1

# Glutathione reductase gsr-1 is an essential gene required for Caenorhabditis elegans early embryonic development

3

2

- 4 José Antonio Mora-Lorca<sup>a,b</sup>, Beatriz Sáenz-Narciso<sup>c</sup>, Christopher J. Gaffney<sup>d</sup>, Francisco José
- 5 Naranjo-Galindo<sup>a,1</sup>, José Rafael Pedrajas<sup>e</sup>, David Guerrero-Gómez<sup>a</sup>, Agnieszka Dobrzynska<sup>f</sup>,
- 6 Peter Askjaer<sup>f</sup>, Nathaniel J. Szewczyk<sup>d</sup>, Juan Cabello<sup>c</sup>, Antonio Miranda-Vizuete<sup>a,\*</sup>
- 7
- <sup>a</sup>Instituto de Biomedicina de Sevilla, Hospital Universitario Virgen del Rocío/CSIC/Universidad de
   9 Sevilla, 41013 Sevilla, Spain
   <sup>b</sup>Departamento de Farmacología, Facultad de Farmacia, Universidad de Sevilla, 41012 Sevilla, Spain
- 11 <sup>c</sup>Center for Biomedical Research of La Rioja (CIBIR), 26006 Logroño, Spain
- <sup>d</sup>MRC/ARUK Centre for Musculoskeletal Ageing Research, University of Nottingham and Medical
   School Royal Derby Hospital, DE22 3DT Derby, United Kingdom
- <sup>e</sup>Grupo de Bioquímica y Señalización Celular, Departamento de Biología Experimental, Universidad de
  Jaén, 23071 Jaén, Spain
- <sup>f</sup>Andalusian Center for Developmental Biology (CABD), CSIC/JA/Universidad Pablo de Olavide, 41013
  Seville, Spain
- 18
- 19 \*To whom correspondence should be addressed: Antonio Miranda-Vizuete, Instituto de Biomedicina de
- 20 Sevilla (IBIS), Hospital Universitario Virgen del Rocío, 41013 Sevilla, Spain. Tel.: +34 955 923061; Fax:
- 21 +34 955 923101; E-mail: <u>amiranda-ibis@us.es</u>
- <sup>1</sup>Present Address: Functional Genomics and Proteomics, Department of Biology, KU Leuven, 3000
   Leuven, Belgium
- 24
- Keywords: Caenorhabditis elegans, embryonic development, glutathione reductase, mitochondria,
   redox
- 27
- 28

#### 28 HIGHLIGHTS

- C. elegans gsr-1 gene encodes cytoplasmic and mitochondria isoforms of glutathione reductase.
- 30 31

32

34

37

- gsr-1 is essential for C. elegans embryonic development.
- The lethality of *gsr-1* mutants is due to a specific requirement of GSR-1 protein in the cytoplasm.
- *gsr-1* embryos have a progressive cell division delay and an aberrant distribution of interphasic
   chromatin.
- *gsr-1* worms with maternally contributed GSR-1 are able to reach adulthood but display
   mitochondria-associated phenotypes such as increased fragmentation, decreased mitochondrial
   membrane potential and induction of mitochondrial UPR.
- 41
- 42

#### 43 ABSTRACT

44 Glutathione is the most abundant thiol in the vast majority of organisms and is maintained in its reduced 45 form by the flavoenzyme glutathione reductase. In this work, we describe the genetic and functional 46 analysis of the *Caenorhabditis elegans gsr-1* gene that encodes the only glutathione reductase protein 47 in this model organism. By using green fluorescent protein reporters we demonstrate that gsr-1 48 produces two GSR-1 isoforms, one located in the cytoplasm and one in the mitochondria. gsr-1 loss of 49 function mutants display a fully penetrant embryonic lethal phenotype characterized by a progressive 50 and robust cell division delay accompanied by an aberrant distribution of interphasic chromatin in the 51 periphery of the cell nucleus. Maternally expressed GSR-1 is sufficient to support embryonic 52 development but these animals are short-lived, sensitized to chemical stress and have increased 53 mitochondrial fragmentation and lower mitochondrial DNA content. Furthermore, the embryonic lethality 54 of gsr-1 worms is prevented by restoring GSR-1 activity in the cytoplasm but not in mitochondria. Given 55 the fact that the thioredoxin redox systems are dispensable in *C. elegans*, our data support a prominent 56 role of the glutathione reductase/glutathione pathway in maintaining redox homeostasis in the 57 nematode.

#### 58 **INTRODUCTION**

59 Maintenance of thiol redox homeostasis is crucial for survival and the thioredoxin and glutaredoxin 60 systems are the two main pathways that control the redox status in virtually all organisms [1]. The 61 thioredoxin system is composed of thioredoxin reductase (TrxR) and thioredoxins (Trx) while the 62 glutaredoxin system comprises glutathione reductase (GR), glutathione (GSH) and glutaredoxins (Grx), 63 where thioredoxins and glutaredoxins operate as terminal oxidoreductases using the reducing power of 64 NADPH [2]. These two systems are mechanistically and structurally very similar, regulating the 65 formation of disulfides within and between proteins, with the main difference being the use of the 66 tripeptide glutathione (L-y-glutamyl-L-cysteinyl-glycine) as electron donor for glutaredoxins in the 67 glutaredoxin system. In addition, the glutaredoxin system also catalyzes the formation of disulfides 68 between proteins and GSH, namely glutathionylation, which has been shown to be an important 69 posttranslational modification, modulating the activity of many proteins [3].

70

71 Thioredoxin system



76

GSH is found in cyanobacteria, proteobacteria and some gram-positive bacteria as well as most eukaryotes [4]. Inactivating mutations in the gene encoding γ-glutamylcysteine synthetase, the enzyme that catalyzes the first step in GSH synthesis, are lethal in all organisms studied from yeast to mammals, including plants [5-9] highlighting the physiological relevance of GSH in these organisms Moreover, mutations in the gene encoding glutathione synthetase, that catalyzes the second, final step of GSH synthesis, are also lethal in *Arabidopsis thaliana* [10] and *Mus musculus* [11] while they are 83 viable in Saccharomyces cerevisiae [12] and Drosophila melanogaster [13]. In these latter organisms, it 84 has been proposed that the intermediate  $\gamma$ -glutamyl-cysteine, which accumulates in glutathione 85 synthetase mutants, substitutes for GSH to allow growth. Of note, although the C. elegans glutathione 86 synthetase ortholog, gss-1, has not been yet characterized, a gss-1(tm672) deletion mutant has been 87 reported as lethal/sterile by the NBRP С. elegans Gene Knockout Consortium 88 (http://www.shigen.nig.ac.jp/c.elegans/).

89 The lethal phenotype of GSH synthesis mutants contrasts with the dispensability, in the vast 90 majority of eukaryotes, of the glutathione reductase gene that encodes the enzyme that recycles 91 reduced GSH from its oxidized form GSSG. This inessentiality of glutathione reductase for normal 92 growth and development is explained by the thioredoxin system being able to reduce GSSG, a trait that 93 is conserved from bacteria to mammals [14] and also by either accumulating GSSG in yeast vacuole 94 [15] or by excreting GSSG in mammalian cells [16]. Exceptions to glutathione reductase dispensability 95 are the fission yeast Schizosaccharomyces pombe [17], the Plasmodium berghei parasite in its 96 mosquito oocyst stage but not in the blood stage [18] and the A. thaliana GR2 gene, encoding a 97 chloroplastic/mitochondrial glutathione reductase whose inactivation causes early embryonic lethality 98 [19]. The lethality of A. thaliana GR2 mutants is most likely due to a deficiency in chloroplasts function, 99 as in this organism mitochondrial thioredoxin reductase TRXR2 is able to reduce GSSG in mitochondria 100 (Meyer AJ, personal communication). In addition, *D. melanogaster* (and probably other insects) lacks a 101 bona fide glutathione reductase gene and the reduction of GSSG is performed by the thioredoxin 102 system, which is essential in this organism [20].

103 All eukaryotic organisms relying on glutathione metabolism have two distinct pools of glutathione 104 reductase activity located in cytoplasm and mitochondria, respectively (with the exception of 105 photosynthetic organisms that also have glutathione reductase activity in chloroplasts and peroxisomes 106 [21]). In most cases, one single gene encodes both cytoplasmic and mitochondrial glutathione 107 reductase isoforms by the use of alternative translation initiation sites [22-25]. Whereas in E. coli the 108 thioredoxin and glutathione systems are functionally redundant in maintaining redox homeostasis [26], 109 important differences on their respective contribution to the redox status of the different subcellular 110 compartments have been identified in eukaryotic organisms, mainly through work in yeast and 111 mammals. Hence, the yeast thioredoxin system controls the thiol redox status in the cytoplasm where 112 the glutathione system acts merely as a backup. In contrast, the glutathione system is the dominant 113 system to maintain thiol redox control in yeast mitochondria [27-29]. In mammals, knock-out mice 114 lacking glutathione reductase are viable [30] while inactivation of the components of the cytoplasmic 115 and mitochondrial thioredoxin systems causes embryonic lethality in mice [31-34], initially suggesting 116 that the mammalian thioredoxin system may have a predominant role in maintaining redox

117 homeostasis. However, studies on mice harboring conditional alleles of cytosolic thioredoxin reductase 118 TrxR1 to bypass embryonic lethality demonstrate that TrxR1-null mice and cells are robustly viable, 119 relying on the glutathione pathway for survival [35, 36]. Indeed, mice lacking both TrxR1 and glutathione 120 reductase in all hepatocytes sustain hepatic redox homeostasis and organismal survival, through de 121 novo GSH synthesis via the transulfuration pathway using dietary methionine as cysteine precursor 122 [16]. Together, these data in mammals uphold a more prominent role of the glutathione pathway on 123 maintaining redox homeostasis while the thioredoxin system appears to play key functions during 124 embryonic development.

125 In contrast to mammals, the C. elegans thioredoxin system is dispensable for embryonic and 126 postembryonic development, as mutants lacking both cytoplasmic and mitochondrial thioredoxin 127 reductases, trxr-1 and trxr-2, are viable and reach adulthood indiscernibly from wild type controls [37, 128 38], supporting the idea that the glutathione system is also the main responsible for maintenance of 129 redox homeostasis in worms. Lüersen et al. have shown that the C. elegans gsr-1 gene encodes a 130 functional glutathione reductase protein [39], which is required for survival to paraguat and juglone 131 treatments. In this work, we demonstrate that the C. elegans gsr-1 gene encodes both cytoplasmic and 132 mitochondrial glutathione reductase isoforms and that is required for embryonic development. This 133 lethal phenotype of gsr-1 mutants arises from a specific requirement of the enzyme in the cytoplasm, 134 the subcellular compartment where GSH is synthesized. In contrast, gsr-1 mutants maternally 135 expressing GSR-1 are able to develop normally but are sensitized to stress, are short-lived and have 136 compromised mitochondria.

#### 137 **RESULTS**

138

#### 139 *C. elegans* **GSR-1** is widely expressed and is targeted to both cytoplasm and mitochondria.

140 The gsr-1 locus is organized into 5 exons and 4 introns and expresses two main mRNA variants, 141 gsr-1a and gsr-1b1 (Figure 1A), whose conceptual translation results into two different isoforms, GSR-142 1a and GSR-1b, with the former having an additional 14 amino acid N-terminal extension (Figure 1B). 143 Two other minor mRNA variants have been reported, gsr-1b2 and gsr-1b3, which differ only in the 144 of 5'-UTRs but sequence their respective also generate the GSR-1b isoform 145 (http://www.wormbase.org/). C. elegans GSR-1 is highly homologous through all protein domains to 146 vertebrate glutathione reductases, including the conserved redox active site CVNVGC (Figure 1B-C). 147 The GSR-1a N-terminal extension, mostly encoded by the first gsr-1 exon, displays characteristics of a 148 mitochondrial targeting sequence (MTS) [40], cleavable by the two matrix protease model (Figure 1D) 149 [41]. This predicts that GSR-1a is located in the mitochondrial matrix while GSR-1b, lacking this putative 150 MTS and initiated from a downstream in-frame ATG codon located at the beginning of the second exon 151 (**Figure 1A**), is a cytoplasmic protein.

