Identifying Longevity Associated Genes by Integrating Gene Expression and Curated Annotations

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Summary

Aging is a complex process with poorly understood genetic mechanisms. Recent studies have sought to classify genes as pro-longevity or anti-longevity using a variety of machine learning algorithms. However, it is not clear which types of features are best for optimizing classification performance and which algorithms are best suited to this task. Further, performance assessments based on held-out test data are lacking. We systematically compare five popular classification algorithms using gene ontology and gene expression datasets as features to predict the pro-longevity versus anti-longevity status of genes for two model organisms (*C. elegans* and *S. cerevisiae*) using the GenAge database as ground truth. We find that elastic net penalized logistic regression performs particularly well at this task. Using elastic net, we make novel predictions of pro- and anti-longevity genes that are not currently in the GenAge database.

1 Introduction

Identifying the genetic and molecular basis of aging is a longstanding goal in medical science (Johnson & Lithgow, 1992; Remolina, Chang, Leips, Nuzhdin, & Hughes, 2012). Many studies have investigated whether individual genes are pro-longevity or anti-longevity on a case-by-case basis (Ailion, Inoue, Weaver, Holdcraft, & Thomas, 1999). Typically, an intervention such as a knockout/knockdown or overexpression is applied to a small number of genes in a model organism such as nematode worm (Caenorhabditis elegans) or yeast (Saccharomyces cerevisiae) followed by quantification of lifespan. A gene is considered pro-longevity if its expression is directly related to lifespan — for instance, if overexpression increases lifespan or underexpression decreases lifespan (Tacutu et al., 2018). Conversely, a gene is considered anti-longevity if its expression is inversely related to lifespan. Meanwhile, many genes do not fall clearly into either category, for instance, a gene might have no discernable effect on lifespan. The GenAge database (Tacutu et al., 2018) contains a catalogue of putative pro- and anti-longevity genes based on current evidence.

Pro/anti-longevity genes can be identified by intervening on individual genes, but this is slow and expensive. Alternatively, a common technique is to randomly knock out or disrupt many genes in a population of organisms, screen for the longest living individuals, and then determine which genes were disrupted in these individuals. This screening technique can rapidly identify anti-longevity genes, but systematically identifying pro-longevity genes is less straightforward. Indeed, among the small number of genes annotated as having some impact on longevity in worms and yeast, there are considerably more anti-longevity genes than pro-longevity genes.

To prioritize which genes to investigate and speed up the discovery process, recent studies have sought to computationally predict the effect of gene interventions on aging, using annotations like Gene Ontology (GO) terms (Gene Ontology Consortium, 2019) as predictors. A survey of such efforts is provided by Fabris, de Magalhães, and Freitas (2017). However, these recent studies suffer from several limitations. First, annotations like GO may be biased by the scope of the existing literature (Haynes, Tomczak, & Khatri, 2018). Second, it is difficult to compare results across studies since there is a lack of consistency in the choice of algorithms, feature sets, and predictive target/outcome. Finally, most recent studies do not report predictive performance on a held-out test dataset, leading to possible overestimation of performance.

We address these gaps by systematically assessing the performance of five popular machine learning algorithms on the task of predicting the pro- versus anti-longevity status of genes in *S. cerevisiae* and *C. elegans*. We use a consistent outcome in all comparisons based on GenAge annotations (Tacutu et al., 2018). We compare the efficacy of GO terms versus gene expression profiles as feature sets for prediction. Further, we predict possible pro/anti-longevity genes that are not currently annotated in GenAge to suggest directions for future experimental studies.

2 Results

2.1 Data sources and algorithms

We compare the performance of five machine learning classification algorithms: elastic net penalized logistic regression (pglm) (Friedman, Hastie, & Tibshirani, 2010), support vector machine with radial basis function (svm) (Karatzoglou, Smola, Hornik, & Zeileis, 2004), gradient boosted trees (xgb) (T. Chen et al., 2019), naive Bayes (nb) (Majka, 2019), and k-nearest neighbors (knn) (Schliep & Hechenbichler, 2016).

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We define the outcome (that is, the target of prediction) to be the pro- versus anti-longevity annotation of individual genes from GenAge. After data cleaning, we identified 398 yeast genes and 848 worm genes with unambiguous annotations. Of these, the majority were labeled as anti-longevity (347 for yeast and 565 for worm). For validation and comparison, in yeast, we also consider replicative lifespan (RLS) outcome data for a comprehensive set of 4,698 single-gene deletions (McCormick et al., 2015). (In yeast, it is more common to use replicative lifespan rather than chronological lifespan to study aging (McCormick et al., 2015)).

As features for prediction, we consider using GO terms (Gene Ontology Consortium, 2019) and ARCHS4 gene expression profiles (Lachmann et al., 2018) for both yeast and worm. For yeast only, we also consider using the Deleteome dataset (Kemmeren et al., 2014), which contains gene expression profiles for nearly 1500 single-gene deletions. For worm only, we also consider using the Worm Cell Atlas dataset (Cao et al., 2017), which contains gene expression profiles for around 50,000 cells. We write GXP to signify Deleteome and Worm Cell Atlas for worm and yeast, respectively. Altogether, we compare the performance of five feature sets for each species: (1) ARCHS4 alone, (2) GO alone, (3) GXP alone (Deleteome for yeast, Worm Cell Atlas for worm), (4) GO combined with ARCHS4, and (5) GO combined with GXP. Normalization, filtering, and other preprocessing steps are described in the Methods section.

To predict whether a particular gene g is pro- or anti-longevity, we construct features in the following manner. Each GO term is considered a separate binary feature taking a value of one if gene g is annotated to the term and zero otherwise. For the ARCHS4, Deleteome, and Worm Cell Atlas data each experimental condition (e.g., a perturbation or tissue sample) is considered a feature and its value is given by the expression of gene g under that condition. Note that this is the transpose of how gene expression data are usually investigated. However, by treating experimental conditions as features and genes as observations, this allows us to exploit arbitrary gene expression data for gene g, not just data from when g is perturbed.

