Gamma-1-Syntrophin Mediates Trafficking of Gamma-Enolase towards the Plasma Membrane and Enhances Its Neurotrophic Activity

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Key Words
Gamma-1-syntrophin · Gamma-enolase · Cathepsin X · Intracellular trafficking · Neurotrophic effect

Abstract
Syntrophins are scaffold proteins that can bind several signaling molecules and localize them to the plasma membrane. We demonstrate here that in neuroblastoma SH-SY5Y cells, brain-specific \(\gamma_1\)-syntrophin binds the neurotrophic factor \(\gamma\)-enolase through its PDZ domain, and translocates it to the plasma membrane, as shown by immunoprecipitation, surface plasmon resonance, fluorescence colocalization and flow cytometry. Extensive colocalization of \(\gamma_1\)-syntrophin and \(\gamma\)-enolase was observed in neurite growth cones in differentiated SH-SY5Y cells. Silencing of the \(\gamma_1\)-syntrophin gene by RNA interference significantly reduced the redistribution of \(\gamma\)-enolase to the plasma membrane and impaired its neurotrophic effects. We demonstrated that an intact C-terminal end of \(\gamma\)-enolase is essential for its \(\gamma_1\)-syntrophin-assisted trafficking. The cleavage of two amino acids at the C-terminal end of \(\gamma\)-enolase by the carboxypeptidase cathepsin X prevents binding with the \(\gamma_1\)-syntrophin PDZ domain. Collectively, these data demonstrate that \(\gamma_1\)-syntrophin participates in \(\gamma\)-enolase translocation towards the plasma membrane, a pre-requisite for its neurotrophic activity. By disrupting this \(\gamma_1\)-syntrophin-guided subcellular distribution, cathepsin X reduces \(\gamma\)-enolase-induced neurotrophic signaling.

Introduction
Nervous system function depends upon highly specific connections that are formed between neurons during development [1, 2]. In this process, neurotrophic factors are significant and they have important roles in nerve growth and regeneration, cell differentiation and survival [3, 4]. Among several proteins, a neurotrophic activity has also been demonstrated for the C-terminal part of \(\gamma\)-enolase [5, 6].

Enolase is a glycolytic enzyme that can form homodimers and heterodimers that are composed of \(\alpha\), \(\beta\) and \(\gamma\) subunits [7]. In mammalian brain, there are three enolase isoforms: \(\alpha\alpha\), \(\alpha\gamma\) and \(\gamma\gamma\) [8]. The \(\alpha\gamma\) and \(\gamma\gamma\) isoforms are also known as neuron-specific enolase: they are primarily found in tissues of the central nervous system, and
they have only been localized to neurons and neuroendocrine cells [9]. Previous studies have shown that the C-terminal part of neuron-specific enolase has a neurotrophic activity when it is bound to the plasma membrane of neuronal cells [5, 6, 10]. However, to date, the mechanisms of γ-enolase translocation and membrane association have remained unknown. The neurotrophic activity of γ-enolase can be regulated by cathepsin X [11], a carboxypeptidase that is prominently localized in the lysosomal system, although it is also seen scattered through the cytoplasm in almost all cells in the brain [12]. Cathepsin X has crucial roles in diverse physiological processes, including cell adhesion, phagocytosis and migration, and cytoskeletal rearrangements [13].

The targeting of proteins to specific membrane domains is organised by the scaffold proteins, which are known to enhance the efficiency and specificity of signal transduction [14]. Syntrophins are scaffold proteins with multiple protein interaction domains [15, 16]. The syntrophin family comprises five structurally related isoforms that are encoded by separate genes: α1, β1, β2, γ1 and γ2. Although all of these isoforms share the same modular domain organization, i.e. two tandem pleckstrin homology domains, a single PDZ domain, and a C-terminal region that is unique to syntrophins, each syntrophin has specific tissue and developmental expression patterns [17, 18]. The γ1 isoform of syntrophin is expressed solely in neurons and it binds a unique set of signalling proteins through its PDZ domain [19, 20]. The C-terminal half of γ1-syntrophin remains accessible for interactions with members of the dystrophin family of proteins, including dystrophin, utrophin and dystrobrevin [15, 21].

Using immunoprecipitation, surface plasmon resonance, immunofluorescence and flow cytometry, we demonstrate here that γ1-syntrophin can regulate the subcellular localization of γ-enolase in neuronal cells, and consequently, its neurotrophic activity. Moreover, we implicate the carboxypeptidase cathepsin X in disruption of the γ1-syntrophin/γ-enolase complex that results in impaired translocation of γ-enolase towards the plasma membrane and its decreased neurotrophic activity.

