Selenoprotein H Is a Nucleolar Thioredoxin-like Protein with a Unique Expression Pattern*§

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The human selenoproteome consists of 25 known selenoproteins, but functions of many of these proteins are not known. Selenoprotein H (SelH) is a recently discovered 14-kDa mammalian protein with no sequence homology to functionally characterized proteins. By sensitive sequence and structure analyses, we identified SelH as a thioreodoxin fold-like protein in which a conserved CXXU motif (cysteine separated by two other residues from selenocysteine) corresponds to the CXXC motif in thioredoxins. These data suggest a redox function of SelH. Indeed, a recombinant SelH shows significant glutathione peroxidase activity. In addition, SelH has a conserved RKRK motif in the N-terminal sequence. We cloned wild-type and cysteine mutant forms of SelH either upstream or downstream of green fluorescent protein (GFP) and localized this fusion protein to the nucleus in transfected mammalian cells, whereas mutations in the RKRK motif resulted in the cytosolic protein. Interestingly, the full-length SelH-GFP fusion protein localized specifically to nucleoli, whereas the N-terminal sequence of SelH fused to GFP had a diffuse nucleoplasm location. Northern blot analyses revealed low expression levels of SelH mRNA in various mouse tissues, but it was elevated in the early stages of embryonic development. In addition, SelH mRNA was overexpressed in human prostate cancer LNCaP and mouse lung cancer LCC1 cells. Down-regulation of SelH by RNA interference made LCC1 cells more sensitive to hydrogen peroxide but not to other peroxides tested. Overall, these data establish SelH as a novel nucleolar oxidoreductase and suggest that some functions in this compartment are regulated by redox and dependent on the trace element selenium.

Selenium is an essential trace element with significant biomedical potential and roles in human health (1). As a dietary supplement, selenium has been shown to serve as a cancer chemopreventive agent and has roles in immune function, mammalian development, male reproduction, and in preventing heart disease and other cardiovascular and muscle disorders (for reviews, see Ref. 2). Its biological significance is attributed to the occurrence of this element in proteins in the form of selenocysteine (Sec),4 the 21st amino acid in the genetic code (3). This rare, highly reactive amino acid (4) is encoded by the UGA codon and is incorporated into proteins co-translationally (5–7). Sec-specific incorporation requires the presence of cis- and trans-acting elements. In addition to UGA, a stem-loop structure known as the SECIS (selenocysteine insertion sequence) element operates as the cis element for Sec incorporation (8–10). In eukaryotes, SECIS is located in 3′-UTRs of selenoprotein mRNAs and interacts with trans-acting factors. Recently, an additional recoding element was identified in a subset of eukaryotic selenoprotein genes that is adjacent to Sec-encoding UGA codons (11). In eukaryotes, the trans-acting factors consist of Sec tRNA (12–15), selenocysteyl-tRNA-specific elongation factor (14, 15), and SBP2 (SECIS-binding protein 2) (16, 17). These three components function together to redefine the UGA codon to dictate Sec insertion (6, 18). Ribosomal protein L30 has also been shown to be a component of the eukaryotic Sec recoding machinery (19).

Selenoproteins are present in all kingdoms of life, and the full set of such proteins in a particular organism is known as the selenoproteome. The human selenoproteome consists of 25 proteins, whereas rodents have 24 such proteins (20). About half of these proteins (e.g. glutathione peroxidases (21) and thioredoxin reductases (22)) play roles in various redox reactions, but functions of the remaining selenoproteins are not known. Therefore, to understand the effects of selenium on human health and to explain how these effects are governed by dietary dependence on selenium, the functions of all selenoproteins need to be elucidated.

Selenoprotein H (SelH) is a recently discovered 14-kDa mammalian protein with no homology to functionally characterized proteins (20). However, certain similarities to other selenoproteins can be seen, particularly a conserved Cys-Xaa-Xaa-Sec (CXXU) motif wherein cysteine and Sec are separated

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4 The abbreviations used are: Sec, selenocysteine; SelH, selenoprotein H; GFP, green fluorescent protein; GPx, glutathione peroxidase; tert-BOOH, tert-butyl hydroperoxide; PBS, phosphate-buffered saline; NLS, nuclear localization signal; DAPI, 4′,6-diamidino-2-phenylindole.

