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Hereditary Spastic Paraplegia-Linked REEP1 Modulates ER-Mitochondria Contacts

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Abstract

Objective—Mutations in receptor expression enhancing protein 1 (*REEP1*) are associated with hereditary spastic paraplegias (HSPs). Although axonal degeneration is thought to be a predominant feature in HSP, the role of *REEP1* mutations in degeneration is largely unknown. Previous studies have implicated a role for REEP1 in the ER, whereas others localized REEP1 with mitochondria. We sought to resolve the cellular localization of REEP1 and to further elucidate the pathobiology underlying *REEP1* mutations in patients.

Methods—A combination of cellular imaging and biochemical approaches was used to refine the cellular localization of REEP1. Next, *Reep1* mutations associated with HSP were functionally tested in neuritic growth and degeneration assays using mouse cortical culture. Finally, a novel assay was developed and used with wild type and mutant *Reep1s* to measure the interactions between the ER and mitochondria.

Results—We found that REEP1 is present at the ER-mitochondria interface, and it contains subdomains for mitochondrial as well as ER localization. Knockdown of *Reep1* and the expression of pathological *Reep1* mutations resulted in neuritic growth defects and degeneration. Finally, using our novel split-*R*Luc8 assay, we show REEP1 facilitates ER-mitochondria interactions, a function diminished by disease-associated mutations.

Interpretation—Our data potentially reconcile the current conflicting reports regarding REEP1 being either an ER or a mitochondrial protein. Furthermore, our results connect, for the first time, the disrupted ER-mitochondria interactions to a failure in maintaining health of long axons in HSPs. Finally, the split-*R*Luc8 assay offers a new tool to identify potential drugs for multiple neurodegenerative diseases with ER-mitochondria interaction defects.

The authors declare no competing financial interests.

Author Contributions

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Conflict of Interest Disclosure

Introduction

Hereditary spastic paraplegias (HSPs), also known as spastic paraplegias (SPGs), are a heterogeneous group of disorders that share clinical symptoms of progressive weakness and spasticity mainly in the lower extremities due to axonal degeneration. More than 70 distinct genetic loci have been mapped in HSP, many of which implicate common cellular processes. Among these are genes involved in membrane shaping and trafficking; mitochondrial function; myelination and lipid metabolism; protein folding and ER stress response; and axonal tract development. 1,2,4

REEP1 is believed to encode an ER protein and mutations are associated with autosomal dominant HSP (SPG31) as well as distal hereditary motor neuropathy^{5–7}. There are six members of the REEP proteins, REEP1–6, which can be grouped into two subfamilies, REEP1–4 and REEP5–6, based on structural and sequence homology. They share sequence homology with yeast Yop1 and plant HVA22 protein and all belong to the Yip (Ypt-interacting protein) family⁸. REEP proteins are known to function in ER membrane shaping: each contains REEP homology domain which has been shown to insert into membranes as a hairpin and actively bend membranes into curved structure. This domain also mediates physical interaction with atlastin-1, spastin, and reticulon.^{5,9–11} Although having a defined role in ER shaping, REEP1 has also been reported as a mitochondrial protein¹²; thus its intracellular localization remains controversial. Interestingly, mitochondrial dysfunctions, such as impaired fusion/fission and defective bioenergetics, have been reported in *REEP1*-associated HSP patients.^{13–15} This further suggests a mitochondrial related role for REEP1, although it is also possible that ER-localized REEP1 could affect mitochondrial functions given the tight association between the ER and mitochondria.^{13,16}

The physical interactions between the ER and mitochondria play important roles in many aspects of cellular functions. The contact sites, known as mitochondria-associated ER membranes (MAMs), serve as channels for transporting molecules (e.g. membrane lipids, proteins, and Ca²⁺) between the two organelles and as hubs for mitochondrial fusion/fission and autophagosome formation.^{17–20} Alterations in ER-mitochondrial contact often leads to mitochondrial dysfunction and compromised ER function.^{1,2,4,21} Recent evidence suggests that the deregulation of ER-mitochondrial contacts and impaired Ca²⁺ transfer at such sites is a common pathogenic mechanism in various neurodegenerative diseases, including Parkinson's disease, Alzheimer's disease, and amyotrophic lateral sclerosis.^{5–7,22–27}

In this study we have found that disease-associated *Reep1* mutations lead to developmental defects and neuritic degeneration in mouse primary cortical neurons. We show that REEP1 contains subdomains for both mitochondrial and ER localization and that REEP1 is detected in MAMs. Using a novel split-*R*Luc8 reassembly assay, we demonstrate that REEP1 can facilitate ER-mitochondrial interactions in live cells and that this function is abrogated in the presence of disease-associated mutations. Together these data provide evidence linking altered ER-mitochondria interactions to *REEP1*-mediated HSP and offer new avenues for potential therapeutic intervention for patients with HSP.

Materials and Methods

DNA constructs

The full length clone of *Reep1* was obtained by screening embryonic day 12.5 (E12.5) mouse cDNA library (in pBluescriptII-SK, provided by Doug Epstein), using the partial cDNA of Reep1 from a subtractive hybridization screen.^{8,28} For Reep1 wild type (Reep1^{WT}) and disease mutant constructs (Reep1P19R, Reep1A20E, Reep1I-115, Reep1I-139, and Reep11-176), each DNA sequence was PCRed out using 5' primers containing EcoR I site and CACCATG, and 3' primers containing the stop codon sequence and Mlu I site and cloned into pCAG-IRES-GFP vector. For GFP-fused Reep1 constructs used to determine protein expression, PCR products encoding each disease mutant construct as well as wild type construct were cloned into pcDNA3.1/CT-GFP TOPO in frame (Life Technologies). For Reep1P19R and Reep1A20E, site directed mutagenesis (Stratagene Chameleon Kit) was used to introduce each point mutation. For Reep1 shRNA constructs, the annealed oligonucleotide was subcloned into pLL3.7 vector. The sequences used include the nontargeting negative control (5'-CAGTCGCGTTTGCGACTGG-3'; Dharmacon), target 1 (5'-GGAAGTTTGTTCATCCCAC-3'), target 2 (5'-GGGTGACCTTTACGTCTTC-3'), and target 3 (5'-GGCTAGAGAAATAGTGTTG-3'). For V5-tagged Reep1, the full length Reep1 was PCRed out and cloned into pcDNA3.1/V5-His-TOPO (Life Technologies) in frame. For GFP-tagged Reep1s used for localization study, PCR products encoding amino acid residues (a.a.) 1-201, 1-115, 116-201, and 116-157 of the mouse Reep1 were cloned into pcDNA3.1/CT-GFP TOPO in frame (Life Technologies). Untagged Reep1 constructs were generated in pcDNA3.1 TOPO with the same cloning strategy. For the split-RLuc8, Renilla luciferase 8 (RLuc8) in pBAD/Myc-His^{5,9–11,29} was used for the initial template. The split site of the RLuc8 was selected according to the previous reports, ^{12,30,31} For Mito- $RLuc8^N$ (the N-terminal half of the RLuc8 targeted to the mitochondria), DNA sequences encoding a.a. 1-91 of the RLuc8 were fused to the 3' end of the Flag tag sequence and the previously reported mitochondria-targeting sequence of the mouse AKAP1 (a.a. residues 34–63)^{13–15,32} in pcDNA3.1 TOPO by PCR. For RLuc8^C-ER (the C-terminal half of the RLuc8 targeted to ER), DNA sequences encoding a.a. 92-311 of the RLuc8 were fused to the 5' end of the myc tag sequence and the previously described ER localization sequence of the yeast UBC6^{13,16,32} by PCR in pcDNA3.1 TOPO. For Reep1^{116–157}RLuc8^N used for subcellular localization study, DNA sequences encoding RLuc8^N (a.a. 1–91) was fused to the 3' end of the DNA sequences encoding REEP1^{116–157} by PCR in pcDNA3.1 TOPO. All constructs and mutations were confirmed by sequencing. ER-dsRed (pDsRed2-ER) and GFP-Sec61β (Plasmid #15108) were purchased from Clontech Laboratories and Addgene, respectively.

