

1 **AlphaFold2-Guided Functional Screens Reveal a Conserved Antioxidant Protein**
2 **at ER Membranes**

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24 **Abstract**

25

26 Oxidative protein folding in the endoplasmic reticulum (ER) is essential for all eukaryotic
27 cells yet generates hydrogen peroxide (H_2O_2), a reactive oxygen species (ROS). The
28 ER-transmembrane protein that provides reducing equivalents to ER and guards the
29 cytosol for antioxidant defense remains unidentified. Here we combine AlphaFold2-
30 based and functional reporter screens in *C. elegans* to discover a previously
31 uncharacterized and evolutionarily conserved protein ERGU-1 that fulfills these roles.
32 Deleting *C. elegans* ERGU-1 causes excessive H_2O_2 and transcriptional gene up-
33 regulation through SKN-1, homolog of mammalian antioxidant master regulator NRF2.
34 ERGU-1 deficiency also impairs organismal reproduction and behavioral responses to
35 H_2O_2 . Both *C. elegans* and human ERGU-1 proteins localize to ER membranes and
36 form network reticulum structures. Human and *Drosophila* homologs of ERGU-1 can
37 rescue *C. elegans* mutant phenotypes, demonstrating evolutionarily ancient and
38 conserved functions. In addition, purified ERGU-1 and human homolog TMEM161B
39 exhibit redox-modulated oligomeric states. Together, our results reveal an ER-
40 membrane-specific protein machinery for peroxide detoxification and suggest a
41 previously unknown and conserved mechanisms for antioxidant defense in animal cells.

42 **Introduction**

43

44 In eukaryotic cells, the endoplasmic reticulum (ER) plays a critical role in protein folding,
45 a process essential for cell function but one that generates the reactive oxygen species
46 hydrogen peroxide (H_2O_2)¹⁻³. The ER-residing oxidase Ero1α stoichiometrically
47 produces one molecule of H_2O_2 per disulfide bond introduced to nascent secreted or
48 membrane proteins. Accordingly, the H_2O_2 concentration in the ER lumen is estimated
49 to be approximately 700 nM, whereas cytosolic H_2O_2 concentration is estimated to be
50 2.2 nM at steady levels⁴⁻⁶. Low cytosolic H_2O_2 levels are maintained by various H_2O_2 -
51 buffering systems and enzymes, including H_2O_2 -degrading catalases. Known families of
52 catalase localize to peroxisomes, cytosol and mitochondria, but not ER⁷⁻¹⁰. While H_2O_2
53 can cross membranes by slow diffusion or aquaporin-facilitated transport¹¹, it remains
54 unknown how ER membranes are guarded against the exceptionally high levels of ER
55 H_2O_2 , whose derivatives (e.g. hydroxyl radicals via Fenton reactions) can chemically
56 attack and damage membrane lipids, nucleic acids and proteins.

57

58 In bacteria, a family of transmembrane proteins (DsbD/ScsB)¹² provide reducing
59 equivalents for correct disulfide bond formation and peroxide reduction in the periplasm,
60 which is considered the bacterial equivalent of eukaryotic ER. DsbD and ScsB carry
61 multi-transmembrane segments and employ a series of cysteine pairs with differential
62 redox potentials to relay electron transfer from cytosol to periplasm¹³. Mammalian
63 counterparts of DsbD and ScsB are proposed to exist yet remain unidentified¹⁴. Our
64 study addresses this gap in knowledge by identifying a previously uncharacterized and

65 evolutionarily conserved *C. elegans* protein Y87G2A.13 (named as ERGU-1, ER
66 guardian of oxidative stress) as the long-sought ER membrane-localized antioxidant
67 protein. ERGU-1 protects against elevated H₂O₂ and ensures optimal organismal
68 functions in *C. elegans*. Furthermore, we reveal the structural and functional
69 conservation of ERGU-1 homologs across various animal species, indicating an ancient
70 and fundamental role for ERGU-1 in maintaining cellular redox homeostasis.

71

72 **Results**

73

74 **Computational and genetic discovery of ERGU-1**

75 We first performed BLASTP search to identify potential eukaryotic sequence homologs
76 of DsbD and ScsB. Such amino acid sequence similarity-based queries yielded no
77 apparent broadly conserved eukaryotic homologs, even after adjusting sensitivity and
78 specificity of the search (Fig. S1). We reasoned that DsbD/ScsB and potential
79 eukaryotic counterparts might not share primary protein sequence similarities owing to
80 distant evolutionary divergence. Nevertheless, eukaryotic counterparts of DsbD/ScsB
81 could exhibit structural and functional, rather than protein sequence features, common
82 for DsbD/ScsB and known families of transmembrane oxidoreductases. These features
83 include multi-transmembrane segments to facilitate intramembrane electron transfer,
84 closely spaced cysteine clusters in different transmembrane segments within proximity
85 (5-10 Angstroms), and broad evolutionary conservation in eukaryotic or multicellular
86 organisms. Based on these features, we sought to leverage the availability of predicted
87 protein structures by AlphaFold2 for nearly the entire *C. elegans* proteome¹⁵ and filter

88 through DsbD/ScsB-like candidates for subsequent functional screens and validation in
89 *C. elegans*.

90

91 To identify DsbD/ScsB-like candidates, we conducted a computational search of the *C.*
92 *elegans* proteome based on the functional features common for DsbD/ScsB-like
93 transmembrane oxidoreductases. Using the *C. elegans* UniProt reference proteome as
94 a starting point (version 26, 19,827 proteins), we found all genes with at least 4
95 annotated transmembrane helices (3,177 proteins) and further filtered to those with at
96 least two transmembrane helices, each containing at least two cysteines separated by
97 at most 2 intervening residues (sequence patterns CC, CXC, or CXXC). For the 190
98 proteins meeting these criteria, we examined AlphaFold2 Database-predicted structures
99 and selected those in which cysteine pairs in two helices were no more than 10
100 Angstroms apart (SG to SG atom distance), producing 53 candidate genes (Fig. 1A and
101 Fig. S1). We implemented the computational screen using customized Python scripts
102 (Supplementary Data file) in UCSF ChimeraX¹⁶.

103

104 Among the 53 candidates with AlphaFold2-predicted structures passing the above
105 screening criteria, we focused on the non-GPCR category with 11 hits for functional
106 validation in *C. elegans* (Fig. 1A, B). We performed RNA interference (RNAi) against
107 genes encoding these 11 hits and tested if any can activate the transcriptional *gst-*
108 *4p::GFP* reporter, which has been previously used to monitor excess oxidative stress
109 and elevated levels of peroxide in *C. elegans*¹⁷. We found that RNAi against one gene
110 *Y87G2A.13* strongly activated *gst-4p::GFP* under standard laboratory conditions without

111 exogenous oxidative stress, while RNAi against other genes or empty vector control did
112 not (Fig. 1B). Based on the T777T vector backbone with improved RNAi specificity and
113 efficacy¹⁸, we designed two independent RNAi constructs targeting different coding
114 regions of Y87G2A.13 and obtained similar results (Fig. 1C, D). In addition, we used
115 CRISPR-mediated genetic deletion to generate a null allele of Y87G2A.13 and
116 observed similar constitutive *gst-4p::GFP* activation in Y87G2A.13 null mutants (Fig.
117 1C, E). We also observed similar *gst-4p::GFP* activation in *ergu-1(gk840471)* mutants
118 with a protein-truncating mutation (Fig 1C, Fig. S2A, B). The transcription factor SKN-1
119 is the *C. elegans* ortholog of NRF2, master regulator of antioxidant responses in
120 mammals, and mediates *gst-4p::GFP* activation upon a variety of oxidative stresses^{19–}
121 ²¹. We found that RNAi against *skn-1* abolished the constitutive *gst-4p::GFP* activation
122 in Y87G2A.13 null and protein-truncating mutants (Fig. 1F, Fig. S2C). N-acetyl-cysteine
123 (NAC), a commonly used precursor for glutathione-based antioxidants and ROS/H₂O₂
124 scavenger in *C. elegans*^{22–24}, also strongly suppressed *gst-4p::GFP* activation in
125 Y87G2A.13 null mutants (Fig. 1G). Taken together, these results show that reduced
126 expression by RNAi or genetic deficiency of Y87G2A.13, one of the computationally
127 identified ERGU candidates, constitutively activates oxidative stress response in a
128 manner that involves excess oxidants and activation of SKN-1. We hereafter refer to the
129 Y87G2A.13 protein as ERGU-1 (ER guardian of redox defense), given additional lines
130 of evidence below.

131

132 **Molecular and organismal roles of ERGU-1 in antioxidant defense in *C. elegans***

133

134 As intramembrane oxidoreductases, DsbD and ScsB transfer electrons via cysteine
135 pairs as redox couples to reduce excess protein disulfide bonds and peroxides in
136 bacterial periplasm¹². We thus assessed similar roles of ERGU-1 in ER protein-folding
137 stress and H₂O₂ reduction, using the *hsp-4p::GFP* transcriptional reporter and a specific
138 fluorescent sensor roGFP2-Orp1 for H₂O₂, respectively^{17,25}. We found that *ergu-1*
139 genetic deletion or RNAi caused constitutive activation of *hsp-4p::GFP* in the absence
140 of exogenous ER or protein folding stresses (Fig. 2A, Fig. S3A). Using *C. elegans*
141 strains expressing roGFP2-Orp1 to specifically monitor cytosolic levels of H₂O₂ through
142 ratiometric dual-color fluorescence (Fig. 2B), we found that *ergu-1* genetic deletion or
143 RNAi caused markedly higher ratios of sensor oxidation versus reduction, indicating
144 elevated H₂O₂ levels. An independent assay based on non-fluorescent compound
145 dichlorofluorescein (DCFH) that reacts with H₂O₂ to produce fluorescent 2',7'-
146 dichlorofluorescein (DCF) showed similarly higher levels of peroxide stress in *ergu-1*
147 mutants (Fig. 2C). Furthermore, we assayed effects of *ergu-1* null, protein-truncating
148 mutants or RNAi on a comprehensive panel of stress-responding reporters, cell type-
149 specific markers, and organelle membrane lipid sensors²⁶⁻²⁸, but did not observe
150 apparent phenotypic consequences, except oxidative stress-related reporters *gst-*
151 *4p::GFP*, *hsp-4p::GFP*, roGFP2-Orp1 and Grx1-roGFP2 (Fig. S2D, 3A-3M). These
152 results suggest ERGU-1 normally defends against specific peroxide-related oxidative
153 stress, consistent with its structural features and functional reporter regulation observed.
154
155 The AlphaFold2-predicted structure of ERGU-1 reveals two clusters of cysteine pairs in
156 its two close transmembrane segments (Fig. 1B, 3D). To examine the importance of

157 such cysteine pairs for the antioxidant function of ERGU-1, we obtained CRISPR-
158 mediated knock-in alleles converting cysteine to alanine at each site, and crossed such
159 cysteine mutants separately with reporter strains of *gst-4p::GFP*. We found that these
160 cysteine mutants caused elevated levels of *gst-4p::GFP* (Fig. 2E, F). In addition, we
161 found that the *ergu-1* null mutant phenotypes were rescued by transgenic
162 extrachromosomal arrays of wild-type but not cysteine mutants of *ergu-1* (Fig. S4A).
163 These results suggest that the clusters of cysteine pairs of ERGU-1 functionally
164 contribute to defending against oxidative stress, supporting their roles in relaying
165 electron transfer across ER membranes to reduce peroxide stress and excess H₂O₂
166 from ER.

