Caenorhabditis elegans alg-n-2 Is Critical for Longevity Conferred by Enhanced Nonsense-Mediated mRNA Decay

HIGHLIGHTS
- C. elegans alg-n-2 is a positive regulator of nonsense-mediated mRNA decay (NMD)
- alg-n-2 is downregulated during aging and contributes to maintaining normal lifespan
- alg-n-2 is required for longevity caused by various genetic interventions
- Upregulation of ALGN-2 by inhibition of daf-2 promotes longevity via increasing NMD
Caenorhabditis elegans algn-2 Is Critical for Longevity Conferred by Enhanced Nonsense-Mediated mRNA Decay

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SUMMARY
Nonsense-mediated mRNA decay (NMD) is a biological surveillance mechanism that eliminates mRNA transcripts with premature termination codons. In Caenorhabditis elegans, NMD contributes to longevity by enhancing RNA quality. Here, we aimed at identifying NMD-modulating factors that are crucial for longevity in C. elegans by performing genetic screens. We showed that knocking down each of algn-2/asparagine-linked glycosylation protein, zip-1/bZIP transcription factor, and C44B11.1/FAS apoptotic inhibitory molecule increased the transcript levels of NMD targets. Among these, algn-2 exhibited an age-dependent decrease in its expression and was required for maintaining normal lifespan and for longevity caused by various genetic interventions. We further demonstrated that upregulation of ALGN-2 by inhibition of daf-2/insulin/IGF-1 receptor contributed to longevity in an NMD-dependent manner. Thus, algn-2, a positive regulator of NMD, plays a crucial role in longevity in C. elegans, likely by enhancing RNA surveillance. Our study will help understand how NMD-mediated mRNA quality control extends animal lifespan.

INTRODUCTION
Eukaryotic cells are equipped with mechanisms that maintain proper gene expression and prevent the production of deleterious proteins. Nonsense-mediated mRNA decay (NMD) is an mRNA quality control mechanism that monitors and degrades abnormal transcripts with premature termination codons (PTCs) (He and Jacobson, 2015; Kim and Maquat, 2019; Kurosaki et al., 2019). NMD targets also include mRNAs with long (>1 kb) 3' untranslated regions, upstream open reading frames, or selenocysteine-encoding UGA codons (He and Jacobson, 2015; Kim and Maquat, 2019; Kurosaki et al., 2019). Thus, NMD is crucial for RNA quality surveillance and the maintenance of the correct transcriptome in organisms.

Key evolutionarily conserved components of NMD have been identified by using model organisms, including the budding yeast Saccharomyces cerevisiae and the nematode Caenorhabditis elegans. A genetic screen using C. elegans has identified the main NMD components, smg (suppressor with morphological effect on genitalia)-1 through smg-7 (Hodgkin et al., 1989; Mango, 2001). SMG-1 is a phosphatidylinositol 3-kinase-related kinase that phosphorylates SMG-2/UPF1 RNA helicase (Hodgkin et al., 1989; Mango, 2001). This event activates the NMD machinery and leads to the cleavage of target mRNAs via the endonuclease SMG-6 (Eberle et al., 2009; Huntzinger et al., 2008; Lykke-Andersen et al., 2014). Subsequently, the target mRNAs are degraded by exosomes and exonucleases (Schmid and Jensen, 2008). Additional research has led to the discovery of other NMD components, including smg-8, smg-9, smgl (smg lethal)-1, and smgl-2 (Longman et al., 2007; Yamashita et al., 2009); however, the role of smg-8 in NMD has been challenged by the characterization of smg-8 mutants (Rosains and Mango, 2012). Unlike the abovementioned, extensively characterized NMD components, the upstream regulators of NMD remain underexplored.

In C. elegans, NMD contributes to longevity by influencing the levels of various mRNAs (Son et al., 2017). Specifically, NMD function is crucial for extended lifespan conferred by reduced insulin/IGF-1 signaling (IIS), an evolutionarily conserved aging-regulatory pathway. Reduced IIS increases NMD and subsequently decreases the levels of specific transcripts, including yars-2b.1/tyrosyl-tRNA synthetase isoform b.1, and this, in turn, contributes to longevity. In addition, NMD is upregulated to modulate the splicing of various
gene transcripts that are important for dietary restriction-induced longevity (Tabrez et al., 2017). Despite these initial findings on the roles of the known NMD components in longevity in C. elegans, it remains poorly understood whether and how NMD contributes to extended lifespan and delayed aging.

In the present report, we identified modulators of NMD by employing two genetic screens. We first performed a genome-wide RNAi screen for the modifiers of NMD by using a fluorescent NMD reporter and subsequently validated the results by measuring the level of rpl-7A, an endogenous NMD target transcript. We found that RNAi targeting each of algn-2/asparagine-linked glycosylation protein, zip-1/bZIP transcription factor, and C44B11.1/FAS apoptotic inhibitory molecule (FAIM) increased the levels of the rpl-7A transcript. We also showed that two of the mutants isolated from our mutagenesis screen exhibited decreased rpl-7A transcript levels. We then found that algn-2, the expression of which declined during aging, was required for maintaining the normal lifespan. Furthermore, we showed that knocking down algn-2 significantly decreased the longevity conferred by various genetic interventions, including daf-2/insulin/IGF-1 receptor mutations, dietary restriction mimetic eat-2 mutations, and mitochondrial respiration-defective isp-1 mutations. We further showed that ALGN-2 was upregulated upon the genetic inhibition of the daf-2/insulin/IGF-1 receptor and contributed to a long lifespan in an SMG-2-dependent manner. Overall, our study identified previously unknown modulators of NMD, including algn-2, which plays key roles in RNA quality control and organismal longevity.