Cell culture and transfection

For immunofluorescence studies, HeLa cells were plated at a density of $1.5\text{-}2\times10^4$ cells/well in 8 well-chamber slides (Thermo Scientific). 16 hours after plating, cells were transfected via polyethylenimine (PEI, High Potency Linear, 40 kDa, Polysciences #24765)-mediated transfection with 200 ng of total DNAs per well. 24 hours after transfection, cells were fixed and used for immunofluorescence studies. For Western blot analysis, HEK293T or HeLa cells were plated at a density of 2×10^5 cells/well in 6 well-plates. After overnight

incubation, cells were transfected via PEI-mediated transfection with 2-3 ug of total DNAs per well. For primary cortical cultures, dissociated cell cultures were generated from E15.5 CD1 mouse embryos as described with minor modifications. ^{17–20,33} Briefly, the cerebral cortex was dissected out in Hank's Balanced Salt Solution buffered with 0.1 M HEPES (pH 7.3) and dissociated with 0.25% trypsin-EDTA (Life Technologies) for 20 min at 37°C, followed by trituration. Dissociated cells were initially plated in neural plating medium (MEM, 10% FBS, 0.6% D-Glucose (w/v), and 10 U/mL penicillin-streptomycin) at a density of 5–8×10⁴ cells/well on 24-well plates with coverslips (12 mm, Fisher Scientific) coated with poly-D-lysine (100 µg/mL). Five hours after plating, the medium was replaced with pre-conditioned neural maintenance medium (Neurobasal medium with B-27 supplement, Glutamax-I supplement, and 10 U/mL penicillin-streptomycin), which was prepared from 7–10 day old astroglial culture (1 day incubation). For primary cortical neuron transfections, the Ca²⁺-phosphate mediated method was used on 3-day old cultures as previously described with slight modification. ^{21,34} A total of 0.5–1 µg of DNA was used for 12 mm coverslips in 24 well plates. Twenty min before the transfection, the neural maintenance medium was changed to transfection medium (same as neural maintenance medium but without penicillin-streptomycin). Four to five days after transfection, neurons were fixed with 4% paraformaldehyde and used for immunofluorescence analyses. The transfection rates among the *Reep1* constructs in the primary cells were similar to each other.

RNA in situ hybridization

Whole mount (E9.5) and section (E14.5 and postnatal day 1; P1) RNA *in situ* hybridization using digoxigenin-labeled *Reep1* riboprobes was performed as previously described. ^{22–27,35,36}

Northern Blot analysis

Adult tissue Northern blotting was performed as previously described using mouse multipletissue Northern blot (Clontech), ULTRAhyb (Ambion) and *Reep1* RNA probes.²⁸

Quantification of degenerating neurites and Sholl analysis

For quantification of degenerating neurites, sixteen 20× (objective) fields per sample were blindly picked and imaged using Zeiss Zen Pro software connected with Hamamatsu ORCA-Flash 4.0 camera attached to a Zeiss Observer Z1 inverted microscope. A 5×5 grid (the area of each square in the grid = 5000 µm²) was superimposed on each image with Grid plug-in for Fiji. Neurites in each square of the grid were manually scored as either whole or beaded as previously described, ^{29,37} and values were expressed as a percentage of degenerating neurites (beaded neurites) among total transfected neurites (sum of beaded and whole neurites). Values from each square (total 16 squares) were averaged for each image, and the resulting values from each image (total 10 images) were averaged for each construct (n=160 per construct from two independent experiments). For Sholl analysis with *Reep1* constructs, the entire coverslip (12 mm) was imaged using Zeiss Zen Pro software connected with Hamamatsu ORCA-Flash 4.0 camera attached to a Zeiss Observer Z1 inverted microscope (10× objective). 30 neurons from each sample (10 neurons per coverslip, three

independent experiments) were randomly picked and cropped out, neurites of each neuron were traced using Simple Neurite Trace plug-in for Fiji, and automated analysis was carried out using Sholl Analysis in Fiji. 30,31,38 For Sholl analysis with shRNAs, 30 to 40 neurons from each sample were randomly picked for imaging (Image-Pro software connected with Hamamatsu C5810 camera attached to a Nikon TE-300 microscope), each image was overlaid with concentric circles (25 μ m intervals) generated in Photoshop, and Sholl analysis was performed manually by counting the numbers of neurites that intersected each circle.

Biochemical fractionation of the mouse brain

Embryonic mouse brains (E17.5) were fractionated into cytosol, crude mitochondria, pure mitochondria, ER, and MAM fractions as previously described. 32,39,40

Western blot

HEK293 or HeLa cells were transfected with DNA constructs or shRNA constructs as indicated. At 48 or 72 hrs after transfection, the cells were lysed in lysis buffer (20 mM Tris-HCl [pH 7.4], 150 mM NaCl, 0.5% TritonX-100) containing a protease inhibitor cocktail (Roche). 10–20 µg of total protein was loaded in each lane, separated by 10% or 4–12% SDS/PAGE and blotted onto PVDF membrane. The blots were blocked with 5% milk, followed by incubation with indicated primary antibodies including anti-V5 antibody (Invitrogen, mouse monoclonal, 1: 5000), anti-GFP antibody (Santa Cruz, rabbit polyclonal, 1:1000), anti-REEP1 antibody (Sigma-Aldrich, rabbit polyclonal, 1:1000), or anti-α tubulin antibody (Santa Cruz, mouse monoclonal, 1:1000), and incubation with HRP-conjugated goat anti-rabbit or mouse IgG secondary antibody (1:5000, Jackson Immunology). For subcellular fractionation samples, 30 µg of protein for each fraction was loaded in each lane and separated by 4-12% SDS/PAGE. The primary antibodies used for subcellular fractionation samples include anti-ATP5A (Abcam, mouse monoclonal, 1:1000), calnexin (BD Biosciences, mouse monoclonal, 1:2000), sigma-1R (ProteinTech, rabbit polyclonal, 1:2000), and REEP1 (Sigma-Aldrich, rabbit polyclonal, 1:1000) antibodies. Blots were developed using ECL kit (SuperSignal West Pico- or Femto Chemiluminescent Substrate, Thermo Scientific) and scanned using ChemiDocTMMP imaging system (Bio-Rad). Quantification of REEP1 expression was normalized to GFP expression using Image Lab software (Bio-Rad).