167
168 We next determined organismal roles of ERGU-1. Morphologically, the null and cysteine
169 knock-in *ergu-1* mutants appear grossly normal. Previous studies have implicated roles
170 of SKN-1 activation in several long-lived genetic mutants or by oxidative stress-induced
171 hormesis in promoting longevity¹⁹. Unexpectedly, we found that the *ergu-1* null mutants,
172 with control or *skn-1* RNAi, exhibited largely normal lifespans (Fig. S2H) under standard
173 laboratory conditions at 20 °C fed with OP50 on FuDR-NGM plates. To test genetic
174 interaction of *ergu-1* with other known genes implicated in ER-specific redox regulation,
175 we examined effects of RNAi against genes in the *ero-1*, *gcs-1*, *gsr-1*, *gpx* and *pdi*
176 families and found that *ergu-1* mutants showed increased oxidative stress responses
177 compared with wild type upon RNAi treatment (Fig. 2G). Notably, assay quantification of
178 reproductive capacity revealed a markedly reduced brood size of *ergu-1* null mutants.
179 Consistently, the *ergu-1* cysteine mutants exhibited similar reduction in brood sizes (Fig.

180 2H). This reduced brood size phenotype can be partially alleviated upon 10 mM NAC
181 antioxidant treatment (Fig. 2H). By contrast, treatment with exogenous oxidative
182 stressor paraquat decreased brood size more severely in *ergu-1* mutants than wild type
183 (Fig. S3I). In addition, we measured the locomotion speed change of young adult
184 hermaphrodites upon exogenous H₂O₂ and found that *ergu-1* null mutants exhibited a
185 markedly occluded locomotion-slowing response to H₂O₂, without apparently affecting
186 neuronal development, cytoskeletal structure or baseline neuronal activity (Fig. S3J-
187 S3N). These results indicate that ERGU-1 is critical for maintaining normal specific
188 organismal functions in reproduction and behavioral responses to H₂O₂.

189

190 **Subcellular localization of *C. elegans* ERGU-1 at ER membrane**

191

192 To elucidate the tissue distribution and subcellular localization of ERGU-1, we
193 constructed both a transcriptional (*ergu-1p::GFP*, fusing the *ergu-1* promoter to the
194 green fluorescent protein GFP) and a translational reporter (*ergu-1p::ergu-1::GFP*,
195 fusing GFP to the C-terminus of ERGU-1 under the control of its native promoter) (Fig.
196 3A). We microinjected the constructs and integrated the transgenic extrachromosomal
197 arrays to the genome at low copy number to ensure the faithful recapitulation of ERGU-
198 1's endogenous expression pattern. Both the transcriptional and translational reporters
199 similarly revealed *ergu-1* expression in major metabolic tissues, including the intestine,
200 body wall muscles, and the spermatheca (Fig. 3B, C). As expected, RNAi against *ergu-*
201 *1* abolished GFP signals from the *ergu-1p::ergu-1::GFP* transgene (Fig. S3B). Notably,
202 the tissues expressing *ergu-1* are also known to exhibit high metabolic activities,

203 generating elevated levels of H₂O₂ during protein folding, as indicated by roGFP2-Orp1
204 (Fig. 2B).

205

206 To ascertain ERGU-1's subcellular localization, we crossed the translational reporter
207 with a previously well-characterized ER membrane marker SEL-1(1-
208 79)::mCherry::HDEL^{28,29}. High-resolution dual-fluorescence confocal microscopy and
209 line-scanning analysis revealed marked co-localization of *ergu-1p::ergu-1::GFP* and
210 SEL-1(1-79)::mCherry::HDEL in a net structure-like reticulum pattern (Fig. 3D-3I, Fig.
211 S5A). ERGU-1::GFP showed particularly prominent intensities in anterior and posterior
212 intestinal cells, consistent with effects we observed with roGFP2-Orp1 (Fig. 2B). We
213 confirmed the ER membrane-specific localization of ERGU-1 by imaging ERGU-1::GFP
214 alongside RFP markers specific for other organelles, including the mitochondria,
215 lysosomes, and plasma membrane (Fig. S5B). The ER membrane localization aligns
216 markedly well with ERGU-1's proposed functional role in mitigating the detrimental
217 effects of H₂O₂ produced during protein folding within the ER lumen. To address
218 potential ER membrane-localized changes in oxidative stresses, we generated ER-
219 proximal redox sensor strain GRX1-roGFP-SPSC-1. We observed markedly higher
220 oxidation to reduction ratio of roGFP upon *ergu-1* and *nduf-7* (positive control) RNAi
221 treatment (Fig. 3J, 3K). Furthermore, biochemical purification of *ergu-1p::ergu-1::GFP*
222 using GFP-trap affinity chromatography revealed a redox-sensitive oligomeric
223 organization of ERGU-1 (Fig. 4F), with monomers being prevalent under reducing
224 conditions. The ER membrane-specific localization, redox regulation and antioxidant
225 function support ERGU-1 as a guardian against peroxide stress originating from ER.

226

227 **Evolutionarily conserved features of ERGU-1 family proteins**

228

229 We next investigated the evolutionary conservation of ERGU-1. First, we constructed a
230 multiple sequence alignment and a maximum likelihood-based phylogenetic tree
231 encompassing ERGU-1 and its homologs from diverse animal species (Fig. 4A). This
232 analysis revealed a high degree of sequence conservation across the ERGU-1 protein
233 family, suggesting a potentially ancient and functionally important role. For instance, the
234 mouse ERGU-1 homolog *Tmem161b* and human *TMEM161B* exhibit 93.6% amino acid
235 percent identities, while *C. elegans* ERGU-1 and human *TMEM161B* exhibit 27.9%
236 amino acid percent identities. To experimentally test evolutionary conservation of the
237 ERGU-1 protein family, we sought to functionally complement the *C. elegans* *ergu-1*
238 mutant phenotype with homologs from other species. We expressed the *ergu-1*
239 homologs *emei* and *TMEM161B* from *Drosophila* and humans, respectively, under the
240 control of endogenous *C. elegans* *ergu-1* promoter, in the *ergu-1* mutant background
241 carrying the *gst-4p::GFP* reporter (Fig. 4B). Remarkably, we found that transgenic
242 expression of these homologs rescued the constitutively activated *gst-4p::GFP*
243 phenotypes, indicating normalized oxidative stress in *C. elegans* *ergu-1* mutants (Fig.
244 4C). This successful complementation across species supports the functional
245 conservation of ERGU-1 and its homologs within the ER-GUARD system.

246

247 To additionally explore the functional conservation of ERGU-1 homologs, we focused on
248 *TMEM161B*, which appears more closely related to ERGU-1 than its paralog

249 TMEM161A (Fig. 4A). To mirror our *C. elegans* studies, we first examined TMEM161B's
250 subcellular localization in mammalian cells. We expressed V5-tagged TMEM161B
251 cDNA under the control of a CMV promoter in HEK293T and U2OS human
252 osteosarcoma cells. Immunofluorescence staining using an anti-V5 antibody revealed a
253 striking colocalization with established ER organelle markers (Fig. 4D, Fig. S6F).
254 Furthermore, to assess the potential antioxidant function of TMEM161B as *C. elegans*
255 ERGU-1, we employed V5-tagged TMEM161B cDNA expressed in HEK293 human
256 embryonic kidney cells. With the H₂O₂ sensor roGFP2-Orp1, we found that cDNA
257 expression of human TMEM161B led to strong suppression of exogenous H₂O₂-induced
258 oxidation of roGFP2-Orp1 (Fig. 4E, Fig. S6G). We also used the latest AlphaFold3
259 webserver³⁰ to model various ligand interaction with TMEM161B and ERGU-1, and
260 identified heme-binding pockets coordinated by a highly conserved tyrosine residue
261 (Y454) (Fig. S4B). We experimentally confirmed that biochemically purified TMEM161B,
262 but not Y454 mutants, bound to heme directly, while functionally the Y454 mutant
263 ERGU-1 or TMEM161B failed to rescue *ergu-1* null mutants (Fig. S4C). Beyond the
264 redox-sensitive oligomeric organization of ERGU-1 (Fig. 4F), purified human
265 TMEM161B exhibited native oligomerization under normal conditions and dissociation
266 into monomers in a reducing environment, as demonstrated by size exclusion
267 chromatography (Fig. 4G). Interestingly, cryo-electron microscopy analysis revealed a
268 filamentary oligomeric form of TMEM161B under non-reducing conditions, while
269 treatment with DTT induced monomerization (Fig. 4H). These findings indicate that
270 human TMEM161B recapitulates several key features of *C. elegans* ERGU-1, including
271 ER membrane localization, antioxidant functions and redox-modulated oligomeric

272 states, providing further evidence for the evolutionary conservation of ERGU-1 in
273 antioxidant defense across the animal kingdom.

274

275

276 **Discussion**

277

278 In this study, we identify ERGU-1, a previously uncharacterized and evolutionarily
279 conserved protein, in a critical ER-resident antioxidant defense system in animal cells.
280 We used a combination of computational and functional screening to pinpoint ERGU-1
281 among the entire UniProt-defined *C. elegans* proteome. Determining the initial list of
282 ERGU candidates for functional validation exemplifies the power of AlphaFold2, an AI
283 for protein structure prediction. Notwithstanding known caveats of predicted structures
284 by AlphaFold2 as compared to experimentally determined ones in accuracy, predicted
285 structures can guide researchers towards promising avenues for functional studies,
286 leading to advances in our understanding of fundamental biological processes and
287 potentially paving the way for new therapeutic strategies^{30–33}. We designed the filtering
288 criteria for screening ERGU candidates based on empirical knowledge gained from
289 DsbD/ScsB and membrane oxidoreductases. Similar strategy based on modified criteria
290 could lead to discovering other protein functions beyond ERGU-1. More broadly, this
291 approach also highlights the potential of AI in accelerating new biological discovery.