152 To demonstrate this dual subcellular localization *in vivo* as well as to describe in detail the tissue 153 and cellular expression pattern of both GSR-1 isoforms, we first generated transgenic worms 154 expressing a translational GFP fusion spanning a 2.2 kb fragment of the gsr-1 promoter plus the 155 complete gsr-1 genomic ORF (Figure 2A). These transgenic animals showed strong fluorescence in 156 pharynx while weaker labeling was found in hypodermis, intestine, vulva muscle cells, spermatheca, 157 uterine cells, coelomocytes, gonad sheath cells, rectal cells and a couple of unidentified neurons 158 (probably PVPL/R) in the tail (Figure 2B-E). Long first introns have been shown to be important for 159 gene regulation by binding a set of transcription factors different to those binding the upstream promoter 160 [42]. As gsr-1 exons 1 and 2 are separated by a 318 bp intron, considerably larger than the rest of gsr-1 161 introns (Figure 1A), we set to investigate whether the *gsr-1* first intron also has promoter activity. 162 Interestingly, transgenic worms expressing a translational GFP fusion spanning the gsr-1 ORF from the 163 first intron (Figure 2F) show labeling in some neurons in the head, excretory channel, intestine and 164 coelomocytes (Figure 2G-K). This result suggests that the gsr-1 first intron possess intrinsic promoter 165 activity, independent from that of the gsr-1 upstream promoter.

To prove that *C. elegans gsr-1* gene encodes both mitochondrial and cytoplasmic isoforms *in vivo*, we generated transgenic worms expressing the full *gsr-1* translational GFP fusion in which either ATG codon is mutated (**Figure 2L**). Thus, inactivation of the second ATG codon forces translation from the first ATG codon resulting in a dotted GFP pattern, consistent with mitochondrial localization (best seen in the pharynx) [43] (**Figure 2M**). In turn, when forcing expression from the first ATG codon by inactivation of the second ATG codon, a diffuse GFP labeling indicating a cytoplasmic localization was

172 obtained (Figure 2N). Similar diffuse fluorescence labeling was found when removing *gsr-1* first exon. 173 which encodes most of GSR-1a MTS (data not shown). We further demonstrate that the additional 14 174 amino acids of GSR-1a constitute a functional MTS on its own as, when expressed in worm muscle 175 cells, the fusion protein MTSgsr-1::GFP is found in the typical tubular distribution along the cell 176 longitudinal axis, colocalizing with the TOMM-20::mRFP mitochondrial marker [38] (Figure 2Q-T). 177 Together, we conclude that the C. elegans gsr-1 gene encodes both mitochondrial (GSR-1a) and 178 cytoplasmic (GSR-1b) isoforms. In addition, we further refine and expand the gsr-1 tissue and cellular 179 expression pattern previously reported [37, 39].

180

### 181 *gsr-1* is essential for *C. elegans* embryonic development and *gsr-1(m+,z-)* mutants are 182 sensitized to chemical and developmental stresses and have compromised mitochondria.

183 To investigate GSR-1 function, we used a worm strain carrying the gsr-1(tm3574) deletion allele, 184 which spans 383 bp and removes part of *gsr-1* third exon, the third intron and part of the fourth exon 185 (Figure 1A). Sequencing of *qsr-1(tm3574*) cDNA demonstrates that it encodes an in frame truncated 186 protein (GSR-1 $\Delta$ 193-302, abbreviated as  $\Delta$ GSR-1) lacking most of the NADPH domain plus a minor 187 fraction of the second FAD domain (Figure 1B). Using purified recombinant protein expressed in 188 bacteria, we show that wild type GSR-1b is able to reduce GSSG in a dose-dependent manner while 189  $\Delta$ GSR-1b was devoid of enzymatic activity (**Figure 3A**), thus confirming gsr-1(tm3574) as a loss of 190 function allele most likely by failure to bind NADPH required for redox cycling. GSR-1b did not function 191 as thioredoxin reductase when using either yeast TRX3 or worm TRX-2 as substrates (data not 192 shown).

193 The gsr-1(tm3574) allele is reported as lethal/sterile by the NBRP C. elegans Gene Knockout Consortium (http://www.shigen.nig.ac.jp/c.elegans/). Six times backcrossed gsr-1(tm3574) animals 194 195 retained the lethal phenotype, so the tm3574 allele was maintained in heterozygosity using GFP or 196 RFP derivatives of the *qC1* balancer [44]. Homozygous *gsr-1* animals segregating from balanced 197 parents (hereafter referred as qsr-1(m+,z-); m, maternal and z, zyqotic), have a normal embryonic and 198 postembryonic development, reaching reproductive stage indistinguishably of wild type controls. Thus, 199 maternally contributed gsr-1 mRNA and/or GSR-1 protein is enough to allow gsr-1 mutants to 200 accomplish a normal life cycle. In turn, the gsr-1(m,z) embryos generated by these animals invariably 201 arrest during embryogenesis.

RNAi downregulation of *gsr-1* expression has been shown to decrease *C. elegans* lifespan, to sensitize worms to the prooxidant juglone and the superoxide anion generator paraquat [39], to impair worm molting in the absence of cytoplasmic thioredoxin reductase *trxr-1* gene [37] and to induce *skn-1* target genes such as *gst-4* or *gcs-1* without an obvious nuclear translocation of a *skn-1* GFP reporter

206 [45]. To determine whether the maternal contribution of gsr-1(m+,z-) worms is functionally equivalent 207 to the decreased gsr-1 mRNA levels caused by RNAi, we tested gsr-1(m+,z-) individuals in all the 208 above mentioned scenarios. Our data indicate that, indeed, this is the case as qsr-1(m+,z-) animals 209 are short-lived (Figure 3B), are highly sensitive to juglone and paraquat (Figure 3C), have a fully 210 penetrant larval arrest phenotype in a trxr-1(sv47) mutant background (Figure 3D) and induce GST-4 211 and GCS-1 reporters without significant SKN-1B/C::GFP nuclear translocation (Figure 3E). Therefore, 212 maternally provided gsr-1 mRNA and/or GSR-1 protein is enough to allow gsr-1(m+,z-) mutants to 213 reach reproductive stage under non-stressed growth conditions but it is insufficient to afford effective 214 protection against stress or developmental constrains.

215 The extreme sensitivity of gsr-1(m+,z-) worms to chemicals that impair mitochondrial function 216 prompted us to assess mitochondrial status in these animals. We found that gsr-1(m+,z-) worms have 217 a clear mitochondrial fragmentation phenotype in muscle cells (Figure 4A) while muscle sarcomere 218 structure is preserved (Figure 4B). In addition, the mitochondrial membrane potential and total 219 mitochondria content measured by incorporation of the fluorescent dye JC-10 was significantly 220 decreased (Figure 4C). Moreover, gsr-1(m+,z-) worms display a strong induction of the mitochondrial 221 UPR reporter HSP-6::GFP [46] (Figure 4D), overall indicating that decreased gsr-1 levels cause 222 mitochondrial stress and dysfunction. As a whole, these results validate the use of gsr-1(m+,z-) worms 223 as a tool to investigate the role of GSR-1 in postembryonic development.

224

## 225 *gsr-1(m-,z-)* embryos arrest at the pregastrula/gastrula stage displaying progressive cell 226 division delay and aberrant interphasic chromatin distribution.

227 Next, we moved to study in detail the gsr-1(m,z) embryonic arrest phenotype. For this purpose, 228 we video-recorded several qsr-1(m-z-) embryos (n=26) and found that they all arrest at the pregastrula/gastrula stage ranging from 17 to 103 cells (Figure 5A-C and Movies 1-3). This arrest 229 230 interval is roughly coincidental with the time at which C. elegans embryos start expressing a Pgsr-231 1:: *qfp* transcriptional construct (**Figure 5D**). Arrested gsr-1(m,z) embryonic cells appear normal for at 232 least the length of normal embryogenesis with no signs of necrosis or apoptosis, as it has been 233 reported for most embryonic lethal mutants [47] (Figure 5A-C and Movies 1-3). However, cell lineage 234 analysis of gsr-1(m,z) embryos identifies a dramatic delay of cell divisions timing, already detectable 235 at the first embryonic divisions that is progressively enhanced until complete arrest occurs (Figure 6A 236 and Supplemental Figure 1A-C).

We next explored the possible causes of the progressive delay of cell division timing of *gsr-1(m-,z-)* embryos. Among other traits, mutations in genes impairing mitochondrial function or genes encoding chromatin regulators have been shown to retard cell divisions in *C. elegans* embryos [48, 49]. Therefore, we set to investigate whether *gsr-1* mutants impact these two pathways given that

mitochondria-associated phenotypes are found in gsr-1(m+,z-) animals and because histone 3 glutathionylation destabilizes nucleosome structure and GSH depletion decreases DNA synthesis rate [50].

244 To evaluate the integrity of the mitochondrial network in gsr-1(m,z) embryos, we attempted 245 labeling these organelles with rhodamine 6G followed by in vivo image analysis [51]. Despite 246 rhodamine 6G efficiently stained embryo mitochondria, no major differences in the dynamics of the 247 mitochondrial network were observed between young wild type and gsr-1(m,z) embryos (Movie 4). 248 Reduced cell size of older embryos precluded analysis of differentiated embryonic cells. Regarding the 249 other possible cause of cell division delay, we examined chromatin dynamics in gsr-1(m-z-) embryos 250 by time-lapse confocal microscopy. During early development, interphasic chromatin occupied the 251 entire nuclear volume whereas mitotic chromosomes condensed in the nuclear interior (Figure 6B 252 and Movie 5; 1:10). However, concomitantly with cessation of cell divisions in gsr-1(m,z) embryos, a 253 progressive aberrant distribution of chromatin at the nuclear periphery was observed (Figure 6B and 254 Movie 5; 2:20-5:00). Collectively, these results suggest that gsr-1 deficiency impacts nuclear 255 chromatin dynamics pointing to a nuclear dysfunction as a possible cause of gsr-1(m-,z-) embryonic 256 arrest.

257

258 Cytoplasmic, but not mitochondrial, GSR-1 expression restores viability of *gsr-1(m-,z-)* 259 embryos.

260 The fully penetrant embryonic arrest phenotype of gsr-1(m-,z-) embryos is rescued by 261 transgenes expressing wild type GSR-1 (vzEx105 transgene) as well as GSR-1::GFP fusion protein 262 (vzEx93 transgene) (Figure 7A), demonstrating that the lethality is due to a gsr-1 deficiency. In 263 contrast, no rescue was obtained when gsr-1(m+,z-) worms and their gsr-1(m-,z-) embryos were 264 maintained on an effective external source of different GSH donors or precursors either in solid or 265 liquid medium [5 mM GSH, 40 μM S-linolenoyl-GSH [52]; 83 mg/ml liposomal GSH [53]; 20 μM 266 glutathione reduced ethyl ester [54]; 10 mM N-acetyl-L-cysteine [55]; 10 mM L-cystine, 10 mM L-267 methionine [56]) or 5mM DTT (data not shown)]. This lack of rescue indicates that qsr-1(m+,z-)268 worms do not provide these compounds to their oocytes (or they are rapidly exhausted) and also 269 reflects the inability of these chemicals to cross the gsr-1(m-z) embryo eqgshell, which is non-270 permeable to most solutes [57].

As mentioned above, both wild type GSR-1 and GSR-1::GFP fusion protein rescue the *gsr-1* embryonic lethal phenotype. In *gsr-1; vzEx105 [Pgsr-1::gsr-1]* animals, untagged GSR-1 produced by the *vzEx105* transgene provides enough maternal load to allow the non-transgenic *gsr-1* progeny to reach the reproductive stage (**Figure 7A, dashed bar**), similarly to *gsr-1(m+,z-)* worms segregating from *gsr-1/qC1* balanced parents. In contrast, in *gsr-1; vzEx93 [Pgsr-1::gsr-1::gsr-1::gfp]* animals, the GSR-

1::GFP fusion protein produced by the *vzEx93* transgene fails to provide maternal rescue, so only the transgenic progeny carrying the *vzEx93* transgene are viable, while all the non-transgenic siblings arrest at early embryogenesis (difference between eggs laid and adults, **Figure 7A**). The failure of the GSR-1::GFP fusion protein to provide maternal rescue could be explained by a lower enzymatic efficiency and/or a faster turnover of the GSR-1::GFP homodimers compared to wild type GSR-1 homodimers.

