A Splice Isoform of DNedd4, DNedd4-Long, Negatively Regulates Neuromuscular Synaptogenesis and Viability in Drosophila

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Abstract

Background: Neuromuscular (NM) synaptogenesis is a tightly regulated process. We previously showed that in flies, Drosophila Nedd4 (dNedd4/dNedd4S) is required for proper NM synaptogenesis by promoting endocytosis of commissureless from the muscle surface, a pre-requisite step for muscle innervation. DNedd4 is an E3 ubiquitin ligase comprised of a C2-WW(x3)-Hect domain architecture, which includes several splice isoforms, the most prominent ones are dNedd4-short (dNedd4S) and dNedd4-long (dNedd4Lo).

Methodology/Principal Findings: We show here that while dNedd4S is essential for NM synaptogenesis, the dNedd4Lo isoform inhibits this process and causes lethality. Our results reveal that unlike dNedd4S, dNedd4Lo cannot rescue the lethality of dNedd4 null (dNedd4T121FS) flies. Moreover, overexpression of UAS-dNedd4Lo specifically in wildtype muscles leads to NM synaptogenesis defects, impaired locomotion and larval lethality. These negative effects of dNedd4Lo are ameliorated by deletion of two regions (N-terminus and Middle region) unique to this isoform, and by inactivating the catalytic activity of dNedd4Lo, suggesting that these unique regions, as well as catalytic activity, are responsible for the inhibitory effects of dNedd4Lo on synaptogenesis. In accord with these findings, we demonstrate by sqRT-PCR an increase in dNedd4S expression relative to the expression of dNedd4Lo during embryonic stages when synaptogenesis takes place.

Conclusion/Significance: Our studies demonstrate that splice isoforms of the same dNedd4 gene can lead to opposite effects on NM synaptogenesis.

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Introduction

Neuronal precursor cell-expressed developmentally downregulated 4 (Nedd4) is an E3 ubiquitin ligase that belongs to the Hect family [1]. Nedd4 proteins share common domain architecture: a C2 domain, 3–4 WW domains and a ubiquitin ligase Hect domain. The C2 domain is primarily responsible for sub-cellular targeting [2,3], while the WW domains mediate substrate recognition and binding usually by associating with PY motifs (L/PPxY) [4,5,6].

In higher eukaryotes, there are several Nedd4 family proteins, including the closely related Nedd4-1 (Nedd4) and Nedd4-2 (Nedd4L). In mammals, Nedd4-2 is known to regulate stability of ion channels, such as ENaC, which has PY motifs that interact with the Nedd4-2-WW domains to promote ENaC endocytosis [7,8,9,10]. Mutations in ENaC PY motifs found in Liddle syndrome (a hereditary hypertension) result in increased retention of ENaC at the plasma membrane in the kidney [11,12]. The interaction between ENaC and Nedd4-2 can be negatively regulated through the phosphorylation of Nedd4-2 by the Ser/Thr kinase, Sgk1, and its close relative Akt1 [13,14]. In contrast to Nedd4-2, mammalian Nedd4-1 has been implicated in the regulation of cellular and animal growth [15], T cell activation [16] and heart [17] and nervous system [18] development.

Recently, Nedd4, which is expressed in muscles, was found to regulate neuromuscular (NM) synaptogenesis in flies [19] and mammals [20]. Specifically, Drosophila Nedd4 (dNedd4) was shown to regulate endocytosis of commissureless (Comm) from the muscle surface to allow proper initiation of NM synaptogenesis [19]. Comm contains two PY motifs (PPCY and LPSY) and 10 Lys residues (ubiquitin acceptor sites) in its intracellular domain. The WW domains of dNedd4 bind to the PY motifs of Comm, leading to Comm ubiquitylation [21]. In Drosophila, each body wall hemisegment contains 30 muscle fibers that are innervated by ~40 motor neurons in a specific, precisely timed manner [22]. Internalization of Comm (expressed on the muscle cell surface) into the muscle is a prerequisite for proper initiation of NM synaptogenesis [23] and is facilitated by dNedd4 [19]. Of note, the
dNedd4 gene is subject to alternative splicing, with several isoforms that can be generally divided into two groups represented by a short isoform (dNedd4S, or dNedd4) and a long isoform, dNedd4Lo. While dNedd4S (dNedd4) was previously shown to enhance NM synaptogenesis, the function(s) of dNedd4Lo was unknown.

Here we characterized dNedd4Lo and show that in contrast to dNedd4S, dNedd4Lo has a negative function in NM synaptogenesis, leading to defects in neuromuscular synapse formation and abnormal larval locomotion.

**Results**

Our previous work showed that dNedd4 is involved in regulating neuromuscular synaptogenesis, and revealed two isoforms of dNedd4 expressed in the body wall muscle [19]. In accord, our immunoblotting analysis of embryo lysates revealed two major splice isoforms of dNedd4, which we named dNedd4-short (dNedd4S, ~92 kDa) and dNedd4-long (dNedd4Lo, ~112 kDa) (Fig. 1A,B). D Neddle4Lo possesses the same functional domains as dNedd4S: a conserved C2 domain, 3 WW domains, including the high affinity WW3 binding domain [24] and a Hect domain, which is catalytically active, much like dNedd4S (Fig. S1A). In addition, dNedd4Lo contains a unique N-terminal (Nterm) region that includes a putative Akt phosphorylation site, as well as a unique middle (Mid) region (Fig. 1A). Our previous work focused on the function of dNedd4S [19]. Here we investigated the biological role of dNedd4Lo and compared it to that of dNedd4S.

