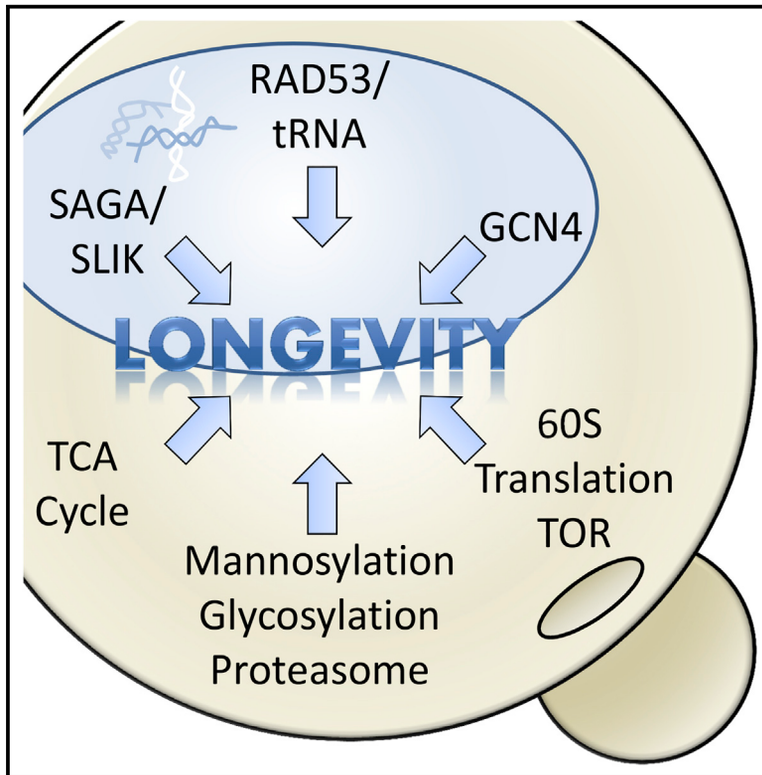


Cell Metabolism

A Comprehensive Analysis of Replicative Lifespan in 4,698 Single-Gene Deletion Strains Uncovers Conserved Mechanisms of Aging

Graphical Abstract



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In Brief

Analysis of 4,698 viable yeast single-gene deletion strains yielded 238 with increased lifespan, in multiple functional groups, showing a significant conservation in longevity pathways between yeast and *C. elegans*. Deletion of tRNA exporter *LOS1* extended lifespan; DNA damage response and mTOR converge on *Los1*-mediated tRNA export to regulate *Gcn4* activity.

Highlights

- 4,698 deletions tested yields the most comprehensive yeast data set on aging
- Longevity clusters center on known, conserved biological processes
- Enrichment of lifespan-extending *C. elegans* orthologs suggests conservation
- Genome-wide information uncovered aging pathways such as tRNA transport

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A Comprehensive Analysis of Replicative Lifespan in 4,698 Single-Gene Deletion Strains Uncovers Conserved Mechanisms of Aging

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SUMMARY

Many genes that affect replicative lifespan (RLS) in the budding yeast *Saccharomyces cerevisiae* also affect aging in other organisms such as *C. elegans* and *M. musculus*. We performed a systematic analysis of yeast RLS in a set of 4,698 viable single-gene deletion strains. Multiple functional gene clusters were identified, and full genome-to-genome comparison demonstrated a significant conservation in longevity pathways between yeast and *C. elegans*. Among the mechanisms of aging identified, deletion of tRNA exporter *LOS1* robustly extended lifespan. Dietary restriction (DR) and inhibition of mechanistic Target of Rapamycin (mTOR) exclude *Los1* from the nucleus in a *Rad53*-dependent manner. Moreover, lifespan extension from deletion of *LOS1* is nonadditive with DR or mTOR inhibition, and results in *Gcn4* transcription factor activation. Thus, the DNA damage response and mTOR converge on *Los1*-mediated nuclear tRNA export to regulate *Gcn4* activity and aging.

INTRODUCTION

Aging has long fascinated and puzzled biologists (Haldane, 1942; Medawar, 1952). In most organisms, a decline in fitness

occurs over time that coincides with increased incidence of several significant causes of morbidity and mortality (Mayeux and Stern, 2012; Simms, 1946). With modern forward and reverse genetic techniques, several groups have shown that single genes can have a dramatic effect on lifespan (Guarente and Kenyon, 2000), and many of these have been found to act together in known signaling pathways (Fontana et al., 2010; Johnson et al., 2013; Kenyon, 2010; Lapierre and Hansen, 2012). Further, the onset of many disadvantageous late life phenotypes is delayed in many long-lived mutant organisms (Garigan et al., 2002; Herndon et al., 2002; Morley et al., 2002).

In yeast, aging can be assayed by at least two methods: chronological lifespan (CLS) and replicative lifespan (RLS) (Kaeberlein, 2010; Longo et al., 2012). The yeast RLS assay measures how many daughter cells a mother can produce before it ceases dividing, and has been proposed as an assay to identify conserved pathways affecting the continued viability of dividing cells such as stem cells in humans (Steinkraus et al., 2008).

A partial genome-to-genome comparison indicated that genes controlling RLS in yeast overlapped with those influencing aging in worms (Smith et al., 2008), which are post-mitotic after adulthood. With the goal of extending this finding to a full genome-to-genome comparison and identifying conserved aging genes, we completed a full genome screen of viable *S. cerevisiae* deletions, identifying 238 long-lived gene deletions, including 189 not previously reported. These cluster into several

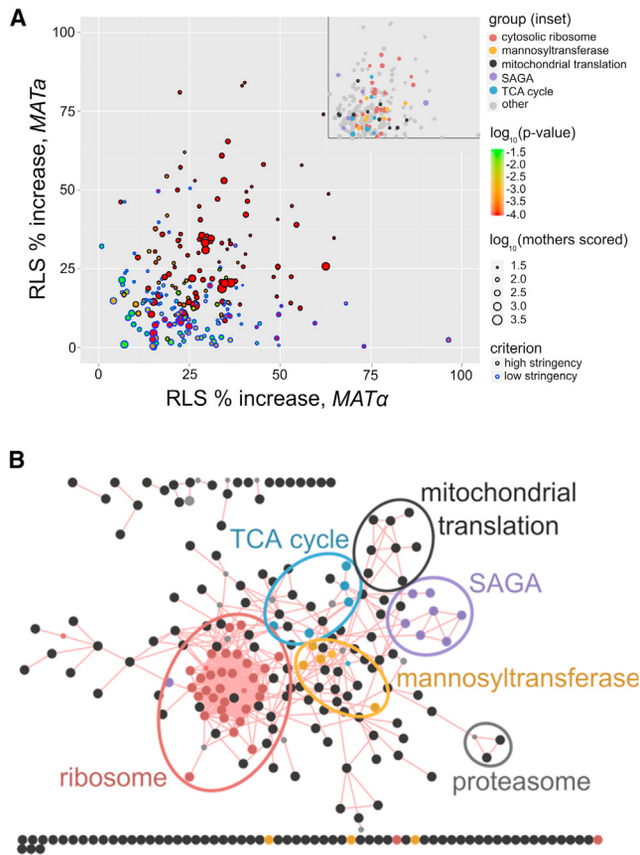


Figure 1. Yeast Gene Deletions that Extend Replicative Lifespan
 (A) Summary of RLS data for long-lived deletion strains. Axes indicate percent increase in RLS relative to control in *MATa* and *MATα*, respectively. Point size is proportional to number of mother cells scored, and point color indicates p value for increased RLS relative to control. Point outline indicates stringency for inclusion: high stringency cutoff was $p < 0.05$ for Wilcoxon rank-sum increased survival independently in both mating types, and low stringency was $p < 0.05$ for pooled data from both mating types with increased RLS shown in each mating type alone.
 (B) Functional clustering of long-lived deletions. Large circles represent long-lived deletions; edges are published physical protein-protein interactions. Overrepresented categories noted in color; $p < 0.05$ with Holm-Bonferroni multiple testing correction.

functional groups, including ribosomal proteins, the SAGA complex, mannosyltransferase genes, and genes involved in TCA cycle metabolism, among others.