Materials and Methods

Cell Culture and Transfections

The human neuroblastoma SH-SY5Y cells were obtained from American Type Culture Collection (CRL-2266, Manassas, Va., USA). The cells were grown in Dulbecco’s modified Eagle’s medium (Sigma, St. Louis, Mo., USA) supplemented with 10% fetal bovine serum (HyClone, Logan, Utah, USA), 2 mM L-glutamine (Sigma), 50 U/ml penicillin and 50 µg/ml streptomycin (Sigma). The cells were maintained at 37°C in a humidified atmosphere containing 5% CO2 and they were grown until 80% confluence. The SH-SY5Y cells were then transfected with the pcDNA3/ cathepsin X construct using lipofectamine 2000 (Invitrogen, Carlsbad, Calif., USA), according to the manufacturer protocol. A stable cell line overexpressing cathepsin X was selected with 400 µg/ml geneticin (Invitrogen, Gibco, Scotland), with upregulation of cathepsin X activity detected using the cathepsin X-specific, intramolecularly quenched fluorogenic substrate Abz-Phe-Glu-Lys(Dnp)-OH [22]. Cell lysates were prepared in lysis buffer (50 mM sodium acetate, pH 5.5, 1 mM EDTA, 100 mM NaCl, 0.25% Triton X-100) with protein concentrations determined using the Bradford method [23], using the Coomassie Plus protein assay reagent (Thermo Scientific, Rockford, Ill., USA). An aliquot of 100 µg of lysate proteins was incubated at 37°C, followed by measurement of cathepsin X activity using Abz-Phe-Glu-Lys(Dnp)-OH at a final concentration of 10 µM. The fluorometric reaction (λex = 320 nm and λem = 420 nm) was quantified at 37°C on a Safire2 (Tecan, Switzerland) microplate reader. Cell lysates were tested in quadruplicates. Silencing of γ1-syntrophin was performed with the Stealth™ RNAi/γ1-Syn system. The single stranded oligos were (RNA)-GUAGCUUGUUGGACCCUCUAUUU and (RNA)-AAUAUGAGGUGCAACACGCU-ACC (Invitrogen Stealth™ RNAi; Invitrogen, Carlsbad). SH-SY5Y cells were transiently transfected with Stealth RNAi/γ1-Syn using Lipofectamine 2000 (Invitrogen, Carlsbad), according to the manufacturer protocol. Silencing of γ1-syntrophin was tested with flow cytometry and Western blotting using γ1-syntrophin antibody (Santa Cruz, USA). The specificity of this antibody to γ1-syntrophin was tested by pretreatment with GST-γ-PDZ fusion protein (online supplementary figure S1, www.karger.com/doi/10.1159/000324292).

Immunoprecipitation and Western-Blotting

To prepare cell lysates for analysis of the protein levels, SH-SY5Y cells were washed with PBS, pH 7.4, and then lysed with 100 µl lysis buffer (400 mM phosphate buffer, pH 6.0, 75 mM NaCl, 4 mM EDTA-Na2, 0.25% Triton X-100) including a protease inhibitor cocktail (Thermo Scientific). The cells were incubated on ice for 30 min and then centrifuged for 10 min at 10,000 g and 4°C. The supernatants were removed and the total protein concentrations were determined by adding lysis buffer, pH 8.0, to a final concentration of 100 µg in 50 µl. Then 5 µg of a mouse anti-γ-enolase monoclonal antibody (Santa Cruz Biotechnology, Santa Cruz, Calif., USA) were added. To purify the immune complexes, 50 µl of this mixture was added to washed protein A-Sepharose beads (Amersham Pharmacia Biotech AB, Uppsala, Sweden) and mixed overnight at 4°C. The beads were then washed with binding buffer (0.14 M phosphate buffer), pH 8.2, and the immune complexes eluted by boiling in SDS sample buffer. The samples were resolved by SDS-PAGE and transferred onto nitrocellulose membranes. The membranes were blocked with 5% skimmed milk powder in PBS, and then incubated with goat anti-γ1-syntrophin or mouse anti-γ-enolase monoclonal antibody (Santa Cruz Biotechnology) in 0.05% Tween in PBS overnight at 4°C. The blots were washed three times for 10 min in...
0.05% Tween in PBS, incubated with the HRP-conjugated secondary antibody (Santa Cruz Biotechnology) for 2 h, and finally washed another three times for 10 min with 0.05% Tween in PBS. The immunoreactive bands were visualized after addition of the Super-Signal chemiluminescent substrate (Thermo Scientific). The band intensities from the Western blotting were quantified using the Sygene's GeneTools Software (Sygene, UK), and expressed as relative values to the control lysate.