292

293 Despite the known generation of H₂O₂ during oxidative protein folding, the identity of the
294 ER membrane-resident protein responsible for its detoxification has remained elusive.
295 Our work bridges this gap by identifying ERGU-1 as such a key player, which harbors

296 functional and structural features characteristic of transmembrane oxidoreductases.
297 Importantly, genetic deletion of ERGU-1 led to constitutive activation of oxidative stress-
298 and ER stress-response reporters, elevated cytosolic H₂O₂ levels, and organismal
299 phenotypes in reproduction and behaviors. These findings strongly suggest ERGU-1's
300 crucial role in mitigating ER-derived oxidative stress, and mechanistically by transferring
301 electrons via cysteine pairs across ER membranes, with ER luminal H₂O₂ as a likely
302 electron acceptor. As loss of ERGU-1 also activated the *hsp-4p::GFP* reporter indicative
303 of unfolded protein stress, it is plausible that ERGU-1 provides reducing equivalents
304 from cytosolic electron donors for additional ER substrates other than H₂O₂, e.g.
305 misfolded proteins and certain peroxide products that ScsB or DscB can act upon to
306 reduce^{12,34,35}. However, it is worth noting that neither ScsB nor DsbD is known to bind to
307 heme, unlike the *C. elegans* and human homologs of ERGU-1. We thus postulate that
308 eukaryotic ER-GUARD may have evolved additional heme-binding capacity that likely
309 enables efficient electron transfer together with cysteine pairs.

310
311 The subcellular localization of ERGU-1 at the ER membrane is well suited for its
312 antioxidant function. By residing at the site close to H₂O₂ generation, ERGU-1 may
313 efficiently neutralize it, safeguarding the ER membrane and nearby cytosol from
314 oxidative damage. Furthermore, the redox-sensitive nature of the tetrameric ERGU-1
315 organization suggests a potential regulatory mechanism for ER-GUARD function. Under
316 more oxidizing conditions, the tetrameric form appears to be more prevalent, potentially
317 enhancing ERGU-1's antioxidant activity to safeguard against excess peroxide stress.
318 Interestingly, our findings on ER-GUARD and its proximity to H₂O₂ generation in the ER

319 align with recent studies suggesting a limited role for mitochondrial ROS in causing
320 global oxidative stress and nuclear DNA damage³⁶. This highlights the ER as a potential
321 major source of cellular oxidative stress impacting nuclear membranes and the genome.
322 By effectively neutralizing ER-derived H₂O₂, ERGU-1 may play a critical role in shielding
323 the genome from oxidative damage. Furthermore, ribosomal RNA, densely packed on
324 ER membranes and crucial for protein synthesis, is another likely target of such
325 oxidative damage. It remains determined whether ERGU-1 protects against oxidative
326 damage in general or more specifically biases towards certain biomolecules. In specific
327 tissues (e.g. intestine), excess peroxide stress caused by loss of ERGU-1 may also
328 activate multiple compensatory antioxidant pathways, including SKN-1/NRF2, and
329 recruit additional mechanisms to safeguard cellular and organismal functions.

330
331 The high degree of sequence conservation observed within the ERGU-1 protein family
332 across diverse animal species underscores its potential evolutionary ancestry and
333 functional importance. The successful complementation of the *C. elegans* *ergu-1* mutant
334 phenotype with homologs from *Drosophila* and humans further strengthens this notion.
335 Although we did not find any apparent orthologues of ERGU-1 in eukaryotic fungi, our
336 findings indicate evolutionary conservation of the ER-GUARD across the animal
337 kingdom, emphasizing its critical and previously undescribed role in maintaining cellular
338 redox balance. Mutations in the *Drosophila* orthologue *emei* impair ER calcium
339 dynamics³⁷, whereas mutations in its vertebrate orthologue *Tmem161B* cause severe
340 pathological cardiac arrhythmias in mice and brain polymicrogyria in humans^{38–41}. Our
341 studies suggest that these previously unexplained phenotypic defects might be

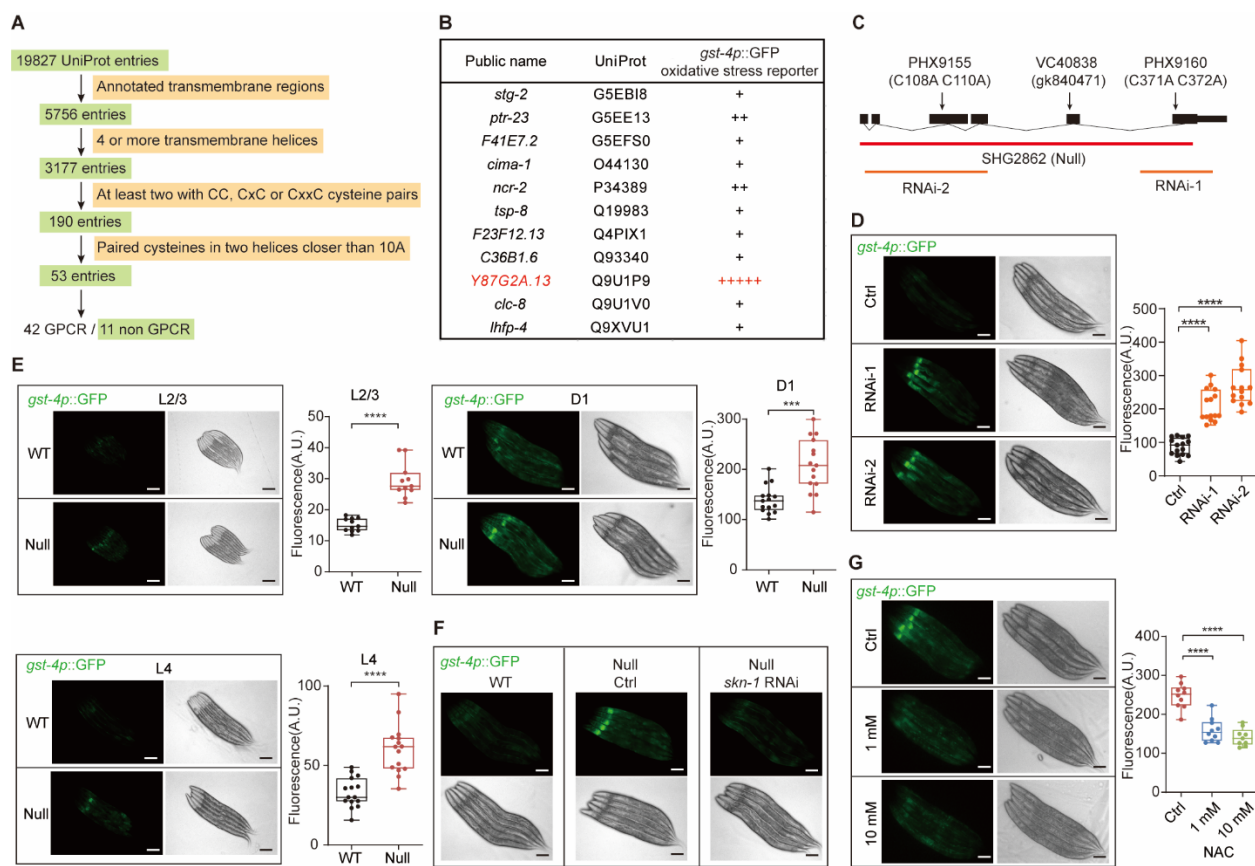
342 mechanically caused by defects in ER-GUARD functions and redox homeostasis.

343 Thus, future studies are warranted to investigate the mechanisms of action, redox

344 regulation of ER-GUARD, and its functional consequences at the cellular and

345 organismal levels that may be broadly important for human physiology and diseases.

346 **Figures**

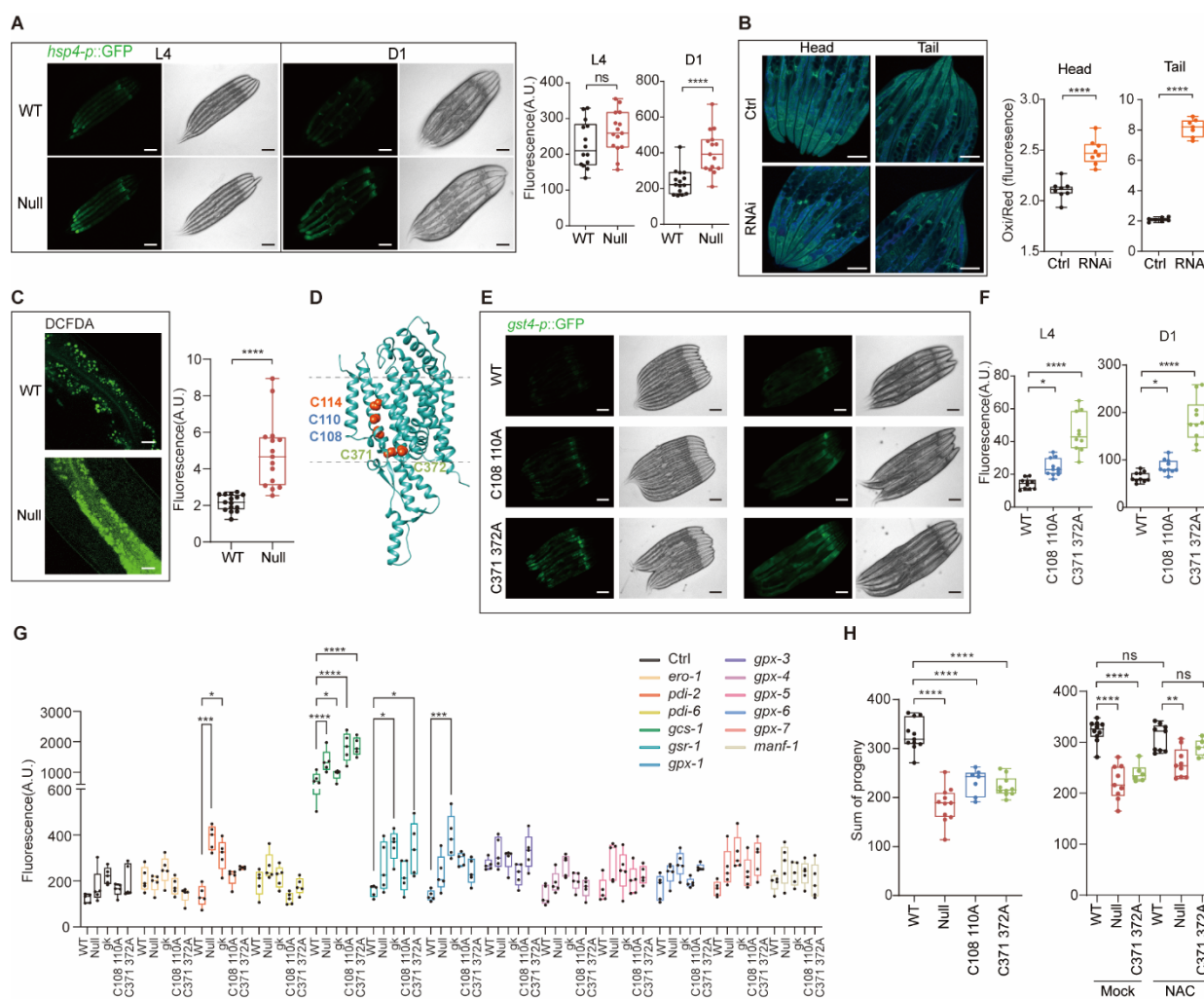


347

348 **Fig. 1. Computational and genetic identification of ERGU-1. (A)** Schematic of
349 AlphaFold2-assisted computational screens. **(B)** Phenotype-driven functional screens
350 based on RNAi activation of *gst-4p::GFP* oxidative stress reporters. **(C)** Gene structure
351 of *ergu-1* showing various alleles. **(D and E)** Representative epifluorescence images
352 showing *gst-4p::GFP* activation in two independent *ergu-1* RNAi (D) and mutants (E)
353 (n=15 for each group). **(F and G)** Representative epifluorescence images showing that
354 *gst-4p::GFP* activation in *ergu-1* null mutants requires ROS and *skn-1* (F), which are
355 suppressed by NAC dose-dependently (G) (n=10 for each group), with quantification of
356 *gst-4p::GFP* fluorescence intensities under conditions indicated. *P* value was

357 determined by an unpaired *t*-test, two-tailed (comparison between two groups) or one-
358 way ANOVA (comparison between multiple groups). Scale bars, 100 μ m.