2.2 Comparative performance of algorithms and feature sets

To assess predictive performance, we use the following cross-validation scheme. For each of the two species, we split the GenAge-annotated genes into five cross-validation folds, and then for each combination of fold, algorithm, and feature set, we compute the area under the receiver-operator curve (AUC). Thus, in total, we compute $2 \times 5 \times 5 \times 5 = 250$ AUC values, 50 for each algorithm (Figures S1,S2).

To summarize the relative performance of the five algorithms, Figure 1 shows how frequently algorithm a has higher AUC than algorithm b for each pair a, b. More precisely, for each pair of algorithms, Figure 1 shows the fraction of times algorithm a has higher AUC than algorithm b across the 50 combinations of species, fold, and feature set. The pglm and svm algorithms consistently outperform the others in terms of AUC. The ranking of algorithms is unchanged when compared using only yeast data. Using only worm data, svm slightly outperforms pglm (0.52 instead of 0.46 in Figure 1), and knn slightly outperforms nb (0.56 instead of 0.34 in Figure 1).

To compare the relative performance of the five different feature sets, Figure 2 shows boxplots of the AUC values over the five cross-validation folds, stratified by species, algorithm, and feature set. For visual clarity, here we only show the results for pglm and svm (the two best algorithms); see Figure S2 for the other algorithms. Generally speaking, using GO terms yields better predictions than gene expression features alone (ARCHS4 or GXP). However, combining GO with gene expression (GO+ARCHS4 or GO+GXP) tends to increase AUC performance compared to GO alone.

Comparing gene expression feature sets, the ARCHS4 features give better performance than GXP (Worm Cell Atlas) for worms, but for yeast, GXP (Deleteome) is superior to ARCHS4. This could be simply due to the fact that the number of features in the worm ARCHS4 data is much larger than in the Worm Cell Atlas data. Alternatively, it could be due to the greater variation in experimental conditions across Deleteome features (which covers a comprehensive set of gene knockouts) compared to Worm Cell Atlas features (which consists of expression profiles of different cell types in normal worms).

Overall, for worms, pglm with GO+ARCHS4 features yields the best performance, whereas for yeast, pglm with GO+GXP is best (Figure 3).

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2.3 Novel predictions of pro/anti-longevity genes

Given the encouraging performance of pglm for predicting pro/anti-longevity genes in GenAge, we applied the algorithm to make novel predictions of pro/anti-longevity genes in *C. elegans* (worm) and *S. cerevisiae* (yeast). To do this, for each species separately, we retrained a pglm model on the full GenAge database, using the combined GO terms plus ARCHS4 gene expression as features (see the Methods section for details on hyperparameter selection). We then used the trained model to generate a predictive score for the pro/anti-longevity effect of each gene not in the GenAge database. Specifically, the predictive score is defined to be the probability that the gene is pro-longevity under the trained model. A score close to 1 indicates that the gene is predicted to be pro-longevity, whereas a score close to 0 indicates that the gene is predicted to be anti-longevity. An intermediate score indicates a gene with unclear pro- or anti-longevity status. Table 1 shows the unannotated genes with the highest confidence levels of being pro- and anti-longevity for worm and yeast, respectively. These genes do not significantly overlap with predictions from the pglm model trained using only GO terms as features (Tables S2-S5), suggesting that these predictions are not simply recapitulating the known biology represented in the GO terms.

To assess the accuracy of the predictions, we looked at the literature to see if there is experimental evidence of pro/anti-longevity effects for these genes. Based on the existing experimental evidence, we find that the model predictions are remarkably good. It turns out that—even though they are not in GenAge yet—there is experimental evidence for the pro/anti-longevity status of most of the predicted genes.

2.3.1 Predicted pro-longevity worm genes

For many of the predicted pro-longevity genes in Table 1, there already exists direct experimental evidence of pro-longevity status. Note that this evidence was not used in making the predictions, implying that the model is producing reliable out-of-sample predictions. We discuss what is known about the top 10 predicted pro-longevity genes: CLEC-196, F44E5.4, CEH-13, LPR-3, HIL-7, W04A8.4, TTH-1, GST-1, F44E5.5, and F20C5.6.

F44E5.4 and F44E5.5 encode members of the hsp70 family of heat shock proteins. The heat shock response is well-known to have strong pro-longevity effects in C. elegans. Indeed, knocking in extra copies of hsp70 extends lifespan (Yokoyama et al., 2002) and knocking down hsp70 via RNAi decreases lifespan and leads to rapid aging phenotypes (Kimura, Tanaka, Nakamura, Takano, & Ohkuma, 2007). GST-1 (Glutathione S-transferase P) is also involved in stress response—particularly, immune response—and GSTs are well-known to be pro-longevity. Overexpression (underexpression) of GSTs has been found to increase (decrease, respectively) lifespan and stress resistance (Avvadevara et al., 2007, 2005). W04A8.4 is an uncharacterized protein that is involved in the pro-longevity effect of metformin on C. elegans (Wu et al., 2016); specifically, knockdown of W04A8.4 leads to metformin resistance. This is intriguing, since metformin treatment has been shown to promote health and extend lifespan in many organisms. Homeobox protein CEH-13 exhibits pro-longevity characteristics based on experimental evidence — specifically, a ceh-13 mutant strain has decreased lifespan compared to wildtype controls (Tihanyi et al., 2010). LPR-3 (LiPocalin-Related protein) is known to be involved in nematode worm locomotion, and appears to mediate the longevity-inducing effect of daf-7 mutation (Hyun, Kim, Dumur, Schroeder, & You, 2016); additionally, expression of lpr-3 is increased in worms fed with $rBm\alpha TX14$, an α -neurotoxin that increases worm lifespan (L. Chen et al., 2016).