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by two other amino acids. This feature is also present in several other mammalian selenoproteins (i.e. SelW, SelIT, SelM, and SelV), but the physiological roles of these selenoproteins are also not known. Structures of SelM (23) and its distant homolog Sep15 (24) have recently been determined (25). Both proteins possess a thioredoxin-like domain and reside in the endoplasmic reticulum, suggesting that these two proteins are thiol-disulfide isomerases with a role in disulfide bond formation.

Among other CXXU-containing selenoproteins, SelW is a better characterized protein, but its function also has not been defined (26). It is the smallest mammalian selenoprotein existing in the form of several isoforms that differ by post-translational modifications (27, 28). SelW is expressed in various tissues and is abundant in muscle. Reduced levels of SelW have been attributed to white muscle disease (29). SelV is a testis-specific distant homolog of SelW having an additional domain of unknown function (20). Here we used computational, molecular, and cell biology techniques to characterize SelH, which revealed that this protein is a nucleolar oxidoreductase with an unusual expression pattern.

EXPERIMENTAL PROCEDURES

Materials—Chemicals were from Sigma, restriction enzymes from Amersham Biosciences, DNA purification kits from Qiagen, and mammalian cell culture reagents from Invitrogen. Primers used in the study are shown in supplemental Table S1.

Multiple Sequence Alignment, Topology Prediction, and Structural Motif Search for SelH—All detected SelH sequences were extracted from GenBank™ nonredundant and EST databases and analyzed by BLAST (30) and ClustalW software (31). Localization of SelH was predicted using PSORT II (32). PSI-BLAST/PHI-BLAST (33) searches with the default setting were carried out to detect conserved protein domains. The HHpred web-based program (34) was used for fold prediction.

Recombinant Protein Expression in Escherichia coli and Antibody Production—Mouse SelH cDNA clone was purchased from Research Genetics. The Sec codon was substituted with that coding for Cys using QuikChange II kit (Stratagene) and for each chimera, the corresponding mutant with disrupted NLS was obtained. Transfection of NIH 3T3 cells was performed using Lipofectamine (Invitrogen) and for each chimera, the corresponding mutant with disrupted NLS was obtained. Transfection of NIH 3T3 cells was performed using Lipofectamine (Invitrogen), and the images were captured using an Olympus FV500 inverted focal microscope at the University of Nebraska-Lincoln Microscopy Core Facility.

Immunohistochemistry—NIH 3T3 cells were seeded on coverslips and transfected with SelH-GFP fusion constructs. After 24 h, the cells were washed with PBS and fixed by flashing with ice-cold methanol containing 0.1% Triton X-100 followed by a 5-min treatment with 4% paraformaldehyde. The cells were then treated with blocking solution (Roche Applied Science) and incubated with rabbit anti-nucleolin antibodies (Novus Biologicals) and mouse monoclonal anti-UB2 antibodies (RDI Research Diagnostics) concurrently followed by washing with PBS and treatment with donkey anti-mouse Cy5-conjugated and monkey anti-rabbit Cy3-conjugated antibodies (Jackson ImmunoResearch Laboratories). The samples were then washed with PBS and distilled water and mounted with Gel/ Mount (Biomedia). Dilutions of all commercial antibodies were as suggested by the manufacturers.