RESULTS
A Genome-wide RNAi Screen Identified Modifiers of NMD
To identify NMD regulators, we first performed a genome-wide RNAi screen in a liquid culture system using an NMD-responsive GFP reporter (Figure 1A). The NMD reporter contains a PTC in the first exon of the GFP-fused lacZ gene, driven by a ubiquitous sec-23 promoter, sec-23p::gfp::lacZ(PTC) (Longman et al., 2007) (Figure 1A). Under normal conditions, this transcript is degraded by NMD, and the worms, therefore, display dim GFP fluorescence (Figure 1A). We used a sensitized loss-of-function mutant background of smg-1, which encodes a kinase that phosphorylates SMG-2/UPF1 and induces NMD (Grimson et al., 2004). With the increased basal GFP expression of the reporter in the smg-1(−/) mutant background, we searched for genes whose knockdown further changed the green fluorescence intensity (Figure 1A). We initially found that the GFP expression levels were further increased by 39 RNAi clones and decreased by 38 RNAi clones (arbitrary cutoff: > 1.5 and −1.5 <, Figure 1B and Table S1). By repeating the experiments six times, we confirmed that 25 and 7 RNAi clones increased and decreased the GFP levels, respectively (cutoff of mean value: > 0.6 and −0.6 <, Figure 1C and Table S1).

RNAi Targeting algn-2, zip-1, or C44B11.1 Increased the Level of an NMD Target Transcript, rpl-7A(PTC)
We then validated the RNAi clones from our liquid culture-based screen by using a solid-medium culture system. We found that 24 of the 25 RNAi clones that increased the GFP levels in liquid culture did the same on solid media (Figures 2A and 2B, Table S1). However, none of the seven RNAi clones that decreased the GFP levels in liquid culture did so on solid media (Figures S1A and S1B, Table S1). We further determined the effects of our hit RNAi clones on an endogenous NMD target gene, rpl-7A(PTC) (Mitrovich and Anderson, 2000) (Figure 2C and Table S1). We found that RNAi knockdown of algn-2, zip-1, or C44B11.1 significantly increased the level of the rpl-7A(PTC) transcript in smg-1(−/) mutants as well as in wild-type animals (Figures 2D–2G, Table S1). Thus, algn-2, zip-1, and C44B11.1 appear to be positive modulators of NMD.

An EMS Mutagenesis Screen Identified Potential Negative Modulators of NMD
Because we were not able to identify RNAi clones that reliably decreased the NMD reporter GFP level from our genome-wide RNAi screen (Figure S1), we sought to identify such genetic inhibition by employing an EMS mutagenesis screen (Figure S3A). From the 83,400 mutagenized haploid genomes of smg-1(−/); sec-23p::gfp::lacZ(PTC) animals, we obtained 11 and 9 fertile mutant strains with decreased and increased GFP levels, respectively (Figure S3A). We then confirmed the reproducibility of the results by quantifying the fluorescence levels of the mutants (Figure S3B). Because of the absence of screened RNAi clones that reproducibly decreased the NMD reporter GFP level, we focused on potential negative NMD regulator mutants. Among the 11 mutants that displayed reduced NMD reporter GFP levels, we found that two mutant alleles, yh47 and yh57, significantly decreased the level of the rpl-7A(PTC) transcript using qRT-
PCR (Figure S3C). These data suggest that genes that are mutated by yh47 and yh57 alleles encode potential negative regulators of NMD.

**algn-2 Whose mRNA Level Decreases with Age Is Required for Maintaining the Normal Lifespan**

We previously reported that NMD function decreases with age and mediates longevity conferred by reduced IIS in *C. elegans* (Son et al., 2017). Thus, in the present study, we tested whether algn-2, zip-1, or C44B11.1 exhibited age-dependent expression changes or contributed to longevity. We found that the mRNA levels of algn-2 and zip-1 decreased with age, whereas that of C44B11.1 exhibited an age-dependent increase (Figures 3A–3C). These data imply a functional association between algn-2, zip-1, and aging in *C. elegans*. We then determined whether RNAi targeting each of algn-2, zip-1, and C44B11.1 affected lifespan. We found that algn-2 RNAi significantly shortened the lifespan of wild-type animals, whereas zip-1 RNAi or C44B11.1 RNAi did not (Figures 3D–3F). Thus, algn-2, which exhibits an age-dependent decrease in mRNA levels, is required for the maintenance of the normal lifespan.
Figure 2. Validation of NMD Modulators Identified from Genome-wide RNAi Screen

(A) Representative RNAi clones targeting potential NMD regulators on solid media. Of 25 potential positive NMD regulators identified from the genome-wide RNAi screen, 24 increased the GFP expression level on solid culture media (scale bar, 100 μm). Ctrl, control RNAi.

