caltech, Pasadena, CA, 91125, USA*

*Corresponding author. Contact: pws@caltech.edu

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Abstract

Abstract goes here

Significance Statement

Introduction

Something something allelic series are important for finding out the properties and potentials of genes.

As case studies, we focus on an allelic series of the *let-60* gene and an allelic series of the *dpy-22* gene in *C. elegans*. *let-60* is the *ras* orthologue in *C. elegans*¹, where it functions to promote the cellular fate of a number of cells during development². *let-60 (ras)* is a GTPase¹ that cycles between GDP- and GTP-binding states. The GTP-binding state is considered to be the signaling competent state. *dpy-22* is the *med-XX* orthologue in *C. elegans*.

In *C. elegans*, *let-60 (ras)* always acts downstream of a receptor. Activated *let-60 (ras)* can activate signaling cascades to transmit information to the eukaryotic nucleus. Often, this signaling cascade consists of *lin-45* (the RAF ortholog)³, *mek-2*⁴ and *mpk-1*⁵. The Ras pathway has been extensively studied in the context of vulval organogenesis, where it acts downstream of *let-23*⁶ to promote vulval induction of the Pn.p cells. In addition to vulval development, *let-60 (ras)* has been implicated in the migration of the Sex Myoblasts (SM), where it acts both cell autonomously and non-autonomously, the formation of the excretory pore, hypodermal fluid homeostasis, as well as germline development and the formation of the male tail⁷. Thus, the Ras pathway is pleiotropic, but its phenotypes are separable and specific.

The intense scrutiny of the Ras pathway has led to the generation of a large number of loss-of-function alleles for *let-60 (ras)*. Null mutations of *let-60 (ras)* are known to be lethal, but reduction-of-function alleles have been useful to carefully dissect the molecular properties of this protein. Gain-of-function mutations of *let-60 (ras)* are also known, although they are much more rare. In particular, a single gain-of-function mutation, *n1046gf*, was famously isolated multiple times in several screens^{8,9,10} leading to a Multivulva (Muv) phenotype. These gain-of-function mutations have a privileged history in the Ras pathway, as they have been used in multiple screens to identify genes that have a Suppressor of Ras (Sur) phenotype such as *sur-1* (now *mpk-1*⁵) or *sur-2*¹¹.

Talk about *dpy-22*.

In this paper, we set out to sequence a weak gain-of-function (*n1046gf*) and a strong loss-of-function (*n2021*) allele of Ras, as well as a weak loss-of-function (*bx93*) and a strong-loss-of-function (*sy622*) allele

of *dpy-22* (*med-12*). In either allelic series we found behaviors that challenge the way we think about alleles and allelic series.

Results

Comparing *ras(gf)* and *ras(lf)* alleles

As a first survey of allelic series, we sequenced two alleles of *let-60*, *let-60(gf)* and *let-60(lf)* because these alleles appear quantitatively different in their functionality when assayed via vulval formation measurements. We predicted that the transcriptomes of these alleles would show the same set of differentially expressed genes (henceforth referred to them as a shared transcriptomic phenotype, STP). Moreover, we predicted genes within this STP would show weaker perturbations on average within one mutant compared to the other. Our results showed that the *let-60(gf)* mutant transcriptome had 3,021 differentially expressed genes, whereas the *let-60(lf)* had 857 differentially expressed genes (see Fig. 1).

Contrary to our expectations, the STP between these two alleles consisted of 31% of the differentially expressed genes in the *let-60(lf)* transcriptome, totalling 269 genes. Moreover, we found that this STP showed a strong positive correlation between the two alleles. In order to assess whether one allele was significantly stronger than another we split the STP into its correlated and anti-correlated components and found the regression line in each component. We measured a correlation coefficient of 1.03 and -1.00 for each component, indicating that the two alleles have effects that are exactly opposite on average. Our results suggest the existence of four phenotypes: A phenotype associated exclusively with *let-60(gf)* (the *let-60(gf)*-specific phenotype), another associated exclusively with *let-60(lf)* (the *let-60(lf)*-specific phenotype), a phenotype that is the same in both alleles and a phenotype that is inverted in one allele relative to the other.

