Torsin Mediates Primary Envelopment of Large Ribonucleoprotein Granules at the Nuclear Envelope

Vahbiz Jokhi,1,4 James Ashley,1,4 John Nunnari,1 Akiko Noma,2 Naoto Ito,3 Noriko Wakabayashi-Ito,3 Melissa J. Moore,2 and Vivian Budnik1,*

1Department of Neurobiology
2RNA and Neuro Therapeutics Institutes, Department of Biochemistry and Molecular Pharmacology, Howard Hughes Medical Institute
University of Massachusetts Medical School, Worcester, MA 01605, USA
3Department of Neurology and Radiology, Massachusetts General Hospital and Program in Neuroscience, Harvard Medical School, Boston, MA 02114, USA
4These authors contributed equally to this work
*Correspondence: vivian.budnik@umassmed.edu
http://dx.doi.org/10.1016/j.celrep.2013.03.015

SUMMARY

A previously unrecognized mechanism through which large ribonucleoprotein (megaRNP) granules exit the nucleus is by budding through the nuclear envelope (NE). This mechanism is akin to the nuclear egress of herpes-type viruses and is essential for proper synapse development. However, the molecular machinery required to remodel the NE during this process is unknown. Here, we identify Torsin, an AAA-ATPase that in humans is linked to dystonia, as a major mediator of primary megaRNP envelopment during NE budding. In torsin mutants, megaRNPs accumulate within the perinuclear space, and the messenger RNAs contained within fail to reach synaptic sites, preventing normal synaptic protein synthesis and thus proper synaptic bouton development. These studies begin to establish the cellular machinery underlying the exit of megaRNPs via budding, offer an explanation for the “nuclear blebbing” phenotype found in dystonia models, and provide an important link between Torsin and the synaptic phenotypes observed in dystonia.

INTRODUCTION

Polarized assembly of cellular complexes often depends on formation of translationally silent RNA transport granules containing mRNAs and associated structural and regulatory components (e.g., proteins and microRNAs). These RNA-protein complexes (RNPs) are shuttled to distinct cellular locales where, upon specific stimuli, the messenger RNAs (mRNAs) are translated into protein building blocks for local cellular architectures and macromolecular complexes (Richter, 2001). Particularly notable is RNP transport in the nervous system, where long-term changes in synaptic structure and function frame key events enabling organisms to respond to their changing environment. A special case of this adaptation is the ability of organisms to learn and remember (Wiersma-Meems et al., 2005). In these processes, localized translation of mRNAs links synaptic-plasticity-inducing stimuli to the synthesis of effector proteins underlying enduring changes in synaptic structure and function (Barco et al., 2008).

Until recently, it was thought that all mRNA export occurred one molecule at a time through the nuclear pore complex (NPC) (Grünewald et al., 2011; Köhler and Hurt, 2007). However, we recently uncovered a mechanism by which large ribonucleoprotein (megaRNP) granules exit the nucleus via nuclear envelope (NE) budding (Speese et al., 2012), a mechanism previously shown to be utilized for the nuclear export of large herpes-type viral capsids (Maric et al., 2011; Mettenleiter et al., 2006). This budding process and the signaling pathway that it initiates are essential for normal synaptic bouton development at the Drosophila larval neuromuscular junction (NMJ) (Ataman et al., 2006; Mathew et al., 2005; Speese et al., 2012). NE budding entails primary envelopment of viral capsids (Mettenleiter et al., 2006) or megaRNPs (Speese et al., 2012) by the inner nuclear membrane (INM); scission of this envelope from the INM creates a membrane-bound particle within the perinuclear space, which subsequently fuses with the outer nuclear membrane (ONM) to allow nuclear escape of the enclosed material. However, the molecular mechanisms required for primary envelopment, INM scission, and fusion were previously unknown. Here, we identify Torsin, an AAA-ATPase that in humans is linked to both dystonia (Breakefield et al., 2008) and herpes virus nuclear egress (Marc et al., 2011), as a major mediator of primary megaRNP envelopment during NE budding, likely functioning to promote INM scission. In torsin mutants, including those mimicking genetic abnormalities in dystonia patients, megaRNPs accumulate within the perinuclear space and the mRNAs contained within fail to reach synaptic sites, preventing normal synaptic protein synthesis and thus proper synaptic bouton development.