Immunofluorescent staining and MitoTracker treatment

For immunofluorescent staining, HeLa cells were fixed with 4% paraformaldehyde for 10 min or with ice-cold methanol for 5 min at room temperature, rinsed with PBS, and treated with 0.1% triton X-100 for 5 min. After blocking in 10% goat serum in PBS, cells were incubated with a primary antibody, including anti-REEP1 (Sigma-Aldrich, rabbit polyclonal, 1:50), calreticulin (abcam, chicken polyclonal, 1:5000), or TOM20 (Santa Cruz Biotechnology, rabbit polyclonal, 1:500) antibodies at 4°C overnight. After rinsing with PBS, cells were incubated with appropriate secondary antibodies conjugated with a fluorescent dye (Alexa-fluor 488 or Alexa-fluor 594; Invitrogen) for 1 hour at the room temperature. After rinsing with PBS, cells were incubated with DAPI (Molecular Probes) for nuclear staining, and mounted with mounting medium (Southern Biotech). For MitoTracker treatment, before the fixation step, cells were incubated with 100 nM of MitoTracker® Red

CMXRos (Invitrogen) for 30 min and then incubated with new medium for 30 min to wash out unincorporated residual MitoTrackers from the cells. Images were captured with Zeiss Zen Pro software using Hamamatsu ORCA-Flash4.0 camera attached to a Zeiss Observer Z1 inverted microscope, or Zeiss LSM 710 confocal microscope.

Split-RLuc8 reassembly assay

HEK293T cells (American Type Culture Collection, Manassas, VA) were cultured in Dulbecco's modified Eagle's medium (Life Technologies) with 10% fetal bovine serum (Life Technologies) at 37°C with 5% CO₂. Cells (4×10^5 /well) were plated in 12-well culture plate the day before transfection. Expression constructs, *Mito-RLuc8*^N and/or *RLuc8*^C-*ER* were co-transfected with the indicated constructs into HEK293T cells using PEI (Polysciences). The cells were split into 96-well plate (4×10^4 /well) at 6 hr post-transfection and further incubated for 24 hrs at which time Enduren live cell substrate (Promega) was added in culture medium for 2.5–3 hrs (30 μ M). The luminescence was measured by POLARstar Omega microplate reader (BMG LABTECH).

Statistical analysis

Prism 6 statistical software (GraphPad) was used to generate charts, and unpaired t-test (two-tailed) with Welch's correction was used to analyze statistical difference. P < 0.05 was considered statistically significant.

RESULTS

Reep1 is expressed in the brain, nerve ganglia, spinal cord, and somites during development

We identified *Reep1* in a previous screen for novel genes involved in mouse forebrain development. ^{28,32} To validate the screen and begin characterizing a function for REEP1, we first analyzed the expression of Reep1 by RNA in situ hybridization. On embryonic day 9.5 (E9.5), mouse *Reep1* was detected in the brain, spinal cord, peripheral nerves, and somites (Fig 1A and B). Its expression in the brain was observed in the telencephalon, diencephalon, and rhombencephalon. Reep1 was also present in the spinal cord and several cranial nerve ganglia including the trigeminal, facio-acustic, glossopharyngeal, and vagal ganglia as well as the dorsal root ganglia (Fig 1A and B). Its expression in the muscle tissues, such as the erector spinae muscle (es mscl) and psoas major muscle (p mscl), were also detected as seen in E13.5 embryos (Fig 1C). By E14.5, Reep1 showed prominent expression in the mantle zone of the telencephalon and diencephalon, where the postmitotic neurons reside, with little or no expression in the proliferative ventricular zone (Fig 1D). In the forebrain of the neonate (P1), Reep1 transcripts were predominantly detected in the neocortex, hippocampus, and thalamus, while weaker labeling was found in the amygdala and hypothalamus (Fig 1E). A Northern blot analysis using adult mouse tissue showed that Reep1 is expressed in the brain as well as in the heart, skeletal muscle and testis (Fig 1F).

HSP-associated Reep1 mutations lead to neuritic degeneration in mouse cortical neurons

Mutations in *REEP1* are known to cause hereditary spastic paraplegia with more than 40 mutations having been identified to date (Figure 1G). Axonal degeneration, predominantly

in cortical motor neurons, is a common pathological feature of HSP.^{33,41} In order to model REEP1-associated degeneration in neurons, mouse primary cortical cultures were transfected with a construct expressing wild type mouse Reep1 or a construct expressing an HSP-associated mutant Reep1 in pCAG-IRES-GFP vector. The HSP-associated mutant Reep1 constructs included two point mutation constructs, Reep1 P19R and Reep1 A20E , and three truncated constructs, $Reep1^{1-115}$, $Reep1^{1-139}$, and $Reep1^{1-176}$, corresponding to the five human mutations, c.56C>G (p.P19R), c.59C>A (p.A20E), c.345C>A (p. Y115X), c. 417+1G>T (p.K139fs), and c.526delG (p.G176fs)(Fig 2A)^{12,13,34,42,43}. Beaded neurites, which are early signs of neurodegeneration, ^{35–37,44} were observed in neurons transfected with mutant Reep1 constructs, as indicated by GFP (transfected cells) and Tuj1 (neuronspecific class III beta-tubulin) expression (Fig 2B). The degree of neurite degeneration was quantified as the percentage of beaded neurites among total transfected neurites (vector, 3.99 \pm 0.52; $Reep1^{P19R}$, 6.32 \pm 0.69; $Reep1^{A20E}$, 11.27 \pm 0.98; $Reep1^{1-115}$, 14.46 \pm 1.46; $Reep1^{1-139}$, 23.19 ± 2.20; $Reep1^{1-176}$, 23.69 ± 1.90; $Reep1^{WT}$, 18.36 ± 2.31; values are in percentage; n=160 per construct) (Fig 2C). Reep1 mutant constructs showed a significant, though varying, increase in the percentage of degenerating neural processes when compared to a vector construct encoding GFP alone. To control for the variation in protein expression levels, we normalized the percentage of degenerating neuronal processes to the level of protein expression, which was quantified by Western blot (Fig 2D, upper panel) (Reep1^{P19R}, 17.35 ± 1.90 ; $Reep1^{A20E}$, 35.52 ± 3.08 ; $Reep1^{l-176}$, 14.05 ± 1.11 ; $Reep1^{WT}$, 11.19 ± 1.40 ; values are in arbitrary unit). The REEP¹⁻¹¹⁵ and REEP1¹⁻¹³⁹ truncated mutant proteins are not recognized by REEP1 antibody due to a deletion of the epitope region, thus normalization was not performed with these constructs. To determine if these two constructs were expressed, we generated GFP fusion constructs, expressed them in cell lines (HEK293T and HeLa) and Western blotted with a GFP antibody (Fig 2D, lower panel). After normalization, the neuritic degeneration rates of the neurons expressing REEP1P19R and REEP1A20E were significantly increased compared to that of the neurons expressing wild type REEP1 (P=0.0187 and P < 0.0001, respectively). Our data provide evidence that disease-associated Reep1 mutations result in neuritic degeneration in mouse cortical neurons, possibly modeling human mutations with a similar degeneration phenotype.

HSP-associated *Reep1* mutations result in defective neurite growth in mouse cortical neurons

Several of the abnormalities observed in HSP patients, including the thin corpus callosum, have been proposed to be developmental in origin and potentially reflect a defect in neurite outgrowth. A,28,45,46 More recently, it has been reported that neurons derived from human pluripotent stem cells (hiPSCs) from HSP patients carrying *SPASTIN* (*SPG4*) mutations have a decreased neuritic complexity. To investigate whether *Reep1* mutations associated with HSP would also result in neuritic growth abnormalities, we transfected cortical neural cultures with wild type or mutant *Reep1* expression constructs. *Reep1* mutant constructs led to reduction in neuritic complexity when compared to that of the wild type or vector construct; similar to what had been observed in neurons derived from *SPG4* hiPSCs (Fig 3A). Individual neuronal images were used to trace neurites semi-automatically (Fig 3B) and measure the level of neuritic complexity by Sholl analysis (See Materials and

Methods for detail) (Fig 3C). Neurons expressing disease-associated *Reep1* mutant constructs had a significantly decreased number of crossings of the proximal neurites compared to those expressing a control vector or wild type *Reep1* (Fig 3C). The effect on distal neurites was not significant, although highly variable between mutations.

