# Genetic Analysis of a Metazoan Pathway using Transcriptomic Phenotypes, Supplementary Information

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# A quality check of the transcriptomic data reveals excellent agreement with the literature

One way to establish whether genes are acting additively or epistatically to each other is to perform qPCR of a reporter gene in the single and double mutants. This approach was used to successfully map the relationships within the hypoxia pathway (see, for example  $^{1,2}$ ). A commonly used hypoxia reporter gene is nhr-57, which is known to exhibit a several-fold increase in mRNA expression when HIF-1 accumulates  $^{2,3,4}$ . Likewise, increased HIF-1 function is known to cause increased transcription of rhy-1 and eql-9 $^5$ .

We can selectively look at the expression of a few genes at a time. Therefore, we queried the changes in expression of rhy-1, egl-9, and nhr-57. We included the nuclear laminin gene lam-3 as a representative negative control not believed to be responsive to alterations in the hypoxia pathway. nhr-57 was upregulated in egl-9(lf), rhy-1(lf) and vhl-1(lf), but remains unchanged in hif-1(lf). egl-9(lf); vhl-1(lf) had an expression level similar to egl-9(lf); whereas the egl-9(lf) hif-1(lf) mutant showed wild-type levels of the reporter expression, as reported previously  $^2$  (see Fig. S1).

We observed changes in rhy-1(lf) expression consistent with previous literature <sup>2</sup> when HIF-1 accumulates. We also observed increases in egl-9 expression in egl-9(lf). egl-9 is known as a hypoxia responsive gene <sup>5</sup>. Although changes in egl-9 expression were not statistically significantly different from the wild-type in rhy-1(lf) and vhl-1(lf) mutants, the mRNA levels of egl-9 still trended towards increased expression in these genotypes. As with nhr-57, egl-9 and rhy-1 expression were wild-type in egl-9(lf) hif-1(lf) and egl-9(lf); vhl-1(lf) mutant showed expression phenotypes identical to egl-9(lf). This dataset also showed that knockout of hif-1 resulted in a modest increase in the levels of rhy-1. This suggests that hif-1, in addition to being a positive regulator of rhy-1 when strongly expressed, also inhibits rhy-1, which constitutes a novel observation. Using a single reporter we would have been able to reconstruct an important fraction of the genetic relationships between the genes in the hypoxia pathway—but would likely fail to observe yet other genetic interactions, such as the evidence for hif-1 negatively regulating rhy-1 transcript levels.

# Weighted Correlations

After we calculated the pairwise correlation within each STP, we weighted the result of each regression by the number of isoforms within the STP and divided by the total number of differentially expressed isoforms present in the two mutant transcriptomes that contributed to that specific STP,  $N_{\text{overlap}}/N_{\text{g}_1\cup\text{g}_2}$ . The weighted regressions recapitulated a module network (see Fig. S3). We identified a strong positive

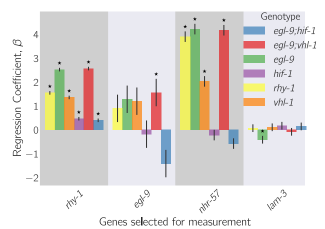
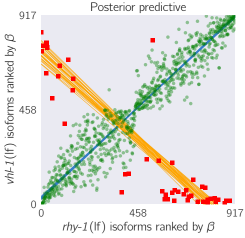


Figure S1. Observed  $\beta$  values of select genes. We selected four genes (rhy-1, egl-9, nhr-57 and lam-3, shown on the x-axis) and plotted their regression coefficients,  $\beta$ , as measured for every genotype (represented by one of six colors) to study the epistatic relationships between each gene. Asterisks above a bar represent a regression coefficient statistically significantly different from 0 ( $q < 10^{-1}$ ) relative to a wild-type control. Error bars show standard error of the mean value of  $\beta$ . nhr-57 is an expression reporter that has been used previously to identify hif-1 regulators  $^{2,1}$ . lam-3 is shown here as a negative control that should not be altered by mutations in this pathway. We measured modest increases in the levels of rhy-1 mRNA when hif-1(lf) is knocked out.