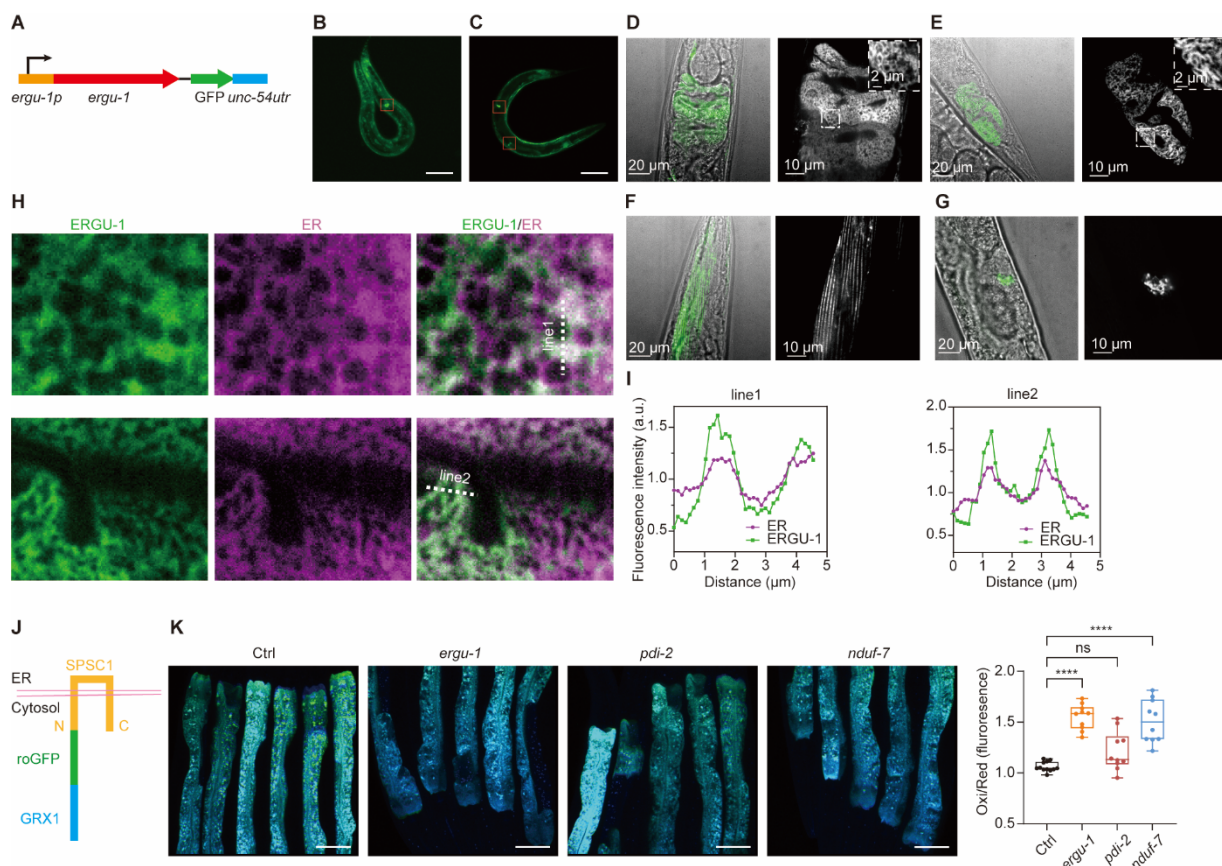
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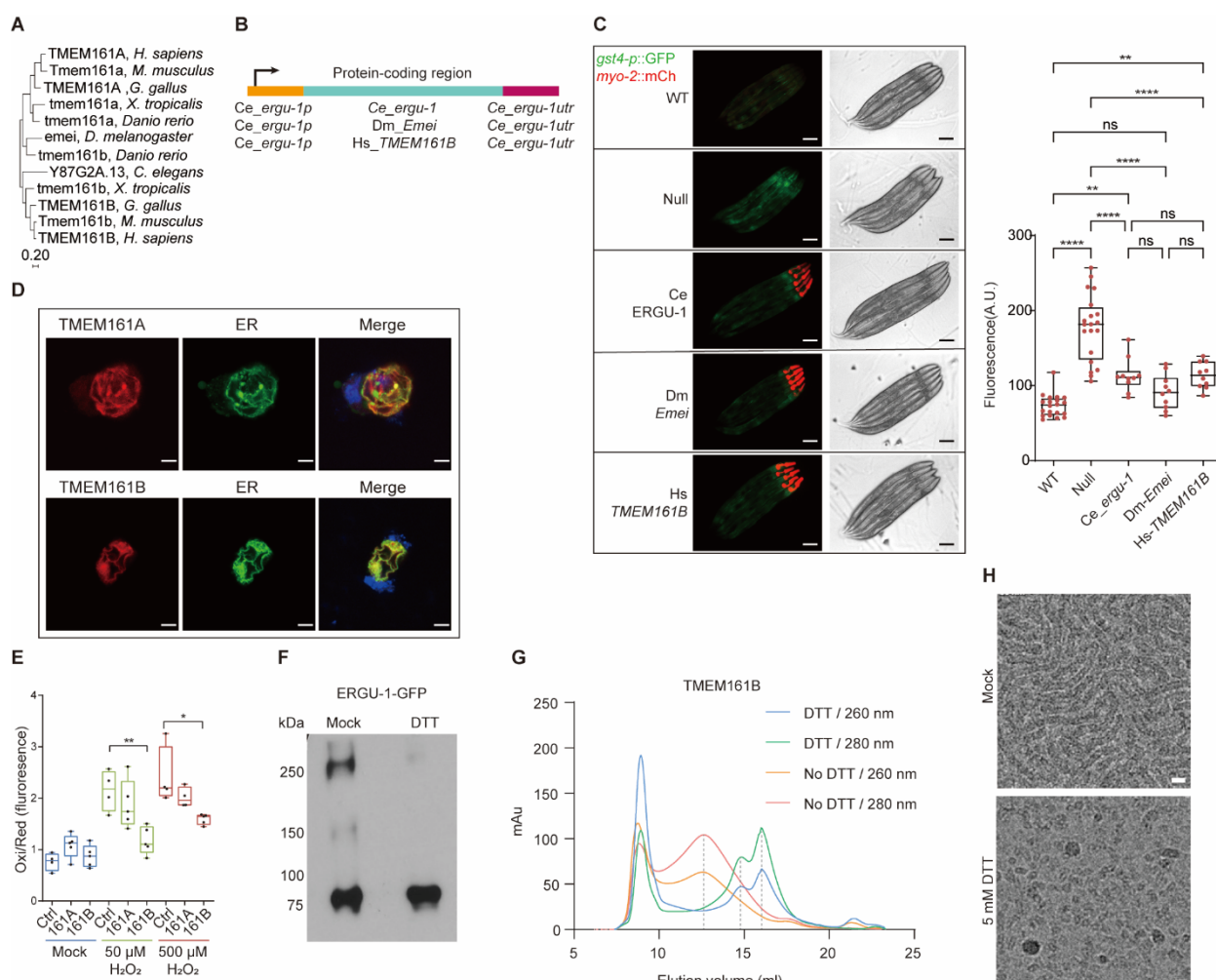
361 **Fig. 2. ERGU-1 defends against H₂O₂ and maintains organismal functions. (A)**
362 Representative epifluorescence images showing increased *hsp-4p::GFP* in *ergu-1* null
363 mutants (n=15 for each group). **(B)** Representative confocal images showing increased
364 oxidation/reduction ratio of Orp1-roGFP (H₂O₂ indicator) with *ergu-1* RNAi. **(C)** DCFDA
365 staining showing increased DCF in null mutants (n=15 for each group). **(D)** AlphaFold2-
366 predicted structure revealing cysteine residues that likely mediate electron transfer

367 across membranes. **(E)** Representative epifluorescence images showing increased *gst-4p::GFP*
368 in two different cysteine pair mutants. **(F)** Quantification of *gst-4p::GFP*
369 fluorescence intensities in WT and two different cysteine pair mutants indicated (n=10
370 for each group). **(G)** Fluorescence intensity of *gst-4p::GFP* in wild type (WT) and *ergu-1*
371 null mutants under control (Ctrl) and RNAi conditions targeting redox regulators,
372 including *ero-1*, *pdi-2*, *pdi-6*, *gcs-1*, *gsr-1*, *gpx-1*, *gpx-3*, *gpx-4*, *gpx-5*, *gpx-6*, *gpx-7*, and
373 *manf-1*. **(H)** Quantification of brood sizes in wild type, *ergu-1* null and cysteine mutants
374 (left panel). Comparison of brood sizes between wild-type and mutant strains under
375 mock and 10 mM NAC treatments (right panel). *P* value was determined by an unpaired
376 *t*-test, two-tailed (comparison between two groups) or one-way ANOVA (comparison
377 between multiple groups). [(A), (B) and (E)] Scale bars, 100 μ m. (C) Scale bars, 20 μ m.



379 Fig. 3. ERGU-1 constitutes a net-like reticulum structure at ER membranes. (A)
380 Schematic of the *ergu-1p*::ERGU-1::GFP translation reporter. **(B and C)** Representative
381 low-mag view of *ergu-1p*::GFP (B) and *ergu-1p*::*ergu-1*::GFP (C) respectively, showing
382 expression in anterior and posterior intestine, body wall muscles and spermatheca.
383 Scale bars, 100 μm. **(D to G)** Representative high-mag confocal view of *ergu-*
384 *1p*::ERGU-1::GFP showing protein localization and expression in anterior (D) and
385 posterior (E) intestinal cells, body wall muscles (F) and spermatheca (G). **(H)**
386 Representative high-mag confocal view of ERGU-1::GFP proteins showing co-
387 localization with the ER membrane marker SEL-1(1-79)::mCherry::HDEL in the
388 intestine, body wall muscles and spermatheca. **(I)** Line scans showing co-localization.
389 Scale bars, 100 μm. **(J)** Schematic design of the ER-proximal H₂O₂ sensor GRX1-

390 roGFP-SPSC-1. (K) Representative ratio metric images and quantification of elevated
391 H₂O₂ levels in the cytosol near the ER under *ergu-1* and *nduf-7* RNAi treatment, as
392 indicated by the ER-proximal H₂O₂ sensor. Scale bars, 100 μm.