(B) Quantification data for panel (A). Error bars represent SEM (two-tailed Student’s t test, *p < 0.05, **p < 0.01, ***p < 0.001, n = 24 from three independent experiments). See Table S1 for the relative GFP level change caused by treating with each RNAi clone.

(C) Schematic diagram showing the alternative splicing of rpl-7A that generates rpl-7A(Normal), a normal transcript, and rpl-7A(PTC), an established NMD target transcript.

(D) Volcano plot showing normalized rpl-7A(PTC) level and p values from RT-PCR in smg-1(tm849) mutants with each of the RNAi clones. p values were calculated using a two-tailed Student’s t test. A dotted line indicates p = 0.05, and black triangles indicate smg-2 RNAi and smg-4 RNAi, which were used as positive controls. See Table S1 for the normalized rpl-7A(PTC) ratio for the indicated RNAi clones and Figure S1 legends for discussion regarding (B and D).
Lastly, we tested whether the transgenic overexpression of algn-2 RNAi (n = 3) (F), and C44B11.1 RNAi (n = 3) (G). Error bars represent SEM (two-tailed Student’s t test, *p < 0.05).

algn-2 Is Crucial for Longevity Conferred by Various Interventions, including daf-2/Insulin/IGF-1 Receptor Mutations that Enhance NMD

Next, we focused our functional analysis on the role of algn-2 in the lifespan. We showed that algn-2 RNAi did not further decrease short lifespan caused by mutations in smg-2 (Figure 4A). Thus, algn-2 appears to act together with SMG-2 to maintain normal lifespan in C. elegans. We have previously reported that SMG-2, an RNA helicase essential for NMD (Kim and Maquat, 2019; Page et al., 1999), is critical for C. elegans longevity resulting from reduced IIS, which increases NMD (Son et al., 2017). We therefore tested whether algn-2 contributed to the longevity conferred by reduced IIS. We found that algn-2 RNAi significantly decreased the long lifespan conferred by two mutant alleles of daf-2/insulin/IGF-1 receptor gene, e1370 and e1368 (Figures 4B and 4C), or a phosphoinositide-3 kinase gene mutation, age-1(hx546) (Figure 4D). In addition, algn-2 RNAi did not further decrease the shortened lifespan of smg-2(–); daf-2(–) mutants (Figure 4E). These results suggest that algn-2 contributes to longevity conferred by reduced insulin/IGF-1 signaling by acting together with SMG-2.

We also tested the effect of algn-2 RNAi on the longevity of dietary restriction mimetic eat-2(–) mutants and mitochondrial respiration-defective isp-1(–) mutants. We found that RNAi knockdown of algn-2 significantly shortened the long lifespan of eat-2(–) mutants (Figure 4F) and isp-1(–) mutants (Figure 4G). These data indicate that algn-2 is generally required for lifespan extension caused by various interventions in C. elegans.

Genetic Inhibition of daf-2 Upregulates ALGN-2 to Increase NMD Function and Lifespan

We next generated transgenic animals expressing GFP-fused ALGN-2 under the promoter of algn-2(2p::algn-2::gfp) and detected ALGN-2::GFP in the intestine and neurons (Figure 5A). We showed that ALGN-2::GFP was localized in the cytoplasm of the cells (Figure 5B), consistent with the localization of mammalian ALG2, the ALGN-2 ortholog, on the cytosolic face of the endoplasmic reticulum (ER) (Dean and Gao, 2014).

We then examined whether the genetic inhibition of daf-2 affected the level of ALGN-2. Interestingly, daf-2 RNAi significantly increased the level of ALGN-2::GFP (Figures 5C and 5D), without affecting that of algn-2 mRNA (Figure S2B), suggesting that reduced insulin/IGF-1 signaling upregulates ALGN-2 at the post-transcriptional level. Next, we determined whether algn-2 affected NMD function in daf-2(–) mutants. Specifically, we measured the level of rpl-12(PTC) transcript, an endogenous NMD target whose level decreases in daf-2(–) mutants and increases with age (Mitrovich and Anderson, 2000; Son et al., 2017) (Figure 5E). We showed that algn-2 RNAi substantially increased the rpl-12(PTC) level in daf-2(–) mutants (Figure 5E). Together, these data suggest that upregulation of ALGN-2 conferred by reduced insulin/IGF-1 signaling enhances NMD in C. elegans.

Lastly, we tested whether the transgenic overexpression of algn-2::gfp affected lifespan. We found that the transgene that expressed algn-2::gfp at the highest level among three lines (Figures 5S and 5Sb) significantly increased lifespan (Figure 5F), whereas the other two did not (Figure 5Sc). Together, these data suggest that upregulation of ALGN-2 can increase lifespan by enhancing NMD.