In order to exclude an overexpression artifact, we also transfected primary neurons with *Reep1* specific shRNAs or control shRNA. Neurons transfected with *Reep1* shRNA1, the most effective knock down construct (Fig 4B), exhibited a decrease in neuritic branch formation (Fig 4A). The Sholl analysis shows that the number of crossings in *Reep1* shRNA1 expressing neurons is significantly reduced compared to the control or shRNA2 (Fig 4C). These data confirm that the endogenous REEP1 is necessary for normal neuritic branching and growth, and that neuritic defects observed with mutant REEP1 are not simply due to an overexpression artifact. These data suggest that mutations in *REEP1* also impair neurite development, a finding that might explain the thin corpus callosum found in some patients.

REEP1 is detected in the ER-mitochondria contact sites known as mitochondriaassociated ER membranes (MAMs)

To explore how *Reep1* mutations might give rise to the observed neuritic defects, we next examined the cellular function of REEP1. We first determined the subcellular localization of REEP1. It was initially reported as a mitochondrial protein, but later studies localized it in the ER. 5,12,3839,40,48 The possibility of it being in the ER-mitochondria interface had been raised but never tested. 13,16,28 Thus, we took a biochemical approach to determine if REEP1 is detected in the ER-mitochondria contact sites known as mitochondria-associated ER membranes (MAMs). Mouse embryonic brain extract (E17.5) was used for subcellular fractionation and assayed for the presence of REEP1 along with known markers in each fraction (Fig 5A). As expected, calnexin, an ER protein known to be enriched at the MAM, ^{19,40,41} was present in the ER and MAM fractions but little in the pure mitochondrial fraction. Conversely, ATP5A, a mitochondrial protein, was predominantly detected in the pure mitochondrial fraction, but none in the ER. For REEP1, it was mainly detected in the ER but also significantly in the MAMs, providing evidence that it is present in the ERmitochondria interface. Low level of REEP1 was also detected in the pure mitochondrial fraction, most likely due to small amounts of ER contamination, although we cannot rule out the possibility that some REEP1 might be present in the mitochondria.

We next sought to determine whether REEP1 can be targeted to the mitochondria. We expressed a series of GFP-tagged *Reep1* constructs (see schematic diagram in Fig 5B) in HeLa cells and examined their subcellular localization. Both full-length REEP1 as well as an N-terminal protein, REEP1¹⁻¹¹⁵, were detected in the ER as determined by overlap with ER-dsRed (Fig 5C) or calreticulin (Fig 8B); both ER markers. In contrast, the C-terminal protein, REEP1^{116–201}, targeted to the mitochondria as demonstrated by its overlap with TOM20, a mitochondrial marker (Fig 5D). With another GFP-tagged deletion construct, *Reep1*^{116–157}, we further honed down the domain required for mitochondrial localization to a.a. 116–157 (Fig 5F). Remarkably, when the amino acids 116–201 or 116–157 of REEP1 were attached to another protein sequence (*Renilla* luciferase 8), it was sufficient to target

the heterologous protein to the mitochondria (data not shown), confirming the specificity of the mitochondrial targeting of the REEP1^{116–201} or REEP1^{116–157} independent of the tag. As the C-terminal domain of REEP1 is known to bind microtubules, we next examined whether REEP1^{116–201} or REEP1^{116–157} localization in the mitochondria is simply due to their alignment with microtubules. Co-staining with α -Tubulin ruled out this possibility as their mitochondrial expression does not overlap with α -Tubulin staining (Fig5E and 5G). Collectively, these data indicate that the N-terminal domain of the REEP1 (a.a. 1–115) is responsible for ER localization, whereas the middle domain (a.a. 116–157) is sufficient for mitochondrial localization. Together our results provide evidence of REEP1 being localized at the ER and mitochondria interface, potentially reconciling contradicting reports on REEP1 subcellular localization. 5,6,12,13,42,43

REEP1, but not HSP-associated mutations, facilitates ER-mitochondrial contacts

Based on our data indicating that REEP1 resides at the ER-mitochondria contact sites, we hypothesized that REEP1 might play a functional role in ER-mitochondrial contact formation. To test our hypothesis, we developed a 'split-Renilla Luciferase 8 (RLuc8) reassembly assay', a tool to quantitatively measure the level of ER-mitochondria interactions in live cells. RLuc8 was split into N-terminal (RLuc8^N) and C-terminal (RLuc8^C) halves based on the previously determined split site, ^{30,37,44} and each half was conjugated with either a mitochondrial or an ER targeting sequence (see Materials and Methods for detail). The resulting constructs were named *Mito-RLuc8*^N and *RLuc8*^C-ER, respectively (Fig 6A). We verified their targeting to the mitochondria or ER in the cells by transfecting these constructs and determining their co-localization with TOM 20 or calreticulin (Fig 6B). Next, these constructs were transfected into HEK293T cells and luciferase activity measured (Fig 6C). Neither mitochondria-targeted N-terminal RLuc8 (Mito-RLuc8^N), or ER-targeted C-terminal RLuc8 (RLuc8^C-ER) alone produced significant luciferase activity (Fig 6C). However, when presenilin2 (PS2) wild type or T122R mutant, known to facilitate ER-mitochondria interactions, 4,22,45,46 was co-transfected with Mito-RLuc8^N and RLuc8^C-ER, a significant increase in luciferase activity was detected when compared to that of an empty vector (Fig 6C). These results validate our split-RLuc8 assay as a method to functionally assess the interaction between the ER and mitochondria.

Using this assay, we next examined if co-transfection of *Reep1* can increase ER-mitochondria interactions. *Reep1* was co-transfected with the split-*R*Luc8 constructs and luciferase activity was measured (Fig 7A). Wild type *Reep1* significantly increased luciferase activity when compared to the empty vector, indicating that it can indeed facilitate the interaction between the ER and mitochondria. When one of the split-*R*Luc8 constructs was expressed in the cytoplasm instead of being targeted to the ER- or mitochondria, the luciferase activity was minimally increased when compared to wild type (Fig 7B). This result supports that the ER and mitochondria specific localizations of the split-*R*Luc8 constructs are required for the restoration of *R*Luc8 activity. Next, our series of disease-associated *Reep1* mutations, *Reep1*^{P19R}, *Reep1*^{A20E}, *Reep1*¹⁻¹¹⁵, *Reep1*¹⁻¹³⁹, and *Reep1*¹⁻¹⁷⁶ were used in our split-RLuc8 assay. None of these induced luciferase activity to the extent observed with wild type *Reep1* (Fig 7A). *Reep1*¹⁻¹¹⁵, which only includes the ER association domain but not the mitochondrial domain, did not activate luciferase above

baseline. Similarly, $Reep1^{P19R}$ and $Reep1^{A20E}$, which possess point mutations in the putative transmembrane sequence within the ER association domain, 5,12,47 also had relatively little effect. The other two truncated mutant constructs, $Reep1^{I-115}$ and $Reep1^{I-139}$, showed some activity but less than a control. Interestingly, $Reep1^{I-176}$ which contains both ER- and mitochondria- association domains, but not the C-terminal domain known to interact with microtubules, 5,48 resulted in the most significant increase in luciferase activity, reaching more than 50% of the wild type. These results indicate that the disease-associated Reep1 mutations impair REEP1's ability to facilitate ER-mitochondria contact. Furthermore, our data also suggest that in addition to the ER- and mitochondria association domains, the microtubule association domain is necessary for its full activity.