RESULTS AND DISCUSSION

In humans, the dystonia-specific Torsin1A (TOR1A) mutation TOR1AΔE302/303 (also known as TOR1AΔAGAG, referred to as TorsinΔE in this paper) at the DYT1 gene locus is linked to
early-onset primary dystonia (Tanabe et al., 2009). Mouse models expressing TOR1A<sup>ΔE302/303</sup> accumulate abnormal vesicular structures at the NE (Goodchild et al., 2005; Naismith et al., 2004). These NE structures show a striking resemblance to the perinuclear megaRNPs we recently reported in <i>Drosophila</i> (Speese et al., 2012), raising the intriguing possibility that these structures could be related. In cultured Schneider-2 (S2) cells and <i>Drosophila</i> larval muscles, megaRNP clusters at the NE can be marked at the light microscopy level by antibodies to the C terminus of the Wnt receptor DFrizzled2 (DFz2C) and the INM-associated protein Lamin C (LamC). DFz2C and LamC partially colocalize at NE-associated foci (DFz2C/LamC foci) (Mathew et al., 2005; Speese et al., 2012). To determine if NE defects observed in <i>TOR1A</i> mutant animal models reflect defects in NE budding, S2 cells were treated with Torsin double-stranded RNA (dsRNA), targeting the sole <i>Drosophila</i> homolog of mammalian <i>TOR1A</i> (Wakabayashi-Ito et al., 2011) (see Figure S1 for Torsin-dsRNA efficiency). This resulted in significant abnormalities in DFz2C/LamC foci at the NE. In untreated S2 cells, NE-DFz2C foci appear as bright immunoreactive spots embedded in a thickening of the lamina, marked by LamC (Speese et al., 2012) or the B-type lamin LamDm0 (Figure 1A). In contrast, Torsin-dsRNA-treated cells displayed small DFz2C-immunoreactive puncta dotting the NE, and thickenings of the lamina were barely visible or absent (Figure 1B; see below for quantification of this phenotype in vivo).

In mammals, Torsin isoforms are derived from four genes: Tor1A, Tor1B, Tor2A, and Tor3A. The DYT1 mutation in <i>Tor1A</i> specifically affects the neuronal NE (Goodchild et al., 2005), consistent with the belief that dystonia is a disease of the nervous system. This neuronal specificity is likely due to compensation by expression of torsinB in nonneuronal tissues, as knockdown of TOR1B in a DYT mutant background caused NE defects in nonneuronal cells (Kim et al., 2010). In <i>Drosophila</i>, there is a single <i>torsin</i> gene, thus overcoming difficulties associated with redundancy. Moreover, we previously showed that NE budding occurs in several cell types, including larval body wall muscle cells wherein the large nuclei are particularly suitable for high-resolution studies (Speese et al., 2012). In addition, the glutamatergic larval NMJ is a powerful model system in which to understand mechanisms of synapse development and function.

To determine the significance of the S2 cell NE phenotype upon Torsin downregulation, DFz2C/LamC foci were examined in <i>torsin</i><sup>KO78</sup>-null mutants (Wakabayashi-Ito et al., 2011) and in
lava in which Torsin was specifically downregulated in muscles by expressing Torsin RNAi using the muscle-specific Gal4 driver C57-Gal4 (Budnik et al., 1996). As in untreated S2 cells, NE DFz2C/LamC foci were observed in wild-type larvae as DFz2C immunoreactive spots surrounded by a thickening of LamC immunoreactivity (Speese et al., 2012) (Figure 1C). In contrast, in larvae expressing Torsin-RNAi in muscles (Figure 1D) or in torsin-null mutants, DFz2C foci were observed as small puncta decorating the NE but lacking any detectable thickening of the lamina. These phenotypes were quantified by determining the percentage of nuclei containing DFz2C spots surrounded by a thickening of the lamina (normal foci; Figure 1G) and the percentage of nuclei containing small NE-associated DFz2C puncta lacking LamC thickening (Figure 1H). There were highly significant differences between wild-type controls and both torsin-null mutants as well as larvae expressing Torsin-RNAi in muscles (Figures 1G and 1H).