**Peptides**

Synthetic peptides corresponding to the C-terminal 30-amino-acid sequence of human brain γ-enolase (γ-Eno: AKYNQLMRIEELGLDEARGHAFGNFRNPSVL), a two-residue-shorter peptide (γ-EnoΔ432–433: AKYNQLMRIEELGLDEARGHAFGNFRNPS), and a peptide corresponding to the C-terminal 15-amino-acid sequence of human α-enolase (α-Eno: KAKFAGRNPRNLAK), were synthesized by Biosynthesis (Lewisville, Tex., USA).

**Expression and Purification of Bacterial Fusion Proteins**

The construct encoding the PDZ domain of γ-syntrophin in the plasmid vector pGEX-5X1 was a generous gift from Stephen H. Gee [19]. The fusion protein was purified as described previously [19]. Briefly, BL21 (DE3)pLys competent cells were transformed with pGEX-5X1 and grown in 2 ml LB medium supplemented with 200 μg/ml ampicillin (LB-Amp) (Sigma) overnight at 37°C in a shaking incubator. The bacteria were pelleted by centrifugation and resuspended in 1 ml fresh LB-Amp. Overnight culturing was used to inoculate 1 liter of bacteria in LB-Amp, which was grown at 37°C for 2 h or until the optical density at 600 nm reached 0.6–1.0. Protein expression was then induced with 1 mM isopropyl-1-thio-β-D-galactopyranoside, and the culture was incubated overnight at 28°C. The expressed fusion protein was recognized by the GST conjugate and was purified on a glutathione-Sepharose column, according to the manufacturer instructions (Novagen, Madison, Wisc., USA). The fusion protein purity was determined by Coomassie blue staining of SDS-polyacrylamide gels. Protein concentrations were determined by the method of Bradford.

**Solid-Phase Binding Assay**

Microtiter plates were coated with 10 μg/ml of the synthetic peptides γ-Eno, γ-EnoΔ432–433 or α-Eno, in 0.01 M carbonate/bicarbonate buffer, pH 9.6, at 4°C overnight. After blocking with 3% BSA in PBS, the expressed and purified GST-γ1-syntrophin (GST-γ1PDZ) was added at different concentrations, for 4 h at 37°C. After washing, a goat anti-GST antibody (Biacore AB, Uppsala, Sweden) was added. Following 2 h of incubation at 37°C, the wells were washed and filled with a secondary donkey anti-goat antibody conjugated with HRP. After a further 2-hour incubation at 37°C, 200 μg/well of the substrate 3,3′,5,5′-tetramethylbenzidine was added. After 15 min, the reaction was stopped by adding 50 μl 2 M H2SO4. Absorbance, which correlated with the binding of GST-γ1-syntrophin, was measured at 450 nm using a Safire 2 microplate reader (Tecan, Switzerland).

**Surface Plasmon Resonance**

The relative binding of γ-enolase C-terminal peptides and the γ1-syntrophin PDZ domain were determined using surface plasmon resonance on a Biacore X apparatus (Biacore AB) at 25°C. The synthetic peptides γ-Eno and γ-EnoΔ432–433 were immobilized at a concentration of 50 μg/ml in sodium acetate buffer, pH 4.5, at a flow rate of 10 μl/min for 10 min (290 RU), onto a CM5 chip (Biacore AB, Uppsala, Sweden). A reference surface was prepared for each peptide in the second flow cell by amine coupling activation followed by immediate deactivation. To measure the relative binding, six different concentrations of GST-γ1PDZ fusion protein in PBST buffer (PBS with 0.1% Tween 20) were injected onto the peptide surfaces for 1 min at a flow rate 10 μl/min. Binding to a reference flow cell was also measured, and used to subtract the non-specific interactions. The association and dissociation rate constants were fitted globally to a two-step binding model using BIA evaluation version 3.2.

**Immunofluorescence Colocalization**

SH-SYSY cells were cultured on glass coverslips at a concentration of 2 × 10⁴/ml in 24-well plates. After 24 h, the cells were fixed with 2% paraformaldehyde at room temperature for 45 min and permeabilized with 0.1% Triton X-100 for 10 min. Non-specific staining was blocked with 3% BSA in PBS, pH 7.4 for 30 min. Then the permeabilized cells were incubated with mouse anti-γ-enolase (10 μg/ml) or goat anti-γ1-syntrophin (10 μg/ml) (Santa Cruz Biotechnology) antibodies in blocking buffer, for 2 h at room temperature. Afterwards, the cells were washed with PBS and incubated with the Alexa-labeled secondary antibodies (Molecular Probes, Eugene, Ore., USA) for an additional 2 h. After washing with PBS, the Prolong Antifade kits (Molecular Probes) were used for mounting the coverslips on glass slides.