In order to gain some insight into the biological meaning each set, we used the WormBase Enrichment Suite to perform Gene, Tissue and Phenotype Ontology enrichment analyses. Analysis of the *let-60(gf)*-specific phenotype revealed enrichment of genes expressed in the early embryo (AB lineage), in the male, the hypodermis and the reproductive system whereas the *let-60(lf)*-specific phenotype showed enrichment of the intestine. Phenotype term enrichment associated the *let-60(gf)*-specific transcriptomic phenotype with the linker cell migration, lipid metabolism and neuropil development, and the *let-60(lf)*-specific transcriptomic phenotype was enriched in mitochondrial alignment. Finally, gene ontology enrichment analysis showed that the *let-60(gf)*-specific transcriptomic phenotype was enriched in regulation of cell-shape, immune system response and side of membrane. The *let-60(lf)*-specific phenotype showed enrichment in terms related to striated muscle, collagen trimer and cell death. Our results suggest that gain- and loss-of-function alleles have separable functions that do not obviously recapitulate the visible phenotypes typically associated with this gene.

A strong and a weak loss-of-function *dpy-22* allele show different transcriptomic profiles

We studied two alleles of *dpy-22* (a Mediator subunit) that previous studies had suggested could be qualitatively distinct. Allele *bx93* (the weak allele) encodes a premature stop codon that removes the terminal 900 amino acids from the protein. *bx93* homozygotic animals are phenotypically wild-type with a very low incidence of male tail defects. Allele *sy622* (the strong allele) encodes a premature stop codon that removes the terminal 1700 amino acids from the protein. *sy622* homozygotes grow slowly, are severely dumpy (Dpy), have a low penetrance multivulva (Muv) phenotype and have a prominent egg-laying defective (Egl) phenotype¹².

We sequenced homozygotes of both alleles. We found that *bx93* homozygotes expressed 434 differentially expressed genes, and *sy622* homozygotes showed 2,821 differentially expressed genes. Of the 434 differentially expressed genes in the *bx93* mutant, 73% were also differentially expressed in the *sy622* alleles, indicating that the impaired functionalities in the *bx93* allele are also impaired in the *sy622* allele. Having established that both alleles affect a shared subset of genes, we proceeded to measure whether the *sy622* allele showed greater perturbations in this subset. We observed that the weak allele, *bx93*, had perturbation magnitudes that were on average 39% weaker than the perturbation magnitudes in the strong allele, *sy622* (see Fig. 2). In summary, the strong allele had more differentially expressed genes than the weak allele, and genes altered

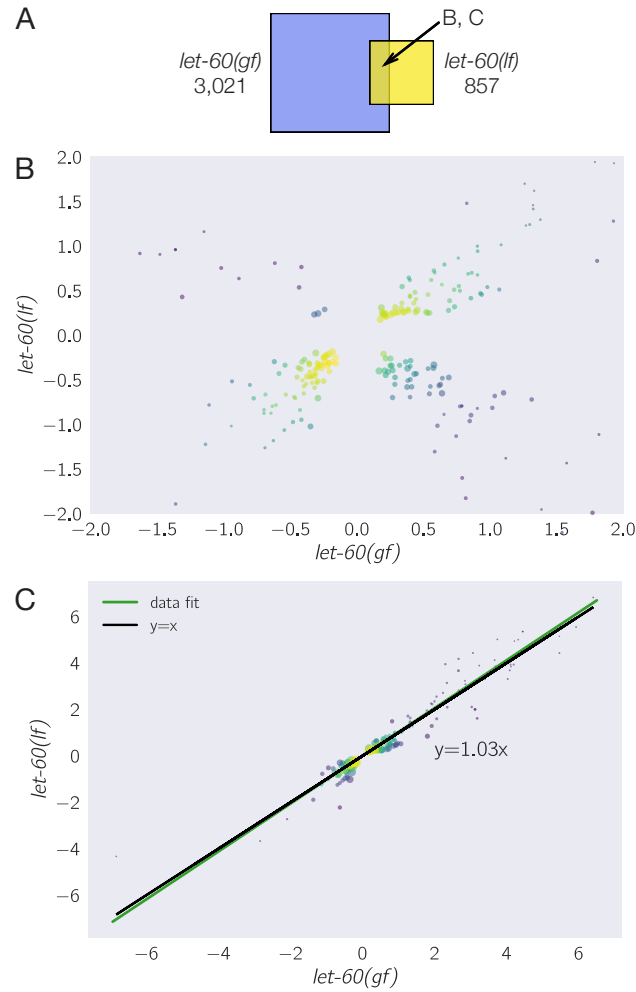


Figure 1. **A.** Venn diagram of the differentially expressed genes in a *let-60(gf)* and a *let-60(lf)* mutant. Diagram is to scale. **B.** β coefficients for isoforms that are differentially expressed in both mutants. **C** When two isoforms change in the same direction in both mutants, the magnitude of the change is also the same.

