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Translation Readthrough Mitigation

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Abstract

A fraction of ribosomes engaged in translation will fail to terminate when reaching a stop codon, yielding nascent proteins inappropriately extended on their C-termini. Although such extended proteins can interfere with normal cellular processes, known mechanisms of translational surveillance are insufficient to protect cells from potential dominant consequences. Through a combination of transgenics and CRISPR/Cas9 gene editing in *C. elegans*, we demonstrate a consistent ability of cells to block accumulation of C-terminal extended proteins that result from failure to terminate at stop codons. 3'UTR-encoded sequences were sufficient to lower protein levels. Measurements of mRNA levels and translation suggested a co- or post-translational mechanism of action for these sequences in *C. elegans*. Similar mechanisms evidently operate in human cells, where we observed a comparable tendency for translated human 3'UTR sequences to reduce mature protein expression in tissue culture assays, including 3' sequences from the hypomorphic "Constant Spring" hemoglobin stop codon variant. We suggest 3'UTRs may encode peptide sequences that destabilize the attached protein, providing mitigation of unwelcome and varied translation errors.

Failure of translation termination to occur at a stop codon can lead to ribosomes translating into a 3'UTR. In some cases translation may proceed through the 3'UTR and into the poly(A) tail, triggering a process termed "nonstop" decay and destabilizing both the mRNA and nascent protein (reviewed in¹). However, for a majority of 3'UTRs a stop codon is encountered prior to the poly(A) tail^{2,3}. Readthrough events that encounter a subsequent termination codon are outside the scope of known translational surveillance pathways including nonstop¹. Depending on the 3'UTR and the frame in which the ribosome enters, the late stop codon can be several, tens, or even hundreds of codons into a 3'UTR, producing variant proteins with potentially problematic C-terminal appendages. This issue is

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Author Contributions

JAA, ESC, and AZF designed *C. elegans* experiments. JAA and ESC conducted *C. elegans* experiments. NJ developed the RNA-seq2 protocol. JAA performed computational analyses. JAA conducted experiments in human cell lines, as designed and aided by JAA, GTH, CHL, MCB, and AZF. JAA and AZF wrote the paper with help from all authors.

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highlighted by several pathologies caused by late frameshifts or stop codon mutations where 3'UTR-encoded C-terminal extensions effect protein mislocalization^{4,5}, aggregation^{6,7}, and instability^{8–12}, with calamitous consequences for organisms. Depending on sequence, genetic background, conditions, and organism, estimates of readthrough efficiency vary from <1% to 10% or more, posing a potential problem of non-trivial magnitude^{10,13}.

We set out to determine whether, and to what extent, 3'UTR translation has an effect on gene expression with a fluorescent reporter system in *C. elegans*. Initially we selected 3'UTRs from three genes: *unc-54* (encoding a muscle myosin), *tbb-2* (a beta tubulin), and *rpl-14* (a ribosomal protein). For each gene, fusion of the 3'UTR to a GFP driven by the *myo-3* promoter resulted in robust fluorescence in body wall muscle (Fig 1a). Next, by mutating stop codons, we created GFP reporters for each gene where translation would read past the normal termination point, terminating instead at a stop codon part-way through the 3'UTR (Fig 1b). In each case the "late stop" reporter accumulated substantially less GFP, with differences in signal of at least 10-fold. As a control, a co-injected mCherry marker robustly expressed in the same cells. We conclude that translation into the 3'UTR can confer substantial loss of protein for at least these three 3'UTRs in *C. elegans*.

To test whether translation into 3'UTRs could confer a loss of protein expression more generally, a two-fluorescent-reporter system with each fluorophore transgene containing an identical 3'UTR was used. Nine genes were chosen arbitrarily to reflect a variety of functions and expression levels: rps-17 (small ribosomal subunit component), r74.6 (dom34/ pelota release factor homolog), hlh-1 (muscle transcription factor), eef-1A.1 (also known as eft-3, translation elongation factor), myo-2 (a pharyngeal myosin), mut-16 (involved in gene/ transposon silencing), bar-1 (a beta catenin), daf-6 (involved in amphid morphogenesis), and alr-1 (neuronal transcription factor). A criterion in choosing these genes was presence (common for *C. elegans* genes, Extended Data 1) of an in-frame stop codon in the 3'UTR at least 30 bases beyond the normal stop but upstream of known poly(A) sites. We fused the 3'UTRs of each gene separately to GFP and mCherry, removing the canonical termination codon in the GFP construct. For each of the nine genes tested, observed GFP signals were extremely faint, with raw GFP/mCherry fluorescence ratios of less than 0.1 (Fig 1c, Extended Data 2). As a control, versions of the GFP reporter with the normal termination codon intact provided robust GFP expression, 10-fold or more higher than the corresponding readthrough constructs (GFP/mCherry fluorescence ratios in the range 0.3–0.9).

Several observations provide hints at how translation into 3'UTRs might reduce protein levels:

- 1. Experiments with specific mutagenesis support a role for the eventual protein sequence. Shortening readthrough peptides (tested for *unc-54* and *tbb-2*) increased GFP expression (Fig 2a). Extending this analysis, an equal-length non-synonymous substitution in the *unc-54* 3'UTR restored GFP expression, whereas synonymous substitution with multiple base differences did not.
- 2. Mutagenesis analysis of constructs using a constant 3'UTR reinforced the inference of peptide sequence as the primary determinant of GFP loss. We

found that the nucleotide sequence between the normal termination codon and the first in-frame termination codon was sufficient to confer GFP loss if inserted at the end of the GFP coding region for *unc-54*, *tbb-2*, *hlh-1*, *daf-6*, *rps-20*, or *rps-30* (Fig 2b). The *rps-30* readthrough region had the weakest effect on GFP, and was the shortest (nine amino acids). We undertook further mechanistic dissection by synonymous variation of readthrough regions from *unc-54*, *tbb-2*, and *rps-20*, with GC contents from 35–60%, in some cases mutating >50% of bases. Each synonymously substituted variant conferred robust loss of GFP.

3.