Gene Expression Analyses—An in silico expression analysis was performed using the SAGE Genie web resource (cgap.ncbi.nlm.nih.gov/SAGE) (36). Pre-made Northern blot membranes (mouse adult tissue blot and mouse conceptus embryonic tissue blot) were purchased from Seegene and probed with 0.3-kb 32P-labeled (Redivue [32P]dCTP, Amersham Biosciences) SelH cDNA. The probe was generated using the Rediprime II random prime labeling system (Amersham Biosciences) in accordance with the manufacturer’s protocol. To analyze the expres-
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In cell lines, total RNA was isolated using the RNAqueous kit (Ambion), separated on an agarose gel, transferred onto a Zeta-Probe blotting membrane (Bio-Rad, protocol from (37)), and probed as described above.

**Western Blot Analyses**—Tissues were extracted from euthanized mice, frozen in liquid nitrogen, and stored at −80 °C until the day of analysis. The tissues were homogenized in PBS containing complete protease inhibitors (Roche Applied Science), and the lysates were normalized with regard to protein concentration. 10% BisTris NOVEX gels (Invitrogen) were used, and each well was loaded with 25 μg of protein. Following SDS-PAGE, the proteins were transferred onto polyvinylidene difluoride membranes (Invitrogen) and probed with anti-SelH antibodies. Secondary horseradish peroxidase-conjugated antibodies were from Amersham Biosciences, and chemiluminescent peroxidase substrate was from Sigma. Lysates from various human cancer cell lines (a generous gift from Dr. Katerina Gurova, Cleveland BioLabs) were examined as described above.

**Cell Culture Labeling, Subcellular Fractionation, and Nucleoli Preparation**—Experiments involving metabolic labeling of selenoproteins were carried out by adding 75Se ([75Se]seleniumic acid; specific activity 1,000 Ci/mmol, Research Reactor Facility, University of Missouri, Columbia, MO) to the culture medium as described previously (38). Separation of cytoplasm and nuclear fractions was performed as described elsewhere (39). In brief, following labeling for 48 h, cells were trypsinized, washed with PBS, homogenized using a tight pestle in 0.25 m sucrose buffer containing complete protease inhibitor mixture (Roche Applied Science), and cytosolic and nuclear fractions were separated by differential centrifugation. The resulting cellular fractions were normalized in regard to protein concentration, subjected to SDS-PAGE, and transferred onto polyvinylidene difluoride membrane, and then 75Se-labeled proteins were visualized with a Storm PhosphorImager system (Amersham Biosciences). Further subfractionation of nuclear compartments to obtain nucleoli and nucleoplasm fractions was conducted as described previously (40) with modifications (41).

**RNA Interference in LCC1 Cell Line and Oxidative Stress**—To knock down SelH gene expression, five separate 21-nucleotide sequences were selected from SelH cDNA as being unique using the online service “siDESIGN” of Dharmacon Research, Inc. These small interfering RNA constructs of pU6-m3 were prepared as described previously (42). The sequences of all five SelH small interfering RNA constructs were confirmed. The five constructs and the pU6-m3 negative control plasmids were separately transfected into LCC1 cells. After stabilizing the transfected cells with 0.8 mg/ml hygromycin, SelH mRNA levels were determined by Northern blot analysis. Based on mRNA levels, the best small interfering RNA target sequences of SelH were identified as 5′-GAA TTG AAG AAG TAC CTT TCA-3′. These small interfering RNA constructs were used in subsequent experiments.

Sensitivity of LCC1 cells to oxidative stress was analyzed by the CellTiter 96 AQeueous 1 Solution cell proliferation assay (Promega). 5 × 103 cells were seeded into each well in 96-well plates and grown overnight in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal calf serum. The cells were washed twice with PBS and incubated in 100 μl of serum/phenol red-free medium containing the indicated (Fig. 6B) concentrations of hydrogen peroxide or 0–1 mM t-BOOH, cumyl hydroperoxide or paraquat for 1 h at 37 °C. To examine sensitivity to menadione, we used 0–30 μM treatment with this compound for 6 h. 20 μl of CellTiter 96 AQeueous One Solution reagent were added to each well and incubated for an additional 4 h. The A490 value was recorded using an ELx808 Ultra microplate reader (Bio-Tek Instruments). Cell viability was calculated as the percentage of untreated controls, and the wells without cells were used as a blank.