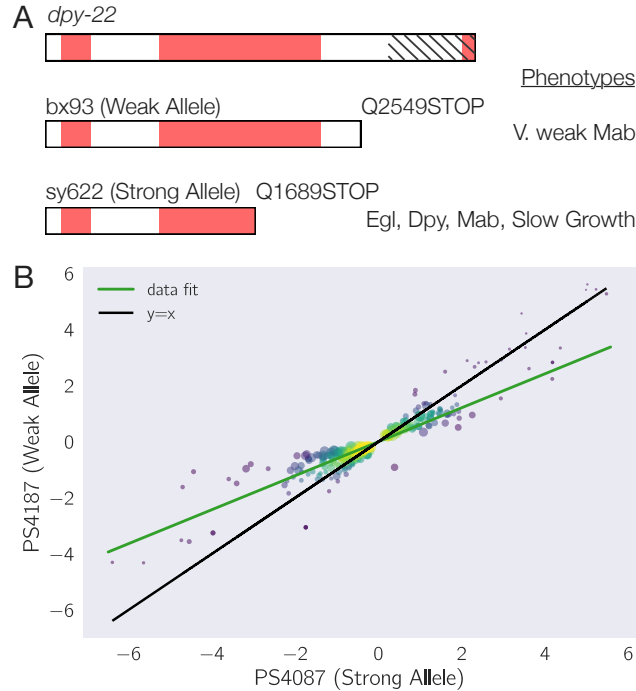


Figure 2. **A** Diagram of the *dpy-22* gene and the *bx93* and *sy622* alleles. **B** Genes that are commonly differentially expressed in both alleles typically change in the same direction, and they tend to change by 30% less in the *bx93* (weak allele) homozygote than in the *sy622* (strong allele) homozygote.

commonly in mutants of both alleles were more perturbed in the strong allele than in the weak allele. However, without a *trans*-heterozygote, it is impossible to tell whether these differences are the result of greater reduction of function in the *sy622* allele relative to the *bx93* allele or whether the *sy622* allele has qualitative differences from *bx93* as a result of additional deleted functional domains.

The *trans*-heterozygote of *dpy-22* strong and weak alleles separates the alleles into four phenotypic classes

A standard method to identify whether two alleles differ quantitatively in their activity levels or whether they are qualitatively different because each allele has inactivated protein domains with separable functions is to generate a *trans*-heterozygote. Theoretically, if two alleles are quantitatively different, the *trans*-heterozygote will have a phenotype intermediate the two homozygote phenotypes. On the other hand, if both alleles are inactivating distinct and separable functions of the protein, then the *trans*-heterozygote will exhibit a wild-type phenotype (intragenic complementation). Finally, if one allele is affecting multiple separable activities whereas the other allele is only affecting one, then the *trans*-heterozygote will exhibit the phenotype of the allele that affects the least number of activities (i.e., one allele will exhibit dominance).

We sequenced a *trans*-heterozygote of the *bx93* and *sy622* alleles with genotype *dpy-10(lf) bx93/+ sy622*. This *trans*-heterozygote appears phenotypically wild-type, resembling the *bx93* mutant morphologically. The *trans*-heterozygote showed 2,930 differentially expressed genes. Using the *trans*-heterozygote, we were able to separate genes into four main phenotypic classes, classifying them by what genotypes caused these genes to become differentially expressed. One phenotypic class consisted of genes that were differentially expressed in the *sy622* homozygote as well as the *trans*-heterozygote, but not in the *bx93* homozygote (989 differentially expressed genes). We called this the *sy622*-associated phenotype. Another phenotypic class consisted of 1,623 genes that were only dysregulated in the *sy622* homozygote, which we called the *sy622*-specific phenotype because it is entirely suppressed by the presence of a single copy of the *bx93* allele. We also found a *trans*-heterozygote-specific phenotype consisting of 1,676 genes which is not present in either homozygote. Finally, we defined a *bx93*-associated phenotype as the set of genes dysregulated in both the *bx93* homozygote and