For detection of the plasma membrane, the cells were transduced with Organelle Lights™ Plasma Membrane-GFP (PM-GFP, Molecular Probes), following the manufacturer instructions a day before the immunofluorescence staining. The N-terminal end of γ-enolase was stained using a goat polyclonal antibody raised against a peptide mapping near the N-terminus of γ-enolase (5 μg/ml) (Santa Cruz Biotechnology).

For actin cytoskeleton localization, F-actin was stained using FITC-labelled phalloidin (500 ng/ml) (Sigma-Aldrich, St. Louis, Mo., USA).

Fluorescence microscopy was performed using a Carl Zeiss LSM 510 confocal microscope (Carl Zeiss, Oberkochen, Germany). Images are presented as single confocal sections and were analyzed using Carl Zeiss LSM image software, version 3.0. The relative colocalization areas, presented in the third quadrant of the images were analyzed for several cells (cell number ≥10).

**Flow Cytometry Analysis**

SH-SYSY cells (2 × 10⁵/ml) were washed with ice-cold PBS and then with 3% BSA in PBS containing 1% sodium azide. Non-specific staining was blocked with 3% BSA in PBS. Next, the cells were incubated with specific antibodies against the N-terminal end of γ-enolase (10 μg/ml) for 45 min at 4°C. After washing with ice-cold PBS, the cells were fixed with 2% paraformaldehyde for 15 min at 4°C. Afterwards, the cells were incubated with an Alexa Fluor-555-labelled secondary antibody (Molecular Probes) for 45 min at room temperature. After additional washing with PBS, comparative analysis of γ-enolase on the cell surface was performed on a FACS Calibur machine (BD Biosciences, San Jose, Calif., USA).

For α-enolase, γ-enolase and γ1-syntrophin staining, cells were first fixed with 2% paraformaldehyde for 15 min at room temperature.
temperature and permeabilized with 0.5% Tween 20 in PBS for 10 min at 4°C. Afterwards, the cells were incubated with specific antibodies against α-enolase, γ-enolase and γ1-syntrophin (Santa Cruz Biotechnology) (10 μg/ml), followed by secondary antibody.

Neurite Outgrowth and Neuronal Proliferation Assay
SH-SY5Y cells were first transfected with Stealth RNAi/γ1-Syn and nontransfected SH-SY5Y cells were used as the control. The next day (24 h after transfection), the cells were plated into 96-well plates (2 × 10^4 cells per well) in the absence or presence of the γ-enolase peptides, γ-Eno and γ-EnoΔ432–433 at concentration of 100 μM. After an additional 24 h, the cells showing neurite outgrowth were quantified (at least 100 cells for each group in one experiment, and in three independent experiments). Cells with one or more neurites for which the length was twice the diameter of the cell body were scored as positive. The cells were observed using an Olympus IX 81 motorized inverted microscope (Olympus Biosystems GmbH, Munich, Germany).

In a similar experiment, cell proliferation was determined with the MTS colorimetric assay (Promega, Madison, Wisc., USA). Cell proliferation was determined using equation (1), where A_test cells and A_control cells were the absorbances of formazan measured for the treated and control cells:

\[
\text{Cell proliferation} = \left( \frac{A_{\text{test cells}}}{A_{\text{control cells}}} \right) \times 100
\]

Statistical Analysis
The SPSS PC software (Release 13.0) was used for the statistical analysis of all of the data. The differences between the groups were evaluated using the non-parametric Mann-Whitney test. p < 0.05 was considered to be statistically significant.

Results

Association of γ-Enolase with γ1-Syntrophin in Neuroblastoma Cells
Scaffold proteins such as the syntrophins may assist to translocate γ-enolase to the plasma membrane. We detected a complex that formed between γ-enolase and the scaffold protein γ1-syntrophin using affinity immunoprecipitation in SH-SY5Y cell lysates. The immunoprecipitates were resolved by SDS-PAGE and immunoblotted. As shown in figure 1a, an antibody to the internal region of γ-enolase co-immunoprecipitated γ1-syntrophin from wild-type SH-SY5Y cell lysates. No effects were seen using an antibody to α-enolase or with control mouse IgG (fig. 1a), demonstrating the specificity of the association between γ-enolase and γ1-syntrophin, although the expression of α-enolase in SH-SY5Y cells was 1.7-fold higher over that of γ-enolase (online suppl. fig. S2). The same pattern was seen in cathepsin-X-overexpressing SH-SY5Y cell lysates with 2.0-fold increased cathepsin X activity (online suppl. fig. S3); however, here the co-immunoprecipitation was decreased by 7.4-fold, in comparison to wild-type cells (fig. 1c). In the reciprocal experiment, γ-enolase was immunoprecipitated using a γ1-syntrophin antibody as well (fig. 1b), where in cathepsin-X-overexpressing cells, the co-immunoprecipitation was 3.0-fold lower (fig. 1d). Together, these data suggest that in neuroblastoma cells, γ1-syntrophin and γ-enolase can form a stable complex that is less stable in cells with up-regulated cathepsin X.