These functional clusters may overlap, or otherwise be linked, especially for a process as complex as aging. One link established by our labs centered around Gcn4; this transcription factor was not only critical for RLS extension by reduced 60S translation, but also for a mitochondrial protease gene deletion *afg3Δ* (Delaney et al., 2013a). Our screen identified another unexpected link, again involving Gcn4. DNA damaging agents such as methyl methane sulfonate (MMS) were shown to inhibit the Los1 tRNA transporter by excluding it from the nucleus, leading to Gcn4 activation. This effect on Los1 required checkpoint response factor Rad53 (Ghavidel et al., 2007). Deletion of *LOS1* extended RLS in our screen, and we chose to further define this mechanism of RLS extension based on the possibility that understand-

ing it might connect DNA damage signaling to translational regulation of lifespan.

RESULTS

Genome-Scale Identification of Single-Gene Deletions that Extend Yeast Replicative Lifespan

We performed a genome-wide analysis of viable *S. cerevisiae* single-gene deletions by measuring the RLS of 5 mother cells in the *MATα* mating type for 4,698 unique strains, based on the approach outlined previously (Kaerberlein and Kennedy, 2005). For each strain that showed a mean RLS increase of $>30\%$ over control, or $p < 0.05$ for increased RLS, we measured RLS for 20 cells in the *MATa* strain carrying the same gene deletion. For all gene deletions that extended RLS significantly in both mating types, at least 20 mother cells total were scored in each mating type. In some cases of divergent mating type RLS, the difference may be due to the selection of slow-growth suppressors in the non-long-lived mating type, as has been observed for ribosomal protein mutants (Steffen et al., 2012). In cases where we have observed a changed RLS upon reconstruction of the strain, only reconstructed data is included. We have observed zero examples in this data where a significant difference between mating types survived reconstruction of the strains, and also note that the very large number of mother cells scored for wild-type *MATa* and *MATα* show no significant difference in RLS. A graphical summary of all long-lived deletions found in this screen is shown in Figure 1A.

Statistical criteria are summarized in the Supplemental Experimental Procedures available online, and tested strains are listed in Table S1, related to Figure 1. A total of 238 long-lived deletion strains are summarized in Table S2, related to Figure 1, and complete survival curves and graphical survival by functional group are shown in Figure S1, related to Figure 1. Mortality analysis for all long-lived strains with over 200 scored mother cells is shown in Figure S2, related to Figure 1.

The more than 780,000 individual manually dissected wild-type yeast daughter cells in this project provide a high resolution for making accurate estimates of false positive and negative rates, allowing us to estimate the total fraction of viable yeast deletions likely to affect RLS. We generated sampling distributions from our wild-type cells (Figure S3, related to Figure 2; Supplemental Experimental Procedures). Using these, we estimated our false positive and false negative rates as a function of the percent increase in RLS and sample size n (Figure 2 and Table S3, related to Figure 2). These results suggest that the estimated total number of additional viable deletions that extend RLS $>50\%$ relative to wild-type is likely <1 . For a 40% increase in RLS, we estimate ~ 10 additional viable deletions, and for a 30% increase, ~ 58 additional viable deletions (Table S3, related to Figure 2). In considering false negative rates, it is worth stating explicitly that there is a class of *S. cerevisiae* genes whose effects cannot be reflected in this work: essential genes. Further, previous work (Curran and Ruvkun, 2007) has suggested that essential genes may be more likely than nonessential genes to have a strong effect on lifespan, implying that the number of essential longevity genes remaining to be discovered could exceed a rough approximation based on extrapolation from nonessential genes.

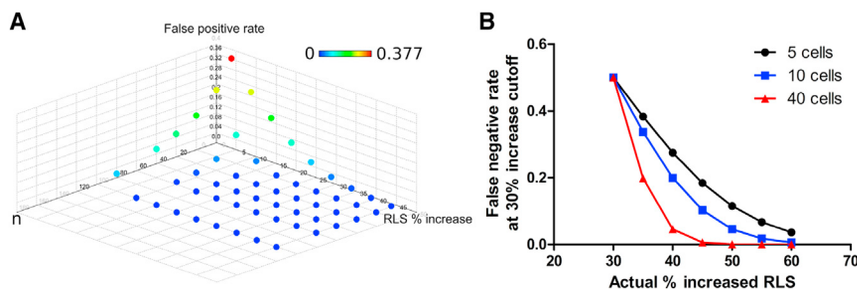


Figure 2. Estimated False Positive and Negative Rates

(A) Estimated false positive rate for given percent increase in RLS and number of mother cells n . x axis indicates percent increase in RLS relative to wild-type control, y axis indicates sample size n , and z axis and point color indicate estimated false positive proportion.

(B) Estimated false negative rate at a threshold of 30% increased RLS, for given actual percent increased RLS and number of mother cells n . x axis indicates actual percent increase expected for the genotype, z axis indicates the estimated false negative proportion using a 30% screening cutoff, and point color indicates sample size n (black, $n = 5$; blue, $n = 10$; red, $n = 40$).

Lifespan-Extending Deletions Are Clustered in Known and Highly Conserved Functional Pathways

A biologically meaningful list of lifespan-extending genes is unlikely to be a random assortment, and the genes identified in our screen are not; many of these genes cluster into known functional groups. All of these groups are conserved from yeast to humans. Genes whose molecular function category is overrepresented among our long-lived deletions are summarized by category (Table 1), and individual categories are described further. A network showing all long-lived deletions as nodes, and published physical protein-protein interactions as edges, shows that many of the genes not only cooperate in related functional processes, but are physically bound in protein complexes such as the cytosolic ribosome, mitochondrial ribosome, and SAGA complex (Figure 1B).

Lifespan Is Altered by Proteins Involved in Both Cytosolic and Mitochondrial Translation, the SAGA Complex, Protein Mannosylation, the TCA Cycle, and Proteasomal Activity

The largest functional category of long-lived deletions is the cytosolic ribosome. These 32 genes include 26 paralogs encoding 20 protein components of the large ribosomal subunit, 4 paralogs encoding 2 proteins in the ribosomal stalk, and 2 paralogs encoding 2 proteins in the small ribosomal subunit. This function is conserved; several components of the large subunit of the cytosolic ribosome have also been shown to affect lifespan in *C. elegans* (Chen et al., 2007; Curran and Ruvkun, 2007; Hansen et al., 2007).

Ribosomal gene deletions were one of the first categories noted to be long-lived during the progress of this project. Four previous publications, three from our group (Kaeberlein et al., 2005; Steffen et al., 2008; Steffen et al., 2012) and one other (Chiocchetti et al., 2007), have investigated the mechanism by which deletion of large subunit genes extends RLS, leading to the identification of the nutrient-responsive transcription factor Gcn4 as an important downstream mediator of longevity (Steffen et al., 2008).

Overall, ribosome and cytosolic ribosome are detected independently as overrepresented categories, due to the fact that alongside the components of the cytosolic ribosome, we identified genes encoding six components of the large subunit of mitochondrial ribosome: *IMG1*, *IMG2*, *MRPL33*, *MRPL40*, *MRPL49*, and *YDR115W*. We also saw increased RLS upon deletion of the genes encoding two mitochondrial tRNA synthetases,

MSK1 and *MSW1*, and three mitochondrial translation control (MTC) genes, *SOV1*, *SUV3*, and *CBS1*, which themselves affect levels of mitochondrial ribosomal proteins. Deletion of these three MTC genes has been shown previously to extend yeast RLS (Caballero et al., 2011). The bias toward the large subunit seen in the RLS-extending cytosolic ribosome deletions is repeated in the mitochondrial ribosome. It will be interesting to learn how much the mechanisms of RLS extension by deletion of cytosolic ribosomal components and mitochondrial ribosomal components have in common.