**D Neddle4S, but not dNedd4Lo, can partially rescue lethality of dNedd4 null mutant flies**

To determine the biological importance of the two splice isoforms of dNedd4, we tested their ability to rescue dNedd4 null mutant flies. The dNedd4 null (dNedd4<sup>T121FS</sup>) homozygote flies contain a frame shift mutation in the dNedd4 gene that truncates the protein products of all the dNedd4 splice isoforms at Thr121, which renders dNedd4 inactive. D Neddle4<sup>T121FS</sup> flies are heterozygous viable and homozygous lethal at the embryonic stage. For rescue experiments, flies containing either a UAS-dNedd4S or UAS-dNedd4Lo transgene under the transcriptional control of the ubiquitous Act-GAL4 driver [25] (Fig. 1C) were crossed into dNedd4<sup>T121FS</sup> homozygote flies. Two independent crosses were performed for each of the UAS-dNedd4S and UAS-dNedd4Lo transgenes, as well as their respective Act-GAL4 driver alone and UAS transgene alone controls. A total of 800 embryos were collected for each UAS line and observed under a fluorescent microscope to follow the survival of rescued larvae, which do not express GFP (non-GFP). Rescue was determined by measuring the percentage of viable dNedd4 null mutant embryos (hatched larvae without GFP expression) out of the expected number of mutant embryos. The expected number of genotype of interest for rescue with dNedd4S (UAS-dNedd4S/X; Act-GAL4/CyO (or Sp); dNedd4<sup>T121FS</sup>/dNedd4<sup>T121FS</sup>) was calculated to be 12.5%, and with dNedd4Lo (+; UAS-dNedd4Lo/CyO (or Sp); dNedd4<sup>T121FS</sup>/dNedd4<sup>T121FS</sup>) was calculated to be 25%. This excludes all other possible genotypes. Our results show that expression of dNedd4S throughout the embryo partially rescued the lethality of dNedd4 null embryos (from 5.5% to 7.25% to 36%), whereas dNedd4Lo did not rescue the lethality (From 5.5% to 3.75% to 4.5%) and larvae died soon after egg hatching (and some did not fully crawl out of their egg shells) (Fig. 1D). These results suggest that the two splice isoforms of dNedd4, both of which are expressed endogenously in embryos (Fig. 1B), have distinct roles during embryo development.

**Overexpression of dNedd4Lo in the muscle reduces fly viability**

Given the different effects of dNedd4S and dNedd4Lo on rescuing dNedd4 null mutant flies, we next examined the effect of overexpression of dNedd4Lo and dNedd4S. Our results show that ubiquitous expression of all transgenic lines of UAS-dNedd4Lo crossed with the Act-GAL4 driver resulted in lethality before the third instar larval stage, while flies expressing UAS-dNedd4S survived to the adult stage. In an attempt to observe the effect of slight variations in expression levels on fly survival, different ubiquitous GAL4 drivers, including da-GAL4 [26], Act-GAL4 and Tubulin-GAL4 [27] were used at 25°C, room temp (RT), ~22°C, or 18°C. All ubiquitous drivers at all temperatures yielded developmental lethality when crossed to UAS-dNedd4Lo, but not UAS-dNedd4S (Table S1). To rule out the possibility that lethality was due to a defect in sub-cellular localization of the protein, we examined the distribution of dNedd4Lo and dNedd4S in salivary glands. The proteins did not form aggregates and properly localized in the cytosol and on the plasma membrane (Fig. S1B), as was previously observed for localization of endogenous dNedd4 [19] [28]. In an attempt to determine the cause of lethality, various tissue-specific GAL4 drivers were used to drive overexpression in the CNS using elav<sup>151-GALA</sup> [29], in motor neurons using D42-GAL4 [29], in muscles using 24B-GAL4 [29] or 5-GALA [19], in eye using GMR-GAL4 [30], in fat body using Ppl-GAL4 [31] and in the epidermal lining of the digestive system and respiratory system using fpl-GAL4 [29]. Each test was performed at 25°C, RT, or 18°C, to determine whether slight variations in expression levels had any effect on fly survival. We found that all UAS-dNedd4Lo transgenic flies died during development only when overexpressed in muscle using the 24B-GAL4 or 5-GALA drivers, while UAS-dNedd4S transgenic flies survived to adulthood (Table S2). Once again, the lethality was not due to a defect in protein sub-cellular localization since we found that dNedd4Lo was properly localized in the muscle (see below).

**Overexpression of UAS-dNedd4Lo in muscle results in aberrant synaptic innervation along the SNb branch from body wall muscle 13 to 12**

Since we previously demonstrated that dNedd4S is involved in neuromuscular (NM) synaptogenesis [19], potential defects in NM synapse formation were analyzed by performing immunofluorescence staining on dissected body wall muscles of third instar larvae from UAS-dNedd4Lo transgenic lines. There are two major categories of innervation defect: pathfinding error and axonal overgrowth/undergrowth error. The abnormal innervations that were identified and scored in this experiment included: backward innervation from muscle 12 onto muscle 13, which belongs to pathfinding error, and increased branching on muscle 12, which belongs to overgrowth error (Fig. 2A). The frequency of abnormalities was calculated as the number of abnormal innervations (backward innervation and/or pathfinding error) that were identified over the total number of neuromuscular junction at muscle 13 and muscle 12 that were scored. By scoring ~100–130 NM synapses at muscles 13 and 12 from each of the UAS-dNedd4Lo and UAS-dNedd4S transgenic lines, the number of normal and abnormal innervation for each UAS-dNedd4 transgenic was compared with that of the muscle driver control (5-GALA muscle driver fly used to drive overexpression of the UAS transgenes) (Fig. 2B). Our results show that the UAS-dNedd4Lo transgenics had a significant amount of abnormal innervation relative to the control (two-tailed Fisher’s exact test, p<0.0001), which was not observed for the UAS-dNedd4S transgenics (p<0.4734).
Figure 1. dNedd4S, but not dNedd4Lo, can rescue the lethality of dNedd4 null mutant flies. (A) Schematic representation of dNedd4S and dNedd4Lo showing their common C2, WW(x3) and Hect domains, putative Akt phosphorylation sites, and N-terminal (Nterm) and middle (Mid) regions unique to dNedd4Lo. (B) Reduction of endogenous dNedd4 protein expression in dNedd4T121FS heterozygote compared to Oregon-R wild type (WT) embryos, detected with an anti-dNedd4 antibody that recognizes both dNedd4S and dNedd4Lo. (C) Protein expression of Flag-tagged UAS-dNedd4Lo/WT, dNedd4S WT and their S→A mutants driven by daughterless(da)-GAL4 in embryo, detected using anti-Flag antibody on western blots (upper panel). Lower panel: actin loading controls. OregonR embryo lysate was included as a negative control, and lysates from mammalian HEK293T cells transfected with dNedd4Lo and dNedd4S were included as positive controls (upper panel). un = untransfected. (D) Expression of dNedd4S, but not dNedd4Lo, driven by Actin-GAL4 throughout the embryo partially rescues dNedd4 null mutant viability. The bar chart represents the percentage of viable dNedd4 null mutant embryos (measured as hatched larvae without GFP expression) out of the expected number of mutant embryos. 800 embryos were scored per genotype.