Decreased expression following translation into the 3'UTR required peptide linkage between the upstream protein and the 3'UTR-encoded segment. To assess the relationship between (i) covalent linkage with the translated C terminal peptide and (ii) the outcome for the larger protein, we took advantage of a picornavirus-derived oligopeptide sequence that causes cleavage and release of the nascent chain, after which ribosomes continue translation of the downstream sequence^{14,15}. Insertion of the T2A peptide (EGRGSLLTCGDVEENPGP) between GFP and the *unc-54* 3'UTR-encoded sequence rescued GFP expression, whereas an uncleavable T2A* point mutant did not (Fig 2b). Restoration of GFP levels by T2A to the level of no-insert controls also argues against mRNA destabilization as a substantial factor in the protein loss observed upon readthrough.

The above results could be explained if GFP was generally incompatible with C-terminal fusions in our system. To address this, we inserted a variety of sequences downstream of GFP: 3xFLAG, 3xHA, three random sequences created *in silico*, and six arbitrary fragments of in-frame coding sequence from *C. elegans* genes, approximately length matched to 3'UTR-encoded sequences (Fig 2b). GFP expression varied between constructs but was generally higher than 3'UTR-encoded sequences: 3xHA, 3xFLAG, 2 of 3 of the random sequences, and 4 of 6 of the coding-derived fragments exhibited GFP:mCherry fluorescence ratios of >0.13, higher than all nine tested 3'UTR-derived C-terminal extensions and significant statistically (p=.004, KS test). Thus the effects of 3'UTR-encoded sequences are not explained by a general intolerance of GFP to C-terminal extensions (see also Methods, Extended Data 3, 4).

It was conceivable that peculiarities of GFP and/or transgene expression systems might underlie the above observations. To establish effects of 3'UTR translation at endogenous genes, we sought loci where: (1) a loss of protein would be detectable phenotypically, (2) Cterminal fusions are known to be functional, (3) the next in-frame stop codon of the endogenous locus is 10 amino acids past the annotated stop codon, yet upstream of annotated poly(A) sites¹⁶, and (4) there is little-or-no autoregulation/feedback. *unc-54* and *unc-22* satisfy all the criteria, and *pha-4*, *unc-45*, and *tra-2* at least the first three points (Methods). For each locus we mutated the stop codon to allow translation into the 3'UTR (Fig 3a)¹⁷. In parallel, we analyzed small insertions/deletions generating late frameshifts for *unc-22* and *unc-54*. Additional controls had length-matched sequences and/or GFP tags at

the C-terminus (Extended Data 5). For each of *unc-22*, *unc-45*, *unc-54*, and *tra-2*, translation into the 3'UTR in at least one frame generated a strong hypomorphic (near null) phenotype specific to each locus. Other C-terminal tags for each gene were well behaved (no loss of expression), although one *tra-2* C-terminal tag did produce a Tra phenotype. The ability to place alternative tags on the C-terminus without obvious phenotypic consequences argues against a general sensitivity of the C-terminus to tagging. For *unc-22* and *unc-54*, that elongation into the 3'UTR in only some frames elicited a hypomorphic phenotype argues against ribosome elongation into the 3'UTR as being detrimental *per se*.

To determine the consequences on gene expression upon translation into 3'UTRs, we analyzed the *unc-54(cc3389)* TAA(Stop)>AAT(Asn) mutation for its effects on RNA, translation, and protein output. We analyzed mRNA expression in *unc-54(cc3389/+)* heterozygotes (phenotypically wild type to avoid complications from an Unc phenotype). RNA-seq revealed the *unc-54(cc3389)* and wild type alleles at approximately equal amounts in the mRNA pool, suggesting 3'UTR translation does not appreciably destabilize the *unc-54* mRNA (Fig 3c). In parallel, we detected a ~20-fold reduction in UNC-54 protein in immunoblots in *unc-54(cc3389)* (Fig 3d). To look for possible alterations in translation for *unc-54(cc3389)*, we examined the distribution of RNase-protected mRNA fragments with ribosome footprint profiling¹⁸. We observed no significant difference in the loading of ribosomes on *unc-54* mRNA (Extended Data 6), nor on the number, distribution, frame, or fragment size of ribosomes in the extended region (Extended Data 7).

A model that arises from these observations is that 3'UTR-encoded peptides mark their resulting products for destruction, either co- or post-translationally. Conceivably this process might operate either in a specific cell/tissue type or in a broad spectrum of different contexts. A broadly-expressed reporter bearing a readthrough extension would be expected to highlight any tissue which failed to destabilize the C-terminal peptide. Using a broadly-expressed promoter (*eef-1A.1*) driving GFP with and without the *unc-54* 3'UTR-encoded peptide, we observed no cells where GFP was robustly retained (data not shown).

We likewise considered the possibility that 3'UTR-encoded peptides might act to limit protein levels in human cells, developing a specific assay using a lentiviral dual fluorescence reporter encoding puromycin N-acetyl-transferase tethered to mCherry-T2A, followed by eGFP and a multiple cloning site (Fig 4a). The resulting reporter expresses both fluorophores from the same mRNA, yet as two disjoint polypeptides, allowing consideration of a peptide tag's effect on eGFP expression independent of effects on mCherry/mRNA expression. We validated the split dual fluorophore approach in K562 cells using tags known to be destabilizing (d1ODC, d4ODC¹⁹) or not (3xFLAG, 3xHA) (Fig 4b). We selected 13 genes of varying expression and function, and inserted the region between the annotated termination codon and first-in-frame termination codon downstream of eGFP. For 9 of 13 genes, the readthrough region reduced the eGFP:mCherry fluorescence ratio between 3 and 30-fold, a stronger reduction than the degron *d4ODC* (Fig 4c). While not universal, the substantial loss of eGFP fluorescence for a majority of readthrough regions opens up the possibility that translation into 3'UTRs may be generally inhibitory to expression across systems.

We hypothesize that a function of 3'UTRs is to minimize the accumulation of extended protein products that could be produced through translational readthrough.