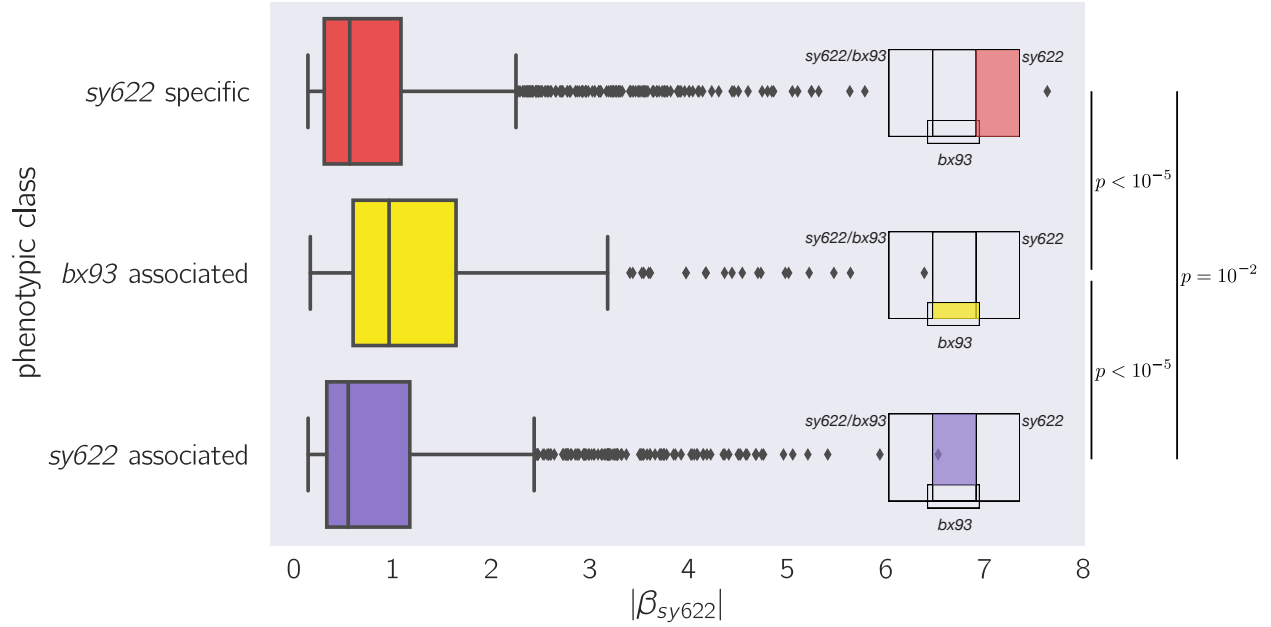


Figure 3. Different phenotypic classes have statistically different perturbation distributions. Genes that are *sy622*-specific have a different perturbation distribution compared to genes that are *bx93*-associated or *sy622*-associated. The lines within the boxes show the 25, 50, and 75 percentiles. Whiskers show the rest of the plot, except for outliers (diamonds). Insets show what genotypes each gene class is expressed in, but the magnitude of the perturbation plotted always corresponds to the *sy622* mutant. The means of each distribution were all statistically different from each other, as assessed by a non-parametric bootstrap test.

the heterozygote, consisting of 310 differentially expressed genes. Having defined these classes, we set out to describe their properties.

We asked whether these classes behaved differently within a homozygote. Specifically, we wanted to know whether the *sy622*-specific, the *sy622*-associated and the *bx93*-associated phenotypes were different in the magnitude of their perturbations or whether these subsets behaved as if they had been randomly selected from the set of differentially expressed genes in the *sy622* homozygote (see Fig. 3). We found that that the *bx93*-associated phenotype had the greatest magnitude of perturbations of the three classes (mean, median and maximum). The *sy622*-associated phenotype had a smaller range of perturbations compared to the *bx93*-associated phenotype (95th percentiles: 3.3 versus 4.2, respectively), and a statistically smaller mean (1.3 vs 0.99, respectively, $p < 10^{-5}$, non-parametric bootstrap). The *sy622*-specific phenotype had the smallest mean of all (0.9, $p < 10^{-5}$ compared with *bx93*-associated phenotype, and $p = 0.01$ compared with the *sy622*-associated phenotype). The medians are almost identical between the *sy622*-specific and the *sy622*-associated phenotypes, which indicates that the small difference in the means of these two distributions is primarily driven by the longer tail of the *sy622*-associated phenotype.