**Evolutionary Analysis**—Amino acid sequence alignment of SelH sequences was generated using the PileUP program. The distances between sequences were calculated from these alignments using the DISTANCE program with the Kimura distance-measuring method, and then the unrooted phylogenetic tree with unscaled distance branches was generated using the GrowTree program with the unweighted pair-group method. All programs were part of the Wisconsin Package, Genetic Computer Group, Madison, WI (43).

**RESULTS AND DISCUSSION**

**SelH Is a Putative Nuclear Oxidoreductase**—Available SelH sequences were collected and assembled into a multiple sequence alignment (Fig. 1A). SelH sequences contain a conserved CXXU motif, which is flanked upstream by a predicted β-strand and downstream by a predicted α-helix. This arrangement is highly similar to thiols/disulfide oxidoreductases of the thioredoxin fold; however, the latter proteins have a CXXC motif instead of CXXU (44, 45). The use of advanced BLAST programs (PSI-BLAST/PHI-BLAST) (33) with several iterations revealed a distant homology to thioredoxin fold proteins. Indeed, further analysis with HHpred identified a thioredoxin-like fold in SelH. Structural alignment places the CXXU motif in SelH in the same place as the CXXC motif in thioredoxin. These data suggest that SelH is an oxidoreductase that uses its CXXU motif as the reversible thiol/selenol/selenenylsulfide redox group.

Sequence analysis also revealed nuclear localization signals (46) in SelH sequences. Remarkably, SelH proteins have a high content of basic residues (18.3% in mouse and 22.2% in zebrafish) due to the high occurrence of arginines and lysines. By using PSORT II, we found six putative NLSs in the SelH sequences of chickens (all monopartite), three in zebrafish (all monopartite), and eight in mice and humans (six monopartite and two bipartite). Comparison of NLS location and conservations using the multiple sequence alignment showed that only two NLSs were conserved, of which one was in the N-terminal and one in C-terminal sequences. Within the N-terminal NLS, the two strictly conserved residues were Arg-6 and Arg-8 (Fig. 1A) (numbering based on the mouse SelH sequence), suggesting that these residues have a critical role in nuclear import as a part of NLS.

**Peroxidase Activity of Recombinant SelH**—To further elucidate whether SelH is an oxidoreductase, we performed several biochemical tests with recombinant mutant forms of the protein in which the Sec residue was substituted with Cys or Ser. Specifically, these mutants were assayed for thioredoxin, per-
oxiredoxin, and GPx activities. In the thioredoxin assay, mammalian thioredoxin reductase 1 was used as the reductant and 5,5'-dithiobis-(2-nitrobenzoic acid) as the oxidant (47, 48), and a separate assay involved insulin reduction, as in the corresponding assay of thioredoxin function (49). We could detect no thioredoxin activity of SelH (data not shown). The peroxiredoxin activity was examined in an assay of protection of glutamine synthetase activity (47). SelH also showed no activity in this assay. Interestingly, SelH had a significant GPx activity (Fig. 1A). This activity was detected only with hydrogen peroxide, whereas t-BOOH was not active as a substrate. The Cys mutant had four times higher activity than the Ser mutant of SelH. We further compared the GPx activity of SelH with that of mammalian GPx1 using mouse liver lysate and quantitative Western blot analysis of GPx1 amounts in the sample (not shown). This activity was detected only with hydrogen peroxide, whereas t-BOOH was not active as a substrate. The Cys mutant was 2,000–4,000-fold lower than that of natural mouse liver GPx1 under the conditions used in the assay. It should be pointed out that, although SelH had a much lower activity than GPx1, we assayed the Cys form of the natural Sec-containing SelH. In addition, the peroxidase activity detected was similar to the activities of peroxiredoxins in the reduction of hydroperoxides. Thus, these data provided further evidence for the oxidoreductase activity of SelH. The natural oxidoreductase activity of SelH, however, remains unknown, and further studies will be needed to identify specific substrates/targets of this protein.