Typical of AAA-ATPases, Torsin contains Walker A and Walker B domains involved in ATP binding and ATP hydrolysis, respectively (Neuwald et al., 1999; Wakabayashi-Ito et al., 2011; Walker et al., 1982), as well as Sensor1 and Sensor2 domains also involved in ATP hydrolysis (Iyer et al., 2004). A conserved amino acid deletion in the Sensor2 domain (TorsinAE; TorsinAE306 in Drosophila) is dominantly linked to dystonia (Ozelius et al., 1997). In addition, an amino acid substitution in the Walker B domain (TorsinE→Q; TorsinE177Q in Drosophila) leads to a Torsin protein that can dominantly bind to its substrate but is unable to hydrolyze ATP and therefore remains bound to this substrate, thus constituting a substrate trap (Goodchild et al., 2005; Wakabayashi-Ito et al., 2011). To determine if TorsinAE or TorsinE→Q transgene expression would also disrupt DFz2C/LamC foci morphology, we expressed these proteins in larval muscles. Expressing TorsinAE mimicked the torsin-null and Torsin-RNAi phenotypes (Figures 1E, 1G, and 1H). In contrast, TorsinE→Q expression resulted in the formation of numerous NE LamC foci, most of which were devoid of DFz2C immunoreactivity (Figures 1F1 and 1G–1I). Careful examination of these depleted LamC foci by confocal microscopy demonstrated that many contained a small DFz2C puncta, but this signal was barely visible (Figure 1F2, arrows). The above phenotypes observed upon expressing TorsinAE and TorsinE→Q were the specific results of the mutations in the Torsin transgenes, as larvae expressing a wild-type Torsin transgene were indistinguishable from wild-type not expressing this transgene (Figures 1G–1I).

To determine the ultrastructural correlates of the above phenotypes, untreated and Torsin-dsRNA-treated S2 cells were examined by transmission electron microscopy (TEM). As previously described (Speese et al., 2012), untreated S2 cells displayed local singlets or clusters of megaRNPs within INM invaginations at discrete regions of the NE (Figures 2A, 2H, and 2I), paralleling light microscopy observations (Figure 1A). In contrast, in Torsin-dsRNA-treated cells, these local megaRNPs at the NE were reduced by ~75% (Figures 2H and 2I), and instead many mega-RNP granules were often observed in rows of singlets lining the perinuclear space (Figures 2B and 2C). In these regions, the perinuclear space appeared distended (green in Figures 2B and 2C) and the ribosome-decorated ONM appeared to evaginate (Figures 2B and 2C). However, NPCs and the rest of the NE appeared normal (Figures S2A–S2D). In addition, the distribution of a number of nuclear proteins, such as the fly Emerin homolog Bocksbeutel (Figures S2E and S2I); dMan1 (Figures S2F and S2J); Otefin, a protein required for NE assembly (Figures S2G and S2K); and the Drosophila homolog of Hsap, Squid, a ribonuclear protein (Figures S2H and S2L), were normally distributed in the mutants. About half of megaRNPs appeared attached to the INM through a collared neck (arrows in Figures 2B, 2C, 2H, and 2I; see Experimental Procedures for definition). Thus, down-regulating Torsin results in abnormal attachment of megaRNPs to the INM, raising the possibility that Torsin could be involved in INM scission after primary megaRNP envelopment.

Corroboration of the above results in vivo was obtained by examining larval body wall muscles of torsin-null mutants. As in S2 cells, megaRNPs tethered to the INM by a collared neck were observed in torsin-null mutant muscles and epithelial cells (Figures 2D, 2E, 2H, and 2I), suggesting this pathway functions in even more tissues than previously characterized. Similarly, muscles expressing TorsinE→Q displayed INM-tethered megaRNPs (Figures 2F, 2H, and 2I). In ~30% of cases, megaRNPs in muscles expressing TorsinE→Q appeared as large (>250 nm), amorphous, dense structures directly apposed to the INM (Figures 2G–2I). Thus, disruption of Torsin function in vivo leads to abnormal megaRNP tethering to the INM.