DISCUSSION

ALGN-2, an NMD Modulator, Contributes to Longevity in C. elegans

NMD is an evolutionarily conserved process that is crucial for the monitoring and maintenance of RNA quality and, therefore, cellular RNA homeostasis. In the present report, we performed a series of genetic screens to identify modifiers of NMD. From our genome-wide RNAi screen, we identified algn-2, zip-1, and C44B11.1/FAIM as NMD modulators that were required for downregulation of NMD target transcripts. Among these, we showed that algn-2 contributed to longevity conferred by daf-2/insulin/IGF-1 receptor mutations in an NMD-dependent manner. Our study highlights the power of genetic screens for identifying factors that regulate NMD function and thereby affect organismal lifespan, possibly through the maintenance of RNA quality.
ALGN-2 May Affect NMD Function by Regulating the ER Stress Response

ALG2, the mammalian ortholog of ALGN-2, is a mannosyltransferase located at the ER membrane (Dean and Gao, 2014). ALG2 catalyzes the second and third mannosylation steps, which convert guanosine diphosphate mannose to core asparagine (N)-glycan (Li et al., 2019). Immature glycosylation causes severe ER stress, resulting in the activation of the unfolded protein response (UPR) (Cherepanova et al., 2016). ER stress can inhibit NMD and causes the NMD complex to be localized to the ER (Sakaki et al., 2012; Usuki et al., 2019).

algn-2 RNAi also increases ER stress in C. elegans (Akiyoshi et al., 2015; Ho et al., 2019) and may therefore decrease NMD by altering ER stress responses. In addition, defects in N-linked glycosylation pathway genes, including ALG2, are linked to congenital myasthenic syndrome (CMS), which involves the impairment of signal transmission at neuromuscular synapses (Cossins et al., 2013; Engel, 2018). Interestingly, NMD is impaired in some types of CMS (Rahman and Nasrin, 2016), suggesting that ALG2-regulated NMD processes play a role in the pathophysiology of CMS. Further studies are required to dissect the mechanisms by which inhibition of algn-2 affects the pathophysiology of the disease and NMD function potentially through affecting ER stress.

ALGN-2 Upregulates NMD in an SMG-2-Dependent Manner

One interesting aspect of our study is that we identified algn-2 from an RNAi screen that used an NMD reporter in a smg-2 mutant background. Interestingly, we found that algn-2 RNAi did not affect the lifespan of smg-2 mutants. Our previous report on NMD and longevity (Son et al., 2017) indicates that different SMG components differentially contribute to NMD function. Specifically, knockdown of smg-1 has an intermediate effect on NMD function among five smg genes that we functionally tested. In contrast, smg-2 RNAi has the biggest effect on NMD function. These data suggest that genetic inhibition of smg-2 causes more severe impacts on NMD function than that of smg-1 does. Thus, we speculate that algn-2 RNAi did not further decrease the lifespan of smg-2(−) animals, while influencing NMD target levels in smg-1(−) mutants, because of their differential effects on NMD functions.

NMD-Regulated RNA Quality Control Is Crucial for Organismal Longevity

Recent reports indicate that NMD-mediated RNA homeostasis is critical for longevity in C. elegans (Son et al., 2017; Tabrez et al., 2017). These studies highlight the importance of NMD for clearing abnormal transcripts and promoting RNA homeostasis, which may help animals maintain health during aging. However, it remained unknown which factor causes a functional decline of NMD during aging. Our data suggest that
reduced levels of factors that regulate NMD, including algn-2, underlie age-dependent decreases in NMD and possibly result in impaired RNA quality control. Our previous reports have also indicated that the NMD in the nervous system is crucial for lifespan extension (Son et al., 2017). Because the brain is the organ that expresses Alg2 at the highest level in young mice (Imamura et al., 2014), Alg2 expression in the brain may decrease in an age-dependent manner, perhaps contributing to the shortening of the lifespan by reducing NMD efficiency. It will be interesting to test whether preserving or increasing the levels of positive NMD modulators, including ALG2/ALGN-2, can be exploited as a strategy for avoiding the adverse effects of age-dependent declines in RNA quality, in particular in the nervous systems.

Figure 4. algn-2 Contributes to Longevity Conferred by Various Interventions, including Reduced Insulin/IGF-1 Signaling by Acting Together with SMG-2

(A) algn-2 RNAi did not decrease the lifespan of smg-2(qd101) [smg-2(–)] mutants. (B–D) RNAi targeting algn-2 significantly decreased the long lifespan of daf-2(e1370) [daf-2(–)] (B), daf-2(e1368) (C), and age-1(hx540) [age-1(–)] (D) mutants. Different from algn-2 RNAi, neither zip-1 RNAi nor C44B11.1 affected the lifespan of daf-2(–) mutants (Figures S4A and S4B). (E) algn-2 RNAi had a small effect on the lifespan of smg-2(–); daf-2(–) double mutants. algn-2 mRNA levels were not changed by smg-2(–) or daf-2(–) mutations (Figure S2B). (F and G) algn-2 RNAi shortened longevity induced by eat-2(ad1116) [eat-2(–)] (F) and isp-1(qm150) [isp-1(–)] (G) mutations. See Table S3 for statistical analysis and additional lifespan data.
Limitations of the Study

Although we identified algn-2 as a positive regulator of NMD, which contributed to longevity in *C. elegans*, the biochemical mechanisms by which ALGN-2 affects NMD functions remain elusive. In addition, global transcriptomic analysis will help identify all the target transcripts of algn-2. Although we showed that decreased NMD function due to *smg-1(C0)* was suppressed by *yh47* and *yh57*, two mutant alleles that we identified from our mutagenesis screen, the molecular identities of the mutations are unknown. Additionally, we did not test whether algn-2 modulated NMD in a cell autonomous or a cell non-autonomous manner. These are important issues that need to be addressed in future studies.