282 We next asked whether the embryonic lethal phenotype is due to a specific GSR-1 requirement 283 in cytoplasm or mitochondria or, instead, to the simultaneous absence of the protein in both 284 compartments. To address this, we performed isoform-specific rescue experiments in *gsr-1* mutants 285 expressing, respectively, the mitochondrial GSR-1a::GFP or cytoplasmic GSR-1b::GFP fusion proteins 286 (Figure 2L-N). As GSR-1::GFP fusions do not provide maternal load, a qualitative determination of the 287 isoform-specific rescue is possible by simply scoring viable versus non-viable progeny. Interestingly, 288 while gsr-1 mutants expressing GSR-1b::GFP exclusively in the cytoplasm (either from the vzEx145 289 [Pgsr-1::gsr-1b(Δ1st exon)::gfp] or the vzEx147 [Pgsr-1::gsr-1b(1st ATG mutated)::gfp] transgenes) 290 rescued the embryonic lethal phenotype, no viable progeny was obtained from gsr-1 mutants 291 expressing the mitochondrial specific GSR-1a::GFP isoform (from the vzEx158 [Pgsr-1::gsr-1a(2nd 292 ATG mutated)::gfp] transgene) (Figure 7A). These data imply that cytoplasmic GSR-1 is sufficient to 293 restore normal embryonic development and that alternative redox systems such as the cytoplasmic 294 thioredoxin system (redundant to the glutathione system in other organisms) are not able to substitute 295 GSR-1 when absent from the cytoplasm.

296 The dispensability of mitochondrial GSR-1a, at least under non-stress normal growth conditions, 297 could be explained by the presence of a redundant GSSG reducing system in mitochondria. In S. 298 cerevisiae and A. thaliana, mitochondrial thioredoxin reductase is able to reduce GSSG in this 299 organelle in the absence of mitochondrial glutathione reductase [58] (Meyer AJ, personal 300 communication). To test if this genetic redundancy is maintained in *C. elegans*, we generated strains 301 expressing only cytoplasmic GSR-1b::GFP (from vzEx145 and vzEx147 transgenes) in a gsr-302 1(tm3574) trxr-2(tm2047) double mutant background. As shown in **Figure 7B**, restoring GSR-1 activity 303 exclusively in the cytoplasm of gsr-1 trxr-2 double mutants also produced viable progeny, arguing 304 against TRXR-2 as a redundant system for mitochondrial GSR-1a in worms. Collectively, these data 305 demonstrate that in *C. elegans*, cytoplasmic GSR-1b is essential for embryonic development and that 306 mitochondrial GSR-1a might be required for a yet to be identified dispensable function in this 307 organelle.

#### 308 **DISCUSSION**

309 Most eukaryotic organisms have one single glutathione reductase gene encoding both 310 cytoplasmic and mitochondrial isoforms by the use of alternative translation initiation sites [22-25]. In

311 this study, we show that this is also the case for C. elegans glutathione reductase gsr-1 gene, where 312 translation from the first ATG codon results in the mitochondrial GSR-1a isoform, with an N-terminal 313 MTS mostly encoded by the first exon, while translation from a second in-frame ATG codon located at 314 the beginning of the second exon generates the cytoplasmic isoform GSR-1b (Figure 2L-T). 315 Furthermore, C. elegans is one of the few examples of an eukaryotic organism with an essential 316 requirement of glutathione reductase for viability. We show here that it is the cytoplasmic isoform of 317 GSR-1 that is essential for viability (Figure 7A). While in many organisms the thioredoxin system 318 backups GSSG reduction in the absence of glutathione reductase [14] this appears not to be the case 319 for C. elegans, despite the fact that both systems have been shown to cooperate in other processes 320 such as worm molting [37]. Therefore, the C. elegans thioredoxin and glutathione systems share 321 common functions but also have specific non-overlapping roles in worm physiology.

322 gsr-1 mutants with maternal load, gsr-1(m+,z-), are viable and reach the reproductive stage 323 indiscernibly from wild type controls. In turn, gsr-1(m,z) animals lacking maternal load arrest during 324 early embryogenesis, displaying a robust progressive cell division delay phenotype (Figure 6A) 325 concomitant with an aberrant distribution of interphasic chromatin (Figure 6B). Several possibilities 326 might account for this lethal phenotype: i) as GSH depletion has been shown to inhibit DNA synthesis 327 and to compromise cell cycle progression in mammalian cells [59, 60], this dependence on GSH supply 328 may also apply to C. elegans embryos. A possible mechanism could involve histone H3 329 glutathionylation that leads to a relaxed histone octamer in which DNA is less tightly packed around the 330 nucleosome, thus opening chromatin [50]. Conversely, deglutathionylation of histone H3 results in a 331 more compact distribution of chromatin ultimately decreasing gene expression and DNA synthesis. 332 Thus, the absence of a GSH regenerating system, in gsr-1(m-z) embryos could compromise nuclear 333 histone H3 glutathionylation, which may explain the aberrant chromatin distribution and cell division 334 delay; ii) related to the this, the thioredoxin and glutathione/glutaredoxin systems are the two enzymatic 335 pathways providing reducing equivalents to the essential enzyme ribonucleotide reductase (RNR), 336 responsible for the supply of deoxyribonucleotides for DNA synthesis and repair [61]. While the 337 glutathione system is much more efficient with the E. coli RNR enzyme, in mammals the thioredoxin 338 system is one order of magnitude more efficient than the glutaredoxin system [62]. If C. elegans RNR is 339 more dependent on the glutathione system, as found in bacteria, failure to reduce RNR could be a 340 cause of the lethality of gsr-1(m,z) worms; iii) mutations in mitochondrial genes have also been shown 341 to cause embryonic arrest in C. elegans [47, 48]. Although we have not been able to detect abnormal 342 mitochondrial network distribution in gsr-1(m,z) embryos (**Movie 4**), the fact that maternally rescued 343 gsr-1 mutants have lower amount of mitochondria, increased mitochondrial fragmentation and induced levels of hsp-6, a mitochondrial stress reporter (Figure 4), it is possible that impaired mitochondrial 344

345 function may also account for the gsr-1(m-,z-) embryonic arrest phenotype; iv) alternatively, it is 346 possible that gsr-1(m,z) embryos could die by GSSG poisoning. In mammalian cell cultures with 347 compromised GSSG reducing activity, GSSG is excreted into the cell culture medium to avoid the 348 toxicity of its intracellular buildup [16]. If such mechanism also happens in C. elegans gsr-1(m-,z-) 349 embryos, excreted GSSG would accumulate within the non-permeable extracellular peri-embryonic 350 space of the embryo [63], where the progressive buildup of GSSG would ultimately cause embryo 351 poisoning. It will be interesting to test this hypothesis by either microinjecting permeable GSH 352 derivatives, GSH precursors or recombinant GSR-1 into the gsr-1(m-,z-) embryos [64] or by 353 transgenically expressing GSR-1 in their peri-embryonic space.

354 As GSH synthesis is restricted to the cytoplasm, the mitochondrial GSH pool is generated by an 355 active import of cytoplasmic GSH via specific carriers. In turn, it is generally accepted that mitochondrial 356 GSSG is not readily exported from this organelle, being therefore reduced by mitochondrial glutathione 357 reductase [65]. C. elegans gsr-1 mutants with transgenic GSR-1 expression restricted to cytoplasm are 358 viable, implying that mitochondrial glutathione reductase is dispensable in C. elegans, at least under 359 normal non-stress conditions and, therefore, that GSSG does not accumulate in mitochondria. As 360 mitochondrial thioredoxin reductase has been shown to substitute for mitochondrial glutathione 361 reductase in yeast and plants [58] and (Meyer AJ, personal communication), we first tested whether this 362 functional redundancy is maintained in C. elegans. However, a double mutant gsr-1(m,z-) trxr-2 (with 363 transgenic GSR-1 expression in the cytoplasm to allow development) is also viable (Figure 7B), 364 suggesting that trxr-2 is not functionally redundant with gsr-1 in worms. We next asked whether 365 mitochondrial GSR-1 is dispensable during normal non-stressed conditions but required under 366 mitochondrial stresses to provide mitochondria with enough GSH to counteract the insults. We tested 367 this possibility and demonstrate that animals lacking mitochondrial GSR-1 are as resistant as wild type 368 controls when exposed to toxic doses of paraguat or juglone, thus ruling out that mitochondrial GSR-1 is 369 needed under mitochondrial stress (Supplemental Figure 2). In this context, the dispensability of 370 mitochondrial gsr-1 (and trxr-2) in C. elegans poses some interesting hypotheses: i) mitochondrial 371 GSSG reduction is carried out by a third, yet unknown, mitochondrial reducing system in the absence of 372 gsr-1 and trxr-2; ii) GSSG can be exported from mitochondria to be reduced in the cytoplasm. Although 373 studies in yeast show that the mitochondrial matrix and intermembrane space glutathione pools are 374 maintained separately [66], a GSSG export activity has been described for the inner mitochondrial 375 membrane ATP-binding cassette transporter ATM3 (in A. thaliana) and Atm1 (in S. cerevisiae) [67]. 376 Thus, it is conceivable that GSSG generated in the mitochondrial matrix could be exported to the 377 intermembrane space by these transporters, from where it can reach the cytoplasm via porins [66] to be 378 reduced by cytoplasmic glutathione reductase as there is no GSSG reducing system in the 379 mitochondrial intermembrane space. The C. elegans abtm-1 gene is the ortholog of ATM3/Atm1 and its 380 mutation causes embryonic lethality. Instead, worms with downregulated levels of abtm-1 by RNAi are 381 viable, although have increased oxidative stress and ferric iron accumulation [68]. This lethality of abtm-382 1 mutants is presumably originated by a failure to transport Fe-S clusters to cytoplasm rather than a 383 GSSG accumulation in mitochondria, as these animals are wild type for *asr-1*. We performed *abtm-1* 384 RNAi in gsr-1(m+,z-) worms and did not observe any compound phenotype, suggesting that GSSG is 385 not accumulating in their mitochondria, even in the absence of this putative GSSG exporter (data now 386 shown); iii) alternatively, mitochondrial GSR-1 could be needed for yet unknown additional non-387 essential roles in this organelle, other than reducing GSSG. An example of a novel additional function of 388 glutathione reductase is the reduction of the redox-regulated mitoNEET protein, an outer mitochondrial 389 membrane protein that has been implicated in energy homeostasis and mitophagy [69, 70].

390 In summary, we describe here the phenotypes that the absence of glutathione reductase causes 391 in the model organism C. elegans. The availability in this organism of cytoplasmic and mitochondrial 392 thioredoxin reductase mutants combined with that of *qsr-1* now characterized offers an ideal scenario 393 where to study the respective contribution of the different compartmentalized redox systems in the 394 context of a multicellular organism. Although our results clearly show that GSR-1 is dispensable in 395 mitochondria, it is yet intriguing why gsr-1(m+,z-) mutants display mitochondria-associated phenotypes. 396 It is plausible that under limiting GSR-1 availability as in gsr-1(m+,z-) mutants, resources are derived to 397 maintain the essential cytoplasmic function thus provoking mitochondrial susceptibility. Future studies 398 aimed to specifically inactivate gsr-1 mitochondrial or cytoplasmic isoforms by CRISPR technology will 399 be instrumental to further refine their respective contribution to cell function and redox homeostasis.

#### 400 MATERIALS AND METHODS

401

#### 402 *C. elegans* strains and culture conditions

The standard methods used for culturing and maintenance of *C. elegans* were as previously described [71]. The strains used in this work are described in **Supplemental Table I**. All experiments were performed at 20°C unless otherwise noted. All VZ strains are 6x backcrossed with N2 wild type.

406

#### 407 *gsr-1* expression constructs, transgenesis and image analysis of transgenic animals.

408 For fluorescent reporter constructs, DNA fragments containing the 2.2 kb upstream promoter 409 region of the gsr-1 gene (including the first 15 nucleotides of the coding region), the 2.2 kb promoter 410 region plus the complete gsr-1 genomic ORF and the gsr-1 genomic ORF except the first exon were 411 amplified from C. elegans wild type genomic DNA and cloned into the PstI and Ball sites of the 412 pPD95.81 vector (Fire Lab C. elegans Vector Kit 1995 (unpublished)) to generate gsr-1 transcriptional and translational GFP reporters, respectively. Mutagenesis reactions (1<sup>st</sup> exon deletion and 1<sup>st</sup> and 2<sup>nd</sup> 413 414 ATG codon substitutions) were performed on the 2.2 kb promoter gsr-1 translational GFP reporter 415 construct using the QuickChange II XL Site Directed Mutagenesis Kit (Stratagene) and confirmed by 416 sequencing. For the reporter construct targeting GFP to muscle cell mitochondria directed by gsr-1 417 MTS, we amplified a 2.4 kb region of the myo-3 gene promoter with primers that included the gsr-1 MTS 418 sequence from the pBN42 plasmid ([Pmyo-3::gfp::3'-UTRunc-54] [72]) and cloned it into the BamHI site 419 of the pPD95.81 vector. Correct orientation of the insert was confirmed by sequencing. For expression 420 of wild type GSR-1, we amplified a fragment containing the gsr-1 2.2 kb promoter region, the genomic 421 ORF and 3'-UTR from C. elegans wild type genomic DNA and cloned it into the Pstl and Spel sites of 422 the pSPARK I vector (Canvax Biotech, Córdoba, Spain). The sequences of all primers used for cloning 423 are available upon request. Transgenic worms were generated by DNA microinjection as previously 424 described [73]. The pRF4 [rol-6(su1006)] [73] and coel::GFP [Punc-122::gfp] [74] plasmids were used 425 as coinjection markers at 50 ng/µl. The gsr-1 expression constructs described above were injected into 426 N2 wild type animals (either as circular plasmids or linear fragments) at varying concentrations ranging 427 from 10 to 50 ng/µl.