393

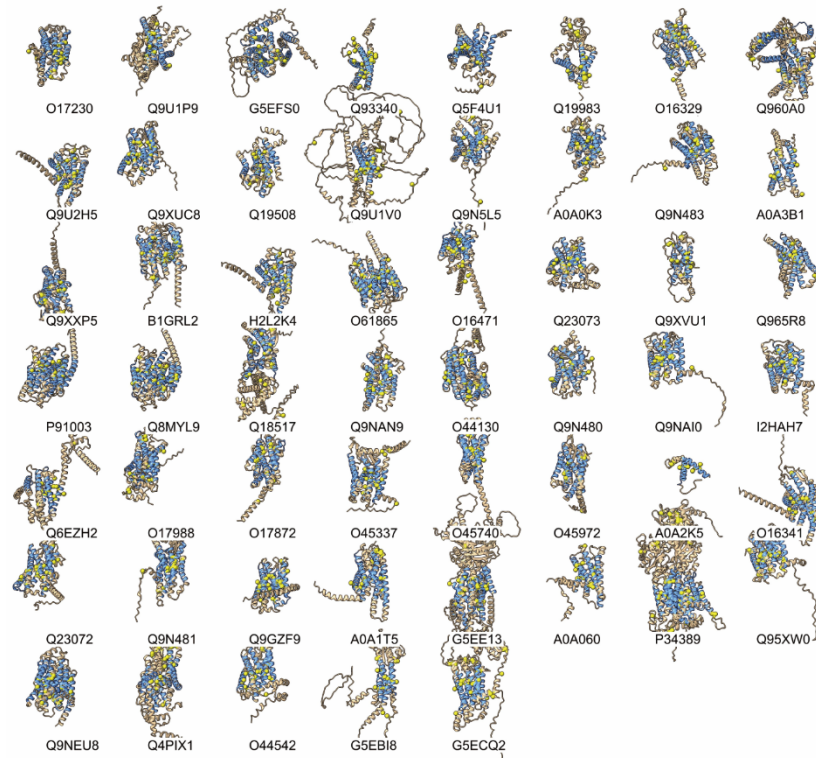
394 **Fig. 4. Conserved ER localization and antioxidant roles of ERGU-1 homologs. (A)**
 395 Maximum likelihood-based phylogenetic tree of ERGU-1 family proteins. (B) Schematic
 396 representation of transgenic constructs and strategy to rescue *ergu-1* null phenotype.
 397 (C) Representative fluorescence images of *gst-4p::GFP* showing rescue of *C. elegans*
 398 *ergu-1* mutants by *Drosophila* and human homologs (n=10 for each rescue group).
 399 Quantification (right). Scale bars, 100 μm. (D) Immunofluorescence images showing
 400 that Human TMEM161A/B localizes to the ER membrane in 293T cells. Scale bars, 10
 401 μm. (E) Reduction of cytosolic H₂O₂ detected by roGFP2-Orp1 in 293T cells
 402 overexpressing human TMEM161A or TMEM161B. (F) Representative SDS-PAGE

403 western blots with antibodies against GFP showing formation of oligomers (likely dimers
404 and tetramers based on the molecular weight) by ERGU-1::GFP and reduction to
405 monomers by 10 mM dithiothreitol (DTT). **(G)** Chromatography plot showing reduction
406 of purified human TMEM161B to smaller size peaks in the presence of 5 mM DTT as
407 analyzed by SEC. **(H)** Reduction of TMEM161B strep-HA from filamentary forms to
408 monomers or dimers in 5 mM DTT as shown by cryo-electron microscopy. Scale bars,
409 10 nm. [(C) and (E)] *P* value was determined by one-way ANOVA.

A

Organism	Blast Name	Score	Number of Hits	Description
Eukaryota	eukaryotes		10	
. . Sar	eukaryotes		2	
. . Reticulomyxa filosa	forams	248	1	Reticulomyxa filosa hits
. . Symbiodinium pilosum	dinoflagellates	95.1	1	Symbiodinium pilosum hits
. . Acer saccharum	eudicots	224	1	Acer saccharum hits
. . Ipomoea batatas	eudicots	151	1	Ipomoea batatas hits
. . Culicoides impunctatus	flies	93.6	1	Culicoides impunctatus hits
. . Idotea baltica	isopods	87.0	2	Idotea baltica hits
. . Friedmanniomyces endolithicus	ascomycete fungi	68.6	3	Friedmanniomyces endolithicus hits

B



410

411 **Fig. S1. AlphaFold2-based computational screen for ERGU candidates. (A)**

412 BLASTP search yielding no apparent broadly conserved eukaryotic homologs of

413 DscB/ScsB based on protein sequence similarities, prompting a search for DscB/ScsB

414 homologs based on functional and structural features using AlphaFold2-predicted

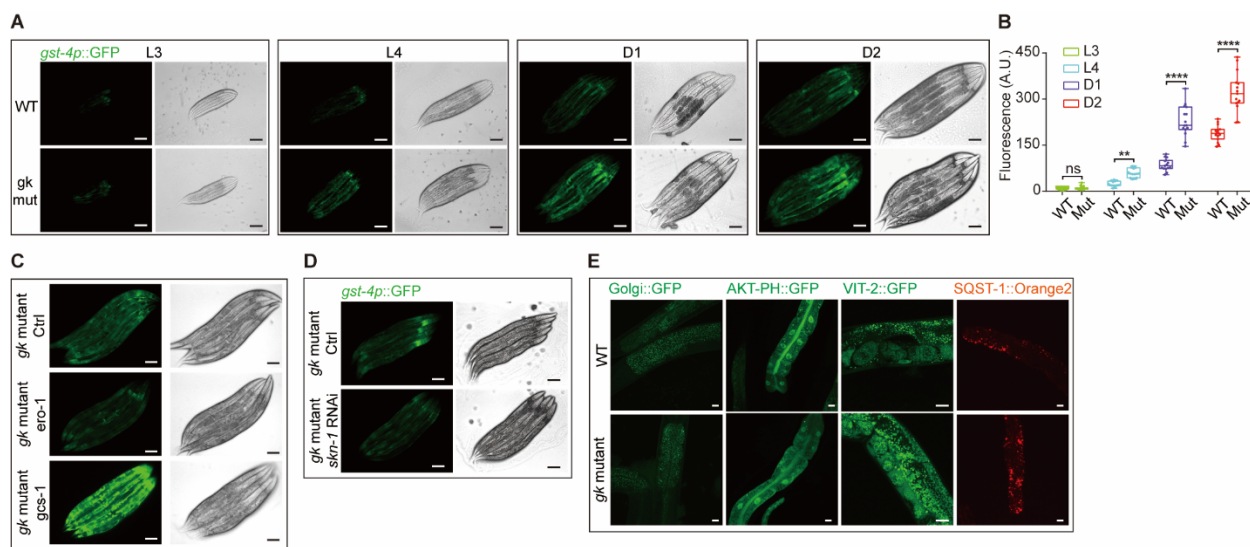
415 structures. **(B)** Predicted structures of the 53 ERGU candidates identified. **(C)** Table

416 listing the annotated functions of the 11 non-GPCR ERGU candidates.

417

C

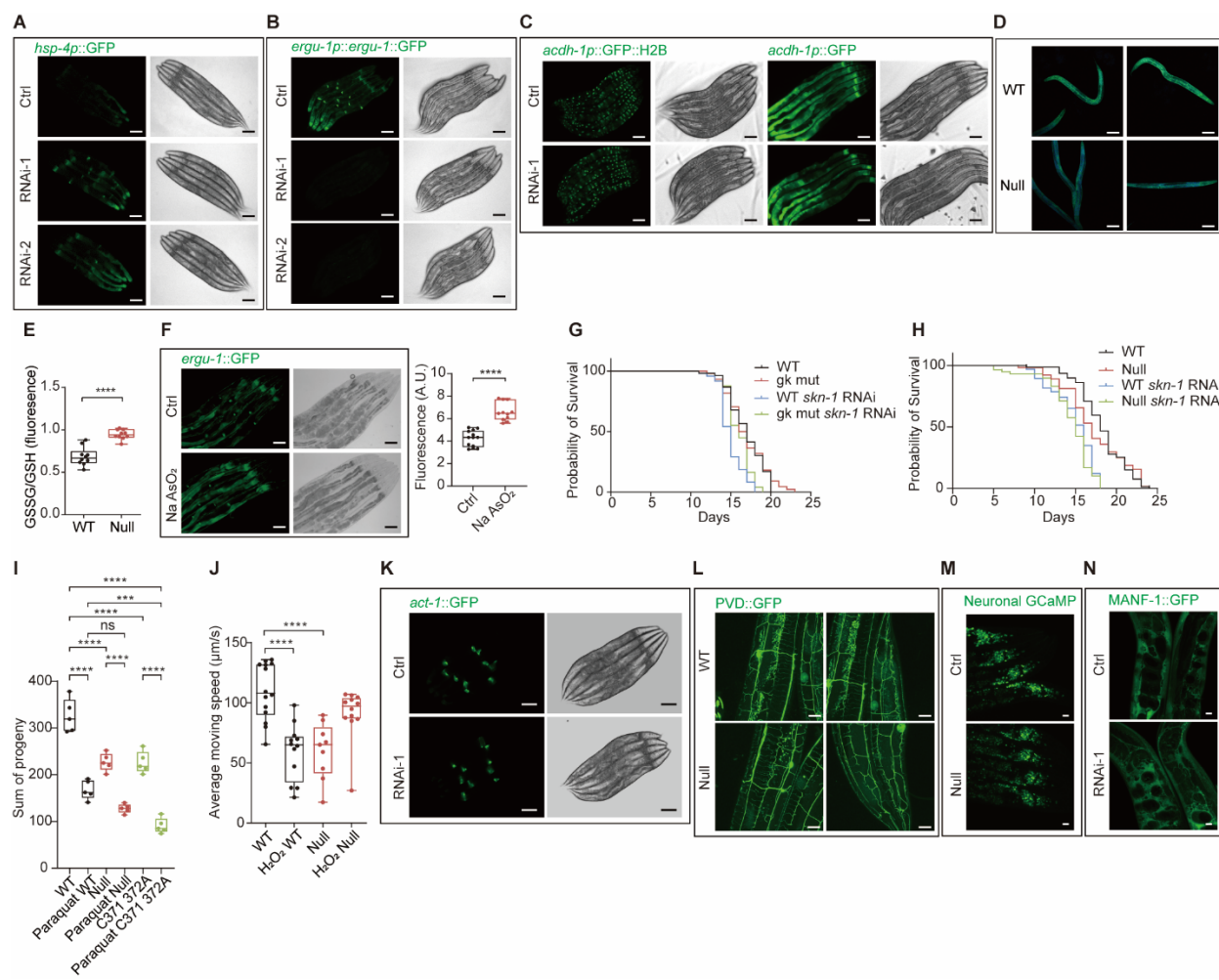
Gene Name	Automated Description
stg-2	Predicted to enable channel regulator activity and voltage-gated calcium channel activity. Involved in adult locomotory behavior and positive regulation of glutamatergic synaptic transmission. Predicted to be located in postsynaptic density membrane. Predicted to be part of AMPA glutamate receptor complex.
ptr-23	Involved in endocytosis and molting cycle. Acts upstream of or within nematode male tail mating organ morphogenesis. Predicted to be located in cytoplasmic vesicle membrane and plasma membrane. Expressed in hypodermis.
F41E7.2	Predicted to enable antiporter activity. Predicted to be involved in proton transmembrane transport. Predicted to be located in membrane. Is an ortholog of human SLC9B1 (solute carrier family 9 member B1) and SLC9B2 (solute carrier family 9 member B2).
cima-1	Predicted to enable sialic acid transmembrane transporter activity. Involved in regulation of protein catabolic process and regulation of synapse organization. Located in lysosome. Expressed in hypodermis and intestine.
ncr-2	Predicted to enable cholesterol binding activity. Involved in cholesterol homeostasis; cholesterol transport; and nematode larval development. Predicted to be located in plasma membrane. Expressed in XXXL; XXXR; gonadal sheath cell; spermathecal; and ventral nerve cord. Used to study Niemann-Pick disease. Human ortholog(s) of this gene implicated in Niemann-Pick disease type C1. Is an ortholog of human NPC1 (NPC intracellular cholesterol transporter 1).
tsp-8	Predicted to be located in membrane.
F23F12.13	Predicted to enable transmembrane transporter activity. Predicted to be involved in transmembrane transport. Predicted to be located in membrane. Is an ortholog of human SLC22A15 (solute carrier family 22 member 15) and SVOPL (SVOPL like).
C36B1.6	Predicted to be located in membrane.
Y87G2A.13	Predicted to be located in membrane. Is an ortholog of human TMEM161A (transmembrane protein 161A).
clc-8	Predicted to be located in membrane.
lhfp-4	Predicted to be involved in sensory perception of sound. Predicted to be located in plasma membrane. Human ortholog(s) of this gene implicated in autosomal recessive nonsyndromic deafness 67. Is an ortholog of human LHFPL4 (LHFPL tetraspan subfamily member 4).