The *bx93* allele is dominant over the *sy622* for the *bx93*-associated phenotype

We explored how expression levels changed in the *bx93*-associated phenotypic class between the homozygotes and the heterozygote. We reasoned that if one allele was dominant over the other in the heterozygote, then plotting the β coefficients in the dominant homozygote versus the heterozygote should lead to a slope of 1. Moreover, since we previously quantified the relationship between the β coefficients in either homozygote in the previous section (the weak allele perturbations are 61% of the strong allele perturbations on average), it should be possible to multiply the recessive allele by this factor such that the relationship between the homozygotes and the heterozygotes is the same. This is important because agreement in both calculations gives increased confidence in the quantitative nature of these results.

We selected the genes within the *bx93*-associated phenotypic class, and plotted the β coefficients of these genes in the *bx93* allele against the coefficients in the heterozygote. The coefficients fell along a line with slope

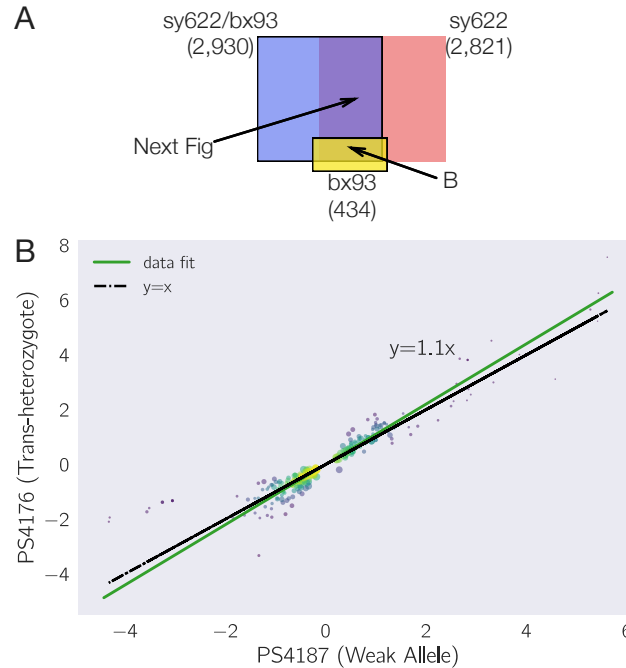


Figure 4. Genes that are differentially expressed in homozygotes of both alleles show expression levels consistent with partial dominance of *bx93* over *sy622* in a trans-heterozygote. **A.** Venn diagram of differentially expressed genes in the *bx93* (weak allele) and the *sy622* (strong allele) homozygotes and the *dpy-6(lf) bx93/+ sy622* trans-heterozygote. Arrows point at the gene subsets used for panel **B** and for Figure 5. **B.** Genes differentially expressed in the *bx93* homozygote have an expression phenotype that is 10% worse on average in the trans-heterozygote.

of 1.1, indicating that the *trans*-heterozygote has a strong resemblance to the *bx93* homozygote, although on average its phenotype is 10% worse (see Fig. 4). As a check, we obtain the β coefficients in the *sy622* allele and multiplied each coefficient by 0.61. This transformation should effectively transform, on average, the β coefficients in the *sy622* to the β coefficients in the *bx93* allele. When we calculate the slope between the transformed *sy622* β coefficients and the coefficients in the heterozygote, the slope is also 1.1. Taken together, this establishes that the *bx93* allele is 90% dominant over the *sy622* allele for the *bx93*-associated phenotypic class.

The *sy622*-associated phenotype is attenuated by the presence of *bx93* in the trans-heterozygote

We also wanted to know whether the *sy622*-associated phenotype showed differences depending on genotypic context. The *sy622*-associated genes are genes that are differentially expressed in the *sy622* homozygote or the trans-heterozygote, but not the *bx93* homozygote. Genes in this group showed a 23% reduction in the magnitude of their perturbations in the heterozygote compared to the homozygote (see Fig. 5). Therefore, these genes are attenuated by the presence of a single copy of the *bx93* allele, but the *sy622* is the dominant allele, showing 77% dominance in the expression level of this phenotypic class. This behavior is qualitatively different from genes in the *bx93*-associated phenotypic class, where *bx93*, not *sy622*, was 90% dominant. Finally, the behaviors of both of these classes are distinct from the behavior of genes in the *sy622*-specific class, which show differential expression in a *sy622* homozygote, but this dysregulation is complemented by the *bx93* allele. This establishes that alleles can have differences in dominance for different phenotypic classes at the gene expression level.