If Torsin is involved in INM scission during megaRNP primary envelopment, then the substrate trap TorsinE→Q should accumulate at the electron-dense collared necks. This prediction was tested by generating wild-type and TorsinE→Q variants fused to a mini-SOG tag (Shu et al., 2011) at their C termini. Mini-SOG is a flavoprotein derived from Arabidopsis Phototropin 2 that when illuminated by blue light produces oxygen species that can convert diaminobenzidine into an electron-dense precipitate (Shu et al., 2011). We first determined if C-terminally tagged Torsin was localized to the NE. Although mini-SOG excitation results in fluorescence emission, its rapid bleaching upon illumination prevented high-resolution acquisition of images. Therefore, we generated Flag-tagged Torsin constructs and expressed them in S2 cells. The wild-type Torsin-Flag signal localized to bright spots coinciding with Lamin foci at the NE; low levels were also observed at the NE and in the cytoplasm (Figure 3A). In contrast, TorsinE→Q-Flag was observed in a punctate pattern lining the NE (Figure 3B). Consistent with the above observations, S2 cells expressing Torsin-SOG displayed an electron-dense signal at sites of megaRNP occurrence in the NE (Figures 3C and 3E; see Figures S3A and S3B for specificity control). An electron-dense SOG-induced signal surrounded each megaRNP (Figures 3C and 3E; see Figures S3A and S3B for specificity control).
that Torsin is present at sites of NE budding. Further, accumulation of the TorsinE−Q substrate trap protein at collared necks of megaRNPs suggests that these necks represent the normal site of Torsin action and provide evidence that Torsin is involved in scission of the INM during primary envelopment.

Our previous study revealed that in Drosophila larval muscles, megaRNPs contain transcripts encoding postsynaptic proteins, including the PDZ scaffolding proteins Par6 and MAGI (Speese et al., 2012). In the case of Par6, interfering with megaRNP formation by inhibiting the Frizzled nuclear import (FNI) pathway or LamC expression results in decreased NMJ localization of par6 mRNA (Speese et al., 2012), decreased postsynaptic Par6 protein levels (Speese et al., 2012), and marked defects in NMJ structure (Ataman et al., 2006, 2008; Packard et al., 2002; Speese et al., 2012). In particular, under these conditions, NMJs fail to expand normally as muscles grow in size during larval development, and a subset of synaptic boutons (called ghost boutons) remain in an immature state. These ghost boutons fail to recruit postsynaptic proteins and to organize postsynaptic specializations, such as the postsynaptic density and subsynaptic reticulum (Ataman et al., 2006, 2008; Packard et al., 2002; Speese et al., 2012).

The above observations support a model in which alterations in Torsin function inhibit nuclear megaRNP exit by slowing or

**Figure 2. Ultrastructural Organization of NE-Associated megaRNPs Is Disrupted in Torsin Mutations**

(A–G) Electron micrographs of nuclear regions in (A–C) S2 cells, (D, F, and G) larval body wall muscles, and (E) larval epithelial cells showing NE-associated megaRNPs. Red, nucleus; blue, cytoplasm; green, perinuclear space. N, nucleus; C, cytoplasm. (A) Untreated S2 cell showing a normal nuclear focus (arrow) containing electron-dense megaRNP granules. (B and C) NE of Torsin-dsRNA-treated S2 cells displaying megaRNPs tethered to the INM by collared necks (arrows), shown at (B) low and (C) high magnification. ribo, ribosome. (D and E) NE in torsin−null mutants also showing megaRNPs tethered to the INM (arrows). (F and G) NE in muscle cells expressing TorsinE−Q showing the presence of (F) a megaRNP (arrow) tethered to the INM and (G) a large, amorphous megaRNP (arrow) tightly apposed to the INM. mi, mitochondria.

