Wnt Signaling in Neuromuscular Junction Development

Kate Koles and Vivian Budnik

Department of Neurobiology, University of Massachusetts Medical School, Worcester, Massachusetts 01605
Correspondence: vivian.budnik@umassmed.edu

Wnt proteins are best known for their profound roles in cell patterning, because they are required for the embryonic development of all animal species studied to date. Besides regulating cell fate, Wnt proteins are gaining increasing recognition for their roles in nervous system development and function. New studies indicate that multiple positive and negative Wnt signaling pathways take place simultaneously during the formation of vertebrate and invertebrate neuromuscular junctions. Although some Wnts are essential for the formation of NMJs, others appear to play a more modulatory role as part of multiple signaling pathways. Here we review the most recent findings regarding the function of Wnts at the NMJ from both vertebrate and invertebrate model systems.

Wnts and their receptors comprise a large family of proteins. (For a description of these families, see He [2012], Willert and Nusse [2012], and www.stanford.edu/group/nusselab/cgi-bin/wnt.) Frizzled receptors form a functional heterocomplex with members of the low-density lipoprotein receptor (LDLR)-related protein (LRP) family, Lrp5 and Lrp6 (Pinson et al. 2000; Tamai et al. 2000) and its Drosophila homolog, Arrow (Wehrli et al. 2000). Furthermore, there are families of proteins belonging to the unconventional Wnt receptors, such as though important roles for Wnt signaling have become known from studies in both the central and peripheral nervous system, this article is concerned with the role of Wnts at the NMJ.

WNT LIGANDS, RECEPTORS, AND WNT SIGNALING PATHWAYS

Wnts and their receptors comprise a large family of proteins. (For a description of these families, see He [2012], Willert and Nusse [2012], and www.stanford.edu/group/nusselab/cgi-bin/wnt.) Frizzled receptors form a functional heterocomplex with members of the low-density lipoprotein receptor (LDLR)-related protein (LRP) family, Lrp5 and Lrp6 (Pinson et al. 2000; Tamai et al. 2000) and its Drosophila homolog, Arrow (Wehrli et al. 2000). Furthermore, there are families of proteins belonging to the unconventional Wnt receptors, such as
members of the ROR (receptor tyrosine kinase-like orphan receptor) (Green et al. 2008) and Derailed (Drl)/Ryk (related to receptor tyrosine kinase) families (Fradkin et al. 2010). Adding to this diversity of ligands and receptors is the number of distinct signaling pathways activated by Wnt ligand/receptor interactions, namely, the canonical Wnt pathway (Logan and Nusse 2004), the divergent canonical Wnt pathway (Ciani et al. 2004), the planar cell polarity (PCP) pathway (Mlodzik 2002), the calcium Wnt signaling pathway (Kuhl et al. 2000), and the Frizzled nuclear import (FNI) Wnt pathway (Speese and Budnik 2007). Given that most of the transduction pathways are described in Cadigan and Peifer (2012), He (2012), Stamos and Weis (2012), and van Amerongen (2012), only a brief description of the pathways is provided in this review, except for the FNI pathway, which so far has only been characterized in the nervous system. In the canonical Wnt pathway, Wnt binding to Frizzled receptors activates Dishevelled (Dvl), which inhibits the phosphorylation of β-catenin by glycogen synthase kinase 3B (Gsk3B), through the disassembly of a destruction complex, which normally triggers the degradation of cytoplasmic β-catenin. Wnt binding to Frizzled receptors disrupts this complex, leading to the stabilization of β-catenin, its import to the nucleus, and the activation of Wnt target genes (Fig. 1A). In the divergent canonical Wnt pathway, Dvl regulates the Gsk3B-dependent phosphorylation of microtubule-associated proteins (MAPs), which control the stability of microtubule networks (Fig. 1B) (Lucas et al. 1998; Ciani et al. 2004). The PCP Wnt pathway regulates tissue and cell polarity through a Dvl-dependent activation of small GTPases, such as Rho, Rac, and Jun amnio-terminal kinase (Jnk) (Fig. 1C). In the Wnt calcium pathway, cell fate decisions and cell movement choices are made via a Dvl-dependent increase in intracellular calcium levels, which leads to the activation of protein kinase C (PKC) and calcium/calmodulin-dependent protein kinase II (CaMKII), the nuclear import of the nuclear factor of activated T-cells (NFAT) and concomitant changes in gene expression (Fig. 1D). Finally, in the FNI Wnt pathway, Frizzled-2 receptors are endocytosed and trafficked toward the nucleus, where a carboxy-terminal fragment of Frizzled-2 is cleaved and imported into the nucleus (Fig. 1E) (Mathew et al. 2005). Within the nucleus, Dfz2C fragments associate with large ribonucleoprotein (RNP) granules, which exit the nucleus through a novel mechanism of nuclear envelope budding, akin to the nuclear egress of Herpes-type viruses (Roller 2008; Speese et al. 2012). The large RNPs contain transcripts encoding postsynaptic proteins, and because these transcripts are also localized at postsynaptic sites, it is presumed that they are involved in local translation (Speese et al. 2012). Indeed, genetic manipulations interfering with the FNI pathway lead to abnormal synaptic growth (Mathew et al. 2005; Speese et al. 2012) (discussed below). A similar pathway of receptor endocytosis, cleavage, and fragment import to the nucleus is observed for the atypical Wnt receptor Ryk (Lyu et al. 2008). However, in this case, the pathway is required for normal cortical neurogenesis, as opposed to synapse development, and the exact role of Ryk receptor fragments within the nucleus has not been investigated. Most importantly, through alternative ligands, receptors, and signaling pathways, Wnts have considerable potential to regulate development differentially.