For image analysis of fluorescent transgenic worms, animals were mounted in a 5 μl drop of 10
mM levamisole on a 3% agarose pad covered with a coverslip. Differential interference contrast (DIC)
and fluorescence imaging was performed on a Zeiss AxioImager M2 with ApoTome Unit fluorescence
microscope and images were captured with the AxioVision 4.8 Software (Zeiss) (Figure 2A-K).

#### 433 Imaging of mitochondria, sarcomeres and measures of mitochondrial function.

434 For Figure 2M-N and 2R-T, animals were picked into 25 ml M9 buffer and were immobilized only 435 by the pressure from the cover slip. Images in M and N were taken using a Nikon H600L microscope 436 (Nikon Corporation, Tokyo, Japan) and a Nikon Digital Sight DS-Fi1 digital camera with proprietary 437 software. Images R - T were taken on a Zeiss AX10 microscope (Carl Zeiss AG, Oberkochen, 438 Germany) with an Axiocam MRC digital camera and Axiovision LE software. Images R – T were taken 439 using GFP, RFP, and triple-pass filter sets, respectively. Images in **Figure 4A** were taken in n = 60 440 animals and Figure 4B-C in n = 20 animals at young adulthood with a fixed exposure of 1 s and the 441 fluorescence was guantified using ImageJ. Corrected total animal fluorescence was calculated using 442 data from integrated density and area, however, mean fluorescence minus background fluorescence 443 produced a similar result. Data were non-parametric and were therefore analysed using a Mann 444 Whitney test and significance was set at P < 0.05.

445

#### 446 **RNA extraction and RT-PCR**

For total RNA extraction, gravid hermaphrodites were washed off the plates with M9 buffer and dissolved in 5M NaOH bleaching solution. Embryos were collected and washed several times with M9 buffer. RNA was extracted from embryos using the NucleoSpin RNA II (Macherey-Nagel) kit following the manufacturer's instructions. Total RNA was DNase-treated using the Amplification Grade Dnase I (Sigma) and 1µg of Dnase-treated RNA was reverse transcribed in a 20 µl reaction mixture. cDNA was generated using the iScript<sup>™</sup> cDNA Synthesis Kit (Biorad). 1µg of cDNA was used for *gsr-1* RT-PCR reactions using MBL-Taq DNA Polymerase (Dominion-MBL).

454

#### 455 **Recombinant protein expression, purification and glutathione reductase enzymatic activity**

456 gsr-1 cDNA from N2 wild type and gsr-1(tm3574) mutants was amplified with the forward primer 457 5'-ACTGCATATGTCTGGCGTCAAG-3' 5´and the reverse primer 458 CATGCTCGAGTTATTATTCCGGCTTCACAC -3' and cloned into the Ndel and Xhol restrictions sites of 459 the pET-15b vector (Novagen) to generate the constructs His-GSR-1b and His- $\Delta$ GSR-1b, respectively. 460 These constructs were used to transform the E. coli BL21(DE3) strain and recombinant protein 461 expression was induced by adding 0.5 mM IPTG to a 500 ml LB medium bacteria culture of 0.5-0.7 OD 462 supplemented with 0.1 mg/ml ampicillin and further incubating the cells at 25°C and 200 rpm during 5 463 hours. Cells were collected by centrifugation, immediately resuspended in 25 ml Tris-HCl 20 mM pH 8, 464 0.1 M NaCl buffer supplemented with 15 mg de lysozyme and 30 mg de Dnase I. After 5 min incubation

465 at room temperature the preparation was sonicated for 30 min on ice and the cell free extract was 466 obtained by centrifugation at 10000 x g during 30 min at 4°C. Recombinant His-GSR-1b and His- $\Delta$ GSR-467 1b proteins were purified from the cell free extract using a TALON column (Clontech) equilibrated with 468 Tris-HCl 20 mM pH 8, 0.1 M NaCl buffer and eluted with 100 mM imidazol. Finally, the purified proteins 469 were dialyzed against the same buffer and concentrated using Centricon YM-10 filter devices 470 (Millipore). The glutathione reductase enzymatic activity was determined as previously described [75]. 471 Briefly, a standard assay mixture (0.5 ml) containing potassium phosphate buffer 0.1 M, pH 7.2, EDTA 1 472 mM, NADPH 200 μM, with different concentrations of His-GSR-1b and His-ΔGSR-1b was prepared and 473 the reaction was initiated by addition of 0.5 mM GSSG. The decrease of absorbance at 340 nm 474 (indicating NADPH consumption) was recorded spectrophotometrically at 25°C.

475

#### 476 Embryonic cell lineage analysis

477 For embryonic cell lineage analysis, 4D-microscopy was carried out using standard live-animal 478 mounting techniques on a Leica DM6000 microscope fitted with DIC optics. The use of DIC optics 479 allows cells tracing without using any dye or fluorescent marker that might alter the cell cycle 480 progression. Embryonic cell lineage was determined as described [76]. In summary, gravid 481 hermaphrodites were dissected and 2- to 4-cell stage embryos were mounted on 4% agar pads in 482 water, and sealed with Vaseline. Imaging was performed at 25°C. The multi-focal time-lapse 483 microscopy of the samples was controlled with the open source software Micro-manager (www.micro-484 manager.org). Pictures on 30 focal planes (1micron/section) were taken every 30 seconds for 12 h. 485 Embryo lineages were analyzed with the software SimiBiocel (SIMI GmbH, www.simi.com).

486

#### 487 **Brood size determination**

L4 larvae were singled onto OP50 seeded plates, allowed to lay eggs at 20°C and transferred to new plates every 12 hours until the animals stopped reproducing. Eggs were counted and then incubated further at 20°C to quantify adult progeny four days later.

491

#### 492 Longevity assay

Lifespan assays were performed at 25°C as previously described [77] with slight modifications. Tightly synchronized embryos from gravid adult hermaphrodites were allowed to develop through the L4 larval stage and then transferred to fresh NMG plates in groups of 20 worms per plate for a total of 80 individuals per experiment. The day animals reached the L4 larval stage was used as t = 0. Animals

497 were transferred to fresh plates daily until progeny production ceased and after that they were 498 transferred every second to third day but monitored daily for dead animals. Nematodes that did not 499 respond to gentle prodding and displayed no pharyngeal pumping were scored as dead. Animals that 500 crawled off the plate or died due to internal hatching or extruded gonad were censored and incorporated 501 as such into the data set.

502

#### 503 Paraquat and Juglone acute treatment

Plates containing the appropriate amount of the respective chemical were prepared freshly prior the experiment. To minimize the number of animals crawling off the agar, OP50 bacteria (from seeded plates) was directly spread in the center of the plates using an inoculating loop. L4 larvae were transferred to plates and survival was determined after 16 hours incubation at 20°C. Nematodes that did not respond to gentle prodding and displayed no pharyngeal pumping were scored as dead. Animals that crawled off the plate were censored.

510

#### 511 Embryonic/larval arrest assays

512 10 gravid *trxr-1(sv47); gsr-1(tm3574)/qC1::rfp* hermaphrodites were allowed to lay eggs during 3 513 hours at 20°C and subsequently removed. Embryos were counted and then allowed to develop for 4 514 days at 20°C after which non RFP *trxr-1(sv47); gsr-1(m+,z-)* adults, larvae and unhatched embryos 515 were quantified.

516

# 517 SKN-1 dependent gene expression quantification and nuclear translocation of SKN-1 fluorescent 518 reporter

519 Worms at their third day of adulthood expressing dvEx166 [Pgst-4::gfp] and svEx741 [Pgcs-1::gfp] 520 extrachromosomal arrays were mounted in a 5 ml drop of 10 mM levamisole on a 3% agarose pad 521 covered with a coverslip. Fluorescence imaging was performed on a Olympus BX61 fluorescence 522 microscope equipped with a Olympus DP72 camera and images were captured with the 523 CellSensDimension 1.12 Software (Olympus). All micrographs were taken with identical image capture 524 settings and quantification of GFP expression (measured as the fluorescence mean of 25 worms 525 divided by the selected area and normalized by the background adjacent to the selected worm in the 526 same image) was performed using the ImageJ Software (NIH). When needed, equal adjustment of 527 brightness and contrast on control and matched problem images was implemented using Adobe 528 Photoshop 10 Software (Adobe Systems). To visualize SKN-1B/C::GFP nuclear translocation, young

adults were washed from plates, anesthetized with 1 mM levamisole and mounted on 2% agarose pads. Visualization and imaging was performed using an Olympus IX81 automated inverted microscope and Slidebook (version 5.0) software. SKN-1B/C::GFP localization quantification, the percent intestinal SKN-1 nuclear localization was categorically scored as follows: none: no localization, low: posterior or anterior intestinal localization, medium: posterior and anterior intestinal localization, high: localization throughout the entire intestine [78].

535

#### 536 Mitochondrial related phenotypes

537 To determine if gsr-1 deficiency had any effect on mitochondria related phentoypes, age-538 synchronized, young adult animals (n = 60) were assessed for mitochondrial networking in gsr-1(m+z-); 539 ccls4251 [Pmyo-3::mito::gfp] versus control ccls4251 [Pmyo-3::mito::gfp] animals. There was 540 significantly greater mitochondrial fragmentation in the gsr-1 animals than controls at 20°C (P < 0.001) 541 and 25°C (P <0.05). Determination of sarcomere structure in gsr-1(m+,z-); jls01[Pmyo-3::myo-3::gfp] 542 were compared to control *jls01[Pmyo-3::myo-3::gfp]* animals (n = 20 per group) both at 20°C and at 543 25°C. There was no significant difference between groups (P >0.05). For mitochondrial content 544 determination, we loaded animals with JC-10, a dye that accumulates proportionally to a negative 545 mitochondrial membrane potential as previously described [79]. Wild type control animals had 546 significantly greater fluorescence of the gut mitochondria than did gsr-1(m+,z-) animals (P <0.001). 547 Images were taken with a fixed exposure of 1 s on the Nikon H600L microscope and the fluorescence 548 was quantified using ImageJ. Corrected total animal fluorescence was calculated as follows: Integrated 549 density – area of the worm x mean fluorescence of three background readings. Data were analyzed 550 using a Mann-Whitney test as data were non-parametric.

551

#### 552 **Embryonic chromatin dynamics analysis.**

553 BN323 animals were maintained at 16°C and shifted to 25°C 2 hours prior to microscopy. 554 Heterozygous and homozygous *gsr-1* mutants were dissected and early embryos were mounted 555 together in M9 buffer on 2% agarose pads. Coverslips were placed over the embryos and sealed with 556 VALAP (1:1:1 mixture of Vaseline, lanolin and paraffin). Confocal epifluorescence and DIC images from 557 three focal planes were acquired at 25°C every 10 minutes on a Nikon A1R microscope through a Plan 558 Apo VC 60x/1.4 objective (Nikon, Tokyo, Japan) using a pinhole of 39.7 μm (1.2 Airy Units).