(H) Percentage of megaRNP granules present in INM invaginations (black), with collared necks (blue) and being large and amorphous (red).

(I) Average number of megaRNP granules in INM invaginations (black), with collared necks (blue), being large and amorphous (red), or per focus (gray). Number of granules/foci: [159:36,166:88,207:33,166:88,181:88]. Error bars represent ±SEM; *p < 0.05; **p < 0.001; ***p < 0.0001). Calibrations are 0.5 μm (A, B, and D–G) and 0.2 μm (C). See also Figure S2.
blocking INM scission during primary envelopment. As a consequence, such alterations should result in abnormal transcript localization both in the nucleus and at synaptic sites as well as decreased synaptic protein synthesis, abnormal NMJ expansion, and an accumulation of ghost boutons. To ascertain the localization of megaRNP transcripts known to be present in megaRNPs at the NE, we carried out fluorescence in situ hybridization (FISH) using par6 and magi RNA probes. As previously described (Speese et al., 2012), in wild-type muscles, par6 and magi mRNAs are enriched at NE foci associated with LamC foci or nuclear folds marked by antibodies to LamC (Figures 4A and 4B, top row). In contrast, in torsin-null mutants, par6 and magi FISH signals appeared as foci that, while associated with the NE, were on the cytoplasmic side of the LamC signal (Figures 4A and 4B, bottom row panels; Figure 4C). This is in agreement with the light and electron microscopy studies, showing that altering Torsin function prevents megaRNP nuclear egress and results in megaRNPs remaining attached to the INM within the perinuclear space.

When we examined FISH signals at the NMJ in wild-type controls, par6 mRNA was concentrated at subsynaptic sites as previously reported (Figure 4D, top panels) (Speese et al., 2012). However, this synaptic par6 FISH signal was virtually eliminated in torsin-null mutants (Figure 4D, bottom panels). Similarly, the synaptic localization of magi mRNA was significantly decreased in the torsin mutants (Figure 4E). The marked decrease in par6 and magi mRNA levels at the NMJ appeared specific for the NE-budding pathway, as no significant differences around the NMJ in torsin mutants were observed upon FISH of discs-large (dlg) RNA, which is not associated with nuclear DFz2C/LamC foci (Figures S4A–S4C) (Speese et al., 2012). Thus, in the absence of Torsin function, synaptic mRNAs known to be present in megaRNPs exhibit reduced localization at the NMJ, but a nonmegaRNP mRNA does not.

We also examined Par6 and MAGI protein levels using antibodies specific to Drosophila Par6 (Ruíz-Canada et al., 2004) and MAGI (this report). In wild-type larvae, Par6 and MAGI immunoreactivity localized primarily to the postsynaptic muscle region of the NMJ (Figures 4F and 4G, top panels). In addition, Par6 immunoreactivity was observed in a diffuse manner at presynaptic boutons (marked by the anti-HRP signal), being particularly prominent at presynaptic microtubule bundles (Ruíz-Canada et al., 2004) (Figure 4F, top panels) and at low levels at the muscle cell cortex (Figure 4F, top panels). MAGI immunoreactivity was also observed at presynaptic compartments, but without noticeable concentration at microtubule bundles (Figure 4G, top panels).
In *torsin* null mutants, postsynaptic localization of Par6 immunoreactivity as well as muscle cell cortex signal was severely reduced (Figure 4F, bottom panels; Figure 4H), while presynaptic localization of Par6 at microtubule bundles appeared normal (Figure 4F, arrows). Similarly, postsynaptic MAGI protein localization was severely reduced in these mutants (Figure 4G, bottom panels; Figure 4I). Unlike Par6 and MAGI, DLG immunoreactivity was not changed in *torsin* mutants (Figure 4J; Figure S4D), suggesting that the defect is not general but affects only a subset of postsynaptic proteins. Thus, disrupting Torsin function prevents normal localization of some synaptic mRNAs and, as a consequence, normal postsynaptic levels of their encoded proteins.