Figure 1. Overview of Wnt signaling pathways. Wnts signal through Frizzled receptors that can form a complex with Lrp-5/6 receptors. With the exception of the FNI pathway, these signaling pathways activate the scaffolding protein Dishevelled (Dvl). The different pathways diverge downstream from Dvl. (A) In the canonical pathway, Dvl disrupts the “destruction complex” composed of Axin, Adenomatous Polyposis Coli (APC), and Gsk3B. This prevents the phosphorylation of β-catenin by Gsk3B, which leads to its stabilization and subsequent nuclear import. In the nucleus, β-catenin associates with the transcription factor T-cell factor/lymphoid enhancer factor 1 (TCF/Lef1) and regulates gene expression. The secreted proteins Dickkopf1 (Dkk1) and secreted Frizzled-related protein (Sfrp) inhibit this pathway. (Legend continues on facing page.)
Figure 1. (Continued) (B) Following Wnt activation of the divergent canonical pathway, Dvl inhibits Gsk3β, which leads to the decreased phosphorylation of microtubule-associated proteins (MAPs), such as Tau, MAP-1B, MAP-2, or the related *Drosophila* protein Futsch. (C) In the planar cell polarity pathway, Dvl activates the small Rho GTPases RhoA and Rac1, and c-Jun–amino-terminal kinase (Jnk), which leads to the regulation of actin and microtubule cytoskeletons. (D) In the Wnt calcium pathway, Dvl activation leads to a rapid increase in intracellular calcium levels and subsequent activation of calcium-sensitive enzymes such as protein kinase C (PKC), calcium/calmodulin-dependent protein kinase II (CaMKII) and the Ca$^{2+}$-activated phosphatase calcineurin (Cn). Calcineurin directly binds to and dephosphorylates nuclear factor of activated T-cells (NFAT), which, in turn, allows their nuclear translocation where they regulate gene expression. (E) In the Frizzled nuclear import pathway (FNI), Wnt binding to Frizzled-2 (DFz2) leads to its endocytosis and Grip-dependent trafficking to the periphery of the nucleus. Here the carboxyl terminus of DFz2 is cleaved and imported. In *Drosophila*, DFz2-C organizes ribonucleoprotein (RNP) particles containing specific transcripts, which then egress from the nucleus and are likely translated in the cytoplasm. A similar nuclear import pathway has also been described for the atypical Ryk receptor during cortical neurogenesis (Lyu et al. 2008).
BRIEF OVERVIEW OF VERTEBRATE NMJ DEVELOPMENT

Vertebrate NMJs are among the longest and best-studied peripheral synaptic units because of their simplicity and accessibility. Motor neuron axons innervate the muscle at the end plate, where axon terminals branch into fine synaptic varicosities, apposed to foldings of the muscle membrane, the junctional folds. The crests of these folds appose presynaptic active zones, the sites of acetylcholine (ACh)–containing vesicle exocytosis at presynaptic terminals, and contain a high density of neurotransmitter receptors. In contrast, the valleys of the junctional folds and their basement membrane are enriched in acetylcholinesterase (AChE), which rapidly degrades acetylcholine, thus terminating ACh-induced responses.