559

560

#### 560 **ACKNOWLEDGMENTS**

561

562 Some strains were provided by the CGC, which is funded by NIH Office of Research 563 Infrastructure Programs (P40 OD010440) and by the Japanese National Bioresource Project. We 564 thank Cristina Cecchi, Amir Shapir, Paul Sternberg, Bart Braeckman, Chris Link, Simon Tuck, 565 Keith Blackwell, José López-Barneo and LivOn Labs for strains and chemicals, Katie McCallum 566 and Danielle Garsin for their help with *skn-1* experiments and Elizabeth Veal and Michel Toledano 567 for critical reading of the manuscript. Prof. Rafael Fernández-Chacón is deeply acknowledged for 568 his continuous support. AMV was supported by grants from the Spanish Ministry of Economy and 569 Competitiveness (BFU2015-64408-P) and the Instituto de Salud Carlos III (PI11/00072, 570 cofinanced by the Fondo Social Europeo) and is a member of the GENIE and EU-ROS Cost 571 Action of the European Union. NJS was supported by a grant from the US National Institutes of 572 Health National Institute for Arthritis and Musculoskeletal and Skin Diseases (AR-054342). CG 573 was funded by a Doctoral Training Studentship provided by the University of Nottingham. PA was 574 supported by the Spanish Ministry of Economy and Competitiveness (BFU2013-42709P). JC is a 575 member of the GENIE Cost action and was funded by Rioja Salud Foundation. 576

577 **FIGURES** 

578

579 Figure 1: gsr-1 mRNA and protein analysis. A) Schematic representation of the two main gsr-1 580 mRNA variants. Boxes represent exons and lines show spliced introns. White boxes indicate 5'-UTR 581 and 3'-UTR, respectively, and grey boxes indicate the ORF. The region of the gsr-1 mRNA missing 582 in the gsr-1(tm3574) deletion allele is underlined. B) Protein domain organization of GSR-1a and 583 GSR-1b isoforms as well as that of the truncated  $\Delta$ GSR-1 protein resulting from translation of the 584 gsr-1(tm3574) allele. Numbers indicate the amino acid residues flanking the different protein domains, based on the domain organization reported for the human GSR orthologue [80]. C) 585 586 Sequence alignment of C. elegans GSR-1 with the corresponding human and zebrafish GSR 587 orthologues. The red rectangle indicates the conserved redox active site CVNVGC. The yellow 588 boxes highlight the residues involved in GSSG binding [81]. The NADPH and FAD binding sites are 589 underlined. Identical residues are shown in black. The numbers on the left indicate the respective 590 amino acid residue. The different domains boundaries are indicated above the sequences and are 591 inferred from comparison with those reported for human GSR protein [80]. D) Sequence of the GSR-592 1a mitochondrial targeting sequence (MTS). Amino acid residues matching the two matrix protease 593 model (in blue. X, any amino acid residue) [41] are highlighted in red. The MTS cleavage site is 594 indicated by the upside-down arrow. The residues encoded by *qsr-1* first and second exons are 595 separated by an hyphen.

596

597 Figure 2: GSR-1 tissue, cellular and subcellular expression analysis. A) Scheme of the GSR-1 598 translational GFP fusion construct used to determine expression from gsr-1 upstream promoter. 599 Exons are in grey boxes and introns in white boxes. B-E) Differential interference contrast and B'-E') 600 fluorescence images of transgenic worms expressing the Pgsr-1::gsr-1::gfp construct: ai, anterior 601 intestine; coe, coelomocytes; gsc, gonad sheet cell; hy, hypodermis; ph, pharynx; rc, rectal cells; sp, 602 spermatheca; tn, tail neurons; uc, uterine cells; vmc, vulva muscle cells. F) Scheme of the GSR-1 603 translational GFP fusion construct used to determine expression from gsr-1 first intron. Exons are in 604 grey boxes and introns in white boxes. G-K) Differential interference contrast and G'-K') 605 fluorescence images of transgenic worms expressing the *gsr-1 first intron::gsr-1::gfp* construct: ai, 606 anterior intestine; coe, coelomocytes; ec, excretory cell; hn, head neurons; int, intestine; pi, posterior 607 intestine. L) Schemes of the GSR-1 translational GFP fusion constructs used to determine 608 subcellular localization of GSR-1a and GSR-1b isoforms. Red X indicates the mutated ATG codon. 609 Exons are in grey boxes and introns in white boxes. M) Dotted fluorescence labeling in pharynx of 610 transgenic worms expressing the *Pgsr-1::gsr-1::gfp* construct with the second ATG codon mutated.

611 N) Diffuse cytoplasmic fluorescence labeling in pharynx of transgenic worms expressing the 612 construct the Pgsr-1::gsr-1::gfp construct with the first ATG codon mutated. Q) Scheme of the 613 construct used to determine GSR-1a MTS functionality when fused to GFP under the control of the 614 *myo-3* muscle promoter. GSR-1a MTS is shown in a grey box. Fluorescence images of muscle cells 615 of transgenic worms expressing R) the Pmyo-3::MTS<sub>ast-1</sub>::gfp construct, T) the Pmyo-3::tomm-616 20::mrfp construct and S) merged picture showing colocalization in muscle tubular mitochondria. 617 Scale bar 20 µm in all images except 100 µm in B. Note Images B-K and M-T were taken with 618 different microscopes and settings (see Materials and Methods).

619

620 Figure 3: Glutathione reductase enzymatic activity and phenotypes of qsr-1(m+,z-) mutants. 621 A) GSSG reductase enzymatic activity of different concentrations of recombinant His-GSR-1b 622 (straight lines) and His- $\Delta$ GSR-1b (dashed lines) in the presence of 0.5 mM GSSG and 0.25 mM 623 NADPH. B) The longevity was assayed at 20°C in *E. coli* OP50. Kaplan-Meier plot was used to show 624 the fraction of animals that survive over time. Longevity was performed twice, obtaining similar 625 results, and the composite data is shown. The survival rate of gsr-1(m+,z-) animals was compared to 626 that of wild type using the log-rank (Mantel-Cox) test and the differences were found significant (\*\*\* 627 p < 0.001). C) Sensitivity of gsr-1(m+,z-) animals to acute paraguat and juglone exposure. Data are 628 the mean  $\pm$  SD of four independent experiments with three biological replicates each (n  $\ge$  200; \*\*\* 629 p < 0.001, by unpaired, two-tail Student *t*-test). **D)** Percentage of developmental stages of single and 630 double mutants combining the trxr-1(sv47) and the gsr-1(m+,z-) mutation. Data are the mean of 631 three independent experiments (n = total number of individuals scored). E) Induction of GST-4 and 632 GCS-1 transcriptional GFP reporters and percentage of nuclear localization of a SKN-1B/C 633 translational GFP reporter in a gsr-1(m+,z-) background. For GST-4 and GCS-1 reporters, the data 634 are the mean  $\pm$  SD of 25 individuals for each genotype (\*\*\* p<0.001, \*\* p<0.01 by unpaired, two-tail 635 Student *t*-test). For SKN-1 reporter the data are from two independent experiments (n = total number 636 of individuals scored).

637

Figure 4: *gsr-1(m+,z-)* mutants display mitochondrial phenotypes. A) Increased mitochondrial fragmentation is observed in *gsr-1(m+,z-)* animals expressing the mitochondrial reporter *ccls4251 [Pmyo-3::mito::gfp]* in muscle cells. Data are the total  $\pm$  SEM of three independent experiments with 20 animals per experiment at 20°C and 25°C (n = 60 total at each temperature) (\* p<0.05, \*\*\* *p*<0.001, by a Mann Whitney test). Representative images of each genotype are shown. Scale bar 20 µm. B) Muscle cell integrity is not affected in *gsr-1(m+,z-)* worms as demonstrated by the maintenance of sarcomere myotubular structure visualized using the *jls01 [Pmyo-3::myo-3::gfp]* 

645 reporter. Data are the total ± SEM of one experiment with 20 animals (p>0.05 by a Mann Whitney 646 test). Representative images of each genotype are shown. Scale bar 20 µm. C) Quantification of the 647 total mitochondrial content measured by JC-10 dye incorporation. Data are the total ± SEM of 20 648 individuals per genotype (\*\*\* p < 0.001 by a Mann Whitney test). Representative images of each 649 genotype are shown. Scale bar 100  $\mu$ m. **D)** Mitochondrial UPR is induced in gsr-1(m+,z-) worms 650 expressing the reporter *zcls13* [*Phsp-6::gfp*]. Data are the mean  $\pm$  SD of 40 individuals per genotype 651 (\*\*\* p<0.001, by unpaired two-tail Student *t*-test). Representative images of each genotype are 652 shown. Scale bar 200 µm.

653

654 Figure 5: Identification of the cell-stage time of arrest of gsr-1(m-,z-) embryos and 655 determination of GSR-1 expression initiation. A-C) Differential interference contrast still 656 images of developing wild type and *asr-1(m-.z-)* embryos analyzed by 4D-microscopy. Three 657 representative independent experiments are provided, illustrating the cases at which gsr-1(m-z-)658 embryos arrest at the earliest, the latest and an intermediate cell-stage. A) Arrest at 17 cell-659 stage; B) Arrest at 33 cell-stage; C) Arrest at 103 cell-stage. Gsr-1 in the images refers to gsr-660 1(m,z) embryos. D) Fluorescence and differential interference contrast still images of 661 developing embryos from transgenic animals expressing the transcriptional construct Pasr-662 1::gfp. Three independent embryos are shown. Gsr-1 expression begins approximately at 100 663 min since embryos start developing. In these initial stages, gsr-1 stronger expression is found in 664 E blastomere descendants (intestine precursors) and MS blastomeres descendants (neurons 665 and mesodermic tissues precursors). Scale bar 10 µm. All analyses were performed at 25°C.

666

667 Figure 6: Cell division delay and progressive perinuclear localization of interphasic chromatin 668 in gsr-1(m-,z-) embryos. A) Cell lineage timing analysis of the EMS blastomere descendants of the 669 gsr-1(m-,z-) embryo depicted in Figure 5B compared to that of wild type embryos. For complete cell 670 lineage analysis, see Supplemental Figure 1B. B) gsr-1(m+,z+) and gsr-1(m-,z-) embryos 671 expressing mCherry::HIS-58 (magenta) and GFP::TBB-2 were observed by live confocal microscopy. 672 Time is indicated in hours: minutes from beginning of recording (see Movie 5). At 2:20 and more 673 evidently at 5:00 chromosomes are condensed at the nuclear periphery of qsr-1(m,z) embryos. 674 Scale bar 10 µm.

675

Figure 7: Rescue of *gsr-1(tm3574)* mutants embryonic lethality. A) The *gsr-1(tm3574)*embryonic lethal phenotype is rescued when GSR-1 activity is restored in the cytoplasm (GSR-1b,
transgenes *vzEx145* and *vzEx147*) but not in mitochondria (GSR-1a, transgene *vzEx158*). The

679 dashed bar in *gsr-1; vzEx105 [Pgsr-1::gsr-1]* animals indicates the fraction of viable, maternally 680 rescued, non-transgenic progeny. Data are the mean  $\pm$  SD of the progeny from at least 10 animals of 681 each genotype. **B)** Viable progeny from *gsr-1; trxr-2* double mutant worms expressing GSR-1b::GFP 682 in cytoplasm. Data are the mean  $\pm$  SD of the progeny from at least 10 animals of each genotype.

683

684

#### 685 **MOVIES**

686

687 **Movies 1-3: Movie recordings of** *C. elegans* wild type vs *gsr-1(m-,z-)* embryos. Developmental 688 arrest of the *gsr-1(m-,z-)* embryo occurs at 17-cell stage (movie 1), 33-cell stage (movie 2) and 103-689 cell stage (movie 3). Records were performed at 25°C over 12h as explained in materials and 690 methods. Movie 1 corresponds to **Figure 5A**, Movie 2 corresponds to **Figure 5B** and Movie 3 691 corresponds to **Figure 5C** 

692

693 Movie 4: Movie recording of *C. elegans* wild type vs gsr-1(m-,z-) embryos stained with 694 rhodamine 6G. The day before the recording wild type and gsr-1(m-,z-) L4 worms were placed on 695 NGM agar plates supplemented with rhodamine 6G at a final concentration of 2.5 ug/ml and 696 incubated at 20°C. Two hours before the recording plates were transferred to 25°C. Recordings were 697 performed with a NIKON A1R confocal microscope through a Plan Apo VC 60x/1.4 objective. Time 698 is indicated in hours : minutes from beginning of recording.

699

700 Movie 5: Movie recording of *C. elegans gsr-1(m+,z+)* and *gsr-1(m-,z-)* embryos expressing

701 mCherry::HIS-58 (magenta) and GFP::TBB-2. Time is indicated in hours:minutes from beginning of
 702 recording.Corresponds to Figure 6B.