Identification of Target Proteins—The finding that SelH possesses a thioredoxin fold and has a putative CXXU redox motif suggests analogy with thioredoxin-like proteins that form transient disulfides with target proteins. Mutation of a resolving Cys in the CXXC motif of thioredoxin often results in stabilization of such complexes (51–53). We utilized this approach to trap the target proteins of SelH. Affinity resins were prepared by linking CXXS and SXXC forms of recombinant SelH and incubating with a nuclear fraction prepared from LCC1 cells. A silver-stained gel showing proteins eluted from the resins is shown in supplemental Fig. S1. Proteins enriched on the SelH resins were then identified by tandem mass spectrometry sequencing. Among those detected in the samples were several thiol oxidoreductases (e.g. peroxiredoxins 1 and 2, thioredoxin, and glutaredoxin), which is consistent with the redox nature of interactions between SelH and target proteins. Interestingly, most other identified proteins have previously been described as nucleolar proteins (e.g. pre-mRNA processing protein 8, putative pre-mRNA splicing factor RNA helicase (DEAH box protein 15), nucleolin, 60 S ribosomal protein L18, ribosomal protein S28). It should be noted that the identified proteins should only be viewed as preliminary targets. Further studies will be required to functionally characterize these interactions and determine their specificity.

Localization of SelH-GFP Fusion Constructs—To test whether SelH is a nuclear protein and whether Arg-6 and Arg-8 are responsible for its location, we prepared and transiently expressed various SelH-GFP fusion proteins in NIH 3T3 cells (Fig. 2A). In these experiments, we used a Sec38Cys mutant form of SelH, which was placed upstream (construct 1) and downstream (construct 5) of GFP. In addition, we fused a 3-kDa N-terminal portion of the protein that contains the putative NLS with GFP (construct 3). We also placed a full-length SelH cDNA sequence including a 3'-UTR containing a SECIS element downstream of GFP as described previously (20). For each of these four constructs, we developed mutants in which Arg-6 and Arg-8 were mutated to serines. Western blot analysis (Fig. 2B) of cells transfected with these constructs revealed correspondence of the actual sizes of the

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**FIGURE 1. Redox properties of SelH.** A, multiple sequence alignment of SelH sequences. The locations of CXXU (which corresponds to the CXXC motif in thioredoxins) and NLS motifs are indicated by boxes, respectively. The predicted secondary structure is shown above the sequence. B, glutathione peroxidase activity of SelH. The indicated recombinant forms of SelH (CXXS and CXXC forms) or mouse liver lysate were assayed for GPx activity as described under “Experimental Procedures.” Reactions were initiated by adding 300 μM H₂O₂ or 233 μM t-BOOH as the substrate. μU, milliunits.
FIGURE 2. Localization of SelH. A, organization of SelH-GFP fusion constructs utilized in the localization experiments. SelH was fused either upstream or downstream of GFP, and for each construct, the two conserved arginines in the NLS motif were mutated to serines. In addition, to increase expression levels, Sec was mutated to Cys in several constructs as shown in the figure. In the constructs expressing a selenoprotein form of SelH, a 3'‐UTR was included that contained a SECIS element. B, detection of SelH-GFP fusion proteins by Western blots using anti-SelH antibodies. Migration of expressed proteins corresponded to their predicted masses and is indicated by arrows on the right. Lanes 1–8 contain proteins generated from the corresponding constructs shown in A above. C, confocal microscopy of GFP-SelH fusion protein constructs containing an N-terminal SelH. Three images are shown for each construct. Left panels show green (GFP) fluorescence, middle panels DAPI staining (nuclear marker), and right panels images obtained by merging the left and middle panels. Numbering from 1 to 4 corresponds to A above. D, Same as in C, except that SelH was fused downstream of GFP in the SelH-GFP protein. Numbering from 5 to 8 corresponds to A above.
fusion proteins to the calculated masses, except for the wild-type SelH-GFP fusion proteins (constructs 7 and 8 on Fig. 2A and lanes 7 and 8 on Fig. 2B, respectively), because of significant premature termination at the Sec position. Full-length Sec-containing SelH-GFP fusion proteins were not detectable using available anti-SelH antibodies.