418

419 **Fig. S2. Phenotypes of ERGU-1 protein-truncating *ergu-1(gk840471)* mutants. (A)**
420 Representative epifluorescence images showing *gst-4p::GFP* activation in *ergu-*
421 *1(gk840471)* mutants (n=15 for each group). **(B)** Quantification of *gst-4p::GFP*
422 fluorescence intensities in WT and *ergu-1(gk840471)* mutants indicated (n=15 for each
423 group). **(C)** Representative epifluorescence images showing *gst-4p::GFP* decrease by
424 *ero-1* RNAi and increase by *gcs-1* RNAi **(D)** Representative epifluorescence images
425 showing *gst-4p::GFP* activation in *ergu-1(gk840471)* mutants requires *skn-1*. **(E)**
426 Representative confocal fluorescence images showing no apparent reporter (Golgi)
427 apparatus-marking Golgi::GFP, apical and endosomal membrane-marking AKT-
428 PH::GFP, yolk organelle-marking VIT-2::GFP and autophagosome-marking SQST-
429 1::mOrange2) alteration in *ergu-1(gk840471)* mutants. *P* value was determined by an
430 unpaired *t*-test, two-tailed (comparison between two groups). [(A), (B) and (C)] Scale
431 bars, 100 μ m. (D) Scale bars, 20 μ m.

432

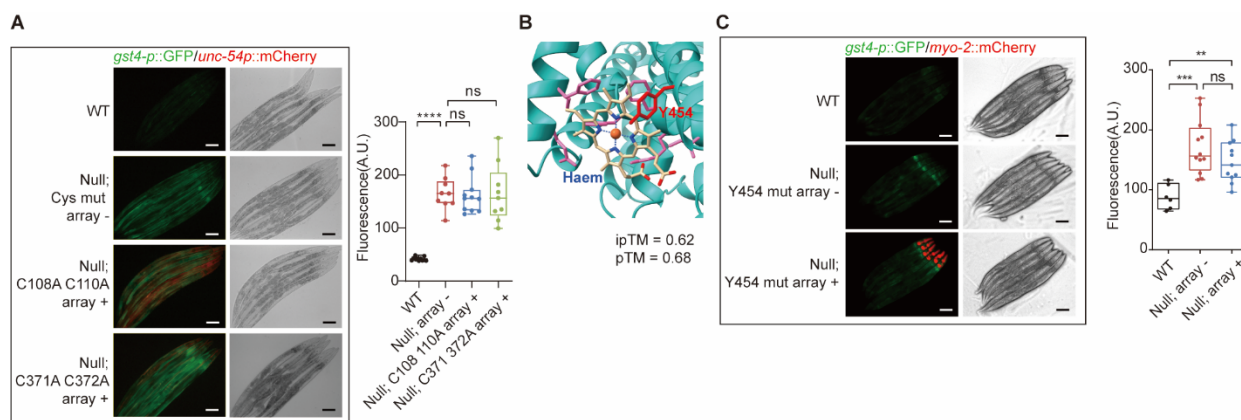


433

434 **Fig. S3. Phenotypic consequences of RNAi or mutant ergu-1. (A)** Representative
435 epifluorescence images showing increased *hsp-4p::GFP* upon *ergu-1* RNAi treatment.
436 **(B)** Representative fluorescence images of *ergu-1p::ergu-1::GFP* showing abolished
437 GFP signals by *ergu-1* RNAi. **(C)** Representative fluorescence images of Vitamin B12
438 stress reporter *acdh-1p::GFP* showing unaltered GFP signals by *ergu-1* RNAi. **(D and**
439 **E)** Representative fluorescence images of glutathione stress GSSG/GSH reporters
440 showing elevated oxidative stress in *ergu-1* null mutants (D) and quantification (n=10)
441 (E). **(F)** Representative fluorescence images of *ergu-1p::ergu-1::GFP* showing reporter
442 up-regulation by oxidative stressors (0.5% NaAsO₂). **(G)** Lifespan curves of wild-type

443 and *ergu-1* null mutants with control or *skn-1* RNAi showing comparable lifespans in
444 wild-type and *ergu-1* null mutants and reduction by *skn-1* RNAi. (H) Lifespan curves of
445 wild-type and *ergu-1* protein-truncating mutants (*gk840471*) with control or *skn-1* RNAi
446 comparable lifespans in wild-type and *ergu-1* (*gk840471*) mutants and reduction by *skn-*
447 1 RNAi ($n \geq 50$). (I) Quantification of brood sizes in wild type, *ergu-1* null and cysteine
448 mutants upon 10 mM paraquat treatment. P value was determined by one-way ANOVA
449 (comparison between multiple groups). (J) Quantification of average moving speed in
450 wild-type, *ergu-1* null and above strain treated with 10 mM H₂O₂. (K) Representative
451 fluorescence images of cytoskeletal actin ACT-1::GFP reporters showing no apparent
452 alteration by *ergu-1* RNAi. (L) Representative confocal fluorescence images showing no
453 apparent reporter (PVD neuron-marking GFP) alteration in *ergu-1* null mutants. (M)
454 Representative confocal fluorescence images showing no apparent reporter (neuronal
455 nuclear GCaMP6) alteration in *ergu-1* null mutants. (N) Representative confocal
456 fluorescence images showing no apparent reporter (secreted MANF-1::GFP) alteration
457 by *ergu-1* RNAi. P value was determined by an unpaired *t*-test, two-tailed (comparison
458 between two groups). [(A) to (J)] Scale bars, 100 μ m. [(K) to (M)] Scale bars, 20 μ m.

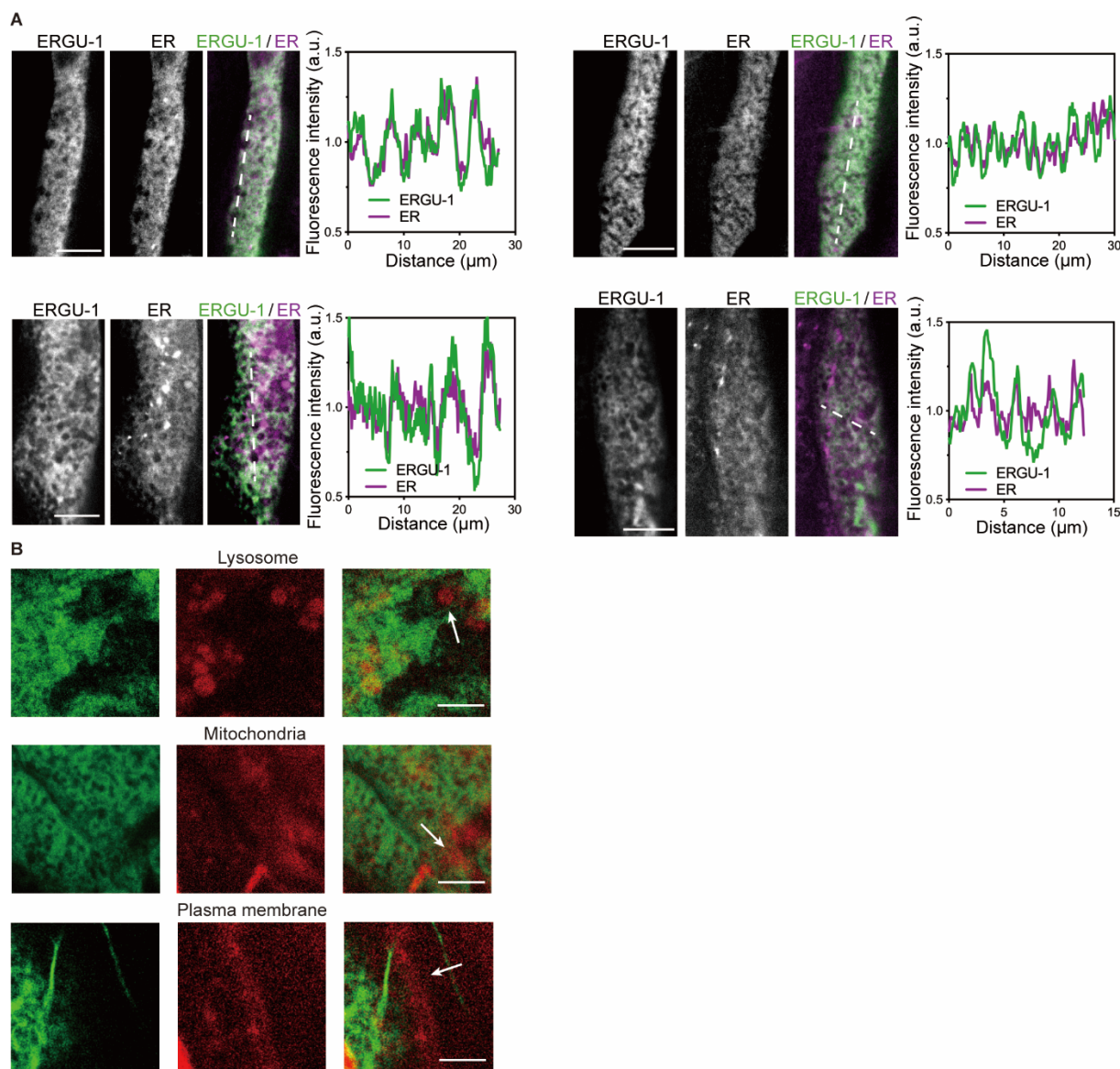
459



461 **Fig. S4. Importance of key cysteine residues for ERGU-1 functions. (A)**

462 Representative fluorescence images of *gst-4p::GFP* showing no rescue of *C. elegans*
463 *ergu-1* mutants by two different cysteine pair mutants of ERGU-1. **(B)** Schematic of key
464 residues for heme binding and interactions. Heme binding pocket is in high confidence
465 region (90 > pIDDT > 70). **(C)** Representative fluorescence images and quantification of
466 *gst-4p::GFP* showing no rescue of *C. elegans* *ergu-1* mutants by *C. elegans* or human
467 Y454 mutants of ERGU-1 homologs. [(A) and (C)] Scale bars, 100 μ m.