- 703
- 704

#### 704 **REFERENCES**

- 705
- [1] Hanschmann, E. M.; Godoy, J. R.; Berndt, C.; Hudemann, C.; Lillig, C. H. Thioredoxins,
  glutaredoxins, and peroxiredoxins--molecular mechanisms and health significance: from
  cofactors to antioxidants to redox signaling. *Antioxid Redox Signal* 19:1539-1605; 2013.
- 709 [2] Meyer, Y.; Buchanan, B. B.; Vignols, F.; Reichheld, J. P. Thioredoxins and glutaredoxins: 710 unifying elements in redev biology. *Annu Bay Const* **42**:225–267: 2000
- unifying elements in redox biology. *Annu Rev Genet* 43:335-367; 2009.
- [3] Xiong, Y.; Uys, J. D.; Tew, K. D.; Townsend, D. M. S-glutathionylation: from molecular
  mechanisms to health outcomes. *Antioxid Redox Signal* 15:233-270; 2011.
- 713 [4] Fahey, R. C.; Sundquist, A. R. Evolution of glutathione metabolism. *Adv Enzymol Relat*714 *Areas Mol Biol* 64:1-53; 1991.
- 715 [5] Grant, C. M.; MacIver, F. H.; Dawes, I. W. Glutathione is an essential metabolite required
- for resistance to oxidative stress in the yeast Saccharomyces cerevisiae. *Curr Genet* 29:511515; 1996.
- [6] Romero-Aristizabal, C.; Marks, D. S.; Fontana, W.; Apfeld, J. Regulated spatial
  organization and sensitivity of cytosolic protein oxidation in Caenorhabditis elegans. *Nat Commun* 5:5020; 2014.
- [7] Fraser, J. A.; Kansagra, P.; Kotecki, C.; Saunders, R. D.; McLellan, L. I. The modifier
  subunit of Drosophila glutamate-cysteine ligase regulates catalytic activity by covalent and
  noncovalent interactions and influences glutathione homeostasis in vivo. *J Biol Chem*278:46369-46377; 2003.
- [8] Cairns, N. G.; Pasternak, M.; Wachter, A.; Cobbett, C. S.; Meyer, A. J. Maturation of
  arabidopsis seeds is dependent on glutathione biosynthesis within the embryo. *Plant Physiol*141:446-455; 2006.
- [9] Dalton, T. P.; Dieter, M. Z.; Yang, Y.; Shertzer, H. G.; Nebert, D. W. Knockout of the mouse
  glutamate cysteine ligase catalytic subunit (Gclc) gene: embryonic lethal when homozygous,
  and proposed model for moderate glutathione deficiency when heterozygous. *Biochem Biophys Res Commun* 279:324-329; 2000.
- [10] Pasternak, M.; Lim, B.; Wirtz, M.; Hell, R.; Cobbett, C. S.; Meyer, A. J. Restricting
  glutathione biosynthesis to the cytosol is sufficient for normal plant development. *Plant J* **53**:999-1012; 2008.
- [11] Winkler, A.; Njalsson, R.; Carlsson, K.; Elgadi, A.; Rozell, B.; Abraham, L.; Ercal, N.; Shi, Z.
  Z.; Lieberman, M. W.; Larsson, A.; Norgren, S. Glutathione is essential for early
  embryogenesis--analysis of a glutathione synthetase knockout mouse. *Biochem Biophys Res Commun* **412**:121–126:2011
- 738 *Commun* **412:**121-126; 2011.
- [12] Grant, C. M.; MacIver, F. H.; Dawes, I. W. Glutathione synthetase is dispensable for
  growth under both normal and oxidative stress conditions in the yeast Saccharomyces
  cerevisiae due to an accumulation of the dipeptide gamma-glutamylcysteine. *Mol Biol Cell*8:1699-1707; 1997.
- 743 [13] Logan-Garbisch, T.; Bortolazzo, A.; Luu, P.; Ford, A.; Do, D.; Khodabakhshi, P.; French, R.
- L. Developmental ethanol exposure leads to dysregulation of lipid metabolism and oxidative
   stress in Drosophila. *G3 (Bethesda)* 5:49-59; 2014.
- 746 [14] Kanzok, S. M.; Schirmer, R. H.; Turbachova, I.; Iozef, R.; Becker, K. The thioredoxin
- 747 system of the malaria parasite Plasmodium falciparum. Glutathione reduction revisited. *J Biol*
- 748 *Chem* **275**:40180-40186; 2000.

- 749 [15] Morgan, B.; Ezerina, D.; Amoako, T. N.; Riemer, J.; Seedorf, M.; Dick, T. P. Multiple
- glutathione disulfide removal pathways mediate cytosolic redox homeostasis. *Nat Chem Biol*9:119-125; 2013.
- [16] Eriksson, S.; Prigge, J. R.; Talago, E. A.; Arner, E. S.; Schmidt, E. E. Dietary methionine can
  sustain cytosolic redox homeostasis in the mouse liver. *Nat Commun* 6:6479; 2015.
- 754 [17] Lee, J.; Dawes, I. W.; Roe, J. H. Isolation, expression, and regulation of the pgr1(+) gene
- encoding glutathione reductase absolutely required for the growth of Schizosaccharomyces
  pombe. *J Biol Chem* 272:23042-23049; 1997.
- 757 [18] Pastrana-Mena, R.; Dinglasan, R. R.; Franke-Fayard, B.; Vega-Rodriguez, J.; Fuentes-
- 758 Caraballo, M.; Baerga-Ortiz, A.; Coppens, I.; Jacobs-Lorena, M.; Janse, C. J.; Serrano, A. E.
- 759 Glutathione reductase-null malaria parasites have normal blood stage growth but arrest 760 during development in the mosquito. *J Biol Chem*; 2010.
- 761 [19] Tzafrir, I.; Pena-Muralla, R.; Dickerman, A.; Berg, M.; Rogers, R.; Hutchens, S.; Sweeney,
- 762 T. C.; McElver, J.; Aux, G.; Patton, D.; Meinke, D. Identification of genes required for embryo 763 development in Arabidopsis. *Plant Physiol* **135**:1206-1220; 2004.
- 764 [20] Kanzok, S. M.; Fechner, A.; Bauer, H.; Ulschmid, J. K.; Muller, H. M.; Botella-Munoz, J.;
- 765 Schneuwly, S.; Schirmer, R.; Becker, K. Substitution of the thioredoxin system for glutathione
- reductase in Drosophila melanogaster. *Science* **291**:643-646; 2001.
- 767 [21] Meyer, Y.; Belin, C.; Delorme-Hinoux, V.; Reichheld, J. P.; Riondet, C. Thioredoxin and
  768 Glutaredoxin Systems in Plants: Molecular Mechanisms, Cross Talks and Functional
  769 Significance. *Antioxid Redox Signal*; 2012.
- 770 [22] Kelner, M. J.; Montoya, M. A. Structural organization of the human glutathione reductase
- gene: determination of correct cDNA sequence and identification of a mitochondrial leader
  sequence. *Biochem Biophys Res Commun* 269:366-368; 2000.
- [23] Tamura, T.; McMicken, H. W.; Smith, C. V.; Hansen, T. N. Gene structure for mouse
  glutathione reductase, including a putative mitochondrial targeting signal. *Biochem Biophys Res Commun* 237:419-422; 1997.
- 776 [24] Outten, C. E.; Culotta, V. C. Alternative start sites in the Saccharomyces cerevisiae GLR1
- gene are responsible for mitochondrial and cytosolic isoforms of glutathione reductase. *J Biol Chem* 279:7785-7791; 2004.
- [25] Song, J. Y.; Cha, J.; Lee, J.; Roe, J. H. Glutathione reductase and a mitochondrial
  thioredoxin play overlapping roles in maintaining iron-sulfur enzymes in fission yeast. *Eukaryot Cell* 5:1857-1865; 2006.
- 782 [26] Prinz, W. A.; Aslund, F.; Holmgren, A.; Beckwith, J. The role of the thioredoxin and
- glutaredoxin pathways in reducing protein disulfide bonds in the Escherichia coli cytoplasm. *I Biol Chem* 272:15661-15667; 1997.
- [27] Kumar, C.; Igbaria, A.; D'Autreaux, B.; Planson, A. G.; Junot, C.; Godat, E.; Bachhawat, A.
  K.; Delaunay-Moisan, A.; Toledano, M. B. Glutathione revisited: a vital function in iron
  metabolism and ancillary role in thiol-redox control. *EMBO J* 30:2044-2056; 2011.
- 788 [28] Toledano, M. B.; Delaunay-Moisan, A.; Outten, C. E.; Igbaria, A. Functions and cellular 789 compartmentation of the thioredoxin and glutathione pathways in yeast. *Antioxid Redox*
- *Signal* **18**:1699-1711; 2013.
- 791 [29] Gostimskaya, I.; Grant, C. M. Yeast mitochondrial glutathione is an essential antioxidant
- with mitochondrial thioredoxin providing a back-up system. *Free Radic Biol Med*; 2016.
- 793 [30] Rogers, L. K.; Tamura, T.; Rogers, B. J.; Welty, S. E.; Hansen, T. N.; Smith, C. V. Analyses of
- 794 glutathione reductase hypomorphic mice indicate a genetic knockout. Toxicol Sci 82:367-
- 795 373; 2004.

- 796 [31] Matsui, M.; Oshima, M.; Oshima, H.; Takaku, K.; Maruyama, T.; Yodoi, J.; Taketo, M. M.
- Farly embryonic lethality caused by targeted disruption of the mouse thioredoxin gene. *Dev Biol* 178:179-185; 1996.
- 799 [32] Conrad, M.; Jakupoglu, C.; Moreno, S. G.; Lippl, S.; Banjac, A.; Schneider, M.; Beck, H.;
- 800 Hatzopoulos, A. K.; Just, U.; Sinowatz, F.; Schmahl, W.; Chien, K. R.; Wurst, W.; Bornkamm, G.
- W.; Brielmeier, M. Essential role for mitochondrial thioredoxin reductase in hematopoiesis,
  heart development, and heart function. *Mol Cell Biol* 24:9414-9423; 2004.
- [33] Jakupoglu, C.; Przemeck, G. K.; Schneider, M.; Moreno, S. G.; Mayr, N.; Hatzopoulos, A. K.;
  de Angelis, M. H.; Wurst, W.; Bornkamm, G. W.; Brielmeier, M.; Conrad, M. Cytoplasmic
  thioredoxin reductase is essential for embryogenesis but dispensable for cardiac
  development. *Mol Cell Biol* 25:1980-1988; 2005.
- 807 [34] Nonn, L.; Williams, R. R.; Erickson, R. P.; Powis, G. The absence of mitochondrial 808 thioredoxin 2 causes massive apoptosis, exencephaly, and early embryonic lethality in 809 homozygous mice. *Mol Cell Biol* **23**:916-922; 2003.
- 810 [35] Mandal, P. K.; Schneider, M.; Kolle, P.; Kuhlencordt, P.; Forster, H.; Beck, H.; Bornkamm,
- 811 G. W.; Conrad, M. Loss of thioredoxin reductase 1 renders tumors highly susceptible to 812 pharmacologic glutathione deprivation. *Cancer Res* **70**:9505-9514; 2010.
- 813 [36] Suvorova, E. S.; Lucas, O.; Weisend, C. M.; Rollins, M. F.; Merrill, G. F.; Capecchi, M. R.;
- Schmidt, E. E. Cytoprotective Nrf2 pathway is induced in chronically txnrd 1-deficient hepatocytes. *PLoS One* **4**:e6158; 2009.
- 816 [37] Stenvall, J.; Fierro-Gonzalez, J. C.; Swoboda, P.; Saamarthy, K.; Cheng, Q.; Cacho-Valadez,
- 817 B.; Arner, E. S.; Persson, O. P.; Miranda-Vizuete, A.; Tuck, S. Selenoprotein TRXR-1 and GSR-1
- are essential for removal of old cuticle during molting in Caenorhabditis elegans. *Proc Natl Acad Sci U S A* 108:1064-1069; 2011.
- [38] Cacho-Valadez, B.; Munoz-Lobato, F.; Pedrajas, J. R.; Cabello, J.; Fierro-Gonzalez, J. C.;
  Navas, P.; Swoboda, P.; Link, C. D.; Miranda-Vizuete, A. The characterization of the
  Caenorhabditis elegans mitochondrial thioredoxin system uncovers an unexpected
  protective role of thioredoxin reductase 2 in beta-amyloid peptide toxicity. *Antioxid Redox Signal* 16:1384-1400; 2012.
- 825 [39] Luersen, K.; Stegehake, D.; Daniel, J.; Drescher, M.; Ajonina, I.; Ajonina, C.; Hertel, P.; 826 Woltersdorf, C.; Liebau, E. The glutathione reductase GSR-1 determines stress tolerance and
- 827 longevity in Caenorhabditis elegans. *PLoS One* **8**:e60731; 2013.
- 828 [40] Habib, S. J.; Neupert, W.; Rapaport, D. Analysis and prediction of mitochondrial 829 targeting signals. *Methods Cell Biol* **80**:761-781; 2007.
- 830 [41] Hendrick, J. P.; Hodges, P. E.; Rosenberg, L. E. Survey of amino-terminal proteolytic 831 cleavage sites in mitochondrial precursor proteins: leader peptides cleaved by two matrix
- b) a cleavage sites in intochondrial precursor proteins: leader peptides cleaved by two matrix
   a) proteases share a three-amino acid motif. *Proc Natl Acad Sci U S A* 86:4056-4060; 1989.
- 833 [42] Fuxman Bass, J. I.; Tamburino, A. M.; Mori, A.; Beittel, N.; Weirauch, M. T.; Reece-Hoyes,
- J. S.; Walhout, A. J. Transcription factor binding to Caenorhabditis elegans first introns reveals lack of redundancy with gene promoters. *Nucleic Acids Res* **42**:153-162; 2014.
- 836 [43] Petit, E.; Michelet, X.; Rauch, C.; Bertrand-Michel, J.; Terce, F.; Legouis, R.; Morel, F.
- 837 Glutathione transferases kappa 1 and kappa 2 localize in peroxisomes and mitochondria,
- respectively, and are involved in lipid metabolism and respiration in Caenorhabditis elegans. *FEBS* / 276:5030-5040; 2009.
- 840 [44] Edgley, M. L.; Baillie, D. L.; Riddle, D. L.; Rose, A. M. Genetic balancers. *WormBook*:1-32; 841 2006.