For the remainder of the genes in Table 1, there is suggestive experimental evidence of pro-longevity status based on associations. C-type Lectin clec-196 expression increases and lifespan increases when hsb-1 is knocked out (Sural, Lu, Jung, & Hsu, 2019). Also, clec-196 is directly adjacent to hsp-1 on chromosome IV, suggesting possible co-involvement, and hsp-1 (heat shock protein) is well-known to be pro-longevity. HIL-7 (Histone H1 Like) gene expression may be associated with Ethosuximide treatment, a drug that increases worm lifespan and affects DAF-16/FOXO target gene expression (X. Chen et al., 2015). TTH-1 (Thymosin beta) is significantly increased in daf-2 mutants, which are very long-lived, suggesting possible pro-longevity status by association (Narayan et al., 2016). F20C5.6 is affected by the well-known longevity genes clk-1 and sir-2.1, as well as by treatment with 1-methylnicotinamide and rotenone, which are well-known for increasing worm lifespan.

This validating evidence from the literature indicates that the model predictions are surprisingly accurate. The predicted pro-longevity genes CLEC-196, HIL-7, TTH-1, and F20C5.6 are candidates for further

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experimental exploration.

2.3.2 Predicted anti-longevity worm genes

Similarly to the predicted pro-longevity genes, there exists experimental evidence of anti-longevity status of most of the predicted anti-longevity genes in Table 1. We discuss what is known about the top 10 predicted anti-longevity genes: MSP-59, Y59E9AR.7, RPL-39, MSP-57, MSP-81, MSP-113, MSP-19, NLP-27, and RPL-11.1.

Major sperm proteins appear to be anti-longevity based on the experimental evidence. A mutation reducing sperm production leads to significantly increased lifespan (Van Voorhies, 1992). Additionally, the expression of sperm-related genes—especially major sperm protein (MSP) genes—is decreased in adult daf-2 mutants, providing further support for an anti-longevity role of MSP genes (Halaschek-Wiener et al., 2005).

RSP-39 and RPL-11.1 are 60S ribosomal proteins. RNAi knockdown of genes encoding ribosomal proteins consistently increases lifespan in *C. elegans*, both in the case of 40S and 60S ribosomal proteins (Hansen et al., 2007). This supports the predicted anti-longevity status.

NLP-27 (Neuropeptide-Like Protein) is the only other predicted anti-longevity gene in the top 10 list. Expression of nlp-27, along with other nlp genes, is increased in long-lived daf-2 mutants. Further, nlp-27 expression is reduced in a short-lived mir-71 deletion strain. This indirect evidence by association suggests a possible pro-longevity role of NLP-27—which would contradict the predicted anti-longevity—but direct over/under-expression of nlp-27 would be needed to establish its pro/anti-longevity status.

2.3.3 Predicted pro-longevity yeast genes

Table 1 lists the top 10 predicted pro-longevity yeast genes. Several of these predictions are borne out by direct experimental evidence via single-gene deletions — specifically, deletion of ACS1, ETR1, UBI4, and POR1 leads to decreased lifespan (Marek & Korona, 2013). Marek and Korona (2013) did not find a significant pro- or anti-longevity effect for UBC5, HSP12, or SBA1, and they do not report results for the remainder of the top 10 genes. However, UBC5 is a strong pro-longevity candidate, since it is involved in cellular stress response and mediates selective degradation of short-lived and abnormal proteins (Seufert & Jentsch, 1990). HSP12 (heat shock protein) is required for the lifespan-extending effect of dietary restriction in yeast (Herbert et al., 2012), validating the pro-longevity prediction. SBA1 is also a strong pro-longevity candidate, as a chaperone-binding protein that is involved in heat shock response and is required for telomere length maintenance (Fang, Fliss, Rao, & Caplan, 1998; Toogun, Zeiger, & Freeman, 2007). PRE3 and PRE7 are part of the proteasome, and it is known that increased proteasome capacity extends lifespan (Kruegel et al., 2011), providing indirect validation of their predicted pro-longevity status. PDI1 is a downstream target of the unfolded protein response (UPR), which is well-known to be pro-longevity (Patil & Walter, 2001).

2.3.4 Predicted anti-longevity yeast genes

Table 1 lists the top 10 predicted anti-longevity yeast genes. As in worms, depletion of ribosomes increases lifespan (Steffen et al., 2008), validating the predictions of the ribosome-biogenesis proteins RPS30B, TMA23, RPS29B, and RLP24 as anti-longevity. HOR7 is reported to influence lifespan, but the direction of the effect may be context-dependent: HOR7 deletion increases lifespan (McCormick et al., 2015), whereas Schleit et al. (2013) find that HOR7 deletion decreases lifespan under dietary restricted conditions.

For URA3, COX9, TOM7, MFA1, and TAR1, we do not find pre-existing corroboration of the predicted anti-longevity status in the literature. TOM7 deletion has been reported to decrease chronological lifespan (Garay et al., 2014), and it does not appear to have a strong effect on replicative lifespan (Marek & Korona, 2013). TOM7 is part of the translocase of the outer mitochondrial membrane (TOM) complex, and the mitochondrial membrane is well-known to be important in yeast longevity (Jazwinski, 2005). Marek and Korona (2013) report a pro-longevity effect for COX9, contrary to the model prediction. (Except for COX9, the Marek and Korona (2013) results are inconclusive for all of the genes in Table 1.) Further investigation of URA3, COX9, TOM7, MFA1, and TAR1 might be interesting to pursue.

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2.4 Validation on a secondary dataset

To further evaluate the predictive accuracy of the trained pglm model, we compare the model predictions to actual lifespan measurements from a non-GenAge validation dataset. For this purpose, we use a dataset of replicative lifespan for a comprehensive set of 4,698 single-gene deletions in yeast (McCormick et al., 2015). Since the McCormick et al. (2015) dataset contains lifespan measurements for deletions of many genes that do not appear in GenAge, in principle it should be well-suited as a secondary validation dataset. Using the pglm model trained on the full GenAge database for yeast with the GO+ARCHS4 feature set as predictors, we made predictions of the longevity effect of all 4,698 genes in the McCormick dataset.