Even before nerve innervation, AChR clusters form in a broad band at the muscle midline, a process called AChR prepatterning (Yang et al. 2001). These “preformed” clusters are also called aneural AChR clusters to underscore their formation independently of innervation. Once the motor axon interacts with the muscle, AChR clusters form at postsynaptic regions exactly opposed to the sites of nerve contact. These mature clusters (also called neural AChR clusters) are formed by the recruitment of aneural AChR clusters as well as by the targeted accumulation of newly synthesized AChRs (Sanes and Lichtman 2001).

Despite its relative simplicity, the vertebrate NMJ has provided us with remarkable insight into the mechanisms of synapse development. Adding to the multiple players found to participate in NMJ development is the recent discovery that, not totally unexpectedly, these also include members of the Wnt family.

An important initial discovery in the quest to understand NMJ development was the identification of Agrin, originally isolated from the electric organ of the pacific electric ray (Smith et al. 1987) as a protein that could trigger AChR clustering in cultured myotubes (Wallace 1986). These prolific initial studies also led to the model that Agrin, a heparan sulfate proteoglycan, was secreted by neurons, accumulated in the basal lamina at the synaptic cleft, where it triggered a molecular cascade in the muscle. This signaling cascade led to the phosphorylation of the β-subunit of AChR and the redistribution of AChRs to the NMJs in association with a 43-kDa AChR-associated protein. This 43-kDa AChR-associated protein was subsequently identified as Rapsyn (receptor-associated protein at synapses) (Frail et al. 1988). Muscles and Schwann cells were also known to express Agrin, but only the neuronal isoform was sufficiently potent to induce AChR clustering (GeSEMANN et al. 1995). Interestingly, agrin−/− mice were able to form aneural AChR clusters before innervation, indicating that Agrin was not required for prepatterning (Lin et al. 2001). During NMJ maturation, the preformed AChR clusters outside of the innervated area are dispersed upon acetylcholine release, and from this perspective, Agrin, which prevents the dispersal of neural AChR clusters, can also be considered as an “anti-declustering factor,” in addition to its AChR clustering role (Lin et al. 2005; Kummer et al. 2006).

During NMJ maturation, the preformed AChR clusters outside of the innervated area are dispersed upon acetylcholine release, and from this perspective, Agrin, which prevents the dispersal of neural AChR clusters, can also be considered as an “anti-declustering factor,” in addition to its AChR clustering role (Lin et al. 2005; Kummer et al. 2006).

One of the key milestones after the identification of Agrin was cloning its muscle-specific receptor, aptly named muscle-specific kinase (MuSK) (Jennings et al. 1993; Valenzuela et al. 1995; DeChiara et al. 1996; Glass et al. 1996). Unlike Agrin, MuSK was required for both neural and aneural AChR clustering in mice and zebrafish (Yang et al. 2001; Zhang et al. 2004). Surprisingly, however, during neural AChR clustering, Agrin did not interact directly with MuSK, suggesting the existence of an MASC (myotube-associated specificity component) (Glass et al. 1996).

WNT SIGNALING AT THE VERTEBRATE NMJ

It was around this time that a link between NMJ development and Wnt signaling was suspected, because the newly cloned MuSK contained a conserved cysteine-rich domain (CRD) found in Frizzled receptors, and this domain was known to be required for Wnt ligand binding (Masiakowski and Yancopoulos 1998). However, experimental evidence supporting the ability of Wnts to bind the MuSK CRD domain was
only obtained a decade later (Jing et al. 2009).

During the intervening decade, a more pressing challenge had been the identification of the mysterious MASC.