Resource Availability

Lead Contact

Further request and information for resources and reagents used in this published article should be directed and will be fulfilled by the Lead Contact, Seung-Jae V. Lee (seungjaevlee@kaist.ac.kr).
Materials Availability
All data analyzed and generated in this research are included in this published article and supplemental information.

Data and Code Availability
The published article includes all data generated in this study.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101713.

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AUTHOR CONTRIBUTIONS
E.J.E.K., H.G.S., H.-E.H.P, Y.J., S.K., and S.-J.V.L designed the experiments. E.J.E.K., H.G.S., H.-E.H.P, Y.J., and S.K. performed the experiments. E.J.E.K. and S.-J.V.L. analyzed the data. E.J.E.K. and S.-J.V.L. wrote the manuscript.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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Supplemental Information

Caenorhabditis elegans alg-2 Is Critical for Longevity Conferred by Enhanced Nonsense-Mediated mRNA Decay

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Figure S1

A

B

Ctrl  gfp  rpb-2  ccch-2  F56C11.3
lim-7  rps-10  T09B4.8  Y116A8C.19

Relative GFP intensity (a.u.)
Figure S2

A

Relative algn-2 mRNA level

RNAi: Ctrl algn-2

B

Relative algn-2 mRNA level

WT daf-2(-) smg-2(-) smg-2(-); daf-2(-)

C

Relative zip-1 mRNA level

RNAi: Ctrl zip-1

D

Relative C44B11.1 mRNA level

RNAi: Ctrl C44B11.1

E

Relative algn-2 mRNA level

WT smg-2(-)

F

Relative zip-1 mRNA level

WT smg-2(-)

G

Relative C44B11.1 mRNA level

WT smg-2(-)
Figure S3

A  EMS Mutagenesis

\[ \text{Norm} \text{alized ratio} = \frac{rpl-7A(PTC)}{rpl-7A(Normal)} \]

B  Relative GFP intensity (a.u.)

| Ctrl 42 43 46 47 48 49 50 56 57 58 61 62 63 64 | yh   |
|------------------------------------------------|------|
| ![Graph showing relative GFP intensity](image)  |      |

C  Normalized ratio

| Normalized ratio | Ctrl 42 43 46 47 48 49 50 56 57 | yh   |
|------------------|---------------------------------|------|
| ![Graph showing normalized ratio](image)      |      |
Figure S4

(A) daf-2(-) daf-2(-) zip-1 RNAi

(B) daf-2(-) daf-2(-) C44B11.1 RNAi
Figure S5

A

[Images of C. elegans expressing GFP in different lines]

B

[Graph showing relative GFP intensity (a.u.) for different lines with statistical significance indicated]

C

[Graph showing % Alive over Days of adulthood for Control and alg-2::gfp lines]
Supplemental Information

Supplemental Figure Legends

Figure S1. RNAi clones that decreased GFP levels in liquid culture media did not exert similar effects on solid media, related to Figures 1 and 2. (A) Seven representative RNAi clones that decreased GFP levels from a genome-wide RNAi screen on solid media (scale bar, 100 μm). gfp RNAi was used as a positive control. (B) Quantification data of images shown in panel A. Some of the hit RNAi clones from our screen may include false positives that affect transgene expression rather than NMD function. RNAi targeting each of hlh-28, K12H6.2, ubc-1, W10G11.19, and mrf-5 that increased the levels of the NMD reporter GFP in this study have been reported to increase the expression of other transgenes (Broday et al., 2004; Kapulkin et al., 2005; Lamitina et al., 2006; Marza et al., 2015; Quach et al., 2013). In addition, ccch-2 and rpb-2 RNAi clones that decreased our NMD reporter GFP level have been shown to decrease transgene expression (MacNeil et al., 2015; Winter et al., 2012). Thus, it seems likely that some of our hit RNAi clones altered the GFP levels in the NMD reporter transgenic worms by generally altering transgene expression, independently of NMD function. Therefore, our additional qRT-PCR experiments, which measured endogenous NMD target mRNA levels, validated genuine hit RNAi clones that affected NMD. We also noticed that increases in rpl-7A(PTC) mRNA levels caused by RNAi targeting each of algn-2, zip-1, and C44B11.1 were smaller than those caused by smg-2 RNAi (Fig. 2D); this is different from the similar effects of these three RNAi clones and smg-2 RNAi on the changes in the NMD reporter GFP levels (Fig. 2A-B). SMG-2 is the central component of NMD.
resulted in a striking increase in \textit{rpl-7A(PTC)} mRNA levels as NMD cannot function properly. In contrast, \textit{alg}-2, \textit{zip}-1, and \textit{C44B11.1} appear to be modulators of NMD. Thus, we speculate that the effects of \textit{alg}-2 RNAi, \textit{zip}-1 RNAi, and \textit{C44B11.1} RNAi on \textit{rpl-7A(PTC)} mRNA levels are smaller than that of \textit{smg-2} RNAi. Error bars represent s.e.m (two-tailed Student’s \textit{t}-test, \(*p < 0.05, **p < 0.001, n=24\) from three independent experiments, a.u.: arbitrary unit). See Supplemental Table S1 for the descriptions and scores of indicated RNAi clones from the screen.