Haploinsufficiency or loss-of-function has been suggested for the underlying molecular genetic mechanism of *REEP1* in SPG31.^{6,13,16,49} The differences in luciferase activity observed in point mutations and truncated mutations (Fig 7A), prompted us to test if *Reep1* mutations also function in a dominant negative fashion. Thus, we examined if *Reep1*^{P19R} or *Reep1*^{P19R} or

Finally, previous studies showed that the ER-mitochondrial contact sites have diverse structural features and in some cases the ER membrane completely wraps around the mitochondria. 19,32,40,50 Thus, maintaining the ER membrane as a curved structure is crucial at these contact sites. Given that REEP1 is thought to shape ER membrane into tubular structures^{5,6,12}, we examined whether the REEP1 mutant proteins retain this function. HeLa cells were transfected with wild type or mutant Reep1 constructs and co-immunostained with calreticulin or GFP-Sec61β marker (Fig 8). Cells transfected with the wild type Reep1 showed a tubular ER pattern as revealed by calreticulin (Fig 8A–F), GFP-Sec61β (Fig 8J– L), and α-Tubulin (Fig 8G–I) staining, consistently with previous reports.⁵ However, the Reep1 mutant constructs, Reep1P19R, Reep1A20E, Reep11-115, and Reep11-139 (Fig 8M-R and data not shown) did not change ER into tubular structures, although $Reep1^{1-176}$ had a mild effect (Fig 8S-U). These results suggest that the ER shaping function of Reep1 might be associated with ER-mitochondrial contact formation directly or indirectly, although further studies are required to confirm this observation. Taken together, our data support a model where REEP1 can facilitate physical ER-mitochondria interactions, possibly through its membrane shaping function, and this function is compromised in disease-associated Reep1 mutations.

Discussion

In this study we have found that REEP1 is present at the ER-mitochondria contact sites, where it can facilitate ER-mitochondria interactions. Furthermore, we found that HSP-associated *Reep1* mutations impair these interactions, possibly explaining defects in neuritic

growth and the degeneration observed in cultured cortical neurons. This novel function provides new insights into the pathogenesis of HSP that will potentially inform new therapeutic options for HSP patients with *REEP1* mutations.

REEP1 subcellular localization and its function at the MAM

When *REEP1* was initially identified as an HSP gene, its protein product was reported to localize in the mitochondria^{12,22}. Later studies, however, showed that REEP1 localized to the ER^{5,6}(ref). It is not clear whether this discrepancy is due to the differences in REEP1 antibody specificity or due to different cell types. Our data show REEP1 contains both a mitochondrial localization domain and an ER localization domain, suggesting the possibility that REEP1 associates with mitochondria as well as ER, and potentially reconciles the previous conflicting data,^{5,6,12} although it is not direct evidence.

Our results also suggest that REEP1 is present and functions in the MAMs. How does REEP1 facilitate the ER-mitochondria interactions? At least three possibilities exist. First, it could directly act as a tethering protein, bridging the two organelles through the formation of homodimers, heterodimers or oligomers. Supporting this model, REEP1 is able to form oligomers and is known to interact with HSP related proteins, spastin and atlastin-1, through hydrophobic domains. ^{5,6,49} Based on our data, it is also reasonable to posit that ER- and mitochondria- localized REEP1 can form dimers with each other, thus bridging the two organelles, similar to MFN2. ^{32,50,51}

Second, REEP1 might indirectly facilitate ER-mitochondria interactions through bending ER membranes, making it topologically possible for the ER to wrap around mitochondria. Ultrastructural data indicate that ER-mitochondria contacts are diverse: in most cases ER tubules form single contacts covering approximately 10% of the mitochondrial surface, while in other cases the ER tubules cover one-half or even completely envelope the mitochondria. 5,32,52,53 Thus, it is conceivable that the membrane shaping function of REEP1 could indirectly enable ER to form contacts with mitochondria.

Finally, REEP1 could function as a molecular chaperone or adaptor that stabilizes or distributes other proteins, i.e. tethering proteins, at the MAM, thus indirectly facilitating ER-mitochondria interactions. A molecular chaperone function of REEP1 has been suggested based on its sequence homology to the plant stress-induced chaperone, HVA22s. ^{5,54} In fact, there are multiple molecular chaperones enriched at the MAM, including sigma-1R, BIP, GRP75, and HSP60; REEP1 could function as another. ^{12,18} The three roles described here appear to be most reasonable, although other possible functions cannot be excluded, and future studies will be directed at elucidating the precise function.

Altered ER-mitochondria contacts and neurodegeneration

Physical contacts between the ER and mitochondria have been observed and functionally implicated in axons, dendrites, and neuronal soma. $^{5,6,55-58}$ For example, in the synapse, ER-mitochondrial contact functions in intracellular Ca^{2+} signaling and for maintaining and modulating synaptic activity. 5,6,12,56 More recently, disruptions in ER-mitochondrial contacts have been proposed as a pathogenic mechanism leading to several

neurodegenerative diseases. 5,21 Presenilins, whose mutations are associated with familial forms of Alzheimer's disease (AD), are enriched in MAMs. 51,59 The T122R mutation in presenilin 2 (PS2-T122R) increases ER-mitochondria interactions, resulting in elevated Ca^{2+} transfer from the ER to mitochondria and leading to chronic mitochondrial Ca^{2+} overload, which is believed to damage metabolic function and eventually cause neuronal death. 18,22,32,52,53 In a similar fashion, α -synuclein, parkin, and DJ-1, whose mutations are responsible for familial forms of Parkinson's disease (PD), and VAPB, whose mutations are associated with amyotropic lateral sclerosis (ALS), are also detected in the MAM fraction and their disease mutations alter ER-mitochondria interactions. $^{25,26,54-58,60-62}$

The axonal injury model has been used to investigate how alterations in ER-mitochondria interactions might lead to degeneration. A mechanical axotomy disrupts Ca²⁺ equilibrium in the axon, activating plasma membrane channels and inducing Ca²⁺ release from the axonal ER. Ca²⁺ release can be buffered by ER-associated mitochondria. Increased contacts between them will lead to mitochondrial Ca²⁺ overload, which eventually triggers opening of the mitochondrial permeability transition pore (mPTP). mPTP opening leads to Ca²⁺ overload in the axon, increased generation of reactive oxygen species, and a drop in ATP production. A a result, the direction of the Na⁺/Ca²⁺ antiporter changes, and additional calcium enters from the extracellular space, ultimately leading to axonal degeneration. These data linking ER Ca²⁺ release with mPTP for axonal degeneration are consistent with those obtained in non-neuronal cells leading to cell death, 22,32 and they suggest that a similar cascade might happen in neurodegenerative diseases including *REEP1*-associated HSPs.