First, Weatherbee et al. (2006) reported that NMJ abnormalities in Lrp4−/− mice had striking similarities to those observed in Musk−/− mutants. Lrp4 is an Lrp family member with an extracellular domain resembling that of Lrp5 and 6/Arrow, but that appears to antagonize Wnt signaling during digit formation in mice (Johnson et al. 2005). Lrp4 has also been shown to interact with postsynaptic density protein 95 (PSD-95) in the brain (Tian et al. 2006). Subsequently, two independent groups reported that Lrp4 was the long-sought Agrin binding receptor (Kim et al. 2008; Zhang et al. 2008). These studies showed that Lrp4 was expressed at NMJs in myotubes, where it bound selectively to the neuronal isoform of Agrin and was necessary for Agrin-induced activation of MuSK and AChR clustering. Indeed, recent detailed structural studies by Zhang and colleagues have now revealed that several of the extracellular domains on Lrp4 serve as ligands for MuSK and that Agrin dramatically stimulates this interaction (Gomez and Burden 2011; Zhang et al. 2011).

Following Agrin-stimulated Lrp4 binding to MuSK, the receptor is tyrosine-phosphorylated and recruits Dok-7 (Inoue et al. 2009), a muscle-specific adaptor protein. Like MuSK, Dok-7 is essential for NMJ formation, because in its absence NMJs fail to form, and Dok-7 overexpression dramatically stimulates AChR clustering (Okada et al. 2006). Dok-7 acts through two different pathways determined by its amino-terminal and carboxy-terminal protein interactions. First, Dok-7 dimerizes (Bergamin et al. 2010), and its amino terminus stimulates MuSK activity. However, this activity of Dok-7 only minimally affects postsynaptic differentiation (Hallock et al. 2010). Second, Dok-7, via its carboxyl terminus, also recruits two adaptor proteins, CT10 regulators of kinase, Crk and Crk-L, downstream from MuSK activation (Hallock et al. 2010; Burden 2011), which are required in the muscle for normal presynaptic and postsynaptic NMJ differentiation. In Crk and Crk-L double mutants, motor axon terminals occupied a larger muscle area than controls and had significantly reduced synapse numbers, with several muscle fibers lacking innervation. These mutants die at birth because of the inability to inflate their lungs. Interestingly, many patients with congenital myasthenia have mutations in the carboxyl terminus of Dok-7, and these results provide insight into the possible molecular pathways affected in this disease. Together with Agrin, Rapsyn, MuSK, and Lrp4, Dok-7 represents an indispensable component of the molecular pathway essential for NMJ formation in vertebrates. Numerous additional molecules and pathways involved in the assembly of NMJ have been revealed over the past few years. These are discussed in excellent reviews (Kummer et al. 2006; Wu et al. 2010).

WNTs AS POSITIVE REGULATORS OF NMJ DEVELOPMENT

The first experimental evidence implicating Wnt signaling in NMJ development was provided when a yeast two-hybrid screen geared to identify intracellular binding partners of MuSK led to the identification of Dishevelled1 (Dvl1), a scaffolding protein activated by several Wnt signaling pathways (Luo et al. 2002). Gain- and loss-of-function experiments in vivo and in vitro showed that Dvl1 was required for the Agrin-mediated clustering of AChRs. In addition, the Dvl1-interacting protein p21-activated kinase 1 (PAK1) was identified, and it was shown that its activation by Agrin required Dvl1. MuSK, Dvl, and PAK1 formed a ternary complex required for AChR clustering in muscle. Furthermore, expression of a Dvl1 mutant transgene lacking the DEP (Dishevelled-Egl-10-Pleckstrin) domain in muscle cells of Xenopus nerve–muscle cocultures led to decreased spontaneous synaptic current frequency and amplitude, indicating a potential muscle-derived retrograde signal that could influence presynaptic neurotransmitter release (Luo et al. 2002).

Subsequently, it was shown that the β-subunit but not the α1-, α3-, α7-, or ε-subunits of AChR interacted with APC, and its function was required in Agrin-induced AChR clustering (Wang et al. 2003). APC is a well-known regulator of β-catenin, a key effector of canonical
Wnt signaling (Aoki and Taketo 2007). APC is also known to interact with microtubules, actin filaments, and EB1 (Su et al. 1995), a microtubule plus-end tracking protein that negatively regulates APC–actin interaction (Reilein and Nelson 2005; Moseley et al. 2007). It was suggested that APC supported a cross talk between Agrin- and Wnt-mediated signaling pathways, by coordinating the functions of actin and the microtubule cytoskeleton. Notably, the APC–AChR interaction was not unique to muscle-specific AChRs, because APC also regulated the synaptic localization of neuronal α3-nicotinic AChRs by binding to PSD-93, β-catenin, and EB1 (Temburni et al. 2004; Rosenberg et al. 2008). Furthermore, APC functions presynaptically as well, where it increases the clustering and membrane localization of the α7-nicotinic AChR in response to Wnt-7a signaling (Farias et al. 2007). All of these data implicate APC as a Wnt downstream effector that is crucially involved in the localization of select AChRs both at neuronal and NMJ synapses.