**Figure S2.** The effects of various genetic interventions that we tested in this study on the mRNA levels of NMD targets and regulators, related to Figures 3 and 4. (A) \textit{alg}-2 RNAi significantly decreased the mRNA levels of \textit{alg}-2 in wild-type worms measured by using qRT-PCR (n=3). (B) \textit{alg}-2 mRNA levels were similar among wild-type (WT), \textit{daf}-2(e1370) [\textit{daf}-2(-)], \textit{smg}-2(qd101) [\textit{smg}-2(-)], and \textit{smg}-2(-); \textit{daf}-2(-) mutant animals (n=3). (C-D) \textit{zip}-1 RNAi (C) and \textit{C44B11.1} RNAi (D) significantly decreased the mRNA levels of \textit{zip}-1 and \textit{C44B11.1}, respectively, in WT worms (n=3). (E-G) The mRNA level of \textit{alg}-2 (E), \textit{zip}-1 (F), or \textit{C44B11.1} (G) was not significantly altered in \textit{smg}-2(qd101) [\textit{smg}-2(-)] mutants compared with that in WT animals (n=3). Error bars represent s.e.m (two-tailed Student’s \textit{t}-test, **\(p < 0.01, ***p < 0.001\)).

**Figure S3.** EMS mutagenesis screen for mutants with altered NMD function, related to Figure 2. (A) Our EMS mutagenesis screen identified 11 suppressors and
9 enhancers that decreased and increased the GFP intensity, respectively, in smg-1(-); sec-23p::gfp::lacZ(PTC) animals. See Supplemental Table S2 for isolated mutants and their lineage. (B) Quantification of the GFP fluorescence levels of isolated mutants compared to the control (Ctrl), smg-1(-); sec-23p::gfp::lacZ(PTC) animals. a.u.: arbitrary unit. Error bars represent s.e.m (two-tailed Student’s t-test, *p < 0.05, ***p < 0.001, n ≥ 24). (C) rpl-7A(PTC) levels normalized to rpl-7A(Normal) measured by qRT-PCR with mutants that displayed decreased GFP levels from the EMS mutagenesis screen using smg-1(-); sec-23p::gfp::lacZ(PTC) animals as a control. Error bars represent s.e.m (two-tailed Student’s t-test, *p < 0.05, **p < 0.01, n=3).

Figure S4. zip-1 RNAi or C44B11.1 RNAi did not affect the lifespan of daf-2(e1370) mutants, related to Figure 4. RNAi targeting zip-1 (A) or C44B11.1 (B) did not affect longevity conferred by daf-2(e1370) [daf-2(-)] mutations. We showed that the mRNA level of algn-2 positively correlates with NMD function with age (Fig. 3A). In contrast, the mRNA level of C44B11.1 increased with age but the NMD function decreased with age (Fig. 3C). One can speculate that C44B11.1 expression increases in an age-dependent manner to compensate the impaired NMD in old worms. C44B11.1 RNAi did not affect lifespan (Fig. 3F), whereas algn-2 RNAi significantly shortened lifespan (Fig. 3D). This is likely because algn-2 affects NMD targets that are critical for lifespan regulation, whereas C44B11.1 may regulate a different subset of NMD targets that have a small or no effect on lifespan. See Supplemental Table S3 for statistical analysis and additional lifespan data.
Figure S5. Lifespan assay using algn-2::gfp, related to Figure 5. (A-B) The images of three transgenic lines that expressed algn-2::gfp. yhEx540[algn-2p::algn-2::gfp; ofm-1p::rfp] [Line 1], yhEx541[algn-2p::algn-2::gfp; ofm-1p::rfp] [Line 2], and yhEx542[algn-2p::algn-2::gfp; ofm-1p::rfp] [Line 3] (scale bar, 100 μm). (B) Quantification of ALGN-2::GFP levels in panel A. Error bars represent s.e.m (two-tailed Student’s t-test, ***p < 0.001, n=24 from three independent experiments, a.u.: arbitrary unit.). (C) algn-2::gfp line 2 or algn-2::gfp line 3 did not affect lifespan. See Supplemental Table S3 for statistical analysis and additional lifespan data.