This feature may prove generally significant in the causation of genetic disease. For example, readthrough alleles (e.g. Stop>Gln) of the *HBA2* locus in humans produce a fraction (~1%⁹) of normal *HBA2* protein (alpha globin), causing thalassemia. Translation into the *HBA2* 3'UTR is known to destabilize the *HBA2* mRNA²⁰, but it is unclear what effect the appended C-terminal 31 amino acids have on *HBA2* protein. We considered the possibility that the *HBA2* 3'UTR-encoded peptide might prevent protein expression in humans, contributing to the loss of *HBA2* protein. When appended to eGFP, the *HBA2* 3'UTR-encoded peptide decreased the eGFP:mCherry fluorescence ratio in K562 cells (Fig 4d). Furthermore, eGFP fluorescence was rescued by a self-cleaving (but not an uncleaveable mutant) T2A peptide.

Several observations from the literature corroborate the notion that 3'UTRencoded peptides may be detrimental to expression for more genes and organisms than those assayed here. In *S. cerevisiae*, translation past a point in the *his3*3'UTR confers a substantial loss in protein expression, without detectable effects on mRNA levels¹¹. Similarly, readthrough of the cyclic AMP phosphodiesterase *pde2* stop codon produces a destabilized protein variant, and this has been suggested to explain elevated cyclic AMP levels in *Psi+* yeast¹⁰. Differential stability by polymorphisms in the readthrough peptide of *sky1* has been postulated to explain *[Psi]*-induced strain differences in diamide sensitivity²¹. Particularly intriguing are very recent findings that stop codon mutations at the cFLIP-L locus confer protein instability for this anti-apoptotic factor in mice, leading to embryonic lethality¹². The same study also noted several hereditary human disease alleles where 3'UTR-encoded peptides are destabilizing, conferring marked decreases in protein activity and level (e.g.⁸).

Not every case of stop codon readthrough is destabilizing^{4–7} (Fig 4c), and some readthrough events are functional and regulated to defined levels (e.g.^{22–25}). Understanding the mechanisms by which some readthrough events are recognized and cleared (while others are not) may prove informative for biological contexts and pathological states where inappropriate readthrough occurs. We do not yet know the determinants of a translated 3'UTR sequence that confer loss of protein, though the ability of numerous sequences (including shuffled and randomized 3'UTR variants, Fig 2b, 4c) suggest a highly degenerate sequence is sufficient. Consistent with the idea that readthrough peptides' effects may be mediated via their biophysical characteristics, we observed a significant negative relationship between hydrophobicity and expression for GFP (K562 cells, *C. elegans*) and endogenous loci (*unc-22* and *unc-54*, *C. elegans*) (Extended Data 8, 9,10, Supplementary Information).

Destabilization by 3'UTR-encoded peptides could effectively screen against at least three types of events where a stop codon is inappropriately bypassed: (1) Stop codon misreading (e.g. by suppressor tRNAs). Suppressor tRNAs permit readthrough of a few to upwards of 30% of ribosomes at a stop codon (one of UAA, UAG, or UGA)¹³. While some suppressor tRNAs can be toxic, other cells tolerate even high levels of readthrough^{13,26–28}. Destabilization of readthrough products by C-terminal appendages may effectively buffer

cells from suppressor tRNA-induced proteostatic chaos. (2) A ribosomal frameshift in a coding region which is late enough that no premature termination codon is encountered. In this case, ribosomes would enter the 3'UTR out-of-frame with the coding region. In our manipulations, translation of 3'UTRs in multiple frames was detrimental to expression (Extended Data 5, data not shown), and similar amino acid and hydropathy biases hold for all three 3'UTR frames (Extended Data 9,10). (3) Aberrant RNA processing or ribosome dysfunction could produce a variety of other improperly terminated peptides from which destabilization would provide valuable relief.

Methods

C. elegans Strain Construction and Husbandry

C. elegans were grown at 23C on agar plates with nematode growth medium seeded with *E. coli* strain OP50 as described²⁹. Some strains were provided by the CGC, which is funded by NIH Office of Research Infrastructure Programs (P40 OD010440). A full list of strains used is available in Supplementary Table 1.

Transgenic array-containing strains were generated as follows: PD5102 (*pha-1(e2123ts)I*; *rde-1(ne300)V*) young adult hermaphrodites (grown at 16C) were injected with a mix of 90 ng/ul pC1 (containing a rescuing fragment of *pha-1*), 5 ng/ul of an mCherry-containing vector, and 5 ng/ul of a GFP-containing vector. Unless otherwise indicated, GFP was driven by the *myo-3* promoter to drive expression in the body wall muscle³⁰. Injectants were shifted to 23C to select for F1 progeny animals bearing a transgenic array (selecting for *pha-1(+)* expression³¹). The *rde-1* allele included in this strain avoided a modest degree of secondary siRNA-based silencing observed with many extrachromosomal transgenes³². For transgenic lines generating low levels of GFP, we considered the possibility that the GFP protein was toxic and selected against.

Under this model, one might expect (1) a subset of sick and/or dead GFP-positive F1 animals, (2) muscle defects due to muscle-specific expression of potentially-toxic GFP derivatives, (3) concomitant low levels of mCherry, and/or (4) a decrease in the efficiency with which transgenic lines were obtained³². None of these effects were observed, arguing against any contribution of negative selection to the observed low GFP expression.

For a subset of strains, we deviated from the above protocol to generate *pha-1* arrays as follows: (1) While most transgenic lines were generated from independently injected parents, a handful of strains were possibly generated from siblings of an injected parent (PD6480, 6481, 6482, 6483, 6484, 6485, 6486, 6493, 6494, 6495). In these cases, all injectants were pooled together on the same plate, and independent F1 were picked off to generate transgenic lines. Previous work has demonstrated independent F1 from the same injected parent carry distinct transgenic arrays^{32,33}. (2) During the course of our analyses, we found a handful of strains with an mCherry-negative subpopulation. The mCherry-positive subpopulation was isolated and propagated to generate the strains PD6401, 6450, 6452, 6456, 6457, and 6464.