First, as a sanity check, we observe that among genes in GenAge, the predicted probability of a gene being pro-longevity is clearly inversely related to the change in lifespan after deletion (Figure 4, left panel). This is not surprising since it simply means that the GenAge annotations are roughly consistent with the McCormick data, and the model was able to fit the GenAge-based training data. More interestingly, we see that the model is able to predict which genes have a larger or a smaller effect on lifespan (Figure 4, left panel). For instance, among pro-longevity genes, the genes with predicted probability near 1 do indeed tend to lead to a larger decrease in lifespan. Meanwhile, among anti-longevity genes, the genes with predicted probability near 0 do indeed tend to lead to a larger increase in lifespan. Since the training data contain no information about the magnitude of the effect on lifespan, this indicates that the model is not simply recapitulating the training data, but is indeed making generalizable predictions.

Next, we compare the model predictions to the lifespan data for genes outside the GenAge database. Figure 4 (right panel) shows the change in lifespan versus the predicted probability of a gene being prolongevity, for genes in the McCormick dataset that are not in GenAge. A downward trend in this plot would indicate concordance between model predictions and the validation data. There is an extremely slight but not convincing downward trend; thus, while suggestive, this does not provide a compelling out-of-sample validation of the model predictions. Note that the pglm classifier trained on GenAge has a strong bias toward predicting genes to be anti-longevity; see Figure 4 (right panel) and Figure S3. This bias is due to class imbalance in the training data, since the majority of genes annotated in GenAge are anti-longevity. This is common when the training data are imbalanced, and can easily be addressed by selecting the classification threshold to yield appropriately balanced predictions.

The lack of concordance between the out-of-sample model predictions and the McCormick lifespan data may be attributable to the fact that for many genes, the McCormick data are not in agreement with the GenAge annotations of pro/anti-longevity. Specifically, many putatively pro-longevity genes led to large increases in lifespan when deleted, and many putatively anti-longevity genes led to large decreases in lifespan when deleted (Figure 4, left panel). It is not clear whether this discrepancy is primarily due to limitations of the GenAge database (e.g., bias and relatively small sample size) or limitations of the McCormick assay. Focusing on the latter possibility, recent studies have identified mechanisms by which disruption of a gene through knockout can activate compensatory mechanisms leading to a dramatically different phenotype than disruption of the same gene through knockdown, which reduces but does not eliminate expression (Wilkinson, 2019). If deletion of a single gene activates similar compensatory mechanisms in yeast, then this could explain the lack of concordance, since it would imply that the change in lifespan under a single-gene deletion is not necessarily related to that gene's pro/anti-longevity status. A comprehensive assay of knockdowns (rather than deletions or knockouts) would shed light on this intriguing question.

2.5 Functional interpretation of model predictions

To interpret the biological basis for the model predictions in terms of functional categories, for each species we retrained the pglm model on the full GenAge dataset using only GO terms as features. We extracted the 20 most influential GO terms from the trained model by ranking the regression coefficients from largest to smallest in absolute value (Table 2). Note that in this model, the coefficient is equal to the log-odds ratio (logOR) of a gene being pro-longevity when it is annotated to a GO term versus when it is not annotated to that GO term. If a GO term has a positive logOR value, then genes annotated with that GO term are more likely to be pro-longevity under the model. Conversely, a negative logOR indicates that genes annotated with that GO term are more likely to be anti-longevity.

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2.5.1 Top GO terms for worm

The current literature supports a strong longevity effect for many of the top categories in Table 2. Translation inhibition is known to increase lifespan (Hansen et al., 2007), so a large negative coefficient for the translation and ribosome GO terms makes sense. Protein homeostasis is known to be key to longevity (Sampaio-Marques & Ludovico, 2018), so it makes sense that the model has positive coefficients for protein transport, endoplasmic reticulum membrane, and endoplasmic reticulum. Ubiquitin-mediated proteolysis is known to be important for promoting longevity, implying that a positive coefficient for ubiquitin-dependent protein catabolic process makes sense. Heat shock response is known to extend lifespan, and indeed, the model has a positive coefficient for response to heat. Activation of the mitochondrial unfolded protein response is known to promote longevity (Durieux, Wolff, & Dillin, 2011), so a positive coefficient for protein import into mitochondrial matrix makes sense. Mitochondria are known to be important for longevity (Sun, Youle, & Finkel, 2016), so a large coefficient for mitochondria makes sense; further, inhibition of mitochondrial respiration is known to extend lifespan (B Hwang, Jeong, & Lee, 2012), so a negative sign for the coefficient could make sense. Similarly, the importance of DNA repair makes sense, and surprisingly, in some cases, DNA repair gene knockdown increases lifespan, possibly due to compensatory biological mechanisms (Lans et al., 2013); thus, a negative coefficient is, in fact, consistent with the literature.

2.5.2 Top GO terms for yeast

For yeast, Table 2 shows the top longevity-related GO terms in the model. The importance of these terms is consistent with the current literature, but the appropriate sign of the coefficient is not always clear, since the genes annotated to each GO term may have contradictory pro/anti-longevity effects and further, there may be compensatory relationships between terms due to correlated predictors.

Replicative cell aging, apoptotic process, and cell cycle obviously make sense as related to yeast aging and longevity. Mitochondrial membrane maintenance is known to be important in yeast longevity (Jazwinski, 2005), and other membranes (e.g., the vacuole membrane) may also be important (Carmona-Gutierrez, Hughes, Madeo, & Ruckenstuhl, 2016); thus, large coefficients for mitochondrion, integral component of mitochondrial outer membrane, mitochondrial intermembrane space, membrane, membrane fraction, and transmembrane transport are consistent with the literature. Depletion of ribosomes is known to increase lifespan (Steffen et al., 2008), so a negative coefficient for chromatin silencing at rDNA is appropriate. Telomeres are known to be important in yeast longevity (Austriaco & Guarente, 1997; Liu, Wang, Wang, & Liu, 2019), so a large coefficient for telomere maintenance makes sense. Longevity effects of cellular response to oxidative stress are corroborated in the literature (Postnikoff, Johnson, & Tyler, 2017). Finally, a negative coefficient for zinc ion binding is consistent with experimental evidence that zinc limitation extends chronological lifespan (Shimasaki et al., 2017).