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Taken together, these results demonstrate that muscle-specific overexpression of UAS-dNedd4Lo in a wild type background causes a high frequency of abnormal motor neuron innervation on muscles 13→12, whereas that of UAS-dNedd4S does not cause significant abnormalities.

Overexpression of UAS-dNedd4Lo in muscle results in reduction of larval locomotor activity

To determine the consequences of the abnormal muscle innervation in larvae that overexpress dNedd4Lo in the muscle, we analyzed larval locomotor activity by measuring the total path length travelled by pre-wandering third instar larvae over a 150 sec period. Our results reveal a severe reduction in locomotor activity of larvae that overexpress dNedd4Lo in the muscle (+/UAS-dNedd4Lo; +/24B-GAL4) (73.76±6.38 mm) relative to the +/UAS-dNedd4Lo (120.35±5.05 mm) or +/24B-GAL4 (137.23±5.46 mm) controls (post hoc analysis, p<0.001) (Fig. 2C,D).

Adverse effects of dNedd4Lo are not a result of dAkt regulation

Differences between the two isoforms of dNedd4 include an alternate start codon site resulting in a longer N-terminal region in dNedd4Lo, and an extra exon inserted between the WW1 and WW3 domains (Fig. 1A and 3A). In the unique N-terminal region of dNedd4Lo, there is a putative Akt phosphorylation site (S39) with the consensus sequence, RxRxxS/T. Another putative Akt phosphorylation site was found common to both dNedd4Lo (S645) and dNedd4S (S444) (Fig. 3A). Interestingly, a close relative of dNedd4 in mammals, Ned4-2, also contains a consensus sequence, RxRxxS/T, that can be phosphorylated by the Ser/Thr kinase Sgk1 and its close relative, Akt1 [13]. Therefore, we tested the possibility that the putative Akt phosphorylation site(s) in dNedd4Lo were phosphorylated by Drosophila Akt (dAkt) in Drosophila S2 cells. We found that while the site containing Ser39 in the unique N-terminal region of dNedd4Lo was indeed

Figure 2. Neuromuscular innervation and locomotion defects in larvae overexpressing dNedd4Lo in the muscle. (A) Muscle-specific overexpression of UAS-dNedd4Lo (dNedd4Lo/5-GAL4) leads to aberrant synaptic innervation along the SNb branch from body wall muscles 13→12 of third instar larvae (HRP stain, red). Muscle driver control line (+/5-GAL4) was included (left panel) to show the normal motor neuron innervation pattern on muscle 13→12. Muscles were stained with Phalloidin (green). Scale bars, 10 μm. (B) Quantification of the muscle innervation defects. Numbers in brackets denote the number of muscles scored (n). * p<0.0001 (Two tailed Fisher’s exact test). (C,D) Overexpression of dNedd4Lo adversely affects larval locomotor activity measured by total path length: (C) Representative path travelled by third instar larvae of muscle driver alone (24B-GAL4), UAS-dNedd4Lo alone (UAS-dNedd4Lo), or larvae overexpressing dNedd4Lo in the muscle (UAS-dNedd4Lo/24B-GAL4). Scale bar, 10 mm. (D) Quantification of locomotor activity shown in panel C of 55–60 larvae (per genotype). Numbers in brackets denote the exact number of larvae scored (n). * indicates a significant difference between the UAS-dNedd4Lo/24B-GAL4 line and both control lines (UAS-dNedd4Lo alone or 24B-GAL4 alone) (p<0.001, post hoc analysis).

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Figure 3. DNedd4Lo, but not dNedd4S, is phosphorylated by Akt in Drosophila S2 cells. (A) Schematic representation of dNedd4S and dNedd4Lo showing their consensus Akt phosphorylation sites. (B) Drosophila S2 cells were transfected (Tfxn) with Flag-tagged wild-type (WT) or the phosphorylation mutants of the dNedd4 isoforms: dNedd4S (dNedd4S(WT)), dNedd4S(S444A), dNedd4Lo(WT), dNedd4Lo(S39A), dNedd4Lo(S645A), or dNedd4Lo (S39A,S645A). Transfected cells were lysed and the lysates immunoprecipitated (IP) with anti Flag antibodies and immunoblotted (IB) with anti Phospho-(Ser/Thr) Akt substrate antibodies to detect phosphorylated dNedd4 (upper panel). dNedd4 expression levels in the lysates were verified with anti-Flag antibodies (lower panel).

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phosphorylated, the site containing Ser653 in dNedd4Lo, which is also found in dNedd4S [Ser543], was not phosphorylated by dAkt (Fig. 3B). Ser39->Ala mutation in dNedd4Lo abolished its phosphorylation. This observation led us to ask whether dNedd4Lo's unique function is mediated by dAkt phosphorylation. To test this possibility, we generated UAS transgenic lines that express a dNedd4Lo S39A/S645A double mutant (UAS-dNedd4Lo 2S->A), in which the Ser residue of each putative Akt phosphorylation site was mutated to Ala. A UAS-dNedd4S S444A (UAS-dNedd4S S->A) mutant transgenic line was also generated for comparison (Fig. 3A).