CRISPR/Cas9 genome editing was performed in the VC2010 (PD1074) N2 background as described¹⁷. We selected *pha-4*^{34,35}, *unc-45*^{36,37}, *tra-2*³⁸⁻⁴⁰, *unc-22*²⁹, and *unc-54*^{41,42,38,26} based on the criteria in the text (citations indicated). The statement that *unc-22* and *unc-54* exhibit little-or-no autoregulation/feedback is based on a number of genetic experiments (with heterozygous²⁹, amber-suppressed²⁶, and/or *smg*-suppressed³⁸ alleles) which express either UNC-54 or UNC-22 at stable intermediate levels (between wild type and null). Alleles of *unc-45* were initially generated in the VC2010 background, though the embryonic lethality made *unc-45*(*TerByP*) difficult to maintain. We subsequently remade all alleles in a balanced heterozygote background (*sC1(s2023)* [*dpy-1(s2170)*] *III/+*) and considered non-Dpy segregants for phenotypic analyses.

Human Cell Line Construction

K562 cells (obtained from ATCC) were grown at a density of ~ $0.5-1\times10^{6}$ cells/ml in RPMI supplemented with penicillin/streptomycin, L-glutamine, and 10% FBS. All cell lines were maintained in a humidified incubator (37C, 5% CO2), and checked regularly for mycoplasma contamination. As means of validating K562 cells, we performed RNA-seq on a subset of lines and observed good correlation with published datasets⁴³ (data not shown). Viral particles were produced in HEK293T cells in 6 well dishes, and 1ml of viral supernatant was used to infect ~100,000 K562 cells by spin infection, 10³ rcf for 2 hours. Polybrene was omitted so as to keep the infection rate low (<10%), ensuring a single incorporation event for most cells. After three days of recovery, cells were selected with puromycin at 0.7 ug/ml for at least 3 days. Fluorescence was examined on a BD Accuri C6 flow cytometer, with appropriate gating for live cell events. For each construct examined via puromycin selection in K562 cells, similar eGFP and mCherry fluorescence levels were also observed in transient transfection in HEK293T cells in the absence of puromycin, arguing against a puromycin-selected skew in mCherry fluorescence.

Plasmids

Plasmids were constructed by restriction digest or Gibson cloning as detailed in Supplementary Table 2. pJA138/L3785 and pJA137/pCFJ104 were used as the basis of all *C. elegans* GFP or mCherry-containing vectors, respectively. Portions of pMCB306 and pMCB309 were used to construct pJA291, the parental puro::mCherry::T2A::eGFP::MCS::wPRE vector for experiments in human cells. Plasmids were confirmed by both sequencing and restriction digest, and plasmid concentrations determined with the QuBit dsDNA Broad Range kit (Invitrogen). A handful of plasmids that may be useful have been deposited with Addgene: pJA327 (*C. elegans* superfolder GFP in L3785), pJA291, pJA317 (pJA291 with d10DC insert) and pJA318 (pJA291 with d40DC insert).

GFP fusions were done with a GFP variant that corresponds to wild-type (*Aequora*) GFP with mutations at position 65 (Ser>Thr for human and Ser>Cys for *C. elegans*) known to improve folding and acquisition of fluorescence. Even with these mutations, GFP has a known propensity to misfold under some circumstances, so we examined the effect of a subset of the 3'UTR-encoded sequences (*hlh-1*, *daf-6*, and *unc-54*) downstream of a faster and more robust-folding GFP variant, superfolder GFP⁴⁴. The observed reduction in

superfolder GFP:mCherry ratios was quantitatively similar to that observed with normal GFP (Extended Data 3).

Sequences of FLAG⁴⁵, HA⁴⁶, d1ODC¹⁹, and d4ODC¹⁹ were obtained from the indicated publications. For exact sequences, see Supplementary Tables 1 and 2. T2A was previously shown to function in *C. elegans*¹⁴. Translation elongation through a member of the 2A peptide family (consensus D(V/I)EXNPGP) causes ribosomal pause, then release of the N-terminal peptide up to and including the Gly¹⁵. Translation elongation resumes, with the C-terminal peptide being produced with an N-terminal Pro.

Microscopy

Animals were immobilized by placing on a slide with a coverslip in 5mM EDTA, 50mM NaCl, 1mM levamisole and imaged on a Nikon Eclipse E6000 microscope using a Nikon super high pressure mercury lamp power supply. Filter cubes for fluorescence images were GFP (96342, Nikon Corp), mCherry (96321, Nikon Corp), and Broad (GFP+mCherry, 59022, Chroma Technology Corp). Images were collected with a 3CCD Digital Camera C7780 (Hamamatsu Corp) using HCImage (Version 1.0.2.060107, Hamamatsu Corp). Pictures of PD4251 and one of PD3363/3364 were taken for each imaging session and compared to ensure consistency between days.

For quantification of GFP to mCherry relative fluorescence, animals were imaged using a 4× objective with a broad filter and a 200 millisecond exposure. To avoid image over- or underexposure, a handful of exceptionally bright or dim strains were taken with a decreased or increased (respectively) exposure time (PD1798 40msec, PD3294 500msec, PD3299 50msec, PD3395 500msec, PD6327 50msec, PD6375 50msec, PD1786 50msec, PD1789 50msec, PD1790 50msec, PD6460 500msec, PD6469 50msec, PD6471 50msec, PD6472 50msec, PD6473 50msec, PD6485 100msec, PD1787 40msec, PD6450 100msec, PD6455 40msec, PD6477 40msec, PD6479 50msec, PD6498 40msec, PD6499 40msec, PD6500 40msec, PD6501 40msec, PD6502 40msec, PD6503 40msec, PD6504 40msec).

Raw pixel values for the red and green channels were obtained from image files using the tifffile package in python. Pixels below a threshold distance (200) from the median pixel intensity of the entire image were discarded as background. Pixels above a threshold intensity distance (4000 of a possible 4095) from the origin were discarded as saturated. The median pixel intensity for the entire image (essentially the black background, given the relatively low density of *C. elegans* tissue) was subtracted from the remaining pixels, and the slope of the linear regression line taken as the GFP/mCherry fluorescence ratio. This metric was robust to different exposure times and neutral density filters.