PDZ Domain of γ1-Syntrophin Binds to the Preserved C-Terminal Peptide of γ-Enolase
It is known that PDZ-domain-mediated interactions are based on the recognition of short C-terminal sequences in the targeted membrane proteins [24]. To verify that this represents the binding mode between γ-enolase and γ1-syntrophin, solid-phase binding assays were performed. For these we used a soluble glutathione-S-transferase (GST) fusion protein containing the PDZ domain of γ1-syntrophin (GST-γ1PDZ), and synthetic peptides representing C-terminal end of both α-enolase and γ-enolase, respectively. Of these, the synthetic peptide γ-Eno corresponded to the entire γ-enolase C-terminal end, and the two-residues-shorter peptide γ-EnoΔ432–433 represented the C-terminal end that results from the gradually cleaving by cathepsin X, meanwhile the synthetic peptide α-Eno corresponded to the entire α-enolase C-terminal end (fig. 2a). The PDZ domain of γ1-syntrophin bound to both of C-terminal peptides of γ-enolase and to the C-terminal peptide of α-enolase in a concentration-dependent manner (fig. 2b); however, the binding was significantly lower with the shorter γ-enolase C-terminal peptide (γ-EnoΔ432–433) and more lower with α-enolase C-terminal peptide compared to γ-Eno peptide.

In addition, the binding of γ-enolase and γ1-syntrophin was assessed by surface plasmon resonance analysis. The binding of purified GST-γ1PDZ to the immobilised γ-Eno, the entire C-terminal peptide of γ-enolase, was concentration dependent (fig. 3a), while the binding of GST-γ1PDZ to the shorter C-terminal peptide of γ-enolase, γ-EnoΔ432–433, was not significantly different from the control (fig. 3b). The dissociation constant, K_d, calculated from a Scatchard plot was 24.3 μM (online suppl. fig. S4). Thus, these data confirm direct interactions of γ-enolase with γ1-syntrophin through its PDZ domain, and they stress the importance of the last two amino acids on the C-terminal end of γ-enolase for binding.
Colocalization of $\gamma_1$-Syntrophin and $\gamma$-Enolase in the Peri-Plasma Membrane Region and Neurite Growth Cone of Neuroblastoma Cells

Using confocal immunofluorescence microscopy in wild-type neuroblastoma SH-SY5Y cells, colocalization of $\gamma_1$-syntrophin and $\gamma$-enolase was seen in the peri-plasma membrane region and along the neurites (fig. 4a). This colocalization was further examined in SH-SY5Y cells induced to differentiate by serum deprivation. Shortly, after serum deprivation, the neuroblastoma cells flatten and produce filopodia and lamellipodia around their circumference, and therefore gradually extend neurites over the subsequent 16–24 h [25, 26]. In SH-SY5Y cells differentiated for 24 h, $\gamma$-enolase and $\gamma_1$-syntrophin were highly concentrated in these early processes of neurite outgrowth (fig. 4b, d). In SH-SY5Y cells transfected with GFP-tagged cathepsin X, the colocalization of $\gamma$-enolase with $\gamma_1$-syntrophin was dramatically abolished (fig. 4c, d).
Fig. 2. Specific interactions of C-terminal peptide of γ-enolase and α-enolase with the PDZ domain of γ1-syntrophin. a Amino acid sequences of the C-terminal peptides: α-Eno, γ-Eno (entire) and cathepsin-X-modified γ-EnoΔ432–433. b Binding of GST fusion protein containing the PDZ domain of γ1-syntrophin (GST-γ1PDZ) to these synthetic peptides, α-Eno, γ-Eno and γ-EnoΔ432–433, using the solid-phase binding assay. GST-γ1PDZ binds to all peptides in a concentration-dependent manner (b), although binding was 3.3-fold greater with γ-Eno than γ-EnoΔ432–433 and 4.4-fold greater than α-Eno at 1 μg/ml of GST-γ1PDZ. The values are relative to those for the control recombinant GST. Data are means ± SD of three independent assays (p < 0.05; as indicated).