We performed the same lethality tests using ubiquitous and muscle-specific GAL4 drivers as described above for UAS-dNedd4Lo and UAS-dNedd4S (wildtype, WT). No difference was found between dNedd4Lo WT and its 2S->A mutant, nor between dNedd4WT and its S->A mutant (Table S3). Next, we scored for muscle innervation defects from UAS-dNedd4Lo 2S->A and UAS-dNedd4S S->A transgenic flies as described above for their WT counterpart. Again, no significant difference was found between UAS-dNedd4Lo WT and its 2S->A mutant (p<0.5298) (Fig. 2B). Because removing the dAkt phosphorylation site of dNedd4Lo mutant did not alter the muscle innervation defects, we conclude that the negative role of dNedd4Lo in NM synaptogenesis is not due to dAkt regulation.

Both unique sequences and the catalytic activity of Nedd4 isoforms are involved in its regulation of neuromuscular synaptogenesis

Since the dAkt phosphorylation site in the unique N-terminal region of dNedd4Lo did not explain the functional difference between dNedd4Lo and dNedd4S, we next studied the role of the N-terminal (Nterm) region as well as the unique middle (Mid) region of dNedd4Lo to determine if either region is involved in the adverse function of dNedd4Lo in NM synaptogenesis. Two dNedd4Lo deletion mutants were created: One with the middle unique sequence deleted (dNedd4LoΔMid) and the other, dNedd4LoΔNterm, had the N-terminal unique sequence replaced by the one from dNedd4HS. In addition, a catalytically inactive Cys961->Ala (C->A) mutant of dNedd4Lo was created to test the effect of abolishing the ubiquitin ligase activity of dNedd4Lo on its function in NM synaptogenesis (Fig. 4 A,B). These mutants were analyzed in comparison to dNedd4Lo WT. Ubiquitous overexpression of UAS-dNedd4LoΔNterm crossed with the da-GAL4 driver, Actin-GAL4, or Tubulin-GAL4 in a wild type background at 25°C, RT or 18°C resulted in developmental lethality, while UAS-dNedd4LoΔMid and UAS-dNedd4Lo C->A transgenic flies survived to adulthood (Table S3). Muscle-specific overexpression of UAS-dNedd4LoΔNterm, UAS-dNedd4LoΔMid and UAS-dNedd4Lo C->A crossed with 24B-GAL4 or 5-GAL4 muscle driver at 25°C, RT or 18°C did not result in any lethality (Table S3). Next, we scored for muscle innervation defects from UAS-dNedd4LoΔNterm and UAS-dNedd4LoΔMid transgenic flies as described above for UAS-dNedd4Lo WT and UAS-dNedd4HS WT. We found that while both UAS-dNedd4LoΔMid and UAS-dNedd4LoΔNterm still exhibited abnormalities compared to the control (5-GAL4 muscle driver fly alone), these abnormalities were seen at a much lower frequency than UAS-dNedd4Lo WT (Fig. 4C). Furthermore, there was no significant difference between UAS-dNedd4LoΔMid and UAS-dNedd4LoΔNterm (p=0.3729). These results show that removing either the N terminal or middle unique regions of dNedd4Lo reduces the frequency of abnormal motor neuron innervation on muscles 13->12 that was observed by muscle-specific overexpression of UAS-dNedd4Lo WT. This suggests that the N-terminal and middle unique regions contribute to the innervation defects caused by dNedd4Lo. Furthermore, mutating the catalytic Cys of dNedd4Lo (UAS-dNedd4Lo C->A mutant) abolishes the abnormality in muscle innervation, indicating that the catalytic activity of dNedd4Lo is required for its adverse effect on NM synaptogenesis (Fig. 4C).

The negative role of dNedd4Lo in NM synaptogenesis is not caused by its inhibition of catalytic activity of dNedd4S, nor by interfering with dNedd4S-mediated regulation of Comm

To explain the negative role of dNedd4Lo and positive role of dNedd4S in NM synaptogenesis, as well as if/how they coordinately regulate this process, we investigated the possibility that dNedd4Lo inhibits the function of dNedd4S by interfering with its catalytic activity. Since the N-terminal and middle unique regions of dNedd4Lo appear to be responsible for its negative effect on fly viability and NM synaptogenesis, we investigated the possibility that they inhibit the catalytic activity of dNedd4S. This was tested in an in vitro ubiquitylation and binding assays. Our results show that adding recombinant proteins corresponding to the Nterm or Mid unique regions of dNedd4Lo into the reaction mixture, either alone or together, did not affect ubiquitylation activity of dNedd4S (Fig. 5A and Fig. S2), nor did these proteins directly bind to dNedd4S (Fig. 5B). In contrast, the C-terminal region of Comm, containing the PY motifs, was able to bind well to dNedd4S (as we previously demonstrated, [19]) and hence was used as a positive control for the binding (Fig. 5B).

Another possible way in which dNedd4Lo could negatively regulate NM synaptogenesis is through the regulation of dNedd4S substrates. Since Comm is a known target of dNedd4S in NM synaptogenesis and removal of Comm from the muscle surface is required for initiation of this event [19], the effect of dNedd4Lo overexpression on endocytosis of Comm from the cell surface was studied in Drosophila S2 cells, which endogenously express dNedd4S. Comm-GFP (WT or 2PY->A mutant, which cannot bind dNedd4) was co-expressed with Flag-dNedd4Lo or Flag-dNedd4S. We hypothesized that overexpression of dNedd4Lo WT would interfere with the dNedd4S-mediated endocytosis of Comm, if indeed dNedd4Lo were to oppose the function of dNedd4S towards Comm. Surprisingly, we found that when co-expressed with dNedd4Lo or dNedd4S, Comm was properly internalized from the cell surface in most cells and co-localized with dNedd4Lo or dNedd4S in intracellular vesicles. As controls, we showed that Comm 2PY->A mutant remained on the plasma membrane of most cells when co-expressed with either dNedd4S or dNedd4Lo (Fig. 6A,B). Likewise, staining for Comm and dNedd4 in muscles of third instar larvae revealed proper internalization of Comm from the muscle surface in muscles that overexpress dNedd4Lo, much like those overexpressing dNedd4S (Fig. 6C). Thus, the negative effect of dNedd4Lo on NM synaptogenesis is likely not caused by interference with Comm endocytosis, since dNedd4Lo was able to promote Comm internalization similar to dNedd4S.