Statistics

Statistical tests and p-values are stated throughout the text and figures.

To test statistical significance of C-terminal appendage effects on the GFP/mCherry fluorescence ratio (Fig 2B), we divided the data into two groups: (1) 3'UTR-derived (*unc-54(TerByP), tbb-2(TerByP), daf-6(TerByP), hlh-1(TerByP), rps-20(TerByP), rps-30(TerByP), shuffle1–3*) and (2) non-3'UTR-derived (*rand1–3(A,C,G,T), 3xHA*,

3xFLAG, eef-1A.1(CDS63-83), bar-1(CDS452-492), daf-6(CDS756-782), mut-16(CDS89-101), alr-1(CDS93-126), hlh-1(CDS290-320,syn1)). For each construct, we took the average GFP/mCherry fluorescence ratio of all available lines. We compared the distribution of 3'UTR-derived and non-3'UTR-derived GFP/mCherry fluorescence ratio values by Kolmogorov-Smirnov test (p=0.004).

Ribosome Footprint

Profiling Ribo-seq was performed essentially as described^{18,47}, with a few modifications. Briefly, animals were grown to the ~L4 stage, and harvested by centrifugation and flash freezing in liquid nitrogen. Animals were ground with a mortar and pestle in liquid nitrogen, after which the powder was thawed in excess volume ice-cold polysome lysis buffer (20mM Tris pH 8.0, 140mM KCl, 1.5mM MgCl2, 1% Triton) with cycloheximide 100ug/ml). RNase 1 and sucrose gradient centrifugation was as before⁴⁷. ~2ug of purified, RNase 1digested monosomal RNA was run on a urea 15% polyacrylamide gel, and the entire region from ~15–30nt was excised for library preparation. At this point the protocol continued with T4 PNK (NEB) treatment as with the RNA-seq1 protocol next section).

RNA Sequencing

Two RNA sequencing (RNA-seq) procotols were used in this study. The first RNA-seq protocol (RNA-seq1) was performed on homozygote populations of animals (Extended Data 6). 5ug of total RNA was treated with the RiboZero kit (Illumina). RNA was fragmented at 95C for 30' by addition of an equal volume of 100mM sodium carbonate, 0.5mM EDTA pH 9.3 buffer. RNA fragments were gel purified, then treated with T4 PNK NEB). 3'Ligation with AF-JA-34.2 (/5rApp/NNNNNAGATCGGAAGAGCACACGTCT/3ddC/, Integrated DNA Technologies) and T4 RNA ligase 1 (NEB) was done at room temperature for 4 hours with 20% PEG8000 in 3.3mM DTT, 8.3mM glycerol, 50mM HEPES KOH (pH 8.3), 10mM MgCl2, 10ug/ml acetylated BSA. Unligated AF-JA-34.2 was removed by sequential treatment with 5'deadenylase (M0331S, NEB), then RecJ_f (M0264S, NEB). Reverse transcription was carried out with AF-JA-126 (/5Phos/AGATCGGAAGAGCGTCGTGT/ iSp18/CACTCA/iSp18/GTGACTGGAAGTTCAG ACGTGTGCTCTTCCGATCT, Integrated DNA Technologies) as a primer. Circular ligase treatment and PCR were as previously described⁴⁷.

A second RNA-seq protocol (RNA-seq2) was used to examine RNA levels with small numbers of heterozygote animals (Fig 3C). ~60 L2–L4 mixed gender animals were picked and flash frozen in 50mM NaCl, and RNA extracted with trizol. RNaseH and 94 oligos complementary to ribosomal RNA were used to deplete rRNA from the sample⁴⁸. Briefly, ~250ng of a cocktail of DNA oligos complementary to rRNA (Supplementary Table 3, ordered from Integrated DNA Technologies) was mixed with ~100ng total RNA in 125mM Tris pH7.4, 250mM NaCl in 8ul. The sample was heat denatured at 95C for 2', then cooled at -0.1C/sec to 45C. 1ul of digestion buffer was added (500mM Tris pH7.4, 1M NaCl, 200mM MgCl₂) with 1ul (5 units) Thermostable RNase H (Epicentre), and the sample was incubated at 45C for one hour. DNA oligos were removed by treatment with TURBO DNase (ThermoFisher) at 37C, and RNA was extracted using an equal volume of phenol/

chloroform. An RNA-seq library was prepared using the SMARTer Stranded RNA-Seq kit (Clontech Laboratories, Inc.).

Sequencing

Libraries were sequenced on a MiSeq Genome Analyzer (Illumina, Inc.). Reads were mapped to the *C. elegans* genome (Ensembl70, WBcel215) using STAR (v2.3.1⁴⁹), with the mutated bases of *unc-54(cc3389)* and *unc-54(e1301)* masked. For Ribo-seq and RNA-seq1, reads bearing the same last 6 nucleotides (from NNNNNN, added with AF-JA-34.2) were assumed to be PCR duplicates and collapsed to a single read. For RNA-seq2, multiple reads containing the same start and stop mapping positions were collapsed to a single read count to reduce effects of PCR bias. The removal of PCR duplicates with either protocol only affected ~5–10% of reads and did not adversely impact any of the analyses shown. RNA-seq1 and Ribo-seq were performed once for each strain shown in Extended Data 6,7.

Genomes and Annotations

While we sought to use the latest genome versions and annotations, we found it prudent to take advantage of the care and time with which other researchers annotated and analyzed earlier versions of genomes. For whole genome alignment of nematode species, *C. elegans* UCSC genome ce10/WS220 was used. To examine the length of predicted C-terminal extensions upon readthrough (Extended Data 1), genomes and annotations of each of the indicated species were as follows: *E. coli* Ensembl genome and annotations from assembly GCA_000967155.1.30, *S. cerevisiae* genome S288c (R57-1-1_20071212) and annotations⁵⁰, *C. elegans* UCSC genome (WS190/ce6) and annotations¹⁶, *H. sapiens* Ensembl genome release 83 and annotations from TargetScan v7.0⁵¹.