The yeast SAGA and SLIK chromatin remodeling complexes and the orthologous human complexes STAGA and SALSA are histone-modifying complexes which contain at least two enzymatic activities: a protein acetyltransferase (in yeast, Gcn5) and a deubiquitinase (in yeast, Ubp8) (Samara et al., 2010). The Ubp8 deubiquitinase acts as part of a deubiquitinating module, or DUBm, alongside Sgf73, Sgf11, and Sus1. Deletions of genes encoding three of the four DUBm components, Ubp8, Sgf73, and Sgf11, increase yeast RLS significantly, and *sgf73Δ* has one of the largest RLS increases seen in our screen. These genes were also identified early in our screen, and a parallel project has now looked in more detail at the mechanisms by which disruption of the SAGA/SLIK DUBm may increase yeast RLS (McCormick et al., 2014).

Protein O-mannosylation is an essential modification that occurs at the endoplasmic reticulum and is conserved from yeast to humans (Hutzler et al., 2007). In yeast, three mannosyltransferase subfamilies, PMT1 (consisting of Pmt1 and Pmt5), PMT2 (Pmt2 and its paralog Pmt3), and PMT4, are thought to mannosylate distinct protein targets (Gentzsch and Tanner, 1997). In our screen, *PMT1*, *PMT3*, and *PMT5* all significantly extended yeast RLS. Closer inspection of the remaining individual *PMT* gene deletions revealed additional interesting results: *pmt2Δ* just missed the cutoff for inclusion, exhibiting a 26.3% increase in RLS ($p < 0.038$) in *MATa*, and a 15.6% increase ($p < 0.054$) in *MATα*. *pmt6Δ* and *pmt7Δ* each extended RLS in one mating type when tested (22.6% and 31.3%, respectively), but not in the other mating type, leaving open the possibility that the nonextended mating type in each case may harbor a second mutation which masks the RLS phenotype of the deletion. We also identified the two yeast mannosyltransferases involved in O-linked glycosylation, *MNN1* and *MNN2*; the N-linked mannosyltransferases *YUR1*, *ALG6*, and *ALG12*; and *EOS1*, which has been implicated in protein N-glycosylation.

Table 1. Genes that Comprise Overrepresented Functional Categories among Long-Lived Single Gene Deletions

Cytosolic Ribosome			Mitochondrial Translation			Mannosyltransferase		
Gene	Mean RLS	Percent Increase	Gene	Mean RLS	Percent Increase	Gene	Mean RLS	Percent Increase
<i>RPL12A</i>	30.11	17.7	<i>CBS1</i>	31.60	36.8	<i>ALG12</i>	31.75	16.6
<i>RPL12B</i>	33.80	21.6	<i>IMG1</i>	33.11	22.5	<i>ALG6</i>	30.36	15.3
<i>RPL13A</i>	34.68	36.9	<i>IMG2</i>	29.57	28.0	<i>EOS1</i>	35.03	23.9
<i>RPL13B</i>	34.70	36.6	<i>MRPL33</i>	28.04	16.3	<i>MNN1</i>	30.54	14.7
<i>RPL16B</i>	35.09	22.7	<i>MRPL40</i>	33.27	20.7	<i>MNN2</i>	30.47	15.5
<i>RPL19A</i>	34.26	29.6	<i>MRPL49</i>	34.69	25.2	<i>PMT1</i>	35.08	38.1
<i>RPL19B</i>	32.36	15.7	<i>MSK1</i>	28.49	11.6	<i>PMT3</i>	31.88	26.8
<i>RPL1B</i>	31.13	24.1	<i>MSW1</i>	32.82	26.0	<i>PMT5</i>	29.07	20.6
<i>RPL20A</i>	35.80	44.4	<i>SOV1</i>	31.29	16.8	<i>YUR1</i>	31.18	19.2
<i>RPL20B</i>	36.40	35.0	<i>SUV3</i>	33.82	21.8			
<i>RPL21B</i>	36.93	49.5	<i>YDR115W</i>	30.13	11.7			
<i>RPL22A</i>	36.59	38.4						
<i>RPL23A</i>	34.02	31.9						
			TCA Cycle			SAGA		
Gene	Mean RLS	Percent Increase	Gene	Mean RLS	Percent Increase	Gene	Mean RLS	Percent Increase
<i>RPL26A</i>	31.39	21.6	<i>EMI5</i>	31.66	28.3	<i>CHD1</i>	31.06	13.5
<i>RPL29</i>	32.08	31.9	<i>IDH1</i>	32.68	17.3	<i>SGF11</i>	36.64	25.8
<i>RPL2B</i>	34.30	38.3	<i>IDH2</i>	30.88	15.0	<i>SGF73</i>	40.81	55.1
<i>RPL31A</i>	34.85	30.0	<i>IDP1</i>	28.37	9.0	<i>SPT8</i>	31.50	9.1
<i>RPL34A</i>	34.20	27.4	<i>LPD1</i>	31.55	36.8	<i>UBP8</i>	36.02	34.4
<i>RPL34B</i>	37.80	48.0	<i>SDH1</i>	32.75	24.9			
<i>RPL35A</i>	24.80	41.2	<i>SDH2</i>	32.09	21.3			
<i>RPL37B</i>	29.99	26.6						
<i>RPL43B</i>	32.51	35.0						
<i>RPL6A</i>	35.50	43.1						
<i>RPL6B</i>	31.51	16.6						
<i>RPL7A</i>	32.10	19.0						
<i>RPL9A</i>	31.43	27.1						
<i>RPP1A</i>	37.38	30.7						
<i>RPP1B</i>	32.91	23.4						
<i>RPP2A</i>	26.39	14.9						
<i>RPP2B</i>	34.85	50.2						
<i>RPS12</i>	32.62	38.3						
<i>RPS6A</i>	30.05	10.2						

Recently, Labunskyy et al. reported that yeast lacking *ALG12* and *BST1*, which controls ER-to-Golgi transport of glycosylphosphatidylinositol-anchored proteins (Tanaka et al., 2004), were long-lived because they activate the unfolded protein response, which promotes multiple forms of stress resistance (Labunskyy et al., 2014). It remains to be determined whether other long-lived deletions of mannosyltransferase-related genes enhance RLS by this mechanism.

We identified seven genes encoding proteins central to the tricarboxylic acid pathway: isocitrate dehydrogenase subunits *IDP1*, *IDH1*, and *IDH2*; lipoamide dehydrogenase *LPD1*; and succinate dehydrogenase subunits *SDH1*, *SDH2*, and *EMI5/SDH5*. All of these genes are involved in a very short section of the TCA cycle: the conversion of isocitrate to alpha-ketoglutarate (*IDP1*, *IDH1*, *IDH2*), on to succinyl-CoA (*LPD1*), and from there to succinate and then fumarate (*SDH1*, *SDH2*, *EMI5*). It is interesting to note that this pathway converges on many events linked to longevity, including production of amino acids that

regulate TOR activity, control of metabolite levels including NAD(+), and use of different substrates for ATP production (Imai et al., 2000; Lin et al., 2000; Yang et al., 2007).

Although not overrepresented by automated gene ontology analysis, one of the longest-lived single deletions identified here is that of *UBR2*, a ubiquitin ligase that regulates the turnover of the transcription factor Rpn4, which positively regulates transcription of components of the yeast proteasome. As the extended RLS of *ubr2Δ* depends completely on Rpn4, we concluded that proteasomal activity is central to this phenotype (Kruegel et al., 2011), and we have recently dissected the RLS effects of *UBR2* and *RPN4* in additional detail (Yao et al., 2015).