Fig. 3. PDZ domain of γ1-syntrophin binding to preserved C-terminal peptide of γ-enolase. a, b Representative association and dissociation curves of PDZ domain of γ1-syntrophin for binding to C-terminal peptide of γ-enolase, obtained by surface plasmon resonance. Different concentrations (0–50 μM) of the GST-γ1PDZ fusion protein were injected onto the surface of a sensor chip coated with the peptides γ-Eno or γ-EnoΔ432–433. Binding of GST-γ1PDZ to the peptides was measured in real time as an increase in resonance units (RU). Binding was stronger at higher concentrations of GST-γ1PDZ protein (a), whereas the interaction was abolished when γ EnoΔ432–433 peptide was immobilized on the cell surface (b).
To investigate whether interaction between \(\gamma\)-enolase and \(\gamma_1\)-syntrophin enhances \(\gamma\)-enolase translocation towards the plasma membrane, we used a fluorescent protein marker that is specific for the plasma membrane (PM-GFP; Organelle LightsTM Plasma Membrane-GFP) and the antibody mapping the epitope close to the N-terminus of \(\gamma\)-enolase. \(\gamma\)-Enolase strongly colocalized with the PM-GFP protein marker specific for the plasma membrane (fig. 5a). To confirm that the translocation of \(\gamma\)-enolase towards the plasma membrane depends on its binding to \(\gamma_1\)-syntrophin, we decreased the expression of \(\gamma_1\)-syntrophin using a \(\gamma_1\)-syntrophin-specific small-interfering (si) RNA, the Stealth RNAi/\(\gamma_1\)-Syn system. Reduced \(\gamma_1\)-syntrophin protein level for 68% 24 h or for 60% 48 h after transfection, respectively, as confirmed by flow cytometry and Western blotting (online suppl. fig. S5) abolished the \(\gamma\)-enolase colocalization with the plasma membrane marker (fig. 5b) in comparison to intact neuroblastoma cells (fig. 5a, c). Additionally, the surface of neuroblastoma cells was examined for the presence of \(\gamma\)-enolase by flow cytometry. A single-cell suspension of wild-type SH-SY5Y cells, cells with overexpressed ca-

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Gamma-1-Syntrophin Regulates Trafficking of Gamma-Enolase

Effects of γ₁-Syntrophin Silencing on γ-Enolase-Mediated Cell Proliferation, Neurite Outgrowth and Actin-Based Cytoskeleton Remodeling

Next, we analyzed the effects of silencing the γ₁-syntrophin gene with the Stealth RNAi/γ₁-Syn system, on γ-enolase-induced neurite outgrowth and cell proliferation. The morphology of SH-SY5Y cells was examined 24 h after exposure to a synthetic peptide that corresponded to γ-Eno, the entire γ-enolase C-terminal end and to shorter peptide, γ-EnoΔ432–433. Treatment with γ-Eno peptide increased the number of cells with neurite elongation, as compared to control cells (data not shown). In cathepsin-X-overexpressing SH-SY5Y cells, the signal for the γ-enolase also decreased, as shown in figure 5f, and was 2.3-fold lower compared to wild-type cells (fig. 5g).

Fig. 5. Silencing of γ₁-syntrophin expression abolishes γ-enolase translocation towards the plasma membrane. a, b SH-SY5Y cells were transfected with γ₁-syntrophin-specific siRNA 24 h before transduction with Organelle Lights™ PM-GFP for direct expression of autofluorescent protein that is localized in the plasma membrane. γ-Enolase antibody was added, and then labeled with an Alexa Fluor 555 secondary antibody. In the representative images, the colocalization of γ-enolase with the plasma membrane marker was seen (a), which was abolished in cells with downregulated γ₁-syntrophin (b). c Quantification of the relative colocalization areas in a and b (cell numbers ≥10). Data are means ± SD (p < 0.03, as indicated). Scale bar = 10 μm. d–f Representative flow cytometric analysis of nonpermeabilized SH-SY5Y cells transfected with γ₁-syntrophin-specific siRNA or pcDNA3/cathepsin X. Broken thin line represents an isotype control and thick solid line a plasma membrane localization of γ-enolase, respectively. Quantitative data are means ± SD of at least three independent experiments (g) (p < 0.001; p < 0.003, as indicated).

thepsin X and cells with reduced γ₁-syntrophin was stained with the antibody to the N-terminal end of γ-enolase under nonpermeabilized conditions. As seen in figure 5d and 5e, the signal for the γ-enolase was 1.9-fold greater, compared to cells with reduced expression of γ₁-syntrophin (fig. 5g). The nonspecific IgG as a control provided no positive staining (data not shown). In cathepsin-X-overexpressing SH-SY5Y cells, the signal for the γ-enolase also decreased, as shown in figure 5f, and was 2.3-fold lower compared to wild-type cells (fig. 5g).