Transparent Methods

Strains

The following strains were used in this study: Lee laboratory N2 wild-type (WT), IJ1600 smg-1(tm849) I outcrossed eight times with N2, PTCxi sec-23p::gfp::lacZ(PTC), IJ1601 smg-1(tm849); sec-23p::gfp::lacZ(PTC) obtained by crossing FX17751 smg-1(tm849) and PTCxi, sec-23p::gfp::lacZ, IJ445 smg-2(qd101) I outcrossed four times with N2, CF1041 daf-2(e1370) III outcrossed six times with N2, IJ446 smg-2(qd101) I; daf-2(e1370) III, IJ385 daf-2(e1368) III outcrossed nine times with N2, IJ256 age-1(hx546) II outcrossed four times with N2, IJ173 eat-2(ad1116) II outcrossed four times with N2, CF2172 isp-1(qm150) IV outcrossed three times with N2, IJ1754 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh42) IJ1755 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh43), IJ1756 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh46), IJ1757 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh47),
IJ1758 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh48), IJ1759 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh49), IJ1760 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh50),
IJ1761 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh51), IJ1766 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh56), IJ1767 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh57),
IJ1768 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh58), IJ1762 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh52), IJ1763 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh53),
IJ1764 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh54), IJ1765 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh55), IJ1769 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh59),
IJ1771 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh61), IJ1772 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh62), IJ1773 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh63),
IJ1774 smg-1(tm849); sec-23p::gfp::lacZ(PTC); (yh64), IJ1539 yhEx414[ofm-1p::rfp],
IJ2047 yhEx540[alg-2p::alg-2::gfp; ofm-1p::rfp], IJ2048 yhEx541[alg-2p::alg-2::gfp; ofm-1p::rfp], IJ2049 yhEx542[alg-2p::alg-2::gfp; ofm-1p::rfp]. All strains
were maintained at 20°C.

Genome-wide RNAi screen
A genome-wide RNAi screen was performed based on a previous study (Lee et al., 2010), with some modifications. Gravid adult smg-1(tm849); sec-23p::gfp::lacZ(PTC)
animals were treated with bleach to obtain eggs (Stiernagle, 2006). The collected
eggs were incubated in M9 buffer at 20°C for approximately 18 h. On the same day
as worms were bleached, double-stranded (ds) RNA–expressing HT115 bacteria
from a commercially available C. elegans RNAi library (Source BioScience,
Nottingham, UK) were cultured in Luria broth (LB) containing 100 μg/ml ampicillin in
96-well plates overnight at 37°C. On the following day, isopropylthiogalactoside (IPTG) (Gold biotechnology, St Louis, MO, USA) was added to each well of the bacterial cultures to a final concentration of 4 mM, and the plates were placed in a 37°C incubator for 1 h. Bacterial pellets were collected by spinning down at 870 g for 10 min. The supernatant was discarded and 190 μl liquid nematode growth media (NGM) containing 100 μg/ml ampicillin plus 4 mM IPTG solution was added to the bacterial pellets. Approximately 20 L1 larval worms were added to each well with cultured bacteria and incubated at 20°C in a shaking incubator. Three days later, the plates were scored for GFP intensity by four researchers independently. Semi-quantitative GFP scores ranging from −3 (darkest) to +3 (brightest) were given to each well. Control empty vector RNAi (L4440) and gfp RNAi were used as controls to provide references for scores 0 and −3, respectively. From the primary RNAi screen, arbitrary cutoff scores for positive hits were set (> +1.5 and < −1.5). The liquid-based screen was repeated six times by the four independent researchers using an RNAi sublibrary that included 77 candidate RNAi clones from our initial RNAi screen. The mean scores of the six repeats were calculated and arbitrary cutoff scores were set as the values +0.6 and −0.6.

Microscopy

Fluorescence imaging was performed based on a previous study (Lee et al., 2015), with some modifications. For measuring GFP fluorescence levels in smg-1(tm849) I; PTCxi::GFP animals, the transgenic worms were fed with designated RNAi bacteria from hatching. Synchronized young adult (day 0) worms were placed on a 2%
agarose pad and paralyzed with 2 mM levamisole. Images of the worms were
captured using an AxioCam HRc CCD digital camera connected to a Zeiss Axio
Scope A1 microscope. The fluorescence intensity of the worms was quantified using
ImageJ (http://imagesj.nih.gov/ij/) (Schneider et al., 2012) after subtracting
background fluorescence signals. Confocal fluorescence images of algn-2p::algn-
2::gfp worms were acquired using an inverted LSM880 laser scanning microscope
(Zeiss Corporation, Germany) with Plan Apochromat 20x0.8 M27, Plan-Apochromat
63x1.4 oil DIC M27 objectives. Green fluorescence was detected with the excitation
wavelength at 488 nm and emission wavelength 526 nm.

EMS mutagenesis screen

An EMS mutagenesis screen was performed based on a previous study (Gürel et al.,
2012), with some modifications. smg-1(tm849) I; PTCxi::GFP transgenic worms
(1,150 animals) at the late L4 stage were collected, washed twice with M9 buffer,
and placed in 2 ml M9 buffer. The worms were then treated with 47 mM liquid ethyl
methanesulfonate (EMS, Sigma, St. Louis, MO, USA) and incubated at 20°C for 4 h
with shaking. The worms were then washed twice with M9 buffer and transferred
onto OP50-seeded NGM plates to recover for 4 h. The EMS-treated P₀ worms were
then evenly transferred onto 14 NGM plates and removed from the plates 24 h later,
when F₁ progeny were generated. Gravid 41,700 F₁ adults were then treated with
bleach solution to obtain F₂ eggs. Approximately 500 F₂ eggs were placed on each
NGM plate (400 plates total) and grown to adulthood. The F₂ adult worms that
exhibited greater or lesser GFP intensity than control worms were isolated and
individually cultured to establish mutant lines. Initially, 98 mutants (63 with increased
GFP levels and 35 with decreased GFP levels) were isolated. Among these 98
mutant lines, 35 and 15 mutants that exhibited increased and decreased GFP levels,
respectively, were sterile; the increased and decreased GFP levels of 19 and 9
mutants, respectively, were not reproducible. Finally, 9 and 11 mutant strains that
reproducibly displayed increased and decreased GFP fluorescence, respectively,
relative to control RNAi-treated worms were established.