The effect of disease-associated *Reep1* mutations on ER-mitochondria interactions, neuritic growth and degeneration

While the disease-associated mutant proteins in AD, PD, and ALS increase ERmitochondria contacts, the opposite is observed with REEP1 disease-associated mutations. Thus, it appears that not only an abnormal increase, but also an abnormal decrease, in ERmitochondria communication might result in neurodegeneration. Support for this result comes from the knock down study of either of the two proteins essential for MAM tethering, sigma-1 receptor and phosphofurin acidic cluster sorting protein-2 (PACS-2), which resulted in neuronal degeneration. ^{25,26,39,60–62} In healthy cells, the ER-mitochondria tethering ensures the propagation of inositol 1,4,5-triphosphate receptor (IP3R)-linked Ca²⁺ signals to the mitochondria to coordinate ATP production and to enable mitochondrial Ca²⁺ buffering. In the situation where the ER-mitochondria coupling is reduced, such as in cells with sigma-1 receptor or PACS-2 knocked down, or cells expressing *Reep1* disease mutations, the Ca²⁺ signal propagation to the mitochondria is suppressed and the Ca²⁺-dependent control of mitochondrial metabolism likely becomes at risk. Consistent with this, cells obtained from HSP patients with REEP1 mutations showed reduced mitochondrial respiratory rate and oxygen consumption. 13,14,55,63,64 In contrast, when the coupling is enhanced, as in the cells with mutations in PS2, α-synuclein, parkin and DJ-1, mitochondria are susceptible to Ca²⁺ overloading and may undergo mPTP permeabilization, thus committing the cells to a cell death pathway. Thus, maintaining proper ER-mitochondria

interactions seems to be pivotal for maintaining normal mitochondrial function and ultimately neuronal integrity.

The most common type of *REEP1* mutations reported to date is small frame shift mutations, which generate pre-mature stops. ^{49,55} For this reason, haploinsufficiency was suggested as a major molecular genetic mechanism of SPG31. A recent *Reep1* knock-out mouse study also supported loss-of-function or haploinsufficency model. ^{6,55,63,64} Interestingly, our data with two point mutations, *Reep1* and *Reep1* and *Reep1* suggest that these non-functional mutant REEP1 proteins, which lost the ability to facilitate ER-mitochondria interactions (Fig 7A), interfere with the function of wild type REEP1, giving a dominant negative effect (Fig 7C). Without neuropathological studies of the postmortem brain or spinal cord tissue from REEP1-associated HSP patients with known genotype, it is difficult to correlate our results with human phenotype. Furthermore, the stability and proper folding of the REEP1 point mutants and truncation mutant proteins need to be assessed in future studies through purification of the REEP1 and folding analyses such as circular dichroism or dynamic light scattering.

In addition to the neuritic degeneration, our Scholl analysis indicates that *Reep1* disease mutations lead to a neuritic growth defect, similar to *Reep1* specific knock down. These data suggest REEP1 is also required for normal neuritic development. In agreement with our results, a recent study using neurons derived from human induced pluripotent stem cells from patients with SPG4 mutations reported a decrease in the complexity of neurites. In contrast, no developmental defects were observed in primary neuronal culture from *Reep1* mutant mice. 6,32 This discrepancy between the knock out mouse study and our results could reflect difference in knock down versus knock out approaches. In shRNA-mediated knock down, Reep1 is abruptly depleted, and cells may not be able to acutely adapt to this change. In contrast, mutant mice never express REEP1 and may compensate through unknown mechanisms.

In conclusion, our data provide compelling evidence that mutations in *Reep1* alter ERmitochondria interactions and that disruption in this cellular process may be a new pathogenic mechanism in *REEP1*-linked HSP. As the deregulation of this interaction between the two organelles is a common pathway in several neurodegenerative diseases, the split-*R*Luc8 reassembly assay we developed should provide a valuable tool for potential drug screening in multiple neurodegenerative diseases.

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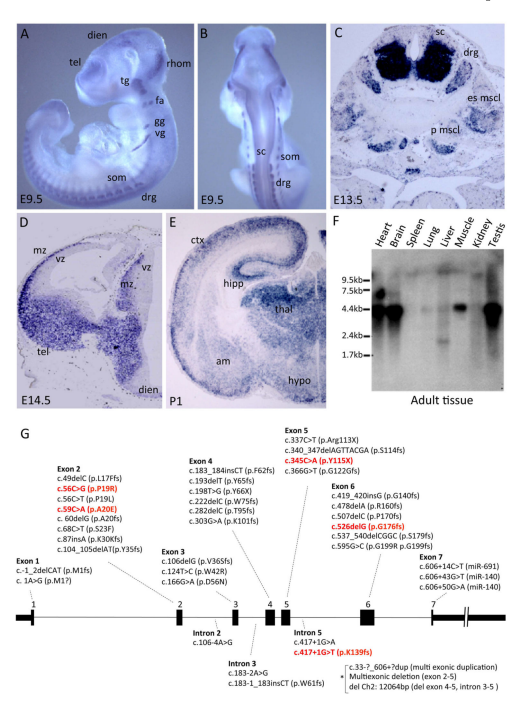


Figure 1. Reep1 expression in the mouse

(A–B) Whole mount RNA *in situ* hybridization of *Reep1* (embryonic day 9.5; E9.5) (A, side view; B, rear view). *Reep1* mRNAs were detected in the brain (telencephalon=tel, diencephalon=dien, rhombencephalon=rhom), spinal cord (sc), peripheral nerve ganglia (trigeminal=tg, facio-acustic=fa, glossopharyngeal=gg, vagal=vg, dorsal root=drg), and somites (som). (C) *Reep1* RNA in situ hybridization on coronal section of the mouse spinal cord (E13.5). *Reep1* is detected in the ventral spinal cord (sc), dorsal root ganglia (drg), erector spinae muscle (es mscl), and psoas major muscle (p muscl) (D–E) *Reep1* RNA in

situ hybridization on coronal sections of the mouse forebrain (E14.5 and P1). *Reep1* is expressed exclusively in the mantle zone (mz) but not in the ventricular zone (vz). Its expression was detected in the neocortex (ctx), hippocampus (hipp), thalamus (thal), amygdala (am), and hypothalamus (hyp). (F) Mouse adult tissue Northern blot analysis detected *Reep1* in the brain as well as in the heart, skeletal muscle and testis. (G) Schematic of the genomic structure (black boxes are exons 1 through 7) with known HSP mutations indicated. ^{12,13,15,39,43,49,65–68} Mutations are distributed over the entire gene (those in red indicate the five *REEP1* mutations modeled in this study). * denotes duplication or deletion mutations spanning more than one exon or intron.

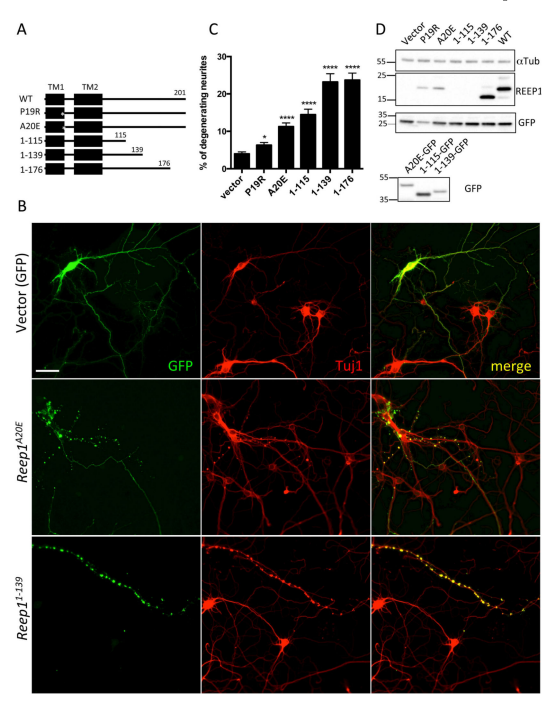


Figure 2. Overexpression of HSP-linked Reep1 mutant constructs leads to neuritic degeneration in primary cortical neurons