Immunobloting

Animals were boiled in $1 \times$ SDS loading buffer (65mM Tris pH 6.8, 10% glycerol, 2% SDS, 2mM PMSF, $1 \times$ Halt Protease Inhibitor (Thermo), 10% 2-mercaptoethanol) and run on a 7.5% Criterion TGX gel (Bio-Rad Laboratories, Inc.). Protein was transferred to a low background fluorescence PVDF membrane (Millipore). The membrane was blocked in 3% nonfat milk in $1 \times$ PBST with 250mM NaCl. The 5–6 antibody was used at a 1:5000 dilution to detect *myo-3*, and 5–8 antibody used at a 1:5000 dilution to detect *unc-54⁵²*. The 5–6 and 5–8 monoclonal antibodies were produced previously by purification of endogenous myosin proteins. Secondary antibody staining was performed with 1:500 Cy3-conjugated affiniPure goat anti-mouse (Jackson Immunoresearch). Imaging was done on a Typhoon Trio (Amersham Biosciences), and quantification done in ImageJ. For the lower blot of Fig 3D, lysates were made from multiple animals, and serial dilutions done to titrate the number of animals per lane.

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Number of Amino Acids Until Next In Frame Stop

Genes with at least one 3'UTR, one Count/Gene Genes with at least one 3'UTR, one Count/Txt Genes with one 3'UTR Nt Shufflings of Genes with one 3'UTR

Extended Data 1. Distribution of C-terminal extensions upon stop codon readthrough

Annotations and genomes were as described in Supplemental Methods. Each 3'UTR was translated starting one codon after the stop codon until the next in-frame stop codon. For metazoans, counting was done three different ways: including only genes for which exactly one 3'UTR was annotated (blue), counting each annotated 3'UTR separately (green), or counting each gene once and splitting gene counts with multiple 3'UTRs equally amongst

the 3'UTR isoforms (red). "Nonstop" indicates 3'UTRs for which no stop codon was encountered prior to the poly(A) tail. For each species the distribution of next in-frame stop codons was calculated for 1,000 nucleotide shufflings of 3'UTR sequences for genes with a single 3'UTR annotated, and 95% confidence interval shown (yellow). A similar "randomized" distribution was obtained upon shuffling 3'UTR sequences and preserving dinucleotide frequency. The frequency of stops immediately after the annotated stop codon (amino acid length 0) is highlighted with a blue arrow in each species. The distribution of peptide lengths follows an exponential decay curve, where the slope is related to the probability of encountering a stop codon at each position. In the simplest model, the probability of encountering a stop codon is constant throughout the 3'UTR, accounting for the roughly linear shape of each plot (previously noted^{2,3}). Notable exceptions are a tendency towards second in-frame stops in *E. coli* (blue arrow), and a tendency towards peptides >60AAs in length in all species. In *E. coli* the enrichment towards longer downstream peptides is at least partially explained by the operonic layout of genes.





Images were taken under a broad excitation/emission filter to allow for simultaneous capture of GFP and mCherry fluorescence. Intensities of each pixel in the red and green channels were extracted in python. Unfiltered pixel intensities are shown as black dots. Pixels were filtered, background subtracted, and linear regression performed (red dots and line, see Methods). For simplicity, the green/red intensities from 1000 random pixels are shown. The GFP/mCherry fluorescence ratio was taken as the slope of the linear regression line.



Extended Data 3. Readthrough regions confer a loss of superfolder GFP fluorescence Each of the indicated TerByP regions were inserted downstream of sfGFP, upstream of the *let-858* 3'UTR. TerByP is the region after the annotated stop codon, up to and including the first in-frame stop codon in the 3'UTR. Quantification was done as described (Extended Data 2).



Extended Data 4. Explanation of "shuffle" sequences

Trinucleotide codons from each TerByP region are color coded by gene (top). Codons were extracted and randomly shuffled in python. A codon was iteratively selected until a stop codon was encountered, defining *shuffle1*. The process was repeated twice more to define *shuffle2* and *shuffle3*. The resulting *shuffle* peptides are a combination of all three TerByP regions. Lengths and color coding of codons for *shuffle1–3* accurately reflect the sequences they are derived from.

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Gene	Genotype	Phenotype	Isolates
pha-4			
	pha-4(TerByP)	wild type	2
unc-22	· · · · · · · · · · · · · · · · · · ·		
	unc-22(TerByP)	wild type	2
	unc-22(-1,TerByP)	twitcher	1
	unc-22(+1,TerByP)	twitcher	1
	unc-22(unc-22::GFP)	wild type	3
unc-45	; ;		
	unc-45(TerByP)	emb lethal	3
	unc-45(3xFLAG::TEV::3xHA)	wild type	2
tra-2			
	tra-2(TerByP)	XX males	2
	tra-2(3xFLAG)	wild type	2
	tra-2(3xFLAG::TEV::3xHA)	XX males	2
unc-54	!		
	unc-54(TerByP)	paralyzed	2
	unc-54(unc-54∷gfp)	wild type	2
	unc-54(+1,TerByP)	paralyzed	1
	unc-54(-1,TerByP)	weak Unc	1

Extended Data 5. Translation into 3'UTRs at endogenous loci tends to yield hypomorphs

CRISPR/Cas9 editing¹⁷ was used to construct the mutations shown. See Supplementary Table 1 for precise nucleotide sequences of all strains. -1/+1 TerByP indicate the loss or gain of one nucleotide relative to the zero frame, generating a frameshift over the stop codon, and translation into the 3'UTR out-of-frame with the coding sequence. For *unc-22*, "wild type" indicates a lack of twitching, even in 1mM levamisole. For *unc-54(R̂1,TerByP)*, "weak Unc" animals were visibly slower than *unc-54(+)*, but faster than *unc-54(TerByP)*. All mutant phenotypes were recessive.