Insights into the physiology of *sy622* homozygotes

Whereas the *sy622* homozygote is strongly phenotypic (see Fig. 2), the *bx93* is almost entirely wild-type. Since the trans-heterozygote also appears grossly wild-type, we hypothesized that the *sy622*-specific pheno-

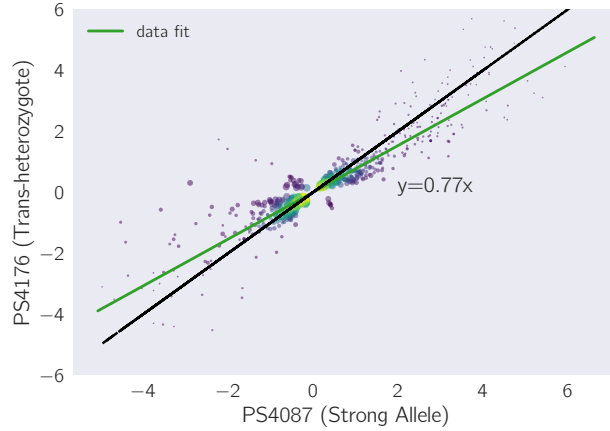


Figure 5. The *sy622/bx93* trans-heterozygote shows a phenotype that is on average 23% weaker than the *sy622* homozygote for genes contained in the *sy622*-associated phenotypic class.

typic class was associated with the macroscopic phenotypes visible in the *sy622* allele. To better understand this phenotypic class, we used the Wormbase Enrichment Suite^{13,14} to query what anatomical, phenotypic or gene ontological terms were enriched in this gene set (see Table 1).

Term	<i>sy622</i> -specific	<i>bx93</i> -associated
Intestine	12	—
Intestinal muscle	4	< 1
PVD	3	< 1
Muscular system	7	< 1
pm3/5	< 1	2
Severe pleiotropic defects early embryo	4	< 1
Rachis absent	4	< 1
Meiosis defective early embryo	3	< 1
Dauer constitutive	< 1	2
Dauer metabolism	< 1	2
Collagen trimers	25	< 1
Muscle cell development	13	< 1
Contractile fibers	14	< 1
Oviposition	12	< 1
Glucuronosyltransferase activity	< 1	9
Monocarboxylic acid catabolic process	< 1	4
Lytic vacuole	< 1	4

Table 1. List of enriched terms for the *sy622*-specific and *bx93*-specific phenotypic classes. The *sy622*-specific and the *bx93*-associated columns show $-\log_{10} q$ for each term. Q-values are calculated using a hypergeometric model and adjusted via a Benjamini-Hochberg algorithm. Q-values shown are rounded to the nearest power of 10 for simplicity. < 1 implies that the q-value did not pass the significance threshold.

The *sy622*-specific phenotypic class was enriched for genes expressed in the intestine and intestinal muscle. We also found enrichment in cell-types that could reasonably be associated with egg-laying defects, namely the PVD neuron, and the muscular system. Phenotype ontology enrichment revealed that the *sy622*-specific phenotypic class was enriched for terms associated with embryonic lethality and small brood size, such as severe pleiotropic defects in the early embryo, oocytes lack nucleus, rachis absent and meiosis defective in the early embryo. Gene ontology enrichment showed that the *sy622*-specific phenotype was enriched in collagen trimers, muscle cell development, contractile fibers and oviposition. In contrast, the *bx93*-specific phenotypic class does not enrich identical terms. Rather, the *bx93*-specific class shows enrichment for genes expressed

in the intestine and pharyngeal muscle cells, pm3 and pm5. It shows enrichment of genes associated with dauer constitutive and dauer metabolism phenotypes, and the gene ontology enrichment primarily reflects terms associated with metabolism, such as glucuronosyltransferase activity, monocarboxylic acid catabolic process, and lytic vacuole.

Conclusions

Gain-of-function and loss-of-function alleles can lead to different transcriptomic states.