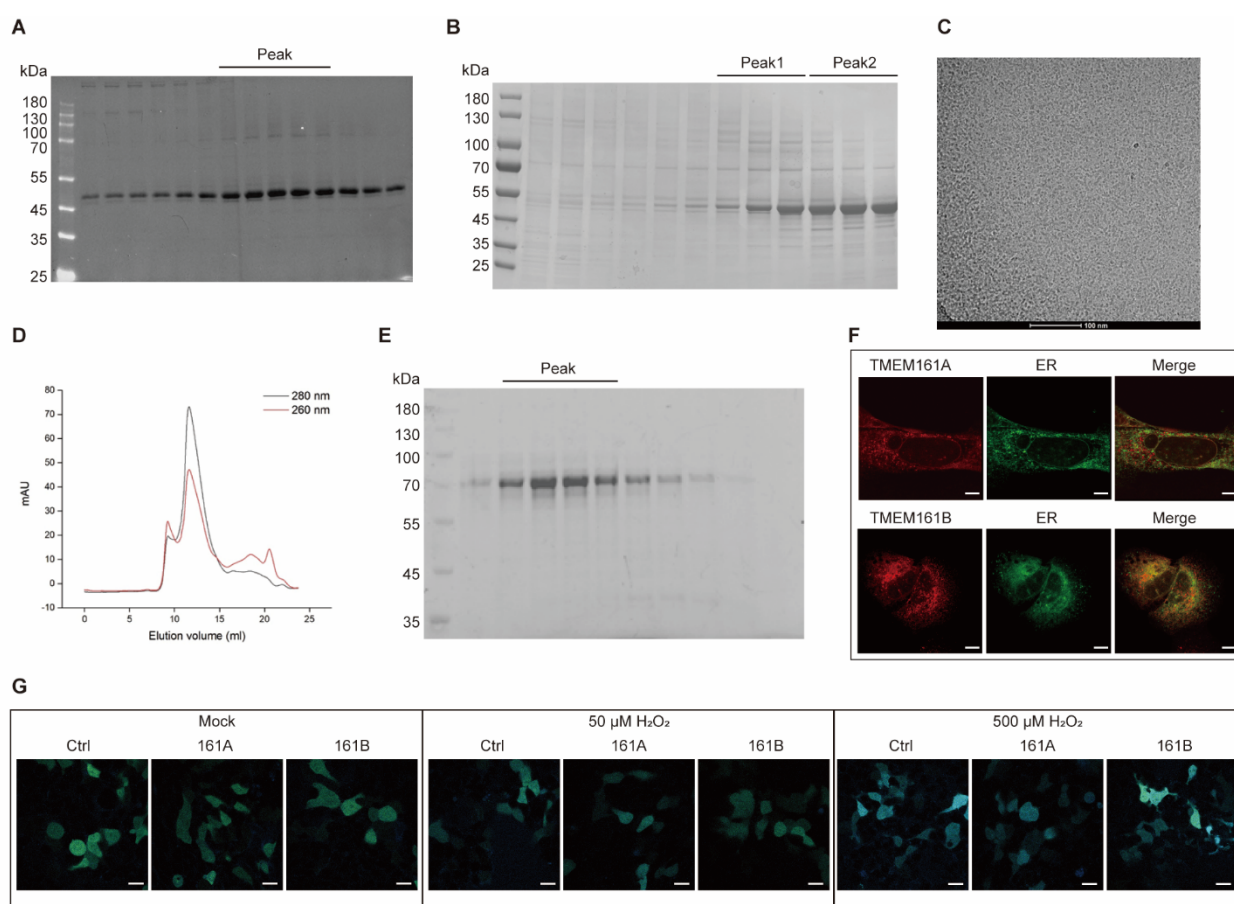
468



469

470 **Fig. S5. Colocalization of ERGU-1::GFP with RFP-marked ER organelle reporters**
471 **but not non-ER organelle reporters.** (A) Representative confocal fluorescence
472 images showing colocalization between ERGU-1::GFP (green) and endoplasmic
473 reticulum-specific reporters (red). Scale bars, 10 μm . (B) Representative confocal
474 fluorescence images showing no colocalization between ERGU-1::GFP (green) and
475 lysosome, mitochondria or plasma membrane-specific RFP reporters (red). Arrow
476 indicates the organelles. Scale bars, 5 μm .

477



478

Fig. S6. Protein characteristics of human TMEM161B, an ERGU-1 ortholog and evolutionarily conserved member of the ERGU protein family. (A) Peak fractions of human TMEM161B-2X Strep-HA protein purified by SEC in a non-reductive environment. (B) Peak fractions of TMEM161B-2X Strep-HA protein purified by SEC in a DTT-containing reductive environment. (C) TMEM161B-GFP in a non-reductive environment shown by Cryo-EM. (D) Peak of TMEM161B-GFP in a non-reductive environment shown by SEC. (E) Peak fractions of TMEM161B-GFP protein purified by SEC in a non-reductive environment. (F) Immunofluorescence images showing that Human TMEM161A/B localizes to the ER membrane in U2OS cells. Scale bars, 10 μm.

488 (G) Representative confocal fluorescence images of roGFP2-Orp1 detecting cytosolic
489 H₂O₂ in 293T cells overexpressing human TMEM161A or B. Scale bars, 20 μm.

490 **Materials and Methods**

491

492 ***C. elegans***

493 *C. elegans* strains were maintained under standard laboratory conditions unless
494 otherwise specified. The N2 Bristol strain was used as the reference wild-type. Feeding
495 RNAi was performed as previously described⁴². Transgenic strains were generated by
496 germline transformation as described⁴³. Transgenic constructs were co-injected (at 10 -
497 50 ng/μl) with dominant *unc-54p::mCherry* or *myo-2p::mCherry*, and stable
498 extrachromosomal lines of fluorescent animals were established for UV-mediated
499 genome integration. Genotypes of strains used are as follows: *ergu-1(ust572, syb9153,*
500 *syb9160)*, *dvls19 [gst-4p::GFP]*, *zcls4 [hsp-4::GFP]*, *jrls10 [unc119(+) rps-0p::roGFP2-*
501 *Orp1]*, *dmals160 [ergu-1p:: ergu-1::gfp; unc-54p::mCherry]*, *dmaEx701[ergu-*
502 *1p::hsTMEM161B::ergu-1utr; myo-2p::mCherry]*, *dmaEx702 [ergu-1p::ergu-1::ergu-1utr;*
503 *myo-2p::mCherry]*, *dmaEx704 [ergu-1p::Dm-161:: ergu-1utr; myo-2p::mCherry]*.

504

505 **AlphaFold-assisted computational screen**

506 A comprehensive computational screen of the *Caenorhabditis elegans* proteome was
507 performed to identify potential DsbD/ScsB-like proteins. The reference proteome utilized
508 for this analysis was the *C. elegans* UniProt reference proteome (version 26),
509 encompassing 19,827 genes. Initial filtering was conducted to select genes encoding
510 proteins with a minimum of four annotated transmembrane helices, resulting in a subset
511 of 3,177 genes. Subsequent filtering criteria focused on the presence of transmembrane

512 helices containing cysteine residues, specifically targeting helices with at least two
513 cysteines separated by no more than two intervening residues, corresponding to the
514 sequence motifs CC, CXC, or CXXC. This filtering narrowed the candidate list to 190
515 genes. The structural analysis was conducted using predicted protein structures from
516 the AlphaFold2 Database. For each of the 190 genes, the predicted structures were
517 examined to determine the spatial proximity of cysteine pairs located within the
518 transmembrane helices. Specifically, structures in which the sulfur atoms (SG) of
519 cysteine pairs were within 10 angstroms of each other were identified. This criterion was
520 met by 53 candidate genes. All computational and structural analyses were executed
521 using custom Python scripts, which are provided in the Supplementary file. Visualization
522 and further structural analyses were performed using UCSF ChimeraX software.

523

524 **RNA interference (RNAi) cloning and screen**

525 RNAi and screen for hits upregulating *gst-4p::GFP* were performed by feeding worms
526 with *E. coli* strain HT115 (DE3) expressing double-strand RNA (dsRNA) targeting
527 endogenous genes. Briefly, dsRNA-expressing bacteria were replicated from the
528 Ahringer library to LB plates containing 100 µg/ml ampicillin (BP1760-25, Fisher
529 Scientific) at 37 °C for 16 hrs. Single clone was picked to LB medium containing 100
530 µg/ml ampicillin at 37 °C for 16 hrs and positive clones (verified by bacteria PCR with
531 pL4440 forward and pL4440 reverse primers) were spread onto NGM plates containing
532 100 µg/ml ampicillin and 1 mM isopropyl β-D-1-thiogalactopyranoside (IPTG, 420322,
533 Millipore) for 24 hrs. Developmentally synchronized embryos from bleaching of gravid

534 adult hermaphrodites were plated on RNAi plates and grown at 20 °C. The worms are
535 collected for imaging at indicated stages.