- 842 [45] Wang, J.; Robida-Stubbs, S.; Tullet, J. M.; Rual, J. F.; Vidal, M.; Blackwell, T. K. RNAi 843 screening implicates a SKN-1-dependent transcriptional response in stress resistance and 844 longevity deriving from translation inhibition. *PLoS Genet* **6**; 2010.
- 845 [46] Yoneda, T.; Benedetti, C.; Urano, F.; Clark, S. G.; Harding, H. P.; Ron, D. Compartment-846 specific perturbation of protein handling activates genes encoding mitochondrial 847 chaperones. *J Cell Sci* **117**:4055-4066; 2004.
- 848 [47] Ahringer, J. Embryonic tissue differentiation in Caenorhabditis elegans requires dif-1, a 849 gene homologous to mitochondrial solute carriers. *EMBO J* **14**:2307-2316; 1995.
- [48] Asencio, C.; Navas, P.; Cabello, J.; Schnabel, R.; Cypser, J. R.; Johnson, T. E.; RodriguezAguilera, J. C. Coenzyme Q supports distinct developmental processes in Caenorhabditis
  elegans. *Mech Ageing Dev* 130:145-153; 2009.
- [49] Kruger, A. V.; Jelier, R.; Dzyubachyk, O.; Zimmerman, T.; Meijering, E.; Lehner, B.
  Comprehensive single cell-resolution analysis of the role of chromatin regulators in early C.
- elegans embryogenesis. *Dev Biol* **398**:153-162; 2015.
- 856 [50] Garcia-Gimenez, J. L.; Olaso, G.; Hake, S. B.; Bonisch, C.; Wiedemann, S. M.; Markovic, J.;
- Basi, F.; Gimeno, A.; Perez-Quilis, C.; Palacios, O.; Capdevila, M.; Vina, J.; Pallardo, F. V. Histone
  h3 glutathionylation in proliferating mammalian cells destabilizes nucleosomal structure.
- 859 Antioxid Redox Sianal **19:**1305-1320: 2013.
- [51] Badrinath, A. S.; White, J. G. Contrasting patterns of mitochondrial redistribution in the
  early lineages of Caenorhabditis elegans and Acrobeloides sp. PS1146. *Dev Biol* 258:70-75;
  2003.
- 863 [52] Cascella, R.; Evangelisti, E.; Zampagni, M.; Becatti, M.; D'Adamio, G.; Goti, A.; Liguri, G.;
- Fiorillo, C.; Cecchi, C. S-linolenoyl glutathione intake extends life-span and stress resistance via Sir-2.1 upregulation in Caenorhabditis elegans. *Free Radic Biol Med* **73**:127-135; 2014.
- 866 [53] Shibamura, A.; Ikeda, T.; Nishikawa, Y. A method for oral administration of hydrophilic
- substances to Caenorhabditis elegans: Effects of oral supplementation with antioxidants on
  the nematode lifespan. *Mech Ageing Dev* 130:652-655; 2009.
- [54] Levy, E. J.; Anderson, M. E.; Meister, A. Transport of glutathione diethyl ester into
  human cells. *Proc Natl Acad Sci U S A* **90**:9171-9175; 1993.
- [55] Yang, W.; Hekimi, S. A mitochondrial superoxide signal triggers increased longevity in
  Caenorhabditis elegans. *PLoS Biol* 8:e1000556; 2010.
- 873 [56] Edwards, C.; Canfield, J.; Copes, N.; Brito, A.; Rehan, M.; Lipps, D.; Brunquell, J.;
- Westerheide, S. D.; Bradshaw, P. C. Mechanisms of amino acid-mediated lifespan extension in
  Caenorhabditis elegans. *BMC Genet* 16:8; 2015.
- 876 [57] Wharton, D. Nematode egg-shells. *Parasitology* **81:**447-463; 1980.
- 877 [58] Trotter, E. W.; Grant, C. M. Overlapping roles of the cytoplasmic and mitochondrial 878 redox regulatory systems in the yeast Saccharomyces cerevisiae. *Eukaryot Cell* **4**:392-400;
- 878 redox regulatory systems in the yeast Saccharomyces cerevisiae. *Eukaryot Cell* 4:392-400;
  879 2005.
  880 [50] Dethleften L. A. Lehmen C. M. Binglew, L. F. Beck, V. M. Tavia effects of equita
- [59] Dethlefsen, L. A.; Lehman, C. M.; Biaglow, J. E.; Peck, V. M. Toxic effects of acute
  glutathione depletion by buthionine sulfoximine and dimethylfumarate on murine mammary
  carcinoma cells. *Radiat Res* 114:215-224; 1988.
- 883 [60] Markovic, J.; Mora, N. J.; Broseta, A. M.; Gimeno, A.; de-la-Concepcion, N.; Vina, J.;
- Pallardo, F. V. The depletion of nuclear glutathione impairs cell proliferation in 3t3
  fibroblasts. *PLoS One* 4:e6413; 2009.
- 886 [61] Sengupta, R.; Holmgren, A. Thioredoxin and glutaredoxin-mediated redox regulation of
- ribonucleotide reductase. *World J Biol Chem* **5**:68-74; 2014.

- 888 [62] Zahedi Avval, F.; Holmgren, A. Molecular mechanisms of thioredoxin and glutaredoxin
- as hydrogen donors for Mammalian s phase ribonucleotide reductase. *J Biol Chem* 284:82338240; 2009.
- 891 [63] Olson, S. K.; Greenan, G.; Desai, A.; Muller-Reichert, T.; Oegema, K. Hierarchical assembly 892 of the eggshell and permeability barrier in C. elegans. *J Cell Biol* **198**:731-748; 2012.
- 893 [64] Brennan, L. D.; Roland, T.; Morton, D. G.; Fellman, S. M.; Chung, S.; Soltani, M.; Kevek, J.
- W.; McEuen, P. M.; Kemphues, K. J.; Wang, M. D. Small molecule injection into single-cell C.
  elegans embryos via carbon-reinforced nanopipettes. *PLoS One* 8:e75712; 2013.
- 896 [65] Ribas, V.; Garcia-Ruiz, C.; Fernandez-Checa, J. C. Glutathione and mitochondria. *Front* 897 *Pharmacol* **5**:151; 2014.
- 898 [66] Kojer, K.; Bien, M.; Gangel, H.; Morgan, B.; Dick, T. P.; Riemer, J. Glutathione redox 899 potential in the mitochondrial intermembrane space is linked to the cytosol and impacts the 900 Mia40 redox state. *EMBO* **/ 31**:3169-3182; 2012.
- 901 [67] Schaedler, T. A.; Thornton, J. D.; Kruse, I.; Schwarzlander, M.; Meyer, A. J.; van Veen, H.
- W.; Balk, J. A conserved mitochondrial ATP-binding cassette transporter exports glutathione
  polysulfide for cytosolic metal cofactor assembly. *J Biol Chem* 289:23264-23274; 2014.
- 904 [68] Gonzalez-Cabo, P.; Bolinches-Amoros, A.; Cabello, J.; Ros, S.; Moreno, S.; Baylis, H. A.;
- 905 Palau, F.: Vazquez-Manrique, R. P. Disruption of the ATP-binding cassette B7 (ABTM-
- 906 1/ABCB7) induces oxidative stress and premature cell death in Caenorhabditis elegans. *J Biol*907 *Chem* 286:21304-21314; 2011.
- [69] Landry, A. P.; Cheng, Z.; Ding, H. Reduction of mitochondrial protein mitoNEET [2Fe-2S]
  clusters by human glutathione reductase. *Free Radic Biol Med*; 2015.
- 910 [70] Lazarou, M.; Narendra, D. P.; Jin, S. M.; Tekle, E.; Banerjee, S.; Youle, R. J. PINK1 drives
- 911 Parkin self-association and HECT-like E3 activity upstream of mitochondrial binding. *J Cell*912 *Biol* 200:163-172; 2013.
- 913 [71] Sulston J, H. J. *Methods. In: The Nematode Caenorhabditis elegans.* . Cold Spring Harbour,
  914 New York: Cold Spring harbour Laboratory Press, Cold Spring Harbour; 1998.
- 915 [72] Morales-Martinez, A.; Dobrzynska, A.; Askjaer, P. Inner nuclear membrane protein LEM-
- 916 2 is required for correct nuclear separation and morphology in C. elegans. *J Cell Sci* 917 **128**:1090-1096; 2015.
- 918 [73] Mello, C. C.; Kramer, J. M.; Stinchcomb, D.; Ambros, V. Efficient gene transfer in
- 919 C.elegans: extrachromosomal maintenance and integration of transforming sequences. *EMBO* 920 / 10:3959-3970; 1991.
- 921 [74] Miyabayashi, T.; Palfreyman, M. T.; Sluder, A. E.; Slack, F.; Sengupta, P. Expression and
- 922 function of members of a divergent nuclear receptor family in Caenorhabditis elegans. *Dev* 923 *Biol* **215**:314-331; 1999.
- 924 [75] Muller, S.; Walter, R. D.; Fairlamb, A. H. Differential susceptibility of filarial and human
  925 erythrocyte glutathione reductase to inhibition by the trivalent organic arsenical melarsen
  926 oxide. *Mol Biochem Parasitol* **71**:211-219; 1995.
- 927 [76] Nieto, C.; Almendinger, J.; Gysi, S.; Gomez-Orte, E.; Kaech, A.; Hengartner, M. O.;
  928 Schnabel, R.; Moreno, S.; Cabello, J. ccz-1 mediates the digestion of apoptotic corpses in C.
  929 elegans. *J Cell Sci* 123:2001-2007; 2010.
- 930 [77] Larsen, P. L.; Albert, P. S.; Riddle, D. L. Genes that regulate both development and 931 longevity in Caenorhabditis elegans. *Genetics* **139**:1567-1583; 1995.
- 932 [78] Inoue, H.; Hisamoto, N.; An, J. H.; Oliveira, R. P.; Nishida, E.; Blackwell, T. K.; Matsumoto,
- 933 K. The C. elegans p38 MAPK pathway regulates nuclear localization of the transcription
- factor SKN-1 in oxidative stress response. *Genes Dev* **19**:2278-2283; 2005.