Expression of dNedd4Lo is suppressed during NM synaptogenesis

Given the adverse effect of dNedd4Lo on NM synaptogenesis and larval locomotion, in contrast to dNedd4S (which positively...
regulates NM Synaptogenesis), it is expected that their timing of expression be tightly regulated. Indeed, our semi-quantitative RT-PCR revealed that expression of dNedd4Lo in the embryo is down-regulated soon after the onset of NM synaptogenesis (13 hrs after egg laying), while expression of dNedd4S remains high (Fig. 7). These results suggest that the negative function of dNedd4Lo has to be suppressed during NM synaptogenesis, and that the constant overexpression of UAS-dNedd4Lo driven by the GAL4 muscle drivers throughout embryogenesis could have contributed to the observed lethality and muscle innervation defects, as well as abnormal locomotor activity.

**Discussion**

We previously showed that dNedd4S is involved in NM synaptogenesis in *Drosophila* [19], most likely by promoting...
Figure 5. The unique regions of dNedd4Lo do not bind to dNedd4S nor inhibit its catalytic activity. (A) In vitro ubiquitylation activity of dNedd4S in the presence of GST alone (control) or GST-tagged dNedd4Lo Nterm or Mid unique regions, detected using anti-ubiquitin antibody on western blots. E1, E2 (UbcH5), E3 (dNedd4S), ubiquitin and ATP were included in the ubiquitylation reactions, as well as increasing concentrations (0.9 μM, 1.8 μM and 3.6 μM) of each potential inhibitor (GST, GST-Nterm, GST-Mid). The blot was stripped and re-blotted with anti-Flag antibody to show equal amount of Flag-dNedd4S (0.27 μM) present in all lanes (upper panel of lower blot), and also re-blotted with anti-GST antibody (lower panel). (B) Lack of in vitro binding of dNedd4S to GST fused dNedd4Lo-Nterm, dNedd4Lo-Mid or GST alone. Flag-tagged dNedd4S expressed in HEK293T cells was precipitated from transfected cells, and immunoblotted for Flag (dNedd4, upper panel) or GST (lower panel). HA-tagged C terminus of Comm (which contains PY motifs) was used as a positive control for the binding to dNedd4S (right side of the blot). doi:10.1371/journal.pone.0027007.g005
Figure 6. Cellular localization of Comm in S2 cells and muscles in the presence of overexpressed dNedd4Lo. (A,B) Comm-GFP (WT or 2PYA mutant, green) was co-expressed with Flag-dNedd4Lo or Flag-dNedd4S (red) in Drosophila S2 cells and their sub-cellular localization (plasma membrane (PM) or intracellular) analyzed by immunofluorescence (A) and quantified (B). In (B), P-values were determined with Fisher’s exact test (two
internalization of Comm from the muscle cell surface (a step necessary for initiation of NM synaptogenesis, [23]). Consistent with this model, knock down of dNedd4 during early muscle development or overexpression of Comm mutants that cannot bind dNedd4 yielded the same defects in NM synaptogenesis [19]. Here, we provide genetic evidence that, in contrast to dNedd4S, the splice isoform dNedd4Lo has a negative role in NM synaptogenesis and embryo development in flies. In accord, expression of dNedd4Lo is reduced (and that of dNedd4S is increased) during synaptogenesis, to permit synaptogenesis to proceed. This negative role of dNedd4Lo does not involve Comm or phosphorylation of dNedd4Lo by Akt, nor does it involve an adverse effect of the unique regions of dNedd4Lo on the catalytic activity of the Hect domain of dNedd4S. Instead, it is likely that the unique Nterm and Mid regions of dNedd4Lo contribute to inhibition of NM synaptogenesis by interacting with other cellular factors or complexes, which are not yet known. Studies to analyze differences in general pattern of ubiquitylation upon overexpression of dNedd4Lo vs. dNedd4S in S2 cells did not reveal overt differences (Fig. S3), most likely due to insufficient sensitivity of the system to detect changes in ubiquitylation of specific substrates among the many ubiquitylated cellular proteins.

Our studies here demonstrate that muscle-specific overexpression of dNedd4Lo causes abnormal motor neuron innervation along the SNb branch on body wall muscles 13–12. The types of defects we found include inappropriate backward innervation from muscles 12–13 and increased number of nerve branches on muscle 12. The backward innervation defect was previously observed for overexpression of Comm 2PY-A (that cannot bind dNedd4) and Comm 10K-R mutants (that cannot become ubiquitylated), as well as dNedd4 RNAi mutants [19]. The other common defect we observed was increased motor nerve branching on muscle 12.

Figure 7. Different expression levels of dNedd4Lo versus dNedd4S during 24 hr embryonic development. (A) Expression of dNedd4S and dNedd4Lo mRNAs were analyzed by semi-quantitative RT-PCR at the indicated embryonic stages. Actin 5c was included as an internal control. (B) The bar chart represents mean expression level ± SD from 4 separate experiments. *Indicates a significant difference between dNedd4S and dNedd4Lo (p<0.01, Student’s t-test).

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It is known that disruption of genes involved in cell adhesion processes cause nerve branching defects, such as position specific (PS) β-integrin, fascinII (FasII), Calcium/Calmodulin dependent Kinase II (CaMKII), and DLG (a PDZ-domain Scaffold protein) [36,37]. They form a post-synaptic complex in the muscle to coordinate regulating desialation of the nerve terminal endings and fine-tune the interaction between motor neurons and their muscle targets. It has been proposed that β-integrin regulates recruitment of the cell adhesion molecule FasII on the muscle surface [36]. Down-regulation of β-integrin and up-regulation of FasII on the muscle surface lead to nerve fasciculation. Whether or not the adverse effects of dNedd4Lo on NM synaptogenesis involve these (or other) proteins is currently unknown.