3 Discussion

We systematically compared the performance of popular machine learning algorithms in classifying genes as pro- or anti-longevity using the GenAge database and combinations of gene expression and gene ontology (GO) feature sets. We identified elastic net penalized logistic regression (pglm) as the most effective classifier and made predictions for unannotated genes. The pglm model fit to GenAge data was only weakly concordant with the McCormick replicative lifespan (RLS) assay, which was based on single gene deletion strains in yeast. This discrepancy could be due to compensatory mechanisms which are known to mitigate the effects of knockouts (Wilkinson, 2019). We suggest future comprehensive longevity assays should consider knockdowns instead of deletions and knockouts. Furthermore, there is a need for increased focus on pro-longevity genes rather than anti-longevity genes, since the latter are much more common in the GenAge database. We offer our predictive probability scores as one possible tool to prioritize future experimental studies which can validate individual genes as pro-longevity mechanistically. We encourage computational researchers to use metrics such as area under receiver-operator curve (AUC) on held-out data from standard databases such as GenAge to assess classification performance and facilitate comparisons across studies. Finally, our approach of combining feature sets to improve predictive performance is generalizable in principle to a wider variety of model organisms as more annotations and datasets become available over time.

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4 Experimental Procedures

Binary pro/anti-longevity annotations were accessed from the GenAge model organisms database (Tacutu et al., 2018), currently available at http://genomics.senescence.info/genes. We used the subset of genes for yeast and worm, and we excluded ambiguous annotations (e.g., if GenAge lists two studies for a gene, one finding it to be pro-longevity and the other finding it to be anti-longevity). GO annotations for all genes were downloaded from the BioMart ENSEMBL database using the biomaRt package in Bioconductor. For both species, gene expression data in the form of RNA-Seq read counts were obtained from the ARCHS4 database (Lachmann et al., 2018), currently available at https://amp.pharm.mssm.edu/archs4/archs4zoo.html. For yeast only, we acquired the Deleteome gene expression microarray dataset (Kemmeren et al., 2014), currently available at http://deleteome.holstegelab.nl. For worm only, we obtained gene expression data from the single-cell RNA-Seq Worm Cell Atlas (Cao et al., 2017), currently available at http://atlas.gs.washington.edu/worm-rna. We reduced the dimensionality of the Worm Cell Atlas data by summing the unique molecular identifier (UMI) counts across all cells within the same tissue, so that each feature is a "pseudobulk" tissue rather than a single cell.

Replicative lifespans (RLS) for 4,698 single-gene deletion yeast strains were obtained from the authors of McCormick et al. (2015). Perturbation genotypes with percent_change greater than 30 and set_lifespan_count less than or equal to 5 were excluded based on the authors' recommendations. We merged results for the same genotype across replicate experiments in the following way. The outcome for each genotype in a single replicate was quantified as the mean of RLS in the perturbation group minus the mean of RLS in the control group. To obtain a single value for the genotype across all replicates, we then computed a weighted average of the outcome values from each replicate, where the weights corresponded to the sample sizes in each group. This ensured that replicates with more observations contributed more to the final value. We refer to this as the McCormick dataset.

All gene expression measurements were normalized to account for sample-specific biases. Specifically, the Deleteome data were already normalized, the ARCHS4 read counts were converted to transcripts-per-million (TPM), and the Worm Cell Atlas UMIs were converted to counts-per-million (CPM). The normalized counts were then log transformed with a pseudocount of one. For Deleteome, genes that were variable in controls and non-responsive mutants were excluded, since these data were likely to contain mostly noise. For each species, we used the subset of genes with no missing values across all feature types (GO features and the two sources of gene expression features), resulting in 703 worm genes (246 pro-longevity, 457 anti-longevity) and 368 yeast genes (46 pro-longevity, 322 anti-longevity). Features with no variation across the included genes were discarded. For yeast, the number of retained features was 3268, 700, and 1390 for ARCHS4, Deleteome, and GO terms, respectively. For worms, the number of features was 2935, 270, and 2051 for ARCHS4, Worm Cell Atlas, and GO terms, respectively. All gene expression features were centered and scaled to have mean zero and standard deviation 0.5 as suggested by Gelman (2008), while binary features (GO) were not centered and scaled. The five sets of features considered for each species were (1) ARCHS4 alone, (2) GO alone, (3) GXP alone (Deleteome for yeast, Worm Cell Atlas for worm), (4) GO combined with ARCHS4, and (5) GO combined with GXP.

To assess predictive performance of different combinations of feature sets, each dataset (consisting of the binary GenAge outcome for a single species matched with one of the five feature sets) was split into 5 external cross-validation (CV) folds. Within each fold, machine learning classifiers were fit to the training data using the caret R package (Kuhn et al., 2019). The same partitioning of the data was preserved across algorithm runs to ensure identical training and test conditions. The algorithms used were k-nearest neighbors (knn), naive Bayes (nb), gradient boosted trees (xgb), support vector machine with radial basis function (svm), and logistic regression with elastic net penalty (pglm). Hyperparameters (Table S1) were selected by grid search using repeated 10-fold internal CV with two repeats within each training fold using the Kappa criterion. Note that this means each algorithm could potentially use different hyperparameter values across the five external CV folds. For all algorithms except naive Bayes, the grid consisted of default caret values. For naive Bayes, the Laplace correction was set to zero, kernel smoothing was always used, and the adjustment to the probabilities was chosen between 0.5 and 1.0. Additionally, for naive Bayes only, many features with near-zero variance caused numerical instabilities and were excluded. Having chosen a final set of hyperparameters for each training fold, the predicted probabilities were computed for the held-out test data and the area under the receiver-operator curve (AUC) was computed to quantify prediction

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performance (discrimination). An AUC value of 1 indicates perfect classification performance, whereas an AUC of 0.5 signifies performance no better than random, or simply always predicting the majority class.