**Figure S2.** A feedback loop can generate transcriptomes that are both correlated and anti-correlated. The vhl-1(lf)/rhy-1(lf) STP shows a cross-pattern. Green large points are inliers to the first regression. Red squares are outliers to the first regression. Only the red small points were used for the secondary regression. Blue lines are representative samples of the primary bootstrapped regression lines. Orange lines are representative samples of the secondary bootstrapped regression lines.

interaction between egl-9(lf) and rhy-1(lf). The magnitude of this weighted correlation derives from the number of genes that are differentially expressed for these mutants (2,549 and 3,005 DEGs respectively) and the size of their STP, which makes the weighting factor considerably larger than other pairs. The weak correlation between hif-1(lf) and egl-9(lf) results from the large effect size of the egl-9(lf) transcriptome coupled with the small STP between both mutants.

The fine-grained nature of transcriptional phenotypes means that these weighted correlations between transcriptomes of single mutants are predictive of genetic interaction.

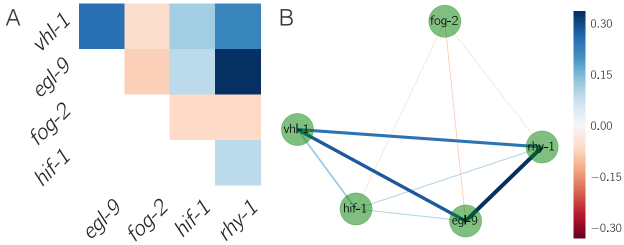
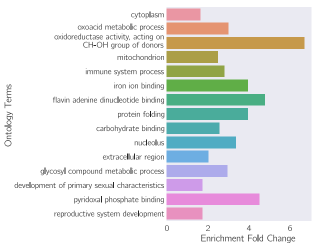


Figure S3. A. Heatmap showing pairwise regression values between all single mutants. B. Correlation network drawn from A. Edge width is proportional to the logarithm of the magnitude of the weighted correlation between two nodes divided by absolute value of the weighted correlation value of smallest magnitude. Edges are also colored according to the heatmap in A. Inhibitors of hif-1 are tightly correlated and form a control module; hif-1 is positively correlated to its inhibitors, albeit weakly; and fog-2, a gene that is not reported to interact with the hypoxia pathway, has the smallest, negative correlation to any gene.

#### Enrichment analysis of the hypoxia response

To validate that our transcriptomes were correct, and to understand how biological functions may vary between them, we subjected each decoupled response to enrichment analysis using the WormBase Enrichment Suite  $^{67}$ .



**Figure S4.** Gene ontology enrichment analysis of genes associated with the main hypoxia response. A number of terms reflecting catabolism and bioenergetics are enriched.

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We used gene ontology enrichment analysis (GEA) on the main hypoxia response program. This showed that the terms 'oxoacid metabolic process'  $(q < 10^{-4}, 3.0 \text{ fold-enrichment}, 24 \text{ genes})$ , 'iron ion binding'  $(q < 10^{-2}, 3.8 \text{ fold-enrichment}, 10 \text{ genes})$ , and 'immune system process'  $(q < 10^{-3}, 2.9 \text{ fold-enrichment}, 20 \text{ genes})$  were significantly enriched. GEA also showed enrichment of the term 'mitochondrion'  $(q < 10^{-3}, 2.5 \text{ fold-enrichment}, 29 \text{ genes})$  (see Fig. S4). Indeed, hif-1(lf) has been implicated in all of these biological and molecular functions  $^{7,8,9,10}$ . As benchmark on the quality of our data, we selected a set of 22 genes known to be responsive to HIF-1 levels from the literature and asked whether these genes were present in our hypoxia response list. We found 8/22 known genes, which constitutes a statistically significant result  $(p < 10^{10})$ . The small number of reporters found in this list probably reflects the conservative nature of our estimates. We studied the hif-1-independent, vhl-1-dependent gene set using enrichment analysis but no terms were significantly enriched.

## HIF-1 in the cellular context

We identified the transcriptional changes associated with bioenergetic pathways in C. elegans by extracting from WormBase all genes associated with the tricarboxylic acid (TCA) cycle, the electron transport chain (ETC) and with the C. elegans GO term energy reserve. Previous research has described the effects of mitochondrial dysfunction in eliciting the hypoxia response  $^{11}$ , but transcriptional control of bioenergetic pathways by HIF-1 has not been studied as extensively in C. elegans as in vertebrates (see, for example  $^{12,13}$ ). We also searched for the changes in ribosomal components and the proteasome, as well as for terms relating to immune response (see Fig S5).