RLS-Extending Single Gene Deletions in Yeast Overlap at High Significance with Known Lifespan-Extending Hypomorphs in *C. elegans*

We examined whether these data would provide further evidence of evolutionary conservation of longevity pathways. To

Table 2. Lifespan Phenotypes of Yeast Deletions Are Significantly Conserved

Worm Orthologs	Yeast Orthologs	Worm Orthologs	Yeast Orthologs
<i>ant-1.1</i>	AAC3	<i>rpl-1</i>	RPL1B
<i>cyc-2.1</i>	CYC1	<i>rpl-19</i>	RPL19A RPL19B
<i>dld-1</i>	LPD1	<i>rpl-23</i>	RPL23A
<i>dod-18</i>	YOR111W	<i>rpl-31</i>	RPL31A
<i>idh-1</i>	IDP1	<i>rpl-6</i>	RPL6A RPL6B
<i>idha-1</i>	IDH2	<i>rpl-7</i>	RPL7A
<i>ifg-1</i>	TIF4631	<i>rpl-9</i>	RPL9A
<i>inf-1</i>	TIF1	<i>rps-6</i>	RPS6A
<i>jmjd-2</i>	RPH1	<i>rsk-1</i>	SCH9
<i>let-363</i>	TOR1	<i>sams-1</i>	SAM1
<i>nac-2</i>	<i>nac-3</i>	<i>spg-7</i>	AFG3
	<i>pos-1</i>	<i>spt-4</i>	SPT4
	<i>raga-1</i>	<i>unc-26</i>	INP53

Listed are genes whose deletion in *S. cerevisiae* was shown to increase RLS in this screen, and whose *C. elegans* orthologs have been published to have increased lifespan upon deletion, hypomorphic mutation, or RNAi knockdown. $p_{\text{overlap}} < 4E-11$; if all ribosomal genes are excluded $p_{\text{overlap}} < 2E-6$.

test this, we asked whether our RLS-extending deletions overlapped significantly with genes whose knockdown or deletion was reported to extend the lifespan of the distantly related nematode *C. elegans*. We compiled a list of 428 genes whose deletion, hypomorphic or null allele mutation, or RNAi knockdown had been reported to extend *C. elegans* lifespan, taking as our starting point previous work where a portion of the yeast genome was similarly examined (Smith et al., 2007; Smith et al., 2008) (Table S4, related to Table 2). We then used paralog-ortholog groups generated by InParanoid (Ostlund et al., 2010) to find genes whose lifespan-extending phenotype was shared between *S. cerevisiae* and *C. elegans*, and calculated the probability of an overlap of the observed magnitude. Long-lived yeast deletions whose hypomorphic lifespan extension has also been reported in *C. elegans* are shown in Table 2. The probability of this overlap absent functional conservation of lifespan phenotypes (i.e., at random) is $p < 4.4E-11$. Because of the large number of ribosomal proteins present in both species' lifespan-altering orthologs, we asked whether they alone were responsible for the significance of the overlap we observed, and found that the remaining list still exhibited a significant overlap ($p < 2.0E-6$) with ribosomal proteins omitted.

We note that the genes affecting both yeast and *C. elegans* lifespan contain a similar proportion of genes from our functionally overrepresented pathways as the screen as a whole. Specifically, 0.255 of the overall screen results fall into the functionally overrepresented groups such as the ribosome, TCA cycle, SAGA, etc., while 0.423 of the conserved genes do so, a higher proportion by gene. Nevertheless, not all five functionally overrepresented pathways we describe here in yeast are represented among the conserved genes; only two (ribosome and TCA cycle) are. We propose at least two possible factors that may contribute to the absence of the other pathways. First, although we here quantify the false negative rate of our screen with high precision, our list of *C. elegans* genes is generated by consideration of aggregate data from multiple studies, most of which are not whole-genome studies, whose aggregate false

negative rate cannot be easily estimated. We suspect that some absences from our list of conserved genes may be because the phenotype has not been observed in *C. elegans* although it may exist. Further, interspecies ortholog prediction is an imperfect science. In the example of the SAGA group, although both the individual genes and the overall protein complexes are conserved at the sequence and functional level between yeast, mice, and humans, we were not able to identify putative *C. elegans* orthologs for any of the RLS-extending SAGA deletions. We do not know whether this is due to sequence divergence, complete loss of this functionality from *C. elegans*, or other reasons, but it may offer an additional explanation for the fact that not every pathway we have uncovered in yeast is represented in the smaller subset of identified conserved genes.

Taken in context, the highly significant overlap between the final results of our screen of viable yeast deletions and published *C. elegans* genes suggests a high degree of general conservation in lifespan genotype-phenotype relationships between very distantly related species. This leaves open the possibility that these genes may be enriched for orthologs that can alter lifespan and aging in other organisms, such as humans.

Nuclear tRNA Accumulation Is Associated with Increased RLS

Among the long-lived deletions identified in our screen, we chose to further examine *los1Δ*, as it had a dramatically enhanced RLS ($p < 0.001$, Figure 3A), and Los1 has a function not previously linked to aging: subcellular localization of tRNA.

Since Los1 is known to function as a nuclear tRNA exporter, and deletion of *LOS1* leads to increased levels of nuclear tRNA (Sarkar and Hopper, 1998), we tested whether altered import of tRNA into the nucleus could also modulate RLS. Mtr10 is a nuclear import receptor required for retrograde transport of tRNAs from the cytoplasm into the nucleus (Shaheen and Hopper, 2005). Deletion of *MTR10* had no effect on RLS ($p = 0.9$, Figure 3B), but overexpression of *MTR10* under its native