For the results in Sections 2.3 and 2.4, the best-performing algorithm (pglm) was retrained on all of the GenAge data for each species with the combined GO plus ARCHS4 feature set. The hyperparameter grid was expanded to 21 alpha values (evenly spaced between zero and one, inclusive), and 97 automatically selected lambda values using five-fold CV. For worm, the optimal alpha was 0.05 (close to an L2 ridge penalty). For yeast, the optimal alpha was 0.5 (an even mix between ridge and the L1 lasso penalty). Using the optimal hyperparameters, predictive probabilities were computed for all genes.

For the results in Section 2.5, for each species the pglm algorithm was retrained on the full GenAge dataset using GO features only. This choice of feature set was used to enable interpretation of regression coefficients. Here, the hyperparameter grid was the same 21 alpha values and 97 automatically selected lambda values with five-fold CV. The optimal alpha values were 0.15 for worm and 0.10 for yeast (both closer to ridge than lasso).

To facilitate reproducibility, all code used to produce the results in this manuscript is publicly available at https://github.com/willtownes/longevity-paper (Townes, F. William & Miller, Jeffrey W., 2020).

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Author Contributions

JWM conceived of the study. FWT assembled the datasets, performed the analyses, and drafted the initial report under JWM's supervision. JWM revised the manuscript and provided biological interpretations of results. Both authors approved of the final manuscript.

Longevity Associated Genes

Tables and Figures

Table 1: Top pro-longevity and anti-longevity genes not in GenAge predicted using GO terms and ARCHS4 gene expression for worm and yeast with the pglm (GLM-Net) algorithm.

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regulation of respiratory metabolism

Table 2: Top GO terms identified by the pglm (GLM-Net) algorithm. logOR: log-odds ratio. OR: odds ratio. Positive logOR indicates a gene annotated to that GO term is more likely to be pro-longevity. BP: biological process, CC: cellular component, MF: molecular function.

Species	ID	logOR	OR	Type	Description
worm	GO:0006412	-0.98	0.38	BP	translation
World	GO:0005634	0.89	2.4	CC	nucleus
	GO:0015031	0.82	2.3	BP	protein transport
	GO:0005789	0.77	2.1	CC	endoplasmic reticulum membrane
	GO:0005840	-0.73	0.48	CC	ribosome
	GO:0009792	0.68	2	BP	embryo development ending in birth or egg hatching
	GO:0006511	0.66	1.9	BP	ubiquitin-dependent protein catabolic process
	GO:0009408	0.66	1.9	BP	response to heat
	GO:0043005	0.65	1.9	CC	neuron projection
	GO:0030150	0.6	1.8	BP	protein import into mitochondrial matrix
	GO:0055120	-0.6	0.55	CC	striated muscle dense body
	GO:0005783	0.6	1.8	CC	endoplasmic reticulum
	GO:0046872	-0.53	0.59	MF	metal ion binding
	GO:0005739	-0.52	0.59	CC	mitochondrion
	GO:0006281	-0.52	0.59	BP	DNA repair
	GO:0035556	-0.52	0.59	BP	intracellular signal transduction
	GO:0045893	0.52	1.7	BP	positive regulation of transcription; DNA-templated
	GO:0008289	0.52	1.7	MF	lipid binding
	GO:0048477	0.5	1.6	BP	oogenesis
	GO:0003824	0.49	1.6	MF	catalytic activity
yeast	GO:0001302	1.8	5.8	BP	replicative cell aging
	GO:0006915	0.87	2.4	BP	apoptotic process
	GO:0016020	-0.82	0.44	$^{\rm CC}$	membrane
	GO:0005634	0.73	2.1	$^{\rm CC}$	nucleus
	GO:0000183	0.72	2.1	BP	chromatin silencing at rDNA
	GO:0005624	0.71	2	$^{\rm CC}$	membrane fraction
	GO:0007049	0.67	1.9	BP	cell cycle
	GO:0005739	0.64	1.9	CC	mitochondrion
	GO:0005515	-0.64	0.53	MF	protein binding
	GO:0003824	0.61	1.8	MF	catalytic activity
	GO:0031307	0.56	1.7	CC	integral component of mitochondrial outer membrane
	GO:0000723	0.55	1.7	BP	telomere maintenance
	GO:0005758	0.53	1.7	CC	mitochondrial intermembrane space
	GO:0055085	0.53	1.7	BP	transmembrane transport
	GO:0017111	0.52	1.7	MF	nucleoside-triphosphatase activity
	GO:0006811	0.51	1.7	BP	ion transport
	GO:0006281	0.5	1.7	BP	DNA repair
	GO:0034599	0.48	1.6	BP	cellular response to oxidative stress
	GO:0008270	-0.48	0.62	MF	zinc ion binding
	GO:0045861	0.47	1.6	BP	negative regulation of proteolysis

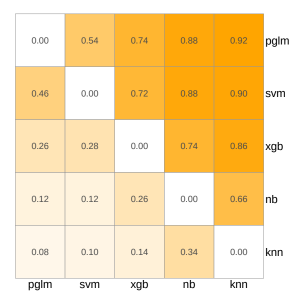


Figure 1: Ranking machine learning algorithms based on AUC. Numeric values indicate the fraction of times the row algorithm has higher classification performance than the column algorithm. pglm: elastic net penalized logistic regression, svm: support vector machine with radial basis function, xgb: gradient boosted trees, nb: naive Bayes, knn: k-nearest neighbors.