We sequenced a weak gain-of-function (*n1046gf*) and a strong loss-of-function (*n2021*) allele of Ras, with the expectation that these alleles would affect the same set of genes in opposite ways, because the gain-of-function allele is thought to signal more than the wild type, whereas the loss-of-function does not signal at all. Contrary to our expectations, we find that these gene sets are substantially disjoint with a relatively small overlap. Within this overlap, we find that genes that are commonly differentially expressed in homozygotes of either allele tend to show changes of exactly the same magnitude and direction. In other words, the effect of loss-of-function and gain-of-function for these genes is exactly the same. Moreover, we find that for a smaller population of genes where loss-of-function and gain-of-function have opposing effects on gene expression, the magnitude of change is also the same. We considered various mechanisms that could generate these patterns.

We considered a scenario where the gain-of-function allele signals constitutively through pathways that are only sparingly used in wild-type animals. This model would predict that the gain-of-function allele mutant would show changes in these pathways as they are flooded with new information through channels that are rarely used. In contrast, the loss-of-function allele mutant should show fewer changes in these pathways, because in this model most of the changes are rarely in use. Thus, genes that are differentially expressed in the gain-of-function allele homozygote would be interpreted as genes where *let-60* (*ras*) is sufficient to trigger changes in expression, and the changes in the loss-of-function allele represent genes where *let-60* (*ras*) is necessary for appropriate expression. Although this scenario begins to explain the difference in size between the number of differentially expressed genes in the gain-of-function allele homozygote and the loss-of-function allele homozygote, it does not explain why there are genes that are differentially expressed in the loss-of-function allele homozygote that are not present in the gain-of-function allele. Moreover, this model does not explain why the STP between the loss-of-function allele and gain-of-function allele homozygotes is predominantly correlated, instead of being entirely anti-correlated. Therefore, it seems unlikely that the transcriptomes of these alleles result from a model where *let-60* (*ras*) is capable of signaling through a large number of pathways but predominantly uses only a subset.

A mechanism that could feasibly generate *let-60(lf)*- and *let-60(gf)*-specific effects as well as correlated and anti-correlated effects would be to postulate that there are four signaling states available to LET-60. In this model, a protein bound to GTP constitutively can signal through a set of proteins, whereas a protein bound to GDP signals through a different, non-overlapping pathway. A third pathway requires GTP-to-GDP cycling at a specific average rate, such that interfering with that cycling by stabilizing either the GTP- or GDP-bound states has the same effect. A similar effect has been described previously for the Sec4 GTPase.

Another way to understand the effects we observed in *let-60(lf)* and *let-60(gf)* mutants is by thinking of the transcriptomes of each mutant as their respective states. Recently, transcriptome profiling has been used to identify novel states in both single cells and whole-organisms¹⁵. If these transcriptomes are states, then these states are the result of continuous *let-60* (*ras*) activity (or inactivity) throughout the lifespan of the animal, and reflect the proximal or immediate effects of altered LET-60 activity as well as compensatory changes due to altered development or life history. This framework has the advantage that it does not *a priori* suggest that the gain-of-function allele homozygote and loss-of-function allele homozygote should have similar, or even overlapping, effects.

Without many more experiments, we cannot definitively point at a mechanism. It is possible that background mutations are contributing to the *let-60(lf)*- and *let-60(gf)*-specific changes, since these mutants were identified in different screens carried out in different labs. A rigorous methodology to exclude background mutations would call for sequencing multiple independent lines containing each allele. As library generation

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and sequencing costs fall, these experiments will become more feasible. Even assuming that background effects are responsible for the lack of overlap between the two mutants, the positive and negative correlations between these two alleles raise important questions about Ras biology. The fact that either correlation has a value with magnitude exactly of 1 suggests the existence of circuits that monitor Ras signaling levels with quantitative accuracy in *C. elegans*.

Loss-of-function allelic series reveal unknown functionality

We sequenced two alleles of *dpy-22* that had been previously studied and reported to have different functionalities. Through transcriptomic profiling, we were able to verify that *sy622* has a transcriptomic phenotype that is quantifiably worse than *bx93*. This worsening manifests as an increase in the number of differentially expressed genes in *sy622* relative to wild-type compared to the number of differentially expressed genes in *bx93*. Moreover, the genes that are commonly dysregulated in both alleles show greater perturbations on average in the *sy622* homozygote relative to the *bx93* allele (see Fig. 2). Unlike the *let-60* (*ras*) alleles we studied, the set of genes differentially expressed in the *bx93* is contained within the *sy622* with few exceptions. We can account for most of these exceptions by invoking a 10% false positive rate (which was the cutoff for our study) and a similar false negative rate.