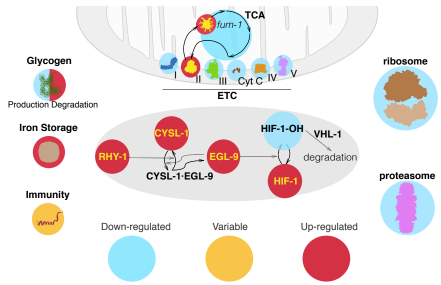


Figure S5. A graphic summary of the genome-wide effects of HIF-1 from our RNA-seq data.

#### Bioenergetic pathways

Our data shows that most of the enzymes involved in the TCA cycle and in the ETC are down-regulated when HIF-1 is induced in agreement with the previous literature <sup>13</sup>. However, the fumarase gene fum-1 and the mitochondrial complex II stood out as notable exceptions to the trend, as they were up-regulated in every single genotype that causes deployment of the hypoxia response. FUM-1 catalyzes the reaction of fumarate into malate, and complex II catalyzes the reaction of succinate into fumarate. Complex II has been identified as a source of reserve respiratory capacity in neonatal rat cardiomyocytes previously <sup>14</sup>. We found two energy reserve genes that were down-regulated by HIF-1. aagr-1 and aagr-2, which are predicted to function in glycogen catabolism <sup>15</sup>. Three distinct genes involved in energy reserve were up-regulated. These genes were ogt-1, which encodes O-linked GlcNac Transferase gene; T04A8.7, encoding an ortholog

of human glucosidase, acid beta (GBA); and T22F3.3, encoding ortholog of human glycogen phosphorylase isozyme in the muscle (PYGM).

### Protein synthesis and degradation

hif-1(lf) is also known to inhibit protein synthesis and translation in varied ways. <sup>16</sup>. Most reported effects of HIF-1 on the translation machinery are posttranslational, and no reports to date show transcriptional control of the ribosomal machinery in C. elegans by HIF-1. We used the WormBase Enrichment Suite Gene Ontology dictionary? to extract 143 protein-coding genes annotated as 'structural constituents of the ribosome' and we queried whether they were differentially expressed in our mutants. egl-9(lf), vhl-1(lf), vhl-1(lf), and egl-9(lf); vhl-1(lf) showed differential expression of 91 distinct ribosomal constituents (not all constituents were detected in all genotypes). For every one of these genotypes, these genes were always down-regulated. In contrast, hif-1(lf) showed up-regulation of a single ribosomal constituent.

Next, we asked whether HIF-1 has any transcriptional effects on the proteasomal constituents; no such effects of HIF-1 on the proteasome have been reported in *C. elegans*. Out of 40 WormBase-annotated proteasomal constituents, we found 31 constituents whose genes were downregulated in at least two out of the four genotypes we studied. Every gene we found was down-regulated in at least two out of the four genotypes we studied.

## A cellular view of hypoxia

In addition to reconstructing the pathway, our dataset allowed us to observe a wide variety of physiologic changes that occur as a result of the HIF-1-dependent hypoxia response. In particular, we observed down-regulation of most components of the TCA cycle and the mitochondrial electron transport chain with the exceptions of fum-1 and the mitochondrial complex II. The mitochondrial complex II catalyzes the reaction of succinate into fumarate. In mouse embryonic fibroblasts, fumarate has been shown to antagonize HIF-1 prolyl hydroxylase domain (PHD) enzymes, which are orthologs of EGL-9<sup>17</sup>. If the inhibitory role of fumarate on PHD enzymes is conserved in C. elegans, upregulation of complex II by HIF-1 during hypoxia may increase intracellular levels of fumarate, which in turn could lead to elevated levels of HIF-1 even after normoxia resumes. The increase in fumarate produced by the complex could be compensated by increasing expression of fum-1. Increased fumarate degradation allows C. elegans to maintain plasticity in the hypoxia pathway, keeping the pathway sensitive to oxygen levels.

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