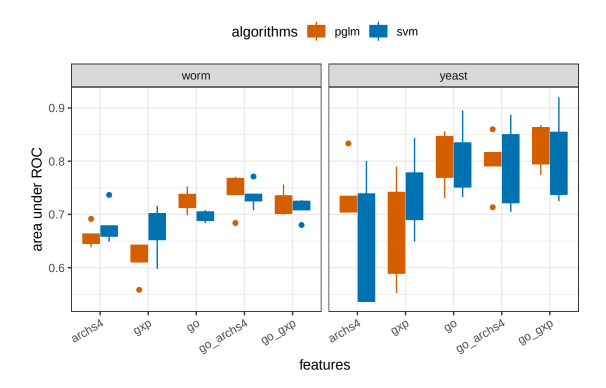


Figure 2: Combining gene expression (archs4, gxp) with gene ontology (GO) features yields improved classification performance in terms of AUC. pglm: elastic net penalized logistic regression, svm: support vector machine with radial basis function. An AUC value of 1 indicates perfect classification, whereas an AUC of 0.5 signifies performance no better than random.

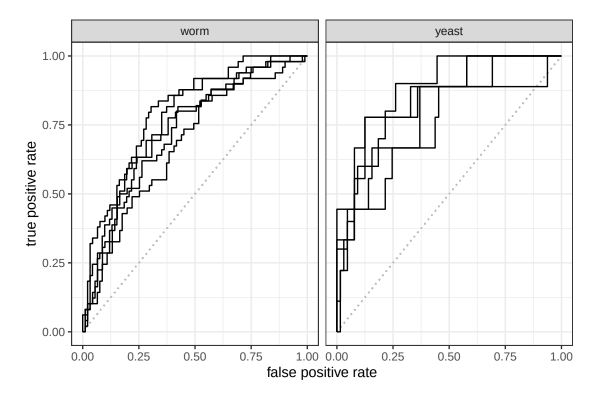


Figure 3: Receiver operator curves (ROC) for the best performing algorithm (pglm: elastic net penalized logistic regression) with the best performing feature sets (GO+GXP for yeast and GO+ARCHS4 for worm). Each curve represents predictive performance on the held-out data from a single cross validation fold. The diagonal gray dotted line indicates the theoretical performance of an untrained random classifier as a baseline.

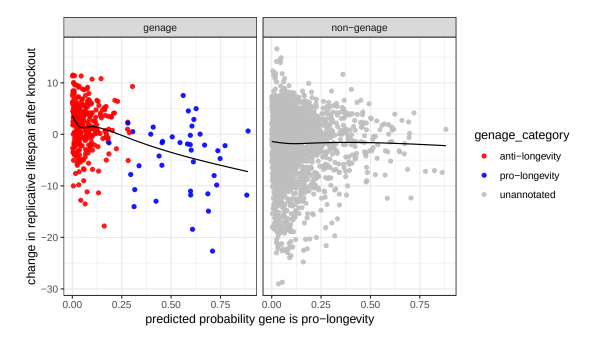


Figure 4: Predicted probability of a gene being pro-aging versus effect of deletion on replicative lifespan (RLS) in yeast. Probabilities are from the pglm classifier trained on the full GenAge dataset. Solid curve is a nonparametric smoother.

Supplemental Figures and Tables

Table S1: Hyperparameters used. pglm: elastic net penalized logistic regression, svm: support vector machine with radial basis function, xgb: gradient boosted trees, nb: naive Bayes, knn: k-nearest neighbors.

algorithm	hyperparameter	values
nb	laplace	0
	usekernel	TRUE
	adjust	0.5; 1
knn	kmax	5; 7; 9
	distance	2
	kernel	optimal
xgb	eta	0.3; 0.4
	\max_depth	1; 2; 3
	gamma	0
	$colsample_bytree$	0.6; 0.8
	$\min_$ child $_$ weight	1
	subsample	0.5; 0.75; 1.0
	nrounds	50; 100; 150
svm	С	0.25; 0.5; 1.0
	sigma	automatic
pglm	alpha	0.1; 0.55; 1.0
	lambda	automatic

Table S2: Top worm pro-longevity genes not in GenAge predicted only using GO terms as features. Compare to Table 1.

Gene	Prob	ID	Description
HLH-2	0.942	WBGene00001949	Helix Loop Helix
NPP-20	0.939	WBGene00003806	Protein SEC13 homolog
HMP-2	0.916	WBGene00001979	Beta-catenin-like protein hmp-2
KGB-1	0.909	WBGene00002187	GLH-binding kinase 1
NPP-13	0.908	WBGene00003799	Nuclear pore protein
DCS-1	0.905	WBGene00000940	m7GpppX diphosphatase
APR-1	0.903	WBGene00000156	Adenomatous polyposis coli protein-related protein 1
UNC-51	0.892	WBGene00006786	Serine/threonine-protein kinase unc-51
EMO-1	0.891	WBGene00001303	Protein transport protein Sec61 subunit gamma
PAS-1	0.891	WBGene00003922	Proteasome subunit alpha type-6

Table S3: Top worm anti-longevity genes not in GenAge predicted only using GO terms as features. Compare to Table 1.

Gene	Prob	ID	Description
DAF-11	0.994	WBGene00000907	Receptor-type guanylate cyclase daf-11
RPS-27	0.986	WBGene00004496	40S ribosomal protein S27
RPL-37	0.985	WBGene00004451	60S ribosomal protein L37
W01D2.1	0.985	WBGene00012179	Ribosomal protein L37
RPL-43	0.985	WBGene00004456	60S ribosomal protein L37a

Table S4: Top yeast pro-longevity genes not in GenAge predicted only using GO terms as features. Compare to

Gene	Prob	ID	Description
SIR2	0.833	YDL042C	Conserved NAD+ dependent histone deacetylase of the Sirtuin family
DNL4	0.774	YOR005C	DNA ligase required for nonhomologous end-joining (NHEJ)
MCM5	0.753	YLR274W	Component of the Mcm2-7 hexameric helicase complex
DOC1	0.702	YGL240W	Processivity factor
POR1	0.7	YNL055C	Mitochondrial porin (voltage-dependent anion channel)
NTG1	0.691	YAL015C	DNA N-glycosylase and apurinic/apyrimidinic (AP) lyase
OYE2	0.635	YHR179W	Conserved NADPH oxidoreductase containing flavin mononucleotide (FMN)
EXO1	0.629	YOR033C	5'-3' exonuclease and flap-endonuclease
MCD1	0.601	YDL003W	Essential alpha-kleisin subunit of the cohesin complex

Table S5: Top yeast anti-longevity genes not in GenAge predicted only using GO terms as features. Compare to Table 1.