One consequence of muscle innervation defects could be abnormal locomotor activity. Indeed, we found a significant reduction in the locomotor activity of larvae that overexpress dNedd4Lo specifically in the muscle. Furthermore, muscle-specific overexpression of dNedd4Lo leads to lethality during development. However, the muscle drivers 24B-GAL4 and 5-GAL4, used in this experiment, drive expression in the entire mesoderm [29,32], which derives into somatic (body wall) muscles for movement [33,34], visceral (gut) muscles for digestion, and cardiac muscles [35]. Thus, while the innervation defects on body wall muscles may contribute to the larval lethality, defects in heart and/or gut muscle functions might contribute as well, since heart activity and feeding are essential for larval survival.

Similar muscle innervation defects were also observed for the mouse homologue of dNedd4, mNedd4 (mNedd4-1) [20]. In mNedd4 mutant embryos, motor nerves defasciculate upon reaching their skeletal muscle targets and the pre-synaptic nerve terminal branches are increased in number. It was also demonstrated that mNedd4 mutants had increased spontaneous miniature endplate potential (mEPP) frequency, which is consistent with the ultra-structural alternation. In addition, β-catenin, a subunit of the cadherin protein complex, was proposed to be a potential substrate for mNedd4 in NM synapse formation and function. β-catenin deficient muscles show similar defects of nerve desialation as mNedd4 mutant [30]. Similarly, molecular manipulation of β-integrins, which is also involved in the cell adhesion process, in muscles of mice also lead to abnormal development of pre-synaptic nerve terminals [39].

Phosphorylation is an important mechanism for the regulation of Nedd4 proteins and other E3 ubiquitin ligases. For example, Nedd4-2 is known to be regulated by Akt/Sgk – mediated phosphorylation, which inhibits its ability to interact with its substrate ENaC [13,14]. However, mutating the dAkt phosphorylation sites (S->A) in dNedd4Lo did not affect the abnormal muscle innervation. Since the dAkt phosphorylation site in the unique N-terminal region of dNedd4Lo did not explain its negative function, we removed the whole unique N-terminal region or the middle region to determine their role in the negative effect of dNedd4Lo on viability and NM synaptogenesis. We demonstrated that removing either the N-terminal or the middle region rescued the lethality and alleviated the muscle innervation defects. Therefore, both regions are involved in the negative function of dNedd4Lo in this event. We thus investigated two possible underlying mechanisms for the negative regulation of dNedd4Lo in NM synaptogenesis. First, we tested inhibition of function of dNedd4S through the unique regions of dNedd4Lo. It is known that the catalytic activity of Nedd4 proteins can be regulated through auto-inhibition mechanism. For example, the WW domains of Nedd4-2 [40] and a close relative of Nedd4, Ich [41], as well as the C2 domain of Smurf 2 [42], were shown to bind to their own Hect domain and inhibit their catalytic activity. However, our data suggest that the unique N-terminal or middle regions of dNedd4Lo do not bind nor inhibit the catalytic activity of dNedd4S in vitro. Second, we investigated the effect of dNedd4Lo overexpression on dNedd4S mediated Comm endocytosis in Drosophila S2 cells and body wall muscles. We hypothesized that if dNedd4Lo acts to inhibit the function of dNedd4S, it would interfere with Comm endocytosis. However, our results show that overexpression of dNedd4Lo did not affect internalization of Comm. Therefore, the unique regions of dNedd4Lo regulate NM synaptogenesis by as yet unknown mechanisms, possibly by targeting other substrates.

Interestingly, differential regulation of substrates is known for isoforms of the E3 ligase Cbl, namely CblL (long) and CblS (short). While the long isoform down-regulates EGFR signaling, the short isoform preferentially controls Notch signaling through regulation of the Notch ligand Delta [43]. DNNedd4 might use a similar mechanism to regulate Drosophila embryo development, particularly NM synaptogenesis. In addition, temporal regulation of expression of dNedd4S and dNedd4Lo differs, allowing NM synaptogenesis to proceed at the appropriate time in development.

Materials and Methods

Constructs

**Generation of UAS-dNedd4S (WT), UAS-dNedd4Lo (WT), UAS-dNedd4S (S444->A), and UAS-dNedd4Lo (S39->A, S645->A) constructs.** dNedd4Lo CDNA was kindly provided by Dr. Guy Tear. PCR-amplified full-length dNedd4S (S444->A) and dNedd4Lo (S39->A, S645->A) mutants were synthesized using the Quick Change Site Directed Mutagenesis Kit (Stratagene). The constructs were subcloned into pUAST, KpnI & BamHI sites with an N-terminal Flag tag to generate transgenic lines, as described [29].

**Generation of UAS-dNedd4Lo Nterm, UAS-dNedd4Lo Mid, and UAS-dNedd4Lo C->A mutants.** UAS-dNedd4Lo Nterm: The DNA fragment flanked by the AarI & Bsp1407I sites of Flag-dNedd4Lo WT in pRmHa3 vector was subcloned to replace the region in the Flag-dNedd4S WT-pRmHa3 flankned by the same restriction sites. UAS-dNedd4Lo Mid: the DNA fragment flanked by the AarI&Bsp1407I sites of Flag-dNedd4S WT in pRmHa3 vector was subcloned to replace the region in the Flag-dNedd4Lo WT-pRmHa3 flankned by the same restriction sites. UAS-dNedd4Lo C->A: The DNA fragment flanked by the XhoI & BamHI sites containing the Cys->A mutation (generated by site-directed mutagenesis) of Flag-dNedd4S C->A in pRmHa3 vector was subcloned to replace the region in the Flag-dNedd4Lo WT-pRmHa3 flankned by the same restriction sites. The constructs were subsequently subcloned into pUAST KpnI & BamHI sites to generate transgenic lines, as described [29].