Extended Data 6. RNA-seq and Ribo-seq from unc-54 mutants

(a) RNA-seq and (b) ribosome footprint profiling (Ribo-seq) library mRNA counts, with summary counts (c) for the indicated strains and mRNAs. Libraries were prepared from L4 animals as described (Methods). "N2" is wild type (PD1074, VC2010⁵⁵). *unc-54(cc3389)* bears a TAA(Stop)>AAT(Asn) mutation, *unc-54(TerByP)*. *unc-54(e1301)* bears GGA(Gly387)>AGA(Arg387), a point mutation that confers a temperature-sensitive Unc phenotype with minimal discernable effects on UNC-54 protein levels. *unc-54(e1301)* was included as a control for the Unc phenotype of *unc-54(cc3389)*, though *e1301* confers a less

severe Unc phenotype than *cc3389*. Values for *unc-54* mRNA (blue) are highlighted throughout, and for comparison, three additional transcripts known to be at least partly expressed in the body wall muscles are also highlighted: *unc-87* (pink), *unc-15* (green), and *unc-22* (red).



Extended Data 7. Ribo-seq of unc-54(cc3389) shows an unexceptional progression of ribosomes in the readthrough region

a. Raw Ribo-seq reads for unc-54(+) (blue) and unc-54(cc3389) (green) animals, plotted as read pile-ups. Mismatched bases are indicated with black bars. Location of the normal stop

codon and the first in-frame stop codon are indicated with "TAA" and dotted lines. The extension in *unc-54(cc3389)* is 30 amino acids.

b. Number of Ribo-seq reads in the last 30 codons, compared to the previous 30 codons, for all mRNAs. Linear regression was performed on all points (solid line), and two-fold difference shown (dashed lines).

c. The distribution of Ribo-seq reads in the last 30 codons (90nts) of *unc-54(cc3389)* is shown in green, and the 95% confidence interval for all open reading frames in dashed lines. d. The fraction of in-frame Ribo-seq reads in the last 30 codons is plotted as a function of read counts in the last 30 codons, and *unc-54(cc3389)* highlighted.

e. The distribution of read lengths in the last 30 codons of *unc-54(cc3389)*, and all open reading frames (95% confidence interval, dashed lines).

For b–d, reads were restricted to 28,29,30 nt lengths. For b–e, a 12 nt offset was done for the ribosomal P-site, and read counts were derived solely from the *unc-54(cc3389)* Ribo-seq library. For c and e, a minimum 15 read counts was imposed to obtain the 95% CI from "all genes".

Extended Data 8. Lack of general conservation of coding potential downstream of stop codons in *Caenorhabditis*

Whole genome alignment of six nematode species with *C. elegans* genome assembly ce10/WS220 was obtained from the UCSC genome browser. For each annotated transcript, the aligned bases from the multiple species alignment were extracted and compared to the

reference (*C. elegans*) genome. The left plot shows summary information of the alignment centered around annotated stop codons; the right plot shows the same centered around the first in-frame stop codon in 3'UTRs. In red is the substitution frequency, i.e. the number of mismatched bases divided by the number of aligned bases at a given position. The enrichment of "wobble" position mutations is apparent as an increase in substitutions at the third position of each codon in the CDS. In green is the synonymous substitution frequency, i.e. for codons beginning at a given position, the fraction of mutations that yield a synonmous substitution divided by all mutations at that position (synonymous+non-synonymous). The tendency to conserve amino acids in the CDS is apparent as a green spike at every in-frame codon. The change in substitution frequency and synonymous substitution frequency about the first in-frame stop codon (right plot) is due to a tendency for NTR codons to be conserved, and for AAN/AGN/GAN codons to not be conserved in 3'UTRs, regardless of frame.

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е

Gene	Genotype	Phenotype	Avg Kyte-Doolitte	Max Kyte-Doolitte (6AA window)		
unc-22						
	unc-22(TerByP)	wild type	0.370	1.950		
	unc-22(-1,TerByP)	twitcher	1.081	3.283		
	unc-22(+1,TerByP)	twitcher	1.685	1.333		
unc-54						
	unc-54(TerByP)	paralyzed	0.928	2.866		
	unc-54(+1,TerByP)	paralyzed	1.391	3.150		
	unc-54(-1,TerByP)	weak Unc	-0.088	1.550		

Extended Data 9. Nucleotide and amino acid composition of readthrough regions (*C. elegans*) Coding sequences (CDS) and 3'UTRs were analyzed for various sequence properties. For simplicity, only genes and 3'UTRs for which a single 3'UTR was annotated were considered. Similar results were obtained with genes with multiple 3'UTRs.

a. Nucleotide frequency of CDS, 3'UTR, and TerByP (region between annotated stop codon and first in-frame stop codon).

b. Frequency of amino acids in all three possible frames for the TerByP region. 3'UTRs were translated one codon past the stop codon of the CDS until the next in-frame stop codon, with

nonstop 3'UTRs ignored. Highlighted are codons with high G content (GGN, Gly) and high T content (TTY, Phe).

c. TerByP regions tend to be hydrophobic, regardless of frame. Kyte-Doolittle score was used as a measure of hydrophobicity⁵⁶. To reduce noise, only TerByP regions at least 10 amino acids long were considered. P-value is for Kolmogorov-Smirnov test comparing CDSs and TerByP sequences (each frame has p-value<10e-293 for this comparison). As the TerByP sequences are shorter than CDSs on average, the distribution of TerByP hydrophobicity scores will tend to have higher variance than CDSs. Random portions of CDSs were taken, length-matched to TerByP frame zero peptide lengths. This was repeated 100 times, and the 95% confidence interval is shown (dashed lines, "CDS rands"). d. Hydrophobicity of the inserts is correlated with a negative effect on GFP fluorescence. The GFP/mCherry fluorescence ratio (Fig 2B) was plotted against the maximum Kyte-Doolittle score in a six amino acid window for each insert. (Similar results were obtained using the Kyte-Doolittle score averaged across the entire sequence.) Mean (circle) and standard deviation (bars) are shown. 3'UTR-derived sequences are in blue, and non-3'UTRderived sequences are in red. So as to avoid redundancy or skewing of the data, in cases where multiple constructs were present with the same peptide sequence (e.g. unc-54(TerByP), unc-54(TerByP,syn1), and unc-54(TerByP,syn2)), only the first of these was used.

e. Hydrophobicity analysis of the TerByP extensions obtained by CRISPR/Cas9 engineering at the *unc-22* and *unc-54* loci. +1/-1 TerByP indicates the gain or loss of a nucleotide, generating a late frameshift and allowing translation to proceed past the annotated stop codon out-of-frame with the upstream ORF. In each case, Kyte-Doolittle hydropathy was used to analyze the C-terminal appendage. In bold is the phenotypically least affected strain of the three.