GFP fluorescence in transfected cells resulting from SelH-GFP fusion proteins was determined by dual wavelength confocal microscopy in parallel with a nuclear marker (DAPI). We found that, when NLS was present in proteins, fluorescence was localized to the nuclear fraction (Fig. 2, C and D; panels 1, 3, 5, and 7), and when the arginines were mutated, the fluorescence signal was confined to the cytosol (Fig. 2, C and D; panels 2, 4, 6, and 8). Thus, SelH is a nuclear protein possessing a RKRK monopartite NLS, and the arginines in this motif are essential for nuclear targeting of the protein.

SelH Is a Nucleolar Protein—As shown in Fig. 2, C and D, fluorescence inside nuclei that reflected localizations of proteins derived from constructs 1, 3, 5, and 7 was heterogeneous. In fact, both the SelH-GFP and GFP-SelH forms appeared to be enriched in nucleoli, whereas when only the NLS-containing N-terminal peptide was fused to GFP (construct 3), fluorescence was diffused throughout the nuclear compartment. Interestingly, in the Sec-containing SelH-GFP fusion protein (construct 7), which migrated as a 310-kDa protein in Western blot assays (Fig. 2B), (indicating significant premature termination at the UGA codon), the fluorescent signal was not diffused but still differed from the typical nucleolar staining. We subjected the cells transfected with constructs 1 and 7 to immunohistochemical analysis and used markers for nucleolar and nucleoplasm staining in parallel. As shown in Fig. 3, upper panel, the full-length SelH co-localizes with nucleolin, a nucleolar marker, indicating that SelH is indeed a nucleolar protein. However, the location of the 3-kDa portion of SelH fused to GFP (construct 7) was surprising, as it appeared to be present in the nucleoplasm. Taken together, these data indicate that SelH is a nucleolar protein and that its correct localization required the full-length folded protein. The nucleolar location of SelH is also consistent with the abundance of nucleolar proteins that were enriched on the SelH resins.

The nucleolus has long been known as a dense subnuclear structure where ribosome biogenesis occurs (for reviews, see Refs. 54 and 55). Unlike the nucleus and other membrane-restricted organelles, there is no evidence for the occurrence of a frontier separating the nucleolus from the surrounding nucleoplasm (56, 57). It is thought that any molecule could migrate in and out of the nucleolus and that targeting of particular cellular components to the nucleolus depends on interactions with the co-called nucleolar binding blocks (58).

Expression of SelH mRNA—To analyze SelH gene expression, we initially carried out an in silico expression analysis using mouse and human expression data that is available in the form of supplemental Figs. S2 and S3. Both mouse and human SelH
mRNA appeared to be moderately expressed in many, if not all, tissues and organs. However, the majority of expressed sequence tags were derived from either embryonic or tumor cells, particularly from carcinomas. Moreover, analysis of SAGE datasets suggested that expression of human SelH is elevated in some tumors, especially thyroid, lung, stomach, and liver cancers. In cell lines, SelH appeared high in embryonic samples, carcinomas, and cell lines derived from bone marrow. In mouse, it appeared to be elevated in developing brain. Interestingly, we previously found that, during zebrafish development, SelH localized within highly proliferative tissues such as the brain ventricular zone, part of the retina and tectum at 24 hpf, and later in branchial arches and pectoral fin buds and the proliferative zone of the retina (59). In addition, zebrafish SelH was identified as essential for embryonic and early larval development (60).

To directly examine SelH mRNA expression, we carried out Northern blot analyses using mouse embryonic and adult tissues and a number of available cell lines (Fig. 4). As expected, we detected SelH mRNA in various tissues. The signal was particularly strong in samples from early stages of embryogenesis and in the adult tissues of thymus, brain, testis, and uterus (Fig. 4, A and B). In cell lines, SelH mRNA expression was elevated in LNCaP, LCC1, and several other cell lines.