- 935 [79] Etheridge, T.; Rahman, M.; Gaffney, C. J.; Shaw, D.; Shephard, F.; Magudia, J.; Solomon, D. 936 E.; Milne, T.; Blawzdziewicz, J.; Constantin-Teodosiu, D.; Greenhaff, P. L.; Vanapalli, S. A.; Szewczyk, N. J. The integrin-adhesome is required to maintain muscle structure, 937 938 mitochondrial ATP production, and movement forces in Caenorhabditis elegans. FASEB J 939 **29:**1235-1246; 2015. 940 [80] Krauth-Siegel, R. L.; Blatterspiel, R.; Saleh, M.; Schiltz, E.; Schirmer, R. H.; Untucht-Grau,
- 941 R. Glutathione reductase from human erythrocytes. The sequences of the NADPH domain and
- 942 of the interface domain. Eur J Biochem 121:259-267; 1982.
- [81] Urig, S.; Lieske, J.; Fritz-Wolf, K.; Irmler, A.; Becker, K. Truncated mutants of human 943
- 944 thioredoxin reductase 1 do not exhibit glutathione reductase activity. FEBS Lett 580:3595-3600; 2006.
- 945
- 946













![](_page_36_Figure_0.jpeg)

FIGURE 7

## Supplemental Table 1: *C. elegans* strains used in this work.

Strain name	Genotype	Reference/Source
N2	Wild type, DR subclone of CB original (Tc1 pattern I)	CGCª
FX3574	gsr-1(tm3574) III	NBRP <sup>b</sup>
JK2533	qC1 [dpy-19(e1259) glp-1(q339) qls26 [Plag-2::gfp; rol- 6(su1006)]] III/eT1 (III;V)	CGC
MT20108	dpy-17(e164) unc-32(e189)/qC1 [dpy-19(e1259) glp-1(q339) nls281 [Pmvo-2::rfp]] []]	CGC
VZ64	gsr-1(tm3574)/qC1 [dpy-19(e1259) glp-1(q339) qls26 [Plag- 2::gfp: rol-6(su1006)]] []]	This study
VZ454	gsr-1(tm3574)/qC1 [dpy-19(e1259) glp-1(q339) nls281 [Pmyo- 2::rfn]] III	This study
VZ291	gsr-1(tm3574) III; vzEx105 [Pgsr-1::gsr-1::3´-UTR <sub>gsr-1</sub> ; Punc- 122::afn]	
VZ292	gsr-1(tm3574) III; vzEx106 [Pgsr-1::gsr-1::3´-UTR <sub>gsr-1</sub> ; Punc- 122::afn]	This study
VZ232	vzEx76 [Pasr-1::afp: rol-6 (su1006)]	This study
VZ233	vzEx77 [Pasr-1::afp: rol-6 (su1006)]	This study
V7234	vzEx78 [Pasr-1::afp: rol-6 (su1006)]	This study
V7257	vzEx93 [Pasr-1::asr-1::afn: rol-6(su1006)]	This study
VZ258	vzEx94 [Pasr-1::asr-1::afp: rol-6(su1006)]	This study
VZ200	asr_1/tm3574) III: vzEv93 [Pasr_1::asr_1::afn: rol_6(su1006)]	This study
VZ3 <del>4</del> 1 V/7307	yzEx117 [asr_1 1 <sup>st</sup> intron: asr_1::afn]	This study
VZ307 VZ309	vzEx117 [951-1 1 1111011951-1919]	This study
VZ300	VZEX110 [951-1 1 1111011951-1910]	This study
VZ309	VZEX119 [951-11 IIIII011951-191]	This study
VZ34Z	gsr-1(tm3574)/qC1[apy-19(e1259)]gip-1(q339)]qis26[Pilag-	i nis study
\/707F	2::gfp; roi-6(su1006)]] III; vzEx119 [gsr-11* Intron::gsr-1::gfp]	<b>T</b> I: ( )
VZ375	$gsr-1(tm_{3574})$ III; VZEX144 [Pgsr-1::gsr-1( $\Delta 1^{a}$ exon)::gfp]	I his study
VZ378	$gsr-1(tm3574)$ III; vzEx145 [Pgsr-1::gsr-1( $\Delta 1^{\circ}$ exon)::gfp]	This study
VZ393	gsr-1(tm3574) III; vzEx146 [Pgsr-1::gsr-1(∆1 <sup>st</sup> exon)::gfp]	This study
VZ395	gsr-1(tm3574)	This study
VZ396	gsr-1(tm3574)	This study
VZ397	gsr-1(tm3574) III; vzEx149 [Pgsr-1::gsr-1(1 <sup>st</sup> ATG->CCT)::gfp]	This study
VZ445	vzEx158 [Pasr-1::asr-1(2 <sup>nd</sup> ATG->GGT)::afp]	This study
VZ446	vzEx159 [Pasr-1::asr-1(2 <sup>nd</sup> ATG->GGT)::afp]	This study
V7447	vzEx160 [Pasr-1"asr-1(2 <sup>nd</sup> ATG->GGT)"afp]	This study
VZ610	asr-1(tm3574)/aC1 [dpv-19(e1259) alp-1(a339) nls281 [Pmvo-	This study
	2"rfp]] []]: vzFx158 [Pasr-1"asr-1/2" ATG->GGT)"afp]	The etday
PS6187	unc-119 (ed3) III; syEx1155 [Pmyo-3::tomm-20::rfp::3xMyc; unc-119(+)]	[1]
V7427	vzEx155 [Pmvo-3::MTSasr_1::afp]	This study
V7428	vzEx156 [Pmvo-3::MTSgsr-1::gfp]	This study
VZ420 VZ420	vzEx150 [Filly0-5WISgsi-1gip]	This study
VZ723 VZ7236	vzEx155 [Pmvo-3::MTSaar 1::arp] vzEx155 [Pmvo-3::MTSaar 1::arp]: avEv1155 [Pmvo-3::tomm	This study
VZ430	20::rfp::3xMyc; unc-119(+)]	
VZ12	trxr-2(tm2047)	
VZ594	gsr-1(tm3574) trxr-2(tm2047)/qC1 [dpy-19(e1259) glp-1(q339) nls281 [Pmyo-2::rfp]] III	This study
VZ603	gsr-1(tm3574) trxr-2(tm2047); vzEx147 [Pgsr-1::gsr-1(1 <sup>st</sup> ATG- >CCT)::gfp]	This study
VZ604	gsr-1(tm3574) trxr-2(tm2047); vzEx145 [Pgsr-1∷gsr-1(∆1 <sup>st</sup> exon)::gfp]	This study

VB1414	trxr-1(sv47) IV	[2]
VZ78	gsr-1(tm3574)/qC1 [dpy-19(e1259) glp-1(q339) qls26 [Plag-	This study
	2::gfp; rol-6(su1006)]] III; trxr-1(sv47) IV	
VZ492	gsr-1(tm3574)/qC1 [dpy-19(e1259) glp-1(q339) nls281 [Pmyo-	This study
	2::rfp]] III; trxr-1(sv47) IV	
CL1166	dvEx166 [Pgst-4::gfp::NLS]	[3]
VZ498	gsr-1(tm3574)/qC1 [dpy-19(e1259) glp-1(q339) nls281 [Pmyo- 2::rfp]] ///: dvEx166 [Past-4::afp::NLS]	This study
VB2317	svFx741 [Pacs-1"afp]	[2]
VZ468	asr-1(tm3574)/aC1 [dpv-19(e1259) alp-1(a339) nls281 [Pmvo-	This study
	2::rfp]] III; svEx741 [Pacs-1::gfp]	
LD001	IdIs007 [Pskn-1::skn-1b/c::gfp; rol-6(su1006)] X	[4]
VZ452	gsr-1(tm3574)/qC1 [dpy-19(e1259) glp-1(q339) nls281 [Pmyo-	This study
	2::rfp]] III; IdIs007 [Pskn-1::skn-1b/c::gfp; rol-6(su1006)] X	-
PD4251	ccIs4251 [Pmyo-3::nucgfp::LacZ::NLS; Pmyo-3::mitogfp); dpy-	CGC
	20(+)] I; dpy-20(e1282) IV	
VZ508	ccIs4251 [Pmyo-3::nucgfp::LacZ::NLS; Pmyo-3::mitogfp); dpy-	This study
	20(+)] l; gsr-1(tm3574)/qC1 [dpy-19(e1259) glp-1(q339) nls281	
	[Pmyo-2::rfp]] []]	
PJ707	jls01[Pmyo-3::myo-3::gfp; rol-6(su1006)]	[5]
VZ507	gsr-1(tm3574)/qC1 [dpy-19(e1259) glp-1(q339) nls281 [Pmyo-	This study
0.14400	2::rfp]] III; JIs01[Pmyo-3::myo-3::gfp; rol-6(su1006)]	101
SJ4100		[6]
VZ495	gsr-1(tm35/4)/qC1 [dpy-19(e1259) glp-1(q339) nIs281 [Pmyo- 2::rfp]] III: zcls13 [Phsp-6::afp] V	This study
BN245	ItIs37 [Ppie-1::mCherry::his-58: unc-119 (+)] IV: oils1 [Ppie-	[7]
	1::gfp::tbb-2; unc-119(+)] V; ltIs24 [Ppie-1::gfp::tba-2; unc-	L. 1
	119(+)]	
BN323	gsr-1(tm3574)/qC1 [dpy-19(e1259) glp-1(q339) qls26 [Plag-	This study
	2::gfp; rol-6(su1006)]] III; ItIs37 [Ppie-1::mCherry::his-58; unc-	5
	119 (+)] IV;	

<sup>a</sup> Caenorhabditis Genetics Center (<u>http://www.cbs.umn.edu/CGC/</u>) <sup>b</sup> National BioResource Project *C. elegans* (http://www.shigen.nig.ac.jp/c.elegans/index.jsp)

#### SUPPLEMENTAL FIGURES

Supplemental Figure 1: Complete cell lineage analysis of *wild type* and *gsr-1(m-,z-)* embryos. Comparison of cell lineage and cell divisions timing of *wild type* control embryos (grey lines) with *gsr-1(m-,z-)* embryos (black lines with red dots) arresting at **A**) 17 cell-stage; **B**) 33 cell-stage; **C**) 103 cell-stage (same as depicted in **Figure 5A-C**). Green dots show cell mitosis.

Supplemental Figure 2: Sensitivity of gsr-1(m-,z-) animals with restricted expression of GSR-1b::GFP in cytoplasm to acute paraquat and juglone exposure. gsr-1(m-,z-) worms with GSR-1 activity restricted to cytoplasm are as resistant as wild type control to paraquat and juglone treatments. Therefore, mitochondrial glutathione reductase isoform is not required for resistance to these chemicals. Data are the mean  $\pm$  SD of three independent experiments with three biological replicates each (n  $\pm$  200).

#### SUPPLEMENTAL REFERENCES

[1] Cacho-Valadez, B.; Munoz-Lobato, F.; Pedrajas, J. R.; Cabello, J.; Fierro-Gonzalez, J. C.; Navas, P.; Swoboda, P.; Link, C. D.; Miranda-Vizuete, A. The characterization of the Caenorhabditis elegans mitochondrial thioredoxin system uncovers an unexpected protective role of thioredoxin reductase 2 in beta-amyloid peptide toxicity. *Antioxid Redox Signal* **16**:1384-1400; 2012.

[2] Stenvall, J.; Fierro-Gonzalez, J. C.; Swoboda, P.; Saamarthy, K.; Cheng, Q.; Cacho-Valadez, B.; Arner, E. S.; Persson, O. P.; Miranda-Vizuete, A.; Tuck, S. Selenoprotein TRXR-1 and GSR-1 are essential for removal of old cuticle during molting in Caenorhabditis elegans. *Proc Natl Acad Sci USA* **108**:1064-1069; 2011.

[3] Link, C. D.; Johnson, C. J. Reporter transgenes for study of oxidant stress in Caenorhabditis elegans. *Methods Enzymol* **353**:497-505; 2002.

[4] An, J. H.; Blackwell, T. K. SKN-1 links C. elegans mesendodermal specification to a conserved oxidative stress response. *Genes Dev* **17**:1882-1893; 2003.

[5] Szewczyk, N. J.; Kozak, E.; Conley, C. A. Chemically defined medium and Caenorhabditis elegans. *BMC Biotechnol* **3**:19; 2003.

[6] Yoneda, T.; Benedetti, C.; Urano, F.; Clark, S. G.; Harding, H. P.; Ron, D. Compartment-specific perturbation of protein handling activates genes encoding mitochondrial chaperones. *J Cell Sci* **117:**4055-4066; 2004.

[7] Morales-Martinez, A.; Dobrzynska, A.; Askjaer, P. Inner nuclear membrane protein LEM-2 is required for correct nuclear separation and morphology in C. elegans. *J Cell Sci* **128**:1090-1096; 2015.

![](_page_41_Figure_0.jpeg)

SUPPLEMENTAL FIGURE 1

![](_page_42_Figure_0.jpeg)

Juglone 150  $\mu$ M, 16 h

![](_page_42_Figure_2.jpeg)

Paraquat 50 mM, 16 h