536 To make the *ergu-1* RNAi-1 and RNAi-2 plasmids, RNAi-1 targeting sequences are

537 PCR- amplified with primers 5'-

538 GGCCCCCCCCTCGAGGTCGACAGGAGACGATGCTATAAACTATGCT -3' and 5'-

539 TCCACCGCGGTGGCGGCCGCGTACGTGTTTCAGCTCG -3' from N2 gDNA and

540 subcloned into the Sall and Notl sites of a digested T777T vector with Promega T4 DNA

541 ligase (Promega, M1801). RNAi-2 targeting sequences are PCR- amplified with primers

542 5'- GGCCCCCCCCTCGAGGTCGACATGCCAAATGAATGCGGGC -3' and 5'-

543 TCCACCGCGGTGGCGGCCGCACGTGGTATGCCTGGTG -3'.

544

545 **Fluorescence microscopy and H₂O₂ sensor imaging**

546 SPE confocal (Leica) and epifluorescence microscopes were used to capture

547 fluorescence images. Animals were randomly picked at the same stage and treated with

548 1 mM levamisole in M9 solution (31742-250MG, Sigma-Aldrich), aligned on a 2% agar

549 pad on a slide for imaging. Identical setting and conditions were used to compare

550 experimental groups with control. For quantification of GFP fluorescence, animals were

551 outlined and quantified by measuring gray values using the ImageJ software. The data

552 were plotted and analyzed by using GraphPad Prism10. For *jrls10* [*unc119*(+) *rps-*

553 *0p::roGFP2-Orp1*] strain, *orp1-roGFP2* was excited sequentially at 405 and 488 nm and

554 emission was recorded at 500-540 nm. Fifteen images were sequentially captured at 1-

555 micrometer z-intervals and subsequently stacked to form a composite image.

556

557 **Western blotting**

558 For SDS-PAGE samples, stage-synchronized animals for control and experiment
559 groups were picked (n = 50) in 60 μ l M9 buffer and lysed directly by adding 20 μ l of 4x
560 Laemmli sample buffer (1610747, Bio-Rad) contain 10% of 2-Mercaptoethanol (M6250-
561 100ML, Sigma (v/v)). Protein extracts were denatured at 95 °C for 10 min and
562 separated on 10% SDS-PAGE gels (1610156, Bio-Rad) at 80 V for ~40 min followed by
563 110 V for ~70 min. The proteins were transferred to a nitrocellulose membrane
564 (1620094, Bio-Rad,) at 25 V for 45 mins by Trans-Blot® Turbo™ Transfer System (Bio-
565 Rad). The NC membrane was initially blocked with 5% non-fat milk and 2% BSA
566 (A4503, Sigma (v/v)) in tris buffered saline with 0.1% Tween 20 (93773, Sigma) (TBST)
567 at room temperature for 1 h. Proteins of interest were detected using antibodies against
568 GFP (A6455, Invitrogen) or V5 (13202T, Cell Signaling Technology) in cold room for
569 overnight. After three washes of 5 min each with tris-buffered saline with 0.1% Tween
570 20, anti-mouse IgG, HRP-linked Antibody (7076, Cell Signaling Technology) was added
571 at a dilution of 1:5000. For DTT treatment, worm lysates were treated with 10 mM DTT
572 and boiled at 95 °C for 10 min.

573

574 **Mammalian cell culture experiments**

575 Human embryonic kidney (HEK) 293T cells and osteosarcoma U2OS cells were
576 maintained in Dulbecco's modified Eagle's medium with 10% inactive fetal bovine
577 serum and penicillin-streptomycin (Gibco, Grand Island, 15140122) at 37 °C supplied
578 with 5% CO₂ in an incubator (Thermo Fisher Scientific) with a humidified atmosphere.
579 Cells were washed once using PBS and digested with 0.05% trypsin EDTA (Gibco) at

580 37 °C for routine passage of the cells. All cells were transfected with 3 μ l LipoD293™
581 (SigmaGen, SL100668) per 1 μ g DNA mixture. HEK293T cells were transfected by the
582 pLX304-TMEM161A-V5 (DNASU, HsCD00441633) or pLX304-TMEM161B-V5
583 (DNASU, HsCD00444935) and collected 2 days after transfection. HEK293T and U2OS
584 cells were transfected by the pHAGE2-TMEM161A-gfp/mCherry or pHAGE2-
585 TMEM161B-gfp/mCherry and collected for imaging 2 days after transfection. pHAGE2-
586 TMEM161A/B-GFP/mCherry was constructed by inserting TMEM161A or TMEM161B
587 into pHAGE2-gfp/mCherry plasmids. TMEM161A was PCR- amplified with the primers
588 5'- TAATTAAACCTCTAGAGCCACCATGGCGGTCCCTCGG -3' and 5'-
589 CTCACCATAGCTCGAGaccagacccGGAGCCTGCCAAGTGC -3' from pLX304-
590 TMEM161A-V5 mentioned before. TMEM161B was PCR- amplified with the primers 5'-
591 TAATTAAACCTCTAGAGCCACCATGGGTGTGATAGGTATACAGC -3' and 5'-
592 CTCACCATAGCTCGAGaccagacccTGCCACAGTCAGATACTGGT -3' from pLX304-
593 TMEM161B-V5 mentioned before.

594

595 **Quantification of brood size**

596 The brood size assay was carried out according to the standard protocol. (1) Briefly,
597 single L4 worms of different stains (N2, ergu-1 null, PHX9155, PHX9160) were
598 individually placed in 60mm Petri plates and kept in the incubator at 20 C. Then, the
599 worms were transferred to a new OP50 containing plate each day at the same time till
600 day 6 of the worms. Progenies were scored after 3 days and plotted for each day and
601 the total sum of progenies. For NAC treatment the plates were supplemented with 10
602 mM of NAC during the entire duration of the assay. For H₂O₂ and paraquat treatment

603 the plates were supplemented with 10 mM of H₂O₂ and paraquat respectively during the
604 assay. Each experiment was repeated 3 times as independent biological replicates with
605 more than 5 animals per group.

606

607 **Behavioral assay**

608 D1 worms (24 hrs post L4) were transferred to a fresh NGM plate seeded with a small
609 and thin OP50 bacterial lawn and allowed to settle for at least ten minutes to recover at
610 room temperature. Moving average speeds, and track lengths of *C. elegans* were
611 monitored for 10 minutes and were analyzed using WormLab. For H₂O₂ treatment, D1
612 worms are treated with 10 mM H₂O₂ for 30 min before assay.

613

614 **Lifespan analysis**

615 For lifespan assays, Animals were cultured under non-starved conditions for at least 2
616 generations before life span assays. For normal NGM life span assay, stage-
617 synchronized L4 stage animals (n ≥ 50) were picked to new NGM plates seeded with
618 OP50 containing 50 µM 5-fluoro-2'-deoxyuridine (FUDR) to prevent embryo growth at
619 20 °C incubator. Animals were scored for survival per 24 hrs. Animals failing to respond
620 to repeated touch of a platinum wire were scored as dead. Three biological replicates
621 were performed, with population sizes larger than 50 in each trial.

622

623 **DCFDA ROS detection**

624 ROS staining in live worms was carried out as in standard protocol with minor
625 modifications. Briefly, day 1 worms were washed three times with M9 and then

626 transferred into 200 μ l of M9 buffer containing 10 mM H2DCFDA (carboxy-H2DCFDA
627 [5-(and-6)-carboxy-2',7'-dichlorodihydrofluorescein diacetate) in an 1.5 ml Eppendorf
628 tube and incubated in the dark for 3 hrs. The worms were randomly selected and
629 treated with 10 mM sodium azide (Sigma-Aldrich) in M9, symmetrically aligned on 2%
630 agar pads on slides for imaging the oxidized dichlorofluorescin (DCF).

631

632 **Expression and purification of TMEM161B for cryo-EM**

633 The complementary DNA encoding human TMEM161B was cloned into a modified
634 pDNA3.1 vector with a C-terminal twin-strep tag and a HA tag. For expression,
635 Expi293F cells (Thermo Fisher Scientific) cultured in Freestyle293 Expression Medium
636 were transfected with the vector DNA/polyethylenimine (1 μ g DNA per ml culture,
637 w/w = 1:3) complex at a cell density of \sim 1.0 \times 10⁶ cells per ml and incubated at 37 °C
638 under 8% CO₂ with agitation at 100 r.p.m for 60 h. Cell pellets were resuspended in
639 buffer A (25 mM Hepes pH 7.5, 150 mM NaCl, protease inhibitor cocktail) and were
640 disrupted by sonication. For TMEM161B monomer purification, an extra incubation with
641 5 mM DTT at room temperature after sonication was required. The cell lysate was then
642 spun at 150,000 \times g for 1h to sediment crude membranes. The membrane pellet was
643 mechanically homogenized in buffer A. The suspension was solubilized in 1% (w/v)
644 1.0% DDM (Anatrance) and 0.1% CHS (Anatrance) for 60 min at 4 °C. The solubilized
645 material was centrifuged at 100,000 \times g for 30 min, and the supernatant was incubated
646 with Strep-Tactin®XT resins (iba) for 2 h at 4 °C. Resins were then washed with 20
647 column volumes of buffer B (25 mM Hepes pH 7.5, 0.025% DDM, 0.0025% CHS,
648 150 mM NaCl). The protein was eluted with buffer B supplied with 50 mM D-biotin,

649 concentrated, and further purified by gel-filtration chromatography on a Superdex 200
650 increase column equilibrated with wash buffer B. The peak fractions of protein were
651 pooled and concentrated to ~5.0 mg/mL. For Cryo-EM observation, 3- μ L aliquots of each
652 sample were applied onto a glow-discharged Quantifoil grid (R1.2/R1.3 300 mesh, Au),
653 blotted for 4.5–5.5 s in 100% humidity at 4 °C, and plunged into liquid ethane using a
654 Vitrobot MkIV (Thermo Fisher Scientific). Cryo-EM micrographs were obtained by using
655 a Talos Arctica G2 microscope (Thermo Fisher Scientific) running at 200 kV and a 300
656 kV Titan Krios microscope (Thermo Fisher Scientific).

657

658 **Statistical analysis**

659 Data were analyzed using GraphPad Prism 9.2.0 Software (Graphpad, San Diego, CA)
660 and presented as means \pm S.D. unless otherwise specified, with *P* values calculated by
661 unpaired two-tailed t-tests (comparisons between two groups), one-way ANOVA
662 (comparisons across more than two groups) and adjusted with Bonferroni's corrections.
663 Box plots are presented as min to max, showing all data points.

664

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674

675 **Author contributions**

676 Z.J. and D.K.M. designed, performed and analyzed the *C. elegans* and cell culture
677 experiments, contributed to project conceptualization and wrote the manuscript. T.P.,
678 B.W., Y.T. and S.H.G. designed, performed and analyzed the *C. elegans* transgenic
679 and CRISPR experiments and editing manuscript. H.B. and Z.J. performed and
680 analyzed the ER imaging experiments. T.D.G conducted the computational screen and
681 contributed to project conceptualization and wrote the manuscript. Z.Y.L., J.X.Y., S.Y.X.
682 contributed to human TMEM161B biochemical analysis. D.K.M, O.D.W. and T.D.G
683 contributed to research materials, funding acquisition and editing manuscript.

684

685 **Competing interests**

686 The authors declare no competing interests.

687 **References**

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