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compared to untreated cells, while the γ-enolase-mediated neuritogenic effects were abolished in γ1-syntrophin down-regulated SH-SY5Y cells (fig. 6a). SH-SY5Y cells with reduced protein level of γ1-syntrophin appeared to undergo little if any γ-enolase-mediated neurite outgrowth, as seen in figure 6b. At the same time, in these siRNA-transfected SH-SY5Y cells γ-enolase-mediated cell proliferation was decreased by 1.4-fold in comparison to nontransfected SH-SY5Y cells (fig. 6c). The same experiments were performed for peptide γ-EnoΔ432–433 represented the C-terminal end that lacks last two amino acids. Nontransfected cells treated with the γ-EnoΔ432–433 did not undergo neurite outgrowth (fig. 6a) as well as γ-EnoΔ432–433 peptide had no significant proliferative ef-

Fig. 6. Neurotrophic effects of γ-enolase are regulated by γ1-syntrophin expression. 

a Inhibition of γ1-syntrophin expression with transient transfection of SH-SY5Y cells with Stealth RNAi/γ1-Syn. After transfection, cells were cultured in the presence or absence of the γ-enolase peptides (γ-Eno and γ-EnoΔ432–433; 100 μM) and after an additional 24 h the frequency of neurite outgrowth was determined. 

b Representative DIC images of γ-Eno-treated nontransfected (left panel) and Stealth RNAi/γ1-Syn transfected SH-SY5Y cells (right panel) are shown. Scale bars = 50 μm. 

c SH-SY5Y cells were treated as in a, and after 24 h the cell proliferation was assessed by the MTS assay. Data are means ± SD of two independent experiments (p < 0.005; p < 0.01; p < 0.0001; p < 0.05; as indicated).
Gamma-1-Syntrophin Regulates Trafficking of Gamma-Enolase

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The present study highlights a key role for \(\gamma_1\)-syntrophin in the regulation of the subcellular distribution and neurotrophic function of \(\gamma\)-enolase. The preserved C-terminus of \(\gamma\)-enolase, which contains a consensus PDZ-binding motif, is necessary for its interaction with the PDZ domain of \(\gamma_1\)-syntrophin. In this way, \(\gamma_1\)-syntrophin aids in the translocation of \(\gamma\)-enolase towards the plasma membrane, where it can then exert its neurotrophic activity. The formation of this signaling complex is abolished by cathepsin X, through its modulation of the C-terminal end of \(\gamma\)-enolase.

Enolase is a glycolytic enzyme that is generally viewed as a soluble cytosolic protein, although several other functions associated with its plasma membrane localization have been reported [28, 29]. For example, \(\alpha\)-enolase can serve as a receptor for the binding of plasminogen to the cell surface, through its exposed C-terminal lysine residues [30]. The neuron-specific enolase has been found in rat brain synapses [31], with half of the amount present in these synapses seen as membrane associated [32]. Here, we demonstrate that \(\gamma\)-enolase is indeed present in neuroblastoma SH-SY5Y cells and that it can be associated with the plasma membrane. Although our results do not directly confirm its membrane integration, the latter may occur via hydrophobic domain in its N-terminal region (32-AAVPSGASTGIY-43) [33] similar to what has been suggested for the hydrophobic domain of \(\alpha\)-enolase (33-AAVPSGASTGIY-44) [34].

Discussion

The present study highlights a key role for \(\gamma_1\)-syntrophin in the regulation of the subcellular distribution and neurotrophic function of \(\gamma\)-enolase. The preserved C-terminus of \(\gamma\)-enolase, which contains a consensus PDZ-binding motif, is necessary for its interaction with the

PDZ domain of \(\gamma_1\)-syntrophin. In this way, \(\gamma_1\)-syntrophin aids in the translocation of \(\gamma\)-enolase towards the plasma membrane, where it can then exert its neurotrophic activity. The formation of this signaling complex is abolished by cathepsin X, through its modulation of the C-terminal end of \(\gamma\)-enolase.
The C-terminal end of γ-enolase contains a consensus PDZ-binding motif (431-SVL-433), but not the C-terminal end of α-enolase (431-LAK-433), that appears to be involved in its cellular redistribution. There are several PDZ-domain-containing proteins that are involved in intracellular signaling pathway events, including localization to the cell membrane [14, 35]. PDZ domains preferentially bind peptides that terminate in hydrophobic amino acids and with a Ser, Thr or Tyr at the −2 position from the C-terminus [36]. Alessi et al. [20] showed that γ1-syntrophin PDZ domain binds a unique set of signaling proteins that are different from those recognized by other syntrophins. We have shown here that in wild-type SH-SY5Y cells, γ1-syntrophin forms a stable complex with γ-enolase; at the same time, α-enolase does not co-purify with γ1-syntrophin. Although both isoforms show high amino-acid sequence homology (82%) [5], there is a significant difference in C-terminal regions [33], as α-enolase lacks the SVL C-terminal consensus PDZ-binding motif. Although we cannot exclude the possibility that the translocation of γ-enolase is due to interactions with PDZ-containing proteins (scaffold proteins) other than γ1-syntrophin, our data clearly show a direct interaction between the PDZ domain of γ1-syntrophin and the C-terminal end of γ-enolase. As the localization of γ-enolase in the periplasma membrane region was diminished after silencing the γ1-syntrophin gene further, this supports the interaction of γ1-syntrophin with γ-enolase and its role in intracellular trafficking.