(A) Diagram of *Reep1* wild type and mutant constructs. Numbers above the line indicate amino acid residues. TM stands for putative transmembrane domain, and asterisks indicate the locations of the point mutations. (B) Representative images of the primary neurons transfected with the indicated *Reep1* constructs or control vector. GFP staining indicates transfected cells and Tuj1 antibody was used for neuron-specific class III beta-tubulin staining. Beaded neurites indicate neuritic degeneration. Scale bar is 50 µm. (C)

Quantification of the beaded neurites of the primary cortical neurons transfected with the indicated DNA constructs. Bars represent mean value \pm SEM (*, P=0.0159; ****, P<0.0001; unpaired t-test; n = 160 per construct). (**D**) A representative Western blot (upper panel) shows expression levels of each Reep1 construct (in pCIG) used to transfect primary culture. As REEP1 antibody does not recognize REEP1¹⁻¹¹⁵ and REEP1¹⁻¹³⁹, GFP-tagged *Reep1* constructs were used to check their expression in Western blot (lower panel). The transfection efficiency of each construct is comparable. α Tub: α -Tubulin

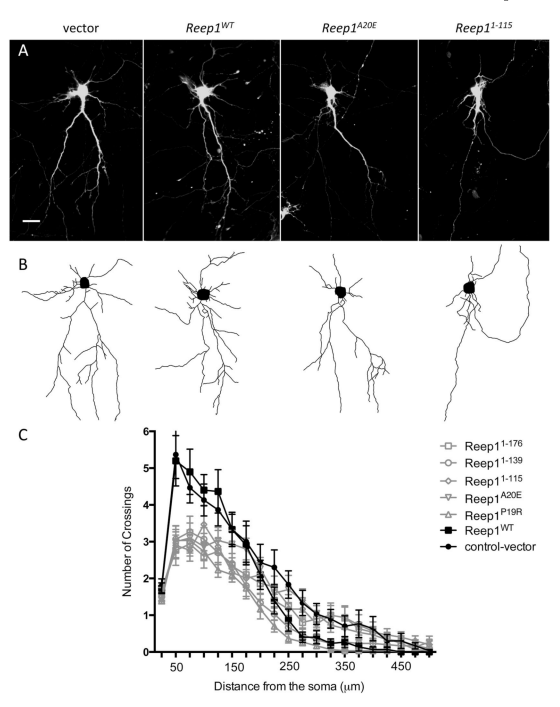


Figure 3. Overexpression of HSP-linked Reep1 mutant constructs leads to defective neurite growth in primary cortical neurons

(A) Representative images of the primary cortical neurons transfected with GFP-tagged control vector, $Reep1^{WT}$, $Reep1^{A20E}$ or $Reep1^{I-115}$, and stained with GFP antibody. (B) Neurite tracings of the corresponding images in (A) using Simple Neurite Tracing plug-in for Fiji. Note some of the GFP positive neurites belonging to adjacent neurons were not traced. (C) Sholl analysis profiles of the primary cortical neurons transfected with the indicated constructs (n=30 neurons per construct, 3 independent experiments). Intersections

were counted at 25 μm intervals from the soma center to a radius of 500 μm . Curves represent mean intersection values \pm SEM. Scale bar in A is 50 μm .

Α

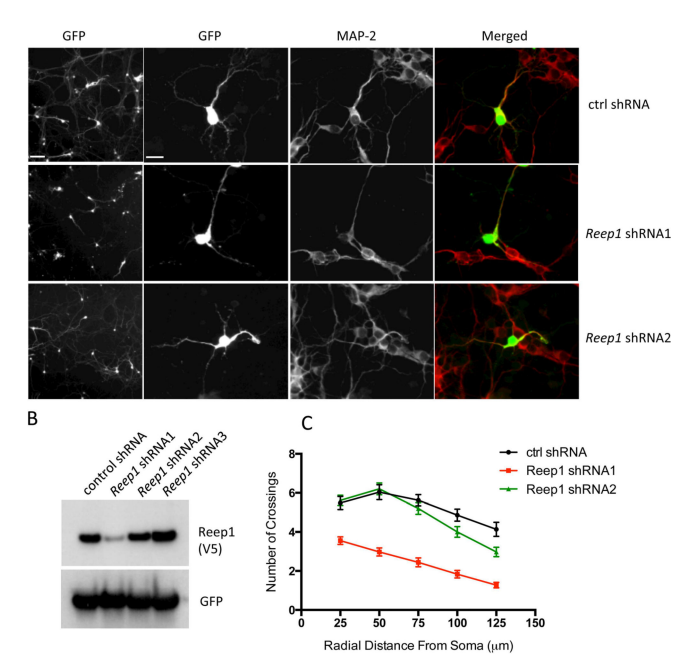


Figure 4. The effect of endogenous REEP1 on neurite growth in primary cortical neurons (A) Representative images of the primary cortical neurons transfected with control shRNA or *Reep1* specific shRNA1 or shRNA2 (see panel B below). Low (left panels) and high magnification (second panels from left) images of GFP fluorescence identify transfected cells. MAP-2 antibody was used for neuron-specific microtubule staining. Neurons transfected with the *Reep1* shRNA1 had more simplified neurite morphology compared to those with the control shRNA. (B) Western blot to detect the level of REEP1 after cotransfecting *Reep1-V5* with the indicated shRNA constructs to HEK293T cells. *Reep1-*

specific shRNA1 was the most effective while the other two shRNAs had little or low activity. (C) Sholl analysis profiles of the primary cortical neurons transfected with the indicated shRNA constructs. Intersections were counted at 25 μ m intervals from the soma center to a radius of 125 μ m. Curves represent mean \pm SEM (n=30 neurons). Scale bars in A are 100 and 50 μ m (from the left).

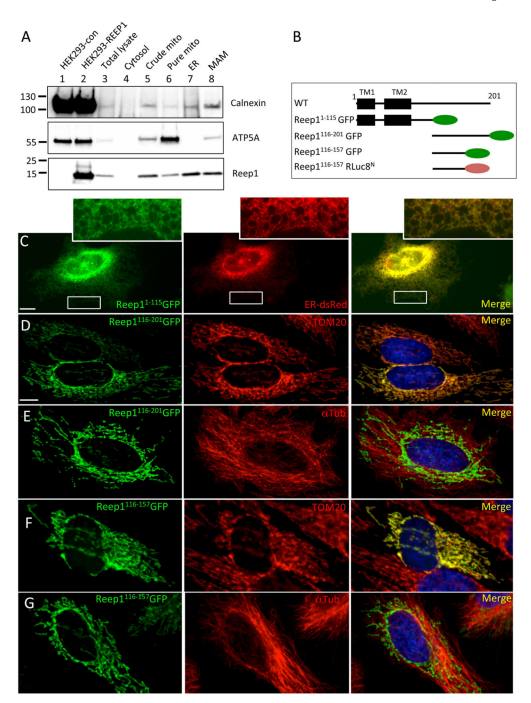


Figure 5. REEP1 is detected in the mitochondria, ER, and ER-mitochondrial contact sites $(M \Delta M)\,$

(A) Western blot using standard subcellular fractionation of mouse brain homogenate (E17.5) into cytosolic, crude mitochondrial, pure mitochondrial, ER, and MAM fractions (ATP-5A, mitochondrial marker; calnexin, ER and MAM marker). REEP1 was mainly detected in the ER and MAM fractions. (B) Schematic diagram of *Reep1* wild type and mutant constructs. Numbers above the line indicate amino acid residues. TM stands for putative transmembrane domain and oval shapes depict GFP (green) or RLuc8^N (brown).