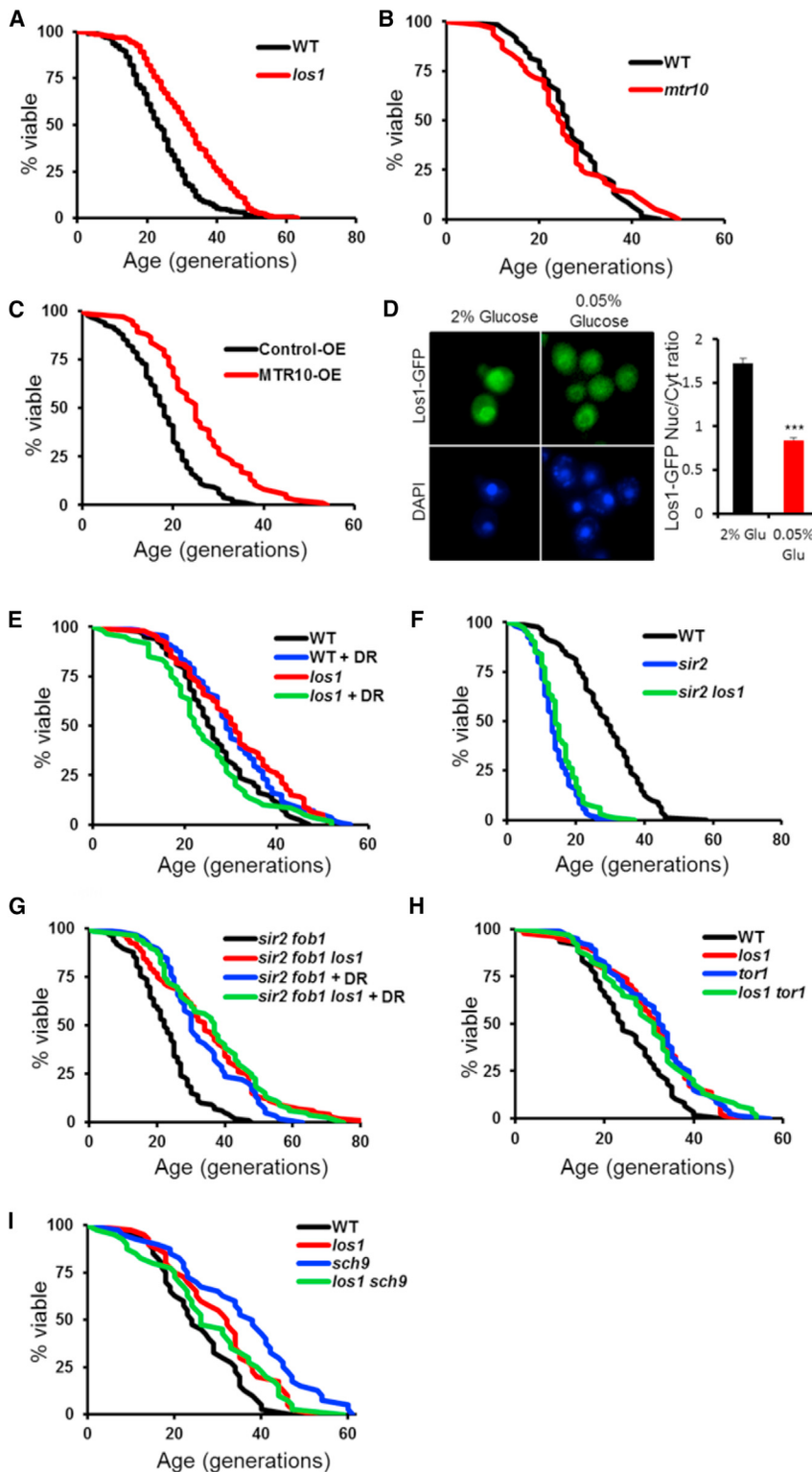


Figure 3. Regulators of Nuclear tRNA Import and Export Influence Lifespan, and DR and *los1* Δ Act in the Same Pathway

(A) Deletion of the nuclear tRNA exporter gene *LOS1* extends RLS.

(B) Deletion of tRNA importer gene *MTR10* does not reduce RLS.

(C) Additional copy of the nuclear tRNA importer *MTR10*, which functions antagonistically to *LOS1*, extends RLS.

(D) Blinded scoring of fluorescence microscopy reveals that DR (0.05% glucose) reduces nuclear *Los1-GFP*, *** $p < 0.001$. Error bars denote the standard error of the mean (SEM).

(E–I) RLS analysis indicates that deletion of *LOS1* acts similarly to DR. Deletion of *LOS1* fails to extend RLS of *sir2* Δ cells, but robustly extends RLS of *sir2* Δ *fob1* Δ cells. Deletion of *LOS1* does not increase RLS additively with DR or with genetic mimics of DR (*tor1* Δ and *sch9* Δ). RLS and microscopy statistics are provided in Table S5.

DR and *Los1* Similarly Modulate RLS

Given prior data showing that *Los1* is excluded from the nucleus upon complete glucose starvation (Quan et al., 2007), we examined whether *Los1* was responsive to DR, by reducing the glucose concentration of the medium from 2% to 0.05%. Relative to control conditions, DR significantly reduced the amount of *Los1-GFP* in the nucleus compared to the cytoplasm ($p < 0.001$, Figure 3D). DR also failed to further increase the RLS of long-lived cells lacking *LOS1* (Figure 3E).

Prior studies have shown that DR fails to increase the RLS of *sir2* Δ single mutant cells lacking the Sir2 histone deacetylase, but robustly increases the RLS of *sir2* Δ *fob1* Δ double mutant cells (Delaney et al., 2011b; Kaeberlein et al., 2004; Lin et al., 2000). Similarly, deletion of *LOS1* did not increase the RLS of *sir2* Δ cells, but significantly increased the RLS of *sir2* Δ *fob1* Δ cells ($p < 0.001$, Figures 3F and 3G). Again, *los1* Δ did not further extend RLS when combined with DR in the *sir2* Δ *fob1* Δ background (Figure 3G).

As an additional test of whether *Los1* acts downstream of DR to modulate lifespan, we combined deletion of *LOS1* with two well-characterized genetic mimetics of DR, deletion of *TOR1* or *SCH9*. As with DR by glucose restriction, deletion of *TOR1* or *SCH9* also failed to further in-

crease RLS in *los1* Δ mother cells (Figures 3H and 3I). Interestingly, *sch9* Δ *los1* Δ cells are slightly shorter lived than *sch9* Δ single mutant cells, and *los1* Δ cells are shorter lived under DR conditions relative to control conditions, similar to the previously observed effect of combining DR with *sch9* Δ (Kaeberlein et al.,

promoter from a low copy plasmid significantly increased RLS relative to a control plasmid ($p < 0.001$, Figure 3C). Taken together, these data indicate that either blocking nuclear export or enhancing nuclear import of tRNA is sufficient to extend yeast RLS.

2005). These data indicate that genetically altering tRNA transport to favor accumulation of nuclear tRNA is sufficient to extend RLS, and support a model in which nuclear sequestration of tRNA via regulation of Los1 localization is one mechanism by which DR increases RLS.

Los1 Is Regulated by Both Rad53 and mTOR Downstream of DR

A previous report showed that Rad53 activation by DNA damage causes Los1 export to the cytoplasm, resulting in nuclear sequestration of tRNA (Ghavidel et al., 2007). Rad53 has been previously implicated in aging (Delaney et al., 2013b; Schroeder et al., 2013; Weinberger et al., 2010) but not linked to the response to DR. Redistribution of Los1 to the cytoplasm in response to DR was prevented in cells lacking *RAD53* ($p = 0.1$, Figure 4A; Table S5, related to Figure 4). Despite the fact that *rad53* Δ cells are extremely short-lived, and in some backgrounds inviable (Zhao et al., 2001), deletion of *LOS1* was still able to significantly extend RLS in this background ($p < 0.001$), consistent with the model that Rad53 acts between DR and Los1 to modulate RLS (Figure 4B).

Since reduced mTOR signaling is believed to mediate RLS extension from DR (Kaeberlein et al., 2005), we attempted to place mTOR genetically within the longevity pathway containing DR, Los1, and Rad53. Rad53 is not required for deletion of *TOR1* to extend RLS ($p = 0.005$, Figure 4C), suggesting that mTOR acts at least partially independently of Rad53. Both *los1* Δ and *tor1* Δ alter the mortality kinetics of *rad53* Δ cells, possibly by reducing a stochastic lifespan-limiting event. Interestingly, Los1-GFP is also enriched in the cytoplasm of *tor1* Δ cells under nutrient rich conditions, albeit to a lesser extent than in wild-type cells subjected to DR ($p < 0.001$, Figure 4D). Unlike DR, however, cytoplasmic localization of Los1-GFP in *tor1* Δ cells does not require Rad53 ($p < 0.001$). Rapamycin, an inhibitor of mTOR complex I that extends lifespan in yeast, worms, fruit flies, and mice (Ramos and Kaeberlein, 2012), also causes Los1-GFP to become enriched in the cytoplasm in a Rad53-independent manner ($p < 0.001$, Figure 4E).

To further explore the role of Rad53 in the DR response, we examined the effect of DR on Rad53 abundance and localization using a strain expressing Rad53-GFP fusion protein under the control of the *RAD53* promoter. DR significantly increased Rad53 nuclear and overall abundance ($p < 0.001$), comparable to treatment with MMS, a known inducer of Rad53 ($p < 0.001$, Figures 4F and 4G). Deletion of *RAD53* prevented RLS extension from DR in both wild-type and *fab1* Δ cells (Figure 4H). Importantly, overexpression of *RAD53* was sufficient to extend RLS in an otherwise wild-type background ($p < 0.001$, Figure 4I), comparable to the effect of *MTR10* overexpression or *LOS1* deletion (Figures 2A and 2C). This RLS extension from *RAD53* overexpression was not associated with a significant change in growth rate either in liquid culture or on agar plates during the lifespan experiment (Figures 4J and 4K), indicating that overexpression of *RAD53* is not sufficient to induce the DNA damage cell-cycle checkpoint.