**Generation of GST-dNedd4Lo N-terminus (Nterm) and GST-dNedd4Lo Middle (Mid) for bacterial expression and purification.** The PCR-amplified DNA fragment corresponding to the N-terminal unique sequence of dNedd4Lo ([bp 1 to 189]) was subcloned into pGEX6P1 BamHI & SalI sites with an N-terminal GST tag. The PCR-amplified DNA fragment corresponding to the middle unique sequence of dNedd4Lo ([bp 912 to 1419]) was subcloned into pGEX6P1 BamHI & EcoRI with an N-terminal GST tag. All fly crosses are summarized in Table S4.

**Rescue of dNedd4 null mutant flies.** The fly lines and final crosses generated to rescue dNedd4null homozygote mutant flies with UAS-dNedd4Lo WT or UAS-dNedd4S WT are listed in the Table S4. Adult flies in final crosses were put on grape agar plates to lay eggs, which were collected and
transferred to new plates after ~16 hrs. Plates were observed under a fluorescent microscope to look for rescued larvae, which did not express GFP (non-GFP).

Semi-quantitative RT-PCR

Oregon-R wild-type flies were put on fruit agar plates to lay eggs, which were collected every 2 hours during the 24-hr embryonic development at ~22°C. Embryos were incubated in 50% bleach for 5 min, collected using embryo collection net (Netwell insert, Corning, Acton, MA), and rinsed in PBS (phosphate-buffered saline). Total RNA was extracted from embryos (Qiagen RNeasy Mini Kit) and reverse transcribed to cDNA (Invitrogen SuperScript III Synthesis System). Forward primer: cgacctgtaaatcagtt and reverse primer: ggtgtctctgaccccgatgtt were used to amplify dNedd4S to give a PCR product of 1124 bp using Platinum Taq DNA Polymerase (Invitrogen). Forward primer: ccatggctgcaataagcctgtgtgtctctgaccccgatgtt were used to amplify dNedd4HS to give a PCR product of 1193 bp using Platinum Taq DNA Polymerase (Invitrogen). For Actin 5C: forward primer: ttggtgcaaggaatgtttgctgtgctcttgatc and reverse primer: cctctccccagaataactcctcaac were used to amplify this internal control, yielding a PCR product of 1193 bp. 50 ng of each cDNA was added to 50 µl PCR mixture. PCR conditions were as follows: 94°C for 30 sec, 55°C for 30 sec, and 68°C for 2 min. A series of amplification cycles from 28 to 42 were tested to determine the exponential phase and 34-cycle was chosen to perform the PCR. Equal volumes of the resulting PCR reactions were analyzed by electrophoresis on a 1.5% agarose gel. Band intensities were measured using Image J software to quantify the expression level of each mRNA at each given embryonic stage.

Biochemical assays

For expression of UAS-dNedd4S (WT and mutants) and UAS-dNedd4Lo (WT and mutants) lines. The UAS-dNedd4S (WT and S444->A [S->A]) or UAS-dNedd4Lo (WT, S39->A, S645->A), ΔNterm, ΔMid, C-'>A)÷daughtersless(da-GAL4) embryos were collected at room temperature (RT). The embryos were dechorionated with 50% bleach for 5 min and lysed with lysis buffer (150 mM NaCl, 50 mM HEPES, 1% Triton-X, 10% glycerol, 1.5 mM MgCl2 and 1.0 mM EGTA) plus protease inhibitors (1 mM PMSF, 1 µg/ml each of aprotinin, leupeptin and pepstatin A) to examine expression by Western blotting.

Antibodies used for this and all other experiments are summarized in Table S5.

Phosphorylation assays

S2 cells were transiently transfected with Flag-tagged dNedd4S [WT, dNedd4HS(S444A)], or dNedd4Lo [WT,dNedd4Lo(S39A), dNedd4Lo(S39A,S645A)] in the pRmHA3 vector and lysed in lysis buffer containing phosphatase inhibitors (10 mM NaF, 10 mM Na pyrophosphate, 10 mM Na-orthovanadate). D Ned4S (WT or mutants) or dNedd4Lo (WT or mutants) was immunoprecipitated (IP) with anti-Flag antibodies and blotted with anti Phospho-(Ser/Thr) Akt substrate antibodies (1:1000,Cell Signaling Technology).

S2 cells immunostaining

Live S2 cells co-expressing Comm [WT or 2PY->A]-GFP [19] and Flag-dNedd4 (S or Lo, WT or C->A) were washed, fixed in 2% PFA (10 min), permeabilized with 0.1% Saponin, stained with M2 anti-Flag antibody and Cy3-conjugated goat anti-mouse antibody (Table S5), and visualized by confocal microscopy.

In vitro ubiquitylation assays

Full-length dNedd4S protein immunopurified from HEK293T cells was incubated in reaction mixtures containing 100 nM human E1 ubiquitin-activating enzyme (BostonBiochem), 200 nM human E2 ubiquitin-conjugating enzyme (UbC-H5; BostonBiochem), 1 µg of ubiquitin (Sigma) and 4 mM ATP in a reaction buffer [25 mM Tris/HCl pH 7.5, 50 mM NaCl, 0.1 mM dithiothreitol and 4 mM MgCl2]. Reactions were incubated for 1 hr at RT. Protein ubiquitylation was detected on western blots. Membranes were blocked and incubated with mouse anti-ubiquitin antibody and HRP-conjugated goat-antimouse antibody to detect ubiquitin (Table S5). To detect Flag-dNedd4HS, membranes were incubated with M2 anti-Flag antibody and HRP-conjugated goat anti-mouse antibody. Mouse anti-glutathione-S-transferase (GST) antibody and HRP-conjugated goat anti-mouse antibody were used to detect bacterially expressed GST-dNedd4Lo Nterm and GST-dNedd4Lo Mid.

In vitro binding assay

Flag-dNedd4S protein immunopurified from HEK293T cells was immobilized on M2 anti-Flag agarose beads (Sigma) and incubated with bacterially expressed GST-dNedd4Lo Nterm or GST-dNedd4Lo Mid at 4°C for 2 hrs. Supernatant was removed and the beads were washed with lysis buffer. Mouse anti-GST antibody and HRP-conjugated goat-antimouse antibody were used to detect GST-dNedd4Lo Nterm or GST-dNedd4Lo Mid (Table S5). Flag-dNedd4HS was detected using M2 anti-Flag antibody and HRP-conjugated goat anti-mouse antibody.