(C) HeLa cells transfected with the $Reep1^{1-115}$ -GFP construct show co-localization with the ER (ER-dsRed), whereas the $REEP1^{116-201}$ -GFP (D, E) or $Reep1^{116-157}$ -GFP (F, G) is colocalized with mitochondria (TOM20) (D, F), but not with microtubules (α Tub) (E, G). Images of each inlet in C are enlarged images of the boxed areas below. α Tub: α -tubulin. Scale bars in C and D are 10 μ m.

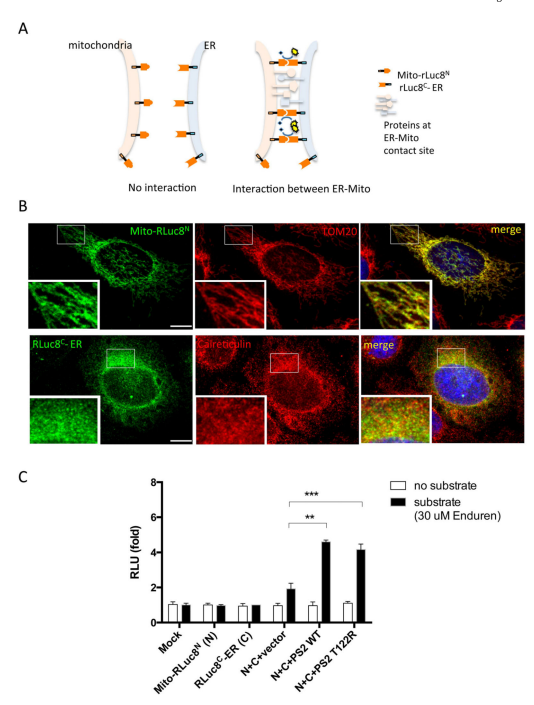
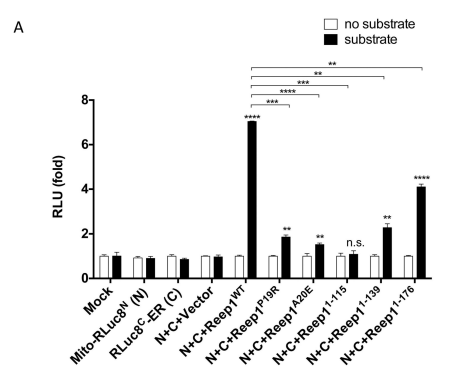


Figure 6. Split-RLuc8 reassembly assay can measure the level of ER-mitochondria interactions (A) A schematic model depicting the split-RLuc8 reassembly assay. Where no interaction between ER and mitochondria exists, Mito-RLuc8 N and RLuc8 C -ER fail to interact and thus do not reconstitute RLuc8 activity. In contrast, where the ER and mitochondria interact, Mito-RLuc8 N moves into proximity of RLuc8 C -ER reconstituting full enzymatic activity, which is detected by the luminescence conversion of substrate. (B) HeLa cells expressing Mito-RLuc8 N were immunostained with Flag (tagged to Mito-RLuc8 N)-antibody together with TOM20. For RLuc8 C -ER, HeLa cells expressing RLuc8 C -ER and ER-dsRed were

detected with Myc (tagged to $RLuc8^C$ -ER)-antibody immunostaining and ER-dsRed fluorescence. (C) Split-RLuc8 reassembly assay performed in HEK293T cells transfected with the indicated constructs in the presence or absence of the substrate. Relative luminescence was measured 24 hrs after transfection (RLU stands for relative luminescence units). Abbreviations: Mito-RLuc8 N , mitochondria targeting sequence- N terminal half of the Renilla luciferase 8; RLuc C -ER, C terminal half of the Renilla luciferase 8-ER targeting sequence; PS2 WT, presenilin 2 wild type; PS2 T122R, presenilin 2 T122R mutant. Bars represent mean \pm SD (**, P = 0.0024; ***, P = 0.0009; unpaired t-test; P = 0.0024; ***



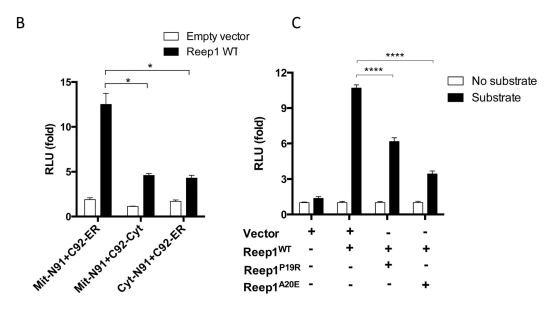


Figure 7. Wild type REEP1, but not REEP1s with HSP-associated mutations, facilitates ER-mitochondria contact formation

(A) Split-RLuc8 reassembly assay performed with the indicated constructs. Wild type Reep1 shows the greatest increase in luminescence, while Reep1s with mutations either have little or no effect. Bars represent mean \pm SEM. The statistical difference was analyzed using unpaired t-test between the vector versus each Reep1 construct (asterisks marked on top of each bar; ****, P < 0.0001; ***, P = 0.0018; **, P = 0.0082; ns, non-significant; **, P = 0.0080; ****, P < 0.0001, from left to right in the graph; n=3), and between the wild type Reep1 and each mutant construct (asterisks marked on top of each bracket; ***, P = 0.001;

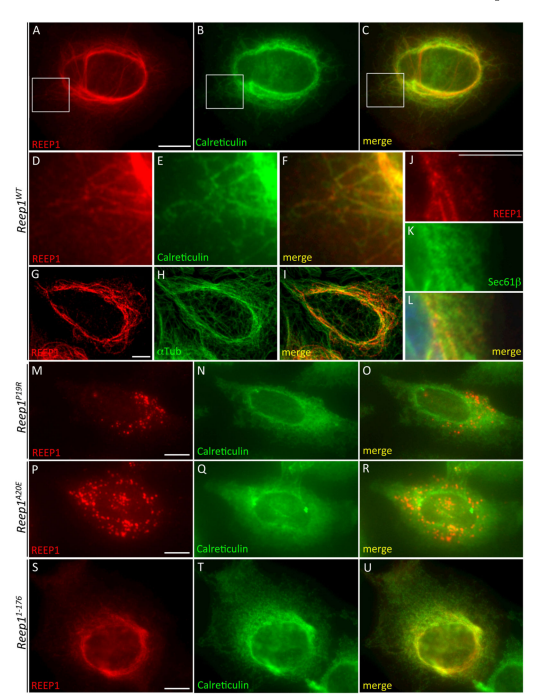


Figure 8. REEP1 disease mutants do not shape ER into tubular pattern unlike wild type REEP1. HeLa cells expressing each indicated construct on the left were immunostained with REEP1 (**A**, **D**, **G**, **J**, **M**, **P**, **S**) and calreticulin (**B**, **E**, **N**, **Q**, **T**) or α -tubulin (**H**), or detected with GFP-Sec61 β fluorescence (**K**). Wild type REEP1 (**A**–**L**) changed ER pattern into tubular whereas mutant REEP1s (M–U) did not, except REEP1^{1–176}, which showed mild changes. Scale bars are 10 μ m.