Deletion of *LOS1* Induces Gcn4 Targets without Reducing mRNA Translation

To further characterize the mechanism by which Los1 regulates RLS, we compared gene expression profiles of wild-type and

los1 Δ cells under nonrestricted conditions. Sixty genes were identified with significantly increased expression in *los1* Δ cells and 31 with significantly reduced expression (Table S6, related to Figure 5). Gene ontology analysis indicated that a majority of the mRNAs upregulated in *los1* Δ cells encoded proteins involved in amino acid biosynthesis and nitrogen metabolism (Table S6, related to Figure 5). Of 60 genes upregulated in *los1* Δ cells, 88% are annotated as targets of the Gcn4 transcription factor on Yeasttract (Teixeira et al., 2006) (Figure 5A; Table S6, related to Figure 5, $p < 0.0001$ enriched for Gcn4 targets compared to downregulated genes). Upregulation of GO term enriched genes was confirmed by qPCR (Figures 5B and 5C; Table S8, related to Figure 5). This reinforces prior evidence that Gcn4 is activated under conditions where tRNA is sequestered in the nucleus (Ghavidel et al., 2007; Qiu et al., 2000), and is consistent with the model that Gcn4 acts downstream of DR and mTOR to promote longevity (Steffen et al., 2008).

Given that *los1* Δ mutants have reduced cytoplasmic tRNA and increased expression of Gcn4 target genes, we hypothesized that loss of *LOS1* may extend RLS by decreasing 60S mRNA translation, similar to *RPL20B* (Steffen et al., 2008). Unexpectedly, we were unable to detect any change in global mRNA translation in *los1* Δ cells, by growth rate in rich media, polysome analysis, or cell cycle progression (Figures 5C–5E). A similar lack of suppression of mRNA translation was also observed in long-lived *tor1* Δ cells, while the *rpl20b* Δ cells had a significant defect. Similar to *tor1* Δ , Gcn4 partially but not completely prevented RLS extension from deletion of *LOS1* (Figure 5F). The magnitude of RLS suppression from *GCN4* deletion in *los1* Δ cells is comparable to the effects in *tor1* Δ cells or cells subjected to DR and cells lacking full 60S translation capacity, such as *sch9* Δ , *afg3* Δ (Delaney et al., 2013a), and *rpl4s* (Figure 5G). Lifespan abrogation is not seen in a DR-independent pathway mutant *ubr2* Δ (Kruegel et al., 2011). These data suggest that Gcn4 is activated in *los1* Δ cells through a mechanism distinct from reduced global mRNA translation, and this Gcn4 activation contributes to increased RLS (Figure 5H).

DISCUSSION

The work summarized here represents one of the most comprehensive ORFeome-wide screens of any model organism for increased lifespan, comprising data from manual dissection of over 2.2 million individual yeast daughter cells. The scale of this screen gives a unique snapshot of the pathways involved in the regulation of lifespan in a single eukaryotic model organism, and has identified multiple significantly overrepresented functional processes. This includes a role for Los1-mediated tRNA transport downstream of dietary restriction, which we have dissected further. Many of the longevity genes identified here are known to act in longevity pathways conserved in multiple species (e.g., components of the cytosolic ribosome), or have greatly extended a functional category which had already shown some genes involved in aging (mitochondrial translation). Others, like the *LOS1* pathway, are largely unstudied (e.g., mannosyltransferase, TCA cycle metabolism). Two gene ortholog groups that modulate the lifespan of yeast and other organisms, mTOR and sirtuins, have already pointed to drugs that extend

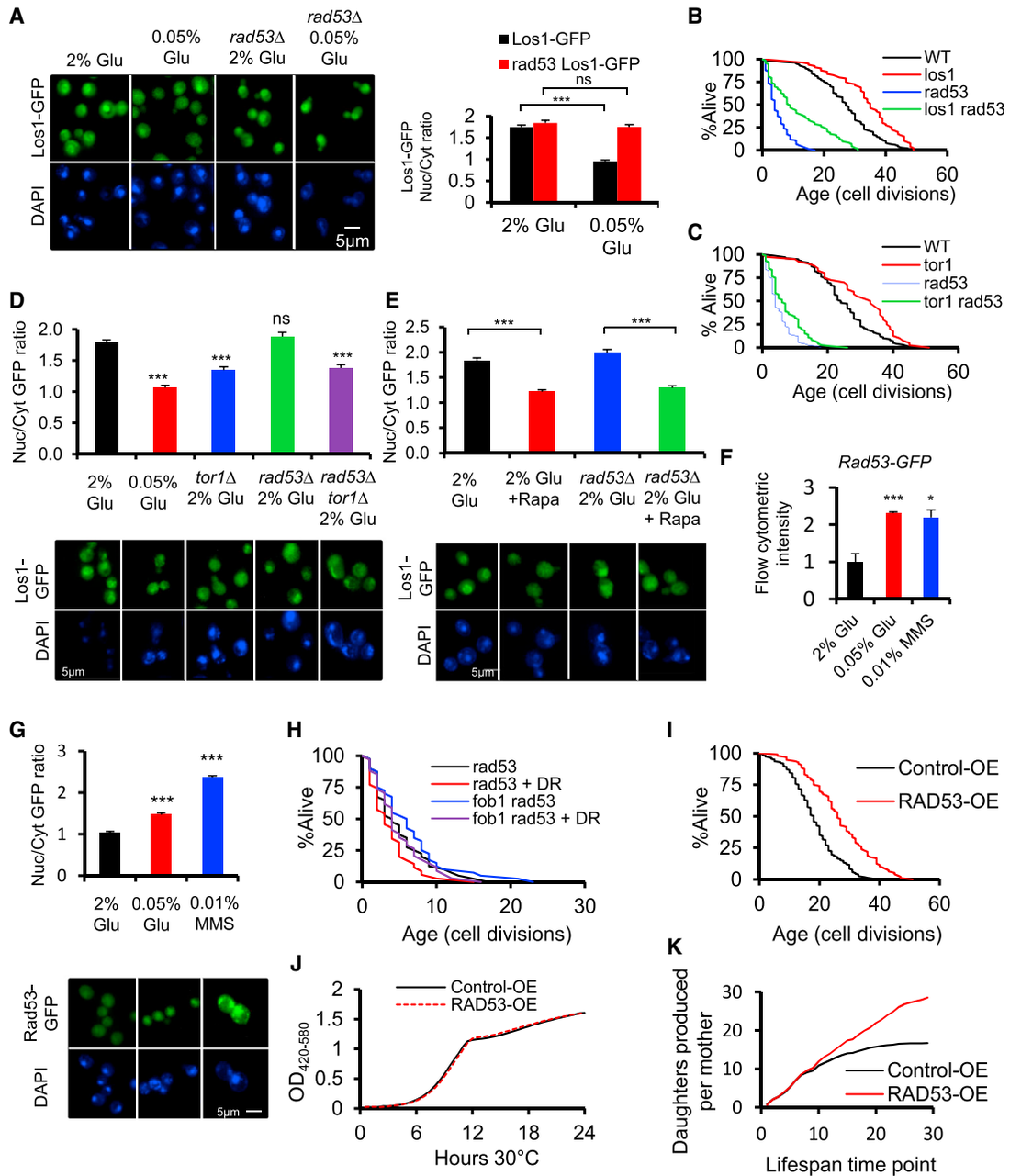


Figure 4. Rad53 Acts Between DR and Los1, but Not Tor1, Is Enhanced by DR, and Modulates Lifespan

(A) Fluorescence microscopy indicates that Rad53 is required for nuclear exclusion of Los1-GFP in response to DR. (B and C) RLS analysis demonstrates that Rad53 is not required for RLS extension by deletion of either *LOS1* or (C) *TOR1*. (D and E) (D) Fluorescence microscopy indicates that Rad53 is not required for nuclear exclusion of Los1-GFP in response to deletion of *TOR1* or (E) by addition of 1 nM rapamycin. (F) Levels of Rad53-GFP in the cell, normalized to 2% glucose growth conditions, as assayed by flow cytometry. (G) GFP-tagged Rad53 is enriched in the nucleus by DR. (H) RLS analysis demonstrates that Rad53 is required for RLS extension from DR. (I) Additional copy of *RAD53* extends RLS. (J) Representative growth curves of *RAD53* overexpression. (K) Mother cell divisions of *Rad53*-OE and wild-type on agar as a function of chronological time. RLS and microscopy statistics are provided in Table S5. *p < 0.05, ***p < 0.001. Error bars denote the standard error of the mean (SEM).