Although a comparison between these two alleles proves fruitful in establishing differences in phenotypic severity, this comparison alone does not allow us to answer whether or not *bx93* and *sy622* act in different ways on subsets of genes. To this end, we sequenced a trans-heterozygote of both alleles, which allowed us to identify four phenotypic classes, which in turn are informative about the biological effects of each allele. We found a *sy622*-specific phenotypic class for which the *bx93* has a dominant wild-type phenotype; we found a *bx93*-associated phenotypic class for which *bx93* also has a dominant phenotype, but this phenotype is definitely not wild-type. We also found a *sy622*-associated class for which *sy622* and *bx93* appear co-dominant. Intriguingly, we also found a phenotype specific to the trans-heterozygote that was not present in either homozygote. A weakness in our study is that we have limited power to study the gene expression changes associated with the trans-heterozygote. This phenotypic class is puzzling because *dpy-22* (*med-12*) is not known to have homotypic interactions, which is a classical explanation for trans-heterozygote-specific phenotypes. Moreover, since this class is specific to a single strain we cannot rule out that this class is actually a result of a strain-specific mutation or set of mutations. In particular, the genotype of the heterozygote includes a mutation at the *dpy-6* locus to balance the *bx93* mutation. One possibility is that the *dpy-6* loss-of-function mutation is not recessive for transcriptomic phenotypes and is responsible for the dysregulation of the new genes observed in the heterozygote. Another possibility is that the *dpy-6* strain had a carrier mutation that fixed in the balanced strain. Finally, it is also possible that the *bx93* and the *sy622* strains had background mutations that did not have effects on their own, but when combined generate a synthetic phenotype with themselves or with one or both of the *dpy-22* alleles. As the cost of sequencing becomes lower, and with improved genetic engineering tools that allow the creation of background-free mutations, it will become increasingly important to rule out these hypotheses by sequencing additional independently derived identical alleles.

Our enrichment analysis of the *sy622*-specific and the *bx93*-associated phenotypic classes revealed that they reflect functionally distinct aspects of *dpy-22* (*med-12*) biology. The *sy622*-specific class contains genes that are associated with severe pleiotropic effects, embryonic lethality and sterility. It also contains genes that are associated with muscle development and function, and there is enrichment of genes expressed in the PVD neuron. Finally, collagen trimers are also overrepresented in this gene class. Taken together, these terms suggest that perturbing this gene class away from the wild-type should lead to an animal that is sickly, has a small brood size and has altered locomotion as well as altered collagen production. Indeed, the *sy622* homozygote is Dpy and Egl and RNAi against *dpy-22* (*med-12*) is known to cause embryonic and larval lethality. The *bx93*-associated class is enriched in a different set of terms, which suggests that transcriptome profiling can be used in conjunction with allelic series to separate genes into distinct phenotypic classes that are biologically relevant. Specifically, for the *sy622* and the *bx93* alleles, we can rule out that these alleles form a qualitative allelic series, since such a series should not exhibit a dosage-dependent phenotype (the *sy622*-associated class). However, it is possible that *sy622* and *bx93* represent a mixed quantitative-qualitative series, because we found phenotypic classes where *bx93* complemented *sy622*, *bx93* was dominant over *sy622* and where *sy622* was co-dominant with or partially dominant over *bx93*.

Allelic series are a cornerstone of genetic analyses. Classically, these series have been important to understand multiple aspects of a gene by comparing and contrasting the properties of different alleles in homozygotes as well as heterozygotes. Due to their sensitivity and quantitative nature, transcriptomic phenotypes represent an exciting new phenotype with which to study allelic series. In this study, we have shown that transcriptomic phenotypes can quickly and easily partition gene sets into phenotypic classes that have different statistical and physiological properties. There is a push for more and more specific sequencing of pure cell populations. This study shows that in addition to sequencing single cells in order to better understand cell-cell heterogeneity, we should sequence a greater allelic diversity in order to understand genotype-to-genotype heterogeneity.

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