It has been proposed that a 30-amino acid C-terminal region of γ-enolase (A404-L433) promotes neurotrophic and neuroprotective effects [5]. As shown by solid-phase binding assays and surface plasmon resonance, the preserved C-terminus of γ-enolase (A404-L433) is also necessary for its interactions with the PDZ domain of γ1-syntrophin. The peptide deduced from the C-terminal of γ-enolase binds to the PDZ domain in the micromolar range (Kd, 24.3 μM); however, a lower affinity for PDZ-mediated interactions might be more suited for the dynamic assembly of PDZ-domain protein complexes [37]. On the contrary, as the product of cathepsin X cleavage of two γ-enolase C-terminal amino acid residues [11], a peptide shorter by two amino acids (A404–S431) showed significantly reduced binding to the PDZ domain of γ1-syntrophin. Similarly, a peptide, corresponding to C-terminal end of α-enolase also exhibited weak binding to the PDZ domain. Collectively, the interactions mediated through the PDZ domain of γ1-syntrophin might be important for the subcellular distribution of γ-enolase, and this regulation can be modulated by cathepsin X.

We favor the idea that γ1-syntrophin serves to sequester the full-length γ-enolase from the cytosol towards the plasma membrane. The colocalization of γ1-syntrophin and γ-enolase mainly at the periplasma membrane region and along neurites in wild-type SH-SY5Y cells supports this idea. On the other hand, in cathepsin-X-overexpressing SH-SY5Y cells, the colocalization of γ-enolase and γ1-syntrophin was abolished. Additionally, co-immunoprecipitation of γ-enolase and γ1-syntrophin from cathepsin X-overexpressing cells was less prominent than it was from wild-type cells. This is thus in agreement with previous reports from our group, where we have shown that the C-terminal of γ-enolase is gradually shorted by two amino acids, V432 and L433, through cleavage by cathepsin X [11], therefore removing the PDZ-binding motif.

The precise subcellular location of this γ-enolase processing by cathepsin X is not clear yet. In neuronal cells, cathepsin X was found in the cytoplasm [12] and can thus interact with its target, γ-enolase. On the other hand, cathepsin X was also shown to colocalize with γ-enolase in perimembrane regions where proteolytic cleavage can also occur [11]. In our previous study, we showed that cathepsin X abolished the neurotrophic activity of γ-enolase and impaired neuritogenesis, linking these effects to the proteolytic cleavage of the C-terminal region of γ-enolase, and consequently with the elimination of its neurotrophic function [11]. In the present study, we have provided another mechanism for the impairment of γ-enolase neurotrophic function, i.e. cathepsin X can regulate this process by eliminating the PDZ-binding motif in the C-terminal region of γ-enolase, preventing the binding to γ1-syntrophin and the translocation towards the plasma membrane.

To our knowledge, this is the first report proposing a mechanism for γ-enolase translocation in neuroblastoma cells. These data are also consistent with the study of Yakubchyk et al. [38], who showed regulation of neurite outgrowth through PDZ-mediated recruitment of diacylglycerol kinase-δ (DGK-δ). They demonstrated that the C-terminal PDZ-binding motif of DGK-δ is necessary for neurite outgrowth after its translocation to the plasma membrane by γ1-syntrophin. For neurite outgrowth dynamic remodeling of the actin-based cytoskeleton is fundamental [1, 39]. Consistent with a potential role of γ1-syntrophin in actin reorganization, we have also shown that γ-enolase colocalizes with actin filaments in the perimembrane region, while this interaction was altered in the absence of γ1-syntrophin. The C-terminal region of syntrophins can bind to members of the dystro-
phin family [15], which, at the same time, are known to bind directly to actin [21, 40]. Therefore, syntrophins might provide a link between γ-enolase and actin cytoskeleton remodeling.

In conclusion, we have identified a mechanism of translocation of γ-enolase towards the plasma membrane in neuroblastoma cells. This arises through the interaction between the PDZ domain of the scaffold protein γ1-syntrophin with the C-terminal region of γ-enolase. We propose that this activity is required for γ-enolase-mediated neurotrophic effects. Within this framework, increased activity of cathepsin X interferes with the formation of this translocation complex thus preventing γ-enolase from exerting its neurotrophic functions, such as enhancement of cell proliferation and inducement of neurite outgrowth.

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