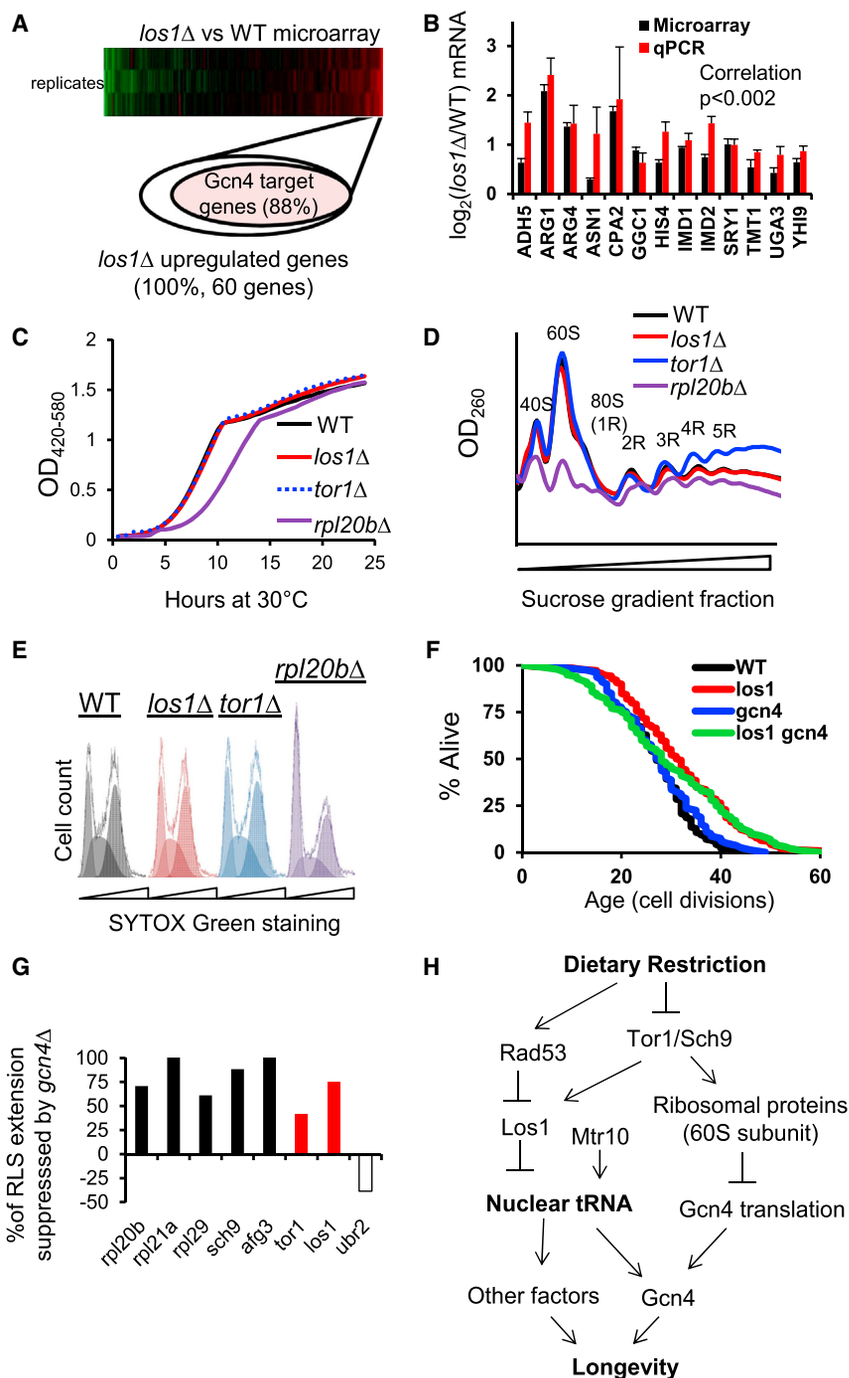


Figure 5. Deletion of *LOS1* Activates *GCN4* without Reducing Global mRNA Translation, and *Gcn4* Is Necessary for RLS Extension by *los1* Δ

(A) Relative proportions of *Gcn4* regulated genes within the subset of genes found via microarray to be upregulated in *los1* Δ yeast.

(B) RT-qPCR verification of a subset *Gcn4* target genes found to be upregulated by microarray analysis of *los1* Δ cells, focused on enriched GO term related genes.

(C–E) (C) Representative growth curves, (D) poly-some profiles, and (E) cell-cycle profiles of *los1* Δ , *tor1* Δ , and WT yeast indicate that neither *los1* Δ nor *tor1* Δ cells have a detectable defect in mRNA translation. For comparison, a translation-deficient *rpl20b* Δ mutant is shown in each case. R values in (E) correspond to the number of ribosomes bound to a particular mRNA.

(F) Deletion of *GCN4* partially blocks RLS extension from deletion of *LOS1*.

(G) Deletion of *GCN4* largely prevents RLS extension in genetic models of DR, but not the DR-independent *Ubr2* pathway. Percent suppression of RLS is calculated by comparing the percent mean RLS extension of the long lived mutant in wild-type and *gcn4* Δ backgrounds. RLS statistics are provided in Table S5.

(H) Diagram summarizing the comprehensive model for DR supported by the data from this study. Error bars denote the standard error of the mean (SEM).

PMT5 mannosyltransferase complex deletions and that of the 60 s subunit-biased ribosomal protein gene deletions.

The detailed analysis here defining a previously unsuspected role for regulation of nuclear tRNA export in modulating longevity provides further insights into linking longevity pathways that may initially appear disparate. Our data demonstrate that *Rad53* and mTOR are both regulated by DR and likely act through parallel mechanisms to influence *Los1* localization (Figure 5). This fits with prior data suggesting that *Rad53* regulates *Los1* in response to DNA damage (Ghavidel et al., 2007), similar to *Ntg1* translocation during mtDNA stress

the lifespan of mice (Harrison et al., 2009; Kaeberlein et al., 2005; Li et al., 2014; Li and Miller, 2014; Mitchell et al., 2014), and each of the pathways described here should be considered with this potential in mind.