Expression of SelH in Mouse Tissues and Human Cancer Cell Lines—To further elucidate the expression of SelH, we carried out Western blot analysis (Fig. 5). SelH was abundant in mouse spleen but was expressed in lower levels in the brain and was not detected in the other organs examined (Fig. 5A). In addition, we analyzed cancer cell lines and found that SelH expression was dissimilar in various cells, but as expected, it was high in LNCaP, LCC1, and several other cell lines (Fig. 5B).

We metabolically labeled LNCaP cells with $^{75}$Se and then prepared nuclear and cytoplasmic fractions (Fig. 5C). A 14-kDa band corresponding in size to SelH was enriched in the nuclear fraction. The nuclear fraction was further fractionated into...
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nucleolar and nucleoplasm fractions. By Western blot analysis, SelH was detected in the nucleolar fraction, in similar manner as the control protein nucleolin (Fig. 5D). Subsequent immunohistochemistry analysis further verified the presence of SelH in nucleoli (Fig. 5E).

Nucleoli are best known as a site for rRNA biosynthesis and modification. However, nucleoli have additional functions, such as virus infection control, maturation of non-nucleolar RNAs and RNPs, regulation of telomerase function and cell cycle, tumor suppressor and oncogene activities, and cell stress sensing and signaling (61–63). Moreover, by proteomic approaches, many uncharacterized proteins have been identified in nucleoli, suggesting additional unknown functions for this compartment (41, 64).

RNA Interference of SelH in LCC1 Cells—Because we found that SelH is a thioredoxin-like protein with significant GPx activity, this protein was further examined for antioxidant functions. We employed RNAi to suppress SelH expression and tested the sensitivity of knockdown cells to oxidative stress. Decreased expression of SelH in LCC1 knockdown cells was verified by both Northern and Western blot analyses (Fig. 6A). We subjected these cells to treatment with hydrogen peroxide and found that the knockdown cells were more sensitive to this form of oxidative stress than control LCC1 cells (Fig. 6B). However, no difference in sensitivity of knockdown and control cells was observed when they were treated with tert-butyl hydroperoxide (t-BOOH), cumyl hydroperoxide, paraquat, and menadione (data not shown). Higher sensitivity of SelH knockdown cells to hydrogen peroxide was consistent with a role of this protein in redox reactions and observed specificity of SelH for hydrogen peroxide compared with t-BOOH.

Nucleoli have been shown to be involved in oxygen-dependent regulatory mechanisms via the hypoxia-inducible factor (HIF-1). This protein is degraded under conditions of normal oxygen tension but is stabilized in hypoxia by VHL (von Hippel-Lindau, a tumor suppressor protein) activating a set of genes implicated in oxygen homeostasis, tumor vascularization, and ischemic preconditioning (65, 66). It has been shown that hypoxia can neutralize the function of VHL by triggering its nucleoli sequestration (67–69). This mechanism underlines the importance of redox status in nucleoli, and we speculate that SelH, being a nucleolar oxidoreductase, may play a related role. Moreover, being a selenoprotein, SelH may mediate certain effects of dietary selenium on nucleolar function.

Evolutionary Analyses—A SelH phylogenetic tree was constructed based on the available SelH sequences. As shown in Fig. 7, various Drosophila species have evolved two additional homologs of the protein, which form separate branches from the selenoprotein forms of the protein. Interestingly, additional fruit fly homologs have neither CXXC nor CXXX motifs. Instead, SelH possesses a CXXR and SelH2 a CXXT motif, the latter more typical of peroxiredoxins and glutathione peroxidases, which are large families of thiol peroxidases involved in various biological processes (50, 70). These data suggest functional differences between SelH and the two Drosophila homologs. It is also possible that various functions of mammalian SelH are carried out by separate proteins in fruit flies.

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REFERENCES

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