Larval Locomotor Activity

Fly crosses generated to analyze larval locomotion are listed in Table S4. 150-sec path length was determined for the following heterozygous genotypes: (1) Driver control line, +/24B-GAL4; (2) UAS control line, +/UAS-dNedd4Lo; and (3) Over-expression line, +/UAS-dNedd4HS; and +/24B-GAL4 to analyze their locomotor activity. All experiments were performed at 24±1°C in a climate-controlled room with 45–50% humidity with a 12/12 hr light/dark cycle. All animals were tested at the pre-wandering physiological stage and the stage was confirmed as previously described [45]. The same number of 3 day old adult flies were put on grape agar plates to lay eggs and transferred to new plates every 24 hrs. After 28 hrs, hatched larvae were cleared from the plate and all newly hatched larvae were collected 2 hrs later. Twenty larvae (3 repetitions per genotype) were separated into vials containing standard fly food medium. Larvae from over-expression lines showed a delay in development, taking 172±4 h after egg-laying to reach the wandering stage, while control larvae reached this stage after 164±4 h. Thus, for all experiments, larvae were aged 168±2 h for the over-expression line and 158±2 h for controls. To measure locomotor activity, larvae were rinsed and placed in a Petri dish containing 3% grape agaroze for 30 sec to adapt to the experimental conditions. The crawling path for each larva was traced for 150 sec on the lid of the dish and scanned for subsequent digital analysis. Image J software (National Institutes of Health) was used for quantification of crawling path length.
**Statistical Analyses**

GraphPad Prism 5.0 software (GraphPad Software, San Diego, CA) and Fisher’s exact test in a 2×2 contingency table were used to analyze the data of motor neuron innervation on body wall muscles, as well as protein localization of dNedd4 and Comm in S2 cells. A p-value of <0.05 was considered significant. For analysis of locomotor activity, the behavioral data were Box-Cox transformed to correct problems of distribution and non-homogeneity of variance before transformation. Normal distribution was confirmed by the Kolmogorov-Smirnov test. The common variance of the three lines was tested by the Levene test ($F_{2, 168} = 2.47, p = 0.087$). One-way ANOVA and subsequent Turkey post-hoc comparisons were performed to evaluate differences among lines. A p-value of <0.01 was considered significant. Statistical analyses were conducted using Statistica 8.0 (StatSoft) software. For analysis using semi-quantitative RT-PCR, Student’s t-test was used to compare the mRNA expression level of dNedd4Lo versus dNedd4S at each given developmental stage during embryogenesis. A p-value of <0.01 was considered significant.

**Supporting Information**

**Figure S1** Catalytic activity of dNedd4S and dNedd4Lo, and normal cellular localization of these isoforms ectopically expressed in salivary glands of third instar larvae. (A) Catalytic activity of dNedd4S and dNedd4Lo: In vitro ubiquitylation assay of wildtype (WT) dNedd4S or dNedd4Lo was performed by incubating E1, E2 (UbcH5), E3 (Flag-tagged dNedd4S or dNedd4Lo), ubiquitin and ATP, and the extent of ubiquitylation (most likely reflecting autoubiquitylation) analyzed by immunoblotting with anti-ubiquitin antibodies. Note the loss of ubiquitylation of the catalytically-inactive C->A mutants of the dNedd4 isoforms. Lower panels: The blot was stripped and re-blotted with anti-Flag antibody to show equal amount of Flag-dNedd4S present in all panels. The blot was also stripped and re-blotted with anti-GST antibody. The catalytically inactive dNedd4S C->A mutant was included as a negative control to demonstrate that the ubiquitylation activity observed was mediated by dNedd4S.

**Figure S2** The unique regions of dNedd4Lo do not inhibit catalytic activity of dNedd4S. In vitro ubiquitylation activity of dNedd4S in the presence of GST alone (control), GST-tagged dNedd4Lo Nterm, Mid, or both unique regions, detected using anti-ubiquitin antibody on western blots. E1, E2 (UbcH5), E3 (dNedd4S), ubiquitin and ATP were included in the ubiquitylation reactions, as well as increasing concentrations (0.9 μM, 1.8 μM and 3.6 μM) of each potential inhibitor (GST, GST-Nterm, GST-Mid, or GST-Nterm+GST-Mid). Middle panel: The blot was stripped and re-blotted with anti-Flag antibody to show equal amount of Flag-dNedd4S present in all lanes. Bottom panel: The blot was also stripped and re-blotted with anti-GST antibody. The catalytically inactive dNedd4S C->A mutant was included as a negative control to demonstrate that the ubiquitylation activity observed was mediated by dNedd4S.

**Table S1** Lethality test for ubiquitous overexpression of dNedd4S (WT and S->A mutant) and dNedd4Lo (WT and S->A mutant) using different GALA enhancer drivers at different temperatures. (DOCX)

**Table S2** Lethality test for overexpression of dNedd4S WT, dNedd4Lo WT and their S->A mutants in different tissues using tissue-specific GALA enhancer drivers. (DOCX)

**Table S3** Lethality test for ubiquitous (Daughterless, Actin, and Tubulin) and muscle-specific (24B and 5) overexpression of UAS-dNedd4LoNterm, UAS-dNedd4LoMid and UAS-dNedd4Lo C->A using different GALA enhancer drivers at different temperatures. (DOCX)

**Table S4** Fly line crosses used for the experiments. (DOCX)

**Table S5** Antibodies used for staining. (DOCX)

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**Author Contributions**

Conceived and designed the experiments: YZ ASK DR. Performed the experiments: YZ ASK KR LN. Analyzed the data: YZ ASK KR. Contributed reagents/materials/analysis tools: GLB. Wrote the paper: YZ ASK KR GI.

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