While many genes identified here cluster into functionally related groups, some genes are alone or unknown in their molecular function, which may yield further unexpected links. For example, the oligosaccharyltransferase complex directly binds to the yeast ribosome at a location near the translocon-binding site, which is on the 60 s ribosomal subunit (Harada et al., 2009). This may link the lifespan extension in the *PMT1/PMT3*/

(Schroeder and Shadel, 2014), as well as evidence that the nutrient responsive hexokinase *Hxk2* and cAMP-dependent protein kinase A (PKA) signaling pathway are important for appropriate regulation of the *Rad53* checkpoint following DNA damage (Searle et al., 2011). Both *Hxk2* and PKA have been previously shown to act within the DR pathway in yeast, further supporting this link (Kaeberlein et al., 2004; Lin et al., 2000). In this regard, it is worth noting that neither DR nor overexpression of *RAD53* is sufficient to induce the DNA damage checkpoint in the absence of DNA damage (Figure S4, related to Figure 5). Thus, the data presented here provide intriguing connections

between DNA maintenance, nutrient response pathways, tRNA transport, and aging. While tRNA synthetases and modifying enzymes have been reported to modulate aging in *C. elegans* (Lee et al., 2003; Yanos et al., 2012), it remains unclear whether the mechanisms are related to those described here.

Some divergence in phenotype was noted for *tor1Δ*, *sch9Δ*, and *los1Δ*. Deletion of *LOS1* did not alter translation, nor does *tor1Δ*, but deletion of *SCH9* reduces translation and slows the cell cycle. RLS extension in a *los1Δ* background was antagonistic in *sch9Δ*, but not in *tor1Δ*. It is currently unclear whether translation rates are the cause for these differences, or if signal transduction downstream of DR causes posttranslational changes of Los1 and/or Tor1. Glucose deprivation may selectively act on Los1 localization via PKA inactivation rather than Snf1/AMPK inactivation (Pierce et al., 2014), which may partially explain the phenotypic differences between these partial DR mimetic mutants.

Based on data presented here and previously, we suggest a model whereby modest reductions in signaling through mTOR can promote longevity by engaging Gcn4 via Los1-dependent nuclear sequestration of tRNA. The branches of this pathway (Rad53, mTOR, ribosomal proteins) ultimately converge on shared Gcn4 targets, explaining the non-additive effects on lifespan when two long-lived mutants within the pathway are combined. Notably, there is accumulating evidence for a role of the GCN4 ortholog, ATF4, in mammalian longevity as ATF4 is upregulated in the liver of Snell dwarf and PAPP-A knockout mice, acarbose and rapamycin treated mice, and calorically restricted and methionine restricted mice, all of which are long-lived (Li et al., 2014; Li and Miller, 2014).

Finally, it is remarkable that so many yeast gene knockouts extend lifespan. This parallels findings from worm genome-wide screens, and raises the question of why it is so easy to extend lifespan genetically. The answer likely lies in evolutionary theories of aging, which posit that organisms are optimized for fitness and not long lifespan. Fitness in yeast likely means a rapid mitotic growth rate when nutrients are available and retention of meiotic capacity, but not the ability to generate 30 daughters instead of 25 when mother cells are swamped by offspring of previous daughters. Since late life vigor lacks selection, there may be many ways to extend lifespan in mammals as well.

EXPERIMENTAL PROCEDURES

Additional information can be found in [Supplemental Experimental Procedures](#).

Strains and Media

All yeast strains were derived from the parent strains of the haploid yeast ORF deletion collections (Winzeler et al., 1999), BY4742 (*MAT α his3 Δ 1 leu2 Δ 0 lys2 Δ 0 ura3 Δ 0*) and BY4741 (*MAT α his3 Δ 1 leu2 Δ 0 met15 Δ 0 ura3 Δ 0*). A list of all strains not contained in either haploid ORF deletion collection is included in [Table S7](#), related to [Figure 3](#). Cells were grown on standard YPD containing 1% yeast extract, 2% peptone, and 2% glucose at 30°C, unless otherwise stated.

Replicative Lifespan

Yeast RLS assays were performed as previously described (Kaerberlein et al., 2004; Steffen et al., 2009). In short, virgin daughter cells were isolated from each strain and then allowed to grow into mother cells while their corresponding daughters were microdissected and counted, until the mother cell could no

longer divide. Statistical significance was determined by calculating p values using the Wilcoxon rank-sum test (Wilcoxon, 1946).

Gene Ontology and Network Analysis

Gene ontology and network analysis was performed using GeneMANIA (Warde-Farley et al., 2010), using all available GeneMANIA published physical protein-protein interaction datasets at the time of the analysis.

Fluorescence Microscopy

GFP labeled cells were grown to log phase under selection in SC-HIS media, and imaged in fresh liquid media on a Teflon printed slide. Pictures were taken on a Zeiss Axiovert 200M, blinded, and analyzed on Zeiss Axiovision microscopy software.

Flow Cytometry

For cell-cycle analysis, all samples were prepared following a protocol previously described (Delaney et al., 2011a).

Growth Curves

All yeast growth rates were analyzed in the OD₄₂₀₋₅₈₀ range in the Bioscreen C automated microbiology growth curve analysis system (Growth Curves USA, Piscataway, NJ, USA) as previously described using the Yeast Outgrowth Data Analyzer (YODA) software (Olsen et al., 2010).

Polysome Profiles

All polysome analysis was carried out as described previously (MacKay et al., 2004; Steffen et al., 2008). Briefly, log phase yeast cultures were chilled with crushed frozen YPD plus 100 μ g/ml cycloheximide. Cells were harvested by centrifugation, washed with 10 ml lysis buffer (25 mM Tris-HCl [pH 7.5], 40 mM KCl, 7.5 mM MgCl₂, 1 mM DTT, 0.5 mg/ml heparin, 100 μ g/ml cycloheximide) and resuspended in 1 ml lysis buffer. Cells were then lysed by vortexing with glass beads. Triton X-100 and sodium deoxycholate were added to a final concentration of 1%, and samples were left on ice for 5 min before the supernatant was clarified by centrifugation. All reagents were ice cold, and all steps were done in a 4°C cold room. For separation on gradients, 1 ml containing 20 A260 units of lysate were loaded onto 11 ml linear 7%–47% sucrose gradients in 50 mM Tris-HCl (pH 7.5), 0.8 M KCl, 15 mM MgCl₂, 0.5 mg/ml heparin, and 100 μ g/ml cycloheximide, and sedimented at 39,000 rpm at 4°C in an SW40 Ti swinging bucket rotor (Beckman) for 2 hr. Gradients were collected from the top and profiles were monitored at 254 nm.

Microarrays

Log phase (exactly 0.5 OD₆₀₀) yeast was harvested and RNA extracted using the QIAGEN RNeasy kit (QIAGEN, Redwood City, CA, USA). Six wild-type cultures and three independently created *los1Δ* isolates were used, with paired cultures on three independent days. Affymetrix GeneChip Yeast Genome 2.0 Arrays (Affymetrix, Santa Clara, CA, USA). Hybridized arrays were scanned with an Affymetrix GeneChip 3000 scanner. Image generation and feature extraction were performed using Affymetrix Gene Chip Command Console software. Raw data was processed and analyzed with Bioconductor. False discovery rates were used at a q value cutoff of 0.05.

ACCESSION NUMBERS

Data were deposited in the GEO public database under accession number GSE37241.

SUPPLEMENTAL INFORMATION

Supplemental Information includes four figures, eight tables, and Supplemental Experimental Procedures and can be found with this article at <http://dx.doi.org/10.1016/j.cmet.2015.09.008>.

AUTHOR CONTRIBUTIONS

M.M., J.R.D., M.K., and B.K.K. conceived the experiments and wrote the manuscript. M.M., J.R.D., A.S., S.S., A.C., and U.A. generated the yeast

strains and contributed to execution and data analysis of all experiments except mouse experiments. J.R.D. and all authors excluding S.J., R.B., T.B., B.O., D.P., R.B.B., X.L., Y.S., Z.Z., and B.K.K. performed the lifespan measurements. D.P. conducted flow cytometry experiments. B.O. assisted in data analysis. R.B. and T.B. executed the microarray analysis. M.K. and B.K.K. provided reagents and support for the studies. All authors discussed the results and commented on the manuscript.

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