Extended Data 10. Nucleotide and amino acid composition of readthrough regions (*H. sapiens*) Similar analysis of hydrophobicity as in Extended Data 9c,d performed in humans.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. Translation into 3'UTRs results in substantial loss of protein expression

a. Dual fluorescence reporter assay to test expression with different 3'UTRs. Transgenic arrays of each GFP construct were created using *pha-1* selection and mCherry (pCFJ104) as a coinjection marker. Broad Filter detects GFP and mCherry signals simultaneously; deviation from yellow towards red or green shows more mCherry or GFP fluorescence, respectively. Three independent transgenic lines were made for each (two for *tbb-2(TerByP)*); transgenic lines with similar mCherry expression are shown. 200 millisecond exposure, $10 \times$ objective.

b. Dual fluorescence reporter assay to test expression of readthrough for different 3'UTRs. The stop codon of each 3'UTR was mutated, allowing translation to proceed into the 3'UTR (Termination ByPass, TerByP). Images were collected as in A. "GFP ($10\times$)" is a 2 second exposure. The dim yellowish fluorescence in "GFP ($10\times$)" for *unc-54(TerByP)* and *tbb-2(TerByP)* is autofluorescence.

c. For each gene, the 3'UTR was fused to mCherry and GFP. GFP expression was tested with the stop codon mutated to a sense codon (*TerByP*). For each of *eef-1A.1*, *rps-17*, *daf-6*, and *hlh-1*, GFP expression was also tested with the normal stop codon in place (*3'UTR*). The ratio of GFP to mCherry fluorescence under a broad fluorescence filter was used as a metric (Extended Data 2, Methods). Each triangle represents an independently-generated transgenic line; mean and standard deviation of n lines shown. Student's t-test two-tailed p-value.

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Figure 2. Identification of determinants for product loss upon translation into the 3'UTR a. Shortening or non-synonymous mutations of the readthrough region can restore GFP expression. Stop codons and/or mutations were inserted into each GFP::3'UTR fusion as diagrammed with stop codons (red stop sign) and poly(A) site (blue arrowhead). [%]Percent indicates same constructs shown in Fig 1. mCherry (pCFJ104) was used as a coinjection marker. "+X AA" indicates amino acids added relative to cognate control ("+0 AA") construct. Constructs and mutated regions drawn to scale, scale bar at top. Mean and standard deviation of n lines shown.

b. 3'UTR-encoded peptides are sufficient to confer GFP loss. Sequences were inserted upstream of the *let-858* 3'UTR. TerByP is the region between the canonical termination codon and first in-frame termination codon in the 3'UTR. "syn" are synonymously-substituted variants. Shuffle1–3 contain shuffled codons of *unc-54*, *tbb-2*, and *rpl-14(VLFL>RSCA)* TerByP regions (Extended Data 4). T2A is a "self-cleaving" peptide which releases the upstream nascent chain; T2A* is a non-cleaving variant^{14,15}. Rand1– 3(A,C,G,T) are random combinations of A, C, G, and T created *in silico*. Each CDSN-M is an arbitrary fragment of the respective gene's Coding DNA Sequence (from amino acid N to M).

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b. Brightfield images of *unc-54* alleles. Arrowhead indicates a "bag of worms", the shell of an egg laying-defective mother consumed by its retained progeny.

c. RNA-seq from *unc-54(TerByP/+)* heterozygotes showed no differential effect on RNA levels. *unc-54(TerByP/+)* heterozygotes were chosen among progeny of *unc-54(+)* males crossed with *unc-54(TerByP)* homozygotes and allele-specific reads identified. Framed inset shows individual allele-specific RNA-seq reads (bars) from *unc-54(TerByP)* (AAT, green) and *unc-54(+)* (TAA, blue). See also Extended Data 6, 7.

d. Quantification of UNC-54 protein levels. Immunoblotting was performed on homozygous populations of the indicated animals. *unc-54(r293)* encodes a nonsensemediated decay allele of *unc-54*, producing <5% of normal UNC-54 protein. *unc-54(r259)* contains a >17kb deletion spanning most of the *unc-54* locus. For the lower blot, the number of animals loaded per lane is indicated. For gel source data, see Supplementary Figure 1.

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Figure 4. Translation into 3'UTRs results in protein loss for several genes in humans a. Lentiviral reporter schematic. A puroR-mCherry fusion was co-translationally cleaved from eGFP-insert by T2A. Constructs in B-D were integrated into K562 cells via lentiviral infection and puromycin selection.

b. Validation of dual fluorescence reporter. Inserts downstream of eGFP were 3xFLAG, 3xHA, and degrons d4ODC ($t_{1/2}$ ~4hr¹⁹), d1ODC ($t_{1/2}$ ~1hr¹⁹).

c. The sequence between the annotated and first in-frame termination codon (TerByP) from each gene was inserted downstream of eGFP (solid line). For comparison, nucleotides of each TerByP region were randomized, producing a length- and nucleotide frequency-matched construct (randTerByP, dashed line). Cells with eGFP lacking an insert and grown a week apart (top, green solid lines) and approx. fluorescence ratio of d4ODC (orange line) are shown.

d. The first 30 amino acids of the *HBA2*3'UTR were inserted downstream of eGFP (orange). Insertion of a self-cleaving T2A peptide restored expression (blue), an uncleavable mutant (T2A*) did not (light blue).