The functional consequence on NMJ structure of reduced Par6 and MAGI mRNA and protein levels at the postsynaptic compartment in *torsin* mutants was assessed by counting the

---

**Figure 4. The Distribution of mRNAs at the NE and Synaptic Sites Is Disrupted in *torsin* Mutants**

(A and B) FISH to body wall muscles showing the nuclear distribution of (A) par6 and (B) magi transcripts in wild-type and *torsin* mutants.

(C) Quantification of FISH signal outside the nucleus. N [nuclei;larvae], [18;6],[18;6],[18;6],[14;6].

(D and E) FISH to body wall muscles showing the distribution of (D) par6 and (E) magi transcript at the NMJ in wild-type and *torsin*-null mutants.

(F and G) Distribution of (F) Par6 and (G) Magi immunoreactivity at the NMJ in wild-type and *torsin* mutants.

(H and I) Quantification of postsynaptic (H) Par6 and (I) Magi immunoreactive signal, normalized to wild-type control. N[NMJs;larvae] is [16;6],[18;6] (H) and [17;6],[15;6] (I).

(J) NMJs in wild-type and *torsin* mutants labeled with anti-HRP and anti-DLG showing reduced size and increased ghost boutons (arrowheads) in *torsin* mutants.

(K and L) Quantification of the number of (K) synaptic boutons and (L) ghost boutons. N[NMJs;larvae], [19;10],[18;10],[19;10],[19;10],[20;10],[19;10],[19;10] for (K) and (L). Error bars represent ±SEM; **p < 0.001; ***p < 0.0001.

Calibration scale are 3 μm (A and B), 10 μm (D–G), and 20 μm (J). See also Figure S4.
number of normal and undifferentiated ghost boutons observable in the last (third-instar) larval stage. Interfering with Torsin function resulted in a significantly reduced number of synaptic boutons (Figures 4J and 4K) and a significantly increased number of undifferentiated ghost boutons (Figure 4J, arrowheads; Figure 4L).

Together, these results demonstrate that inhibiting Torsin function results in megaRNP accumulation at the NE, likely due to a defect in INM scission during primary envelopment. As a consequence, synaptic transcript-containing megaRNPs fail to efficiently exit the nucleus, limiting trafficking of the mRNAs contained within to postsynaptic sites where they are normally enriched. This reduced synaptic mRNA localization results in reduced levels of specific postsynaptic proteins during NMJ expansion and thus in poorly developed NMJs containing fewer synaptic boutons and increased numbers of undifferentiated ghost boutons lacking postsynaptic proteins. These results provide mechanistic insight into the molecular machinery underlying nuclear egress of megaRNPs by NE budding. They also provide a mechanism by which Torsin influences synaptic development as well as important clues as to how torsinA dysfunction might lead to the alterations in synaptic plasticity observed in DYT1 mouse models and human patients.

EXPERIMENTAL PROCEDURES

Fly Strains
Flies were reared on standard Drosophila medium at 25°C. RNAi crosses and controls were performed at 29°C.

Molecular Biology
Torsin dsRNA was prepared by amplifying exon1 by PCR and in vitro transcribed using the Ambion MEGAscript T7 kit.

S2 Cell Culture and dsRNA Treatment
Drosophila SL2-NP2 cells were cultured and treated as described in Koles et al. (2012).

Immunocytochemistry
Third-instar larval body wall muscles were dissected and fixed as described in Budnik et al. (1996).

Antibody Generation
Anti-MAGI was generated by immunizing rats with a bacterially produced Magi peptide (amino acids 337–558).

Fluorescence In Situ Hybridization
Procedures for FISH were as described in Speese et al. (2012).

Transmission Electron Microscopy
TEM was performed as described in Ashley et al. (2005).

Diaminobenzidine conversion of Mini-SOG
Diaminobenzidine (DAB) photoconversion was adapted from Grabham and Goldberg (1997).

Image Acquisition
Confocal images were acquired using a Zeiss LSM700 confocal microscope equipped with a Zeiss ×63 Plan-Apochromat 1.4 NA DIC oil-immersion objective.
REFERENCES