**RT-PCR**

RT-PCR was performed based on a previous study (Son et al., 2017), with some
modifications. dsRNA-expressing HT115 bacteria were cultured in LB containing 50
μg/ml ampicillin overnight at 37°C. The bacteria were then seeded on NGM plates
and incubated overnight at 37°C. IPTG (1 mM) was added onto the bacteria-seeded
plates, which were then kept at room temperature for 1 d. Approximately 500
bleached eggs of *smg-1(tm849)* mutants were placed on the plates and grown until
reaching the young adult stage. Total RNA was extracted from synchronized worms
using RNA IsoPlus (Takara, Shiga, Japan). cDNA was generated using ReverTra
Ace® qPCR RT Master Mix with gDNA remover (Toyobo, Osaka, Japan). *rpl-7A(PTC)*
and *rpl-7A(Normal)* cDNAs were amplified using the primers that were reported
(Rosains and Mango, 2012). PCR products were analyzed by electrophoresis on a
2% agarose gel for 25 min. The band intensity was quantified using ImageJ
(http://imagesj.nih.gov/ij/). The sequences of the primers that were used are as
follows:

*rpl-7A(PTC)-F*: GACAGCCAGTCGGTGG
Quantitative RT-PCR analysis

Quantitative RT-PCR was performed based on a previous study (Jeong et al., 2020), with some modifications. All worms were maintained at 20°C. Levels of rpl-7A(PTC) transcript were measured using the mutant worms that we isolated, and smg-1(tm849); sec-23p::gfp::lacZ(PTC) transgenic animals were used as a control. Total RNA was extracted from synchronized day 1 adult worms using RNA IsoPlus. cDNA was generated using ReverTra Ace® qPCR RT Master Mix with gDNA remover and was used for quantitative PCR to measure mRNA levels. Quantitative PCR with the Power SYBR® Green PCR master mix (Applied Biosystems, Foster City, CA, USA) was performed using a StepOne Real Time PCR System (Applied Biosystems, Foster City, CA, USA). The analysis was performed by using a comparative C₉ method. The ama-1 mRNA level was used as a control for normalization. The sequences of the primers that were used are as follows:

ama-1-F: TGGAACTCTGGAGTCACACC
ama-1-R: CATCCTCCTTCAATGACCG
rpl-7A(PTC)-F: GACAGCCAGTCCCGTGG
rpl-7A(PTC)-R: GTCTAGTTCACCTACAGGAAATG
Lifespan assays

Lifespan assays were performed based on a previous study (Lee et al., 2019), with some modifications. HT115 bacteria that expressed dsRNA were cultured in LB containing 50 μg/ml ampicillin (USB, Santa Clara, CA, USA) at 37°C overnight. The cultured HT115 RNAi bacteria (100 μl) were then seeded on NGM containing 50 μg/ml ampicillin and incubated at 37°C overnight. IPTG (1 mM) was added and incubated for 24 h at room temperature. To prevent progeny from hatching, 5-fluoro-
2'-deoxyuridine (FUDR) was added to the RNAi bacteria-seeded plates at a final concentration of 5 μM. Young (day 1) adult worms were placed on freshly prepared plates and subsequently transferred onto new plates after 1 or 2 d. Worms that did not respond to a gentle touch with a platinum wire pick were scored as dead. Worms that ruptured, burrowed, bagged, or crawled off the plates were censored but included for subsequent statistical analysis. All lifespan assays were performed at 20°C by at least two independent researchers. OASIS (online application of survival analysis, http://sbi.postech.ac.kr/oasis) was used for statistical analysis of the lifespan assay results (Yang et al., 2011). A log-rank (Mantel-Cox method) test was used to calculate p values.

Cloning and generation of transgenic worms

A promoter (~0.3 kb upstream of the start codon) and the coding region of algn-2 (~1.8 kb) were PCR amplified by using C. elegans genomic DNA as a PCR template. pPD95.75 (Fire lab C. elegans vector kit) plasmid was linearized by KpnI restriction enzyme. The PCR product of algn-2 promoter and algn-2 genomic region were inserted into the linearized pPD95.75 using In-fusion HD cloning kit (Takara, Shiga, Japan) to generate algn-2p::algn-2::gfp plasmid. The mixture of algn-2p::algn-2::gfp plasmid (25 ng μl⁻¹) and the control plasmid ofm-1p::rfp (75 ng μl⁻¹) were injected into day 1 adult worms.

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