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Gene	Prob	ID	Description
VPS34	0.997	YLR240W	Phosphatidylinositol (PI) 3-kinase that synthesizes PI-3-phosphate
VPS15	0.996	YBR097W	Serine/threonine protein kinase involved in vacuolar protein sorting
AKR1	0.996	YDR264C	Palmitoyl transferase involved in protein palmitoylation
AKR2	0.995	YOR034C	Ankyrin repeat-containing protein
ENV7	0.995	YPL236C	Vacuolar membrane protein kinase
RPL37A	0.994	YLR185W	Ribosomal 60S subunit protein L37A
RPL43A	0.994	YPR043W	Ribosomal 60S subunit protein L43A
RPS27A	0.993	YKL156W	Protein component of the small (40S) ribosomal subunit
RPS27B	0.993	YHR021C	Protein component of the small (40S) ribosomal subunit
RPS29A	0.993	YLR388W	Protein component of the small (40S) ribosomal subunit

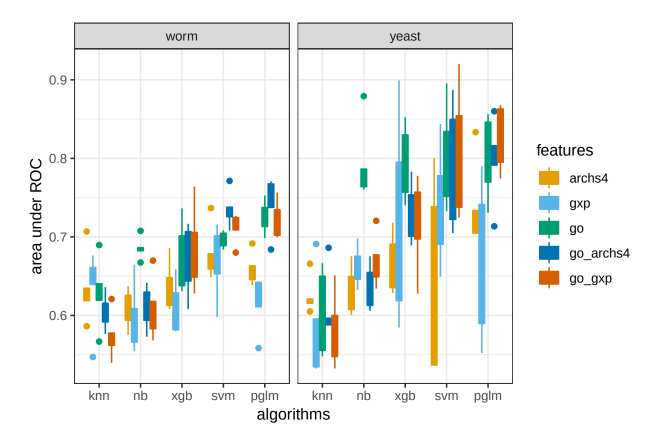


Figure S1: Comparison of predictive performance of machine learning algorithms on classifying genes as pro- or antilongevity. pglm: elastic net penalized logistic regression, svm: support vector machine with radial basis function, xgb: gradient boosted trees, nb: naive Bayes, knn: k-nearest neighbors, gxp: gene expression, ROC: receiver-operator curve.

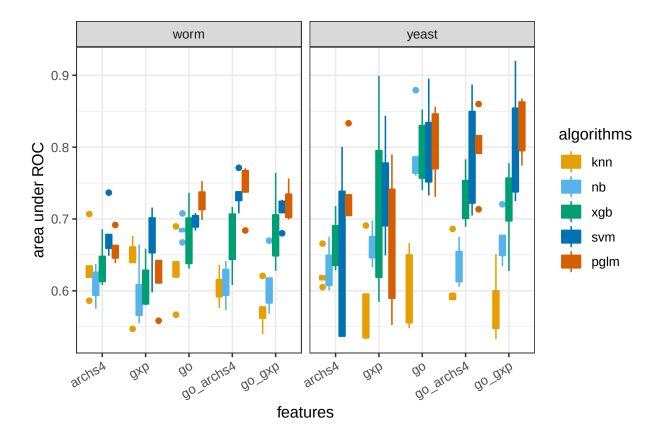


Figure S2: Comparison of predictive performance of different feature sets on classifying genes as pro- or anti-longevity. pglm: elastic net penalized logistic regression, svm: support vector machine with radial basis function, xgb: gradient boosted trees, nb: naive Bayes, knn: k-nearest neighbors, gxp: gene expression, ROC: receiver-operator curve.

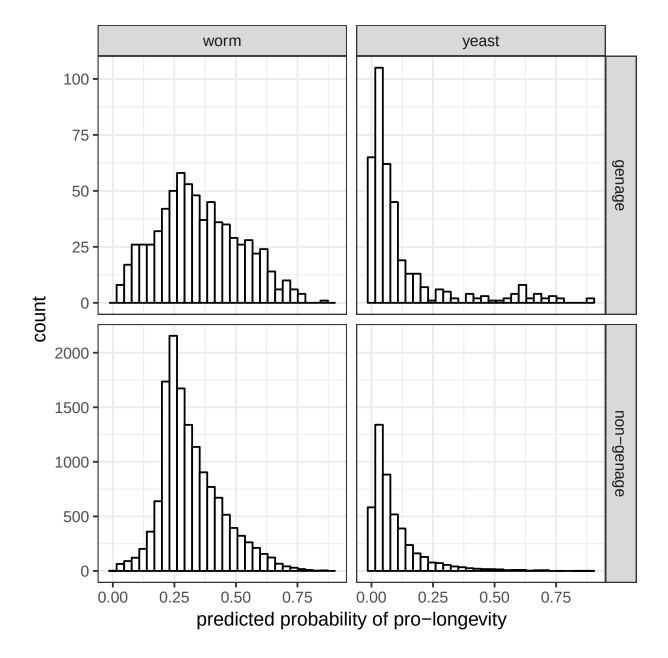


Figure S3: Distribution of predictive probabilities after training elastic net penalized logistic regression (pglm) on the full GenAge dataset with GO terms and ARCHS4 gene expression as features.

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