More recently, it was found that Rapsyn binds to β-catenin (Zhang et al. 2007). Rapsyn has been known to associate with AChRs in an Agrin-dependent manner (Moransard et al. 2003), and, indeed, β-catenin associated with AChR following Agrin treatment. This Agrin-triggered association did not occur in rapsyn−/− mice, indicating that Rapsyn is required for β-catenin association with AChR. Reducing β-catenin levels in C2C12 myotubes by shRNA resulted in decreased AChR clustering following Agrin treatment (Zhang et al. 2007; but see Li et al. 2008). To understand the role of β-catenin in vivo, β-catenin was conditionally knocked out only in the muscles of mice. Surprisingly, in addition to muscle-specific phenotypes, such as the increased size and wider distribution of AChR clusters, primary nerve branches were mislocalized, evoked and spontaneous neurotransmitter release was decreased, and short-term plasticity was abnormal in these mice (Li et al. 2008). In contrast, Agrin-induced AChR clustering was unaffected in cultured muscles from β-catenin−/− mice. Most importantly, β-catenin conditional knockout in neurons did not have any effect on NMJ morphology or function, implying that muscle β-catenin is involved in presynaptic development and function in an as-yet-unidentified retrograde pathway (Fig. 3) (Li et al. 2008).

In addition to Wnt signaling pathway components, some of the Wnt ligands have also been identified. Whereas Wnt1 (Luo et al. 2002), Wnt4, Wnt6, and Wnt7b (Zhang et al. 2007) were found to have no effect on AChR clustering, Wnt3, which is expressed in mouse motor neurons at the time of innervation (Krylova et al. 2002), enhanced AChR clustering in developing chicken wing NMJs and the clustering ability of Agrin in mouse cultured myotubes (Henriquez et al. 2008). Dickkopf-1 (Dkk1), which specifically blocks canonical Wnt signaling (Fig. 1A) (Niehrs 2006), had no effect, whereas secreted Frizzled-related protein 1 (Sfrp1), which blocks both canonical and non-canonical Wnt signaling (Rattner et al. 1997), reduced the number of AChRs induced by Wnt3, indicating that Wnt3 is likely to mediate this effect via a non-canonical Wnt signaling pathway in myotubes, even though Wnt3 can activate canonical signaling (Barrow et al. 2003). It was also found that Wnt3 induced a quick, Agrin-independent transient up-regulation of Rac1 activity, which led to the rapid formation of AChR microclusters (Fig. 2A). These microclusters were subsequently stabilized into full-sized clusters in an Agrin-dependent manner, in a process involving Rho signaling (Fig. 2B) (Weston et al. 2003; Henriquez et al. 2008). These observations led to a model in which Wnt3 released from motor neurons induces the first wave of Rac1 activation and subsequent AChR microclustering, which is then consolidated into mature AChR clusters via Agrin-mediated signaling. Nevertheless, the muscle receptor(s) through which Wnt3 signals this activation remain to be identified.

A role for Wnts in both AChR prepatterning and axon guidance is also provided by studies in zebrafish. As discussed above, before the arrival of motor nerve terminals to the muscle field, AChRs spontaneously cluster in the mid-region of the muscle, where prospective NMJs eventually form, and this prepatterning depends on MuSK (Lin et al. 2001; Yang et al. 2001).
Figure 2. Wnt signaling at vertebrate neuromuscular junctions. (A) Mouse Wnt3 and zebrafish Wnt11r function during prepatterning. Wnt11r signaling through Unplugged/MuSK before innervation leads to aneural AChR clustering and also via a parallel signaling pathway to changes in extracellular matrix modifications (EMMs), possibly via chondroitin sulfate proteoglycans (CSPGs) that eventually influence incoming motor neuron axonal growth cone guidance. Wnt3 signaling leads to the rapid and transient activation of Rac1 and AChR microcluster formation, which (B) in the presence of neural agrin at the time of innervation are stabilized in exact opposition to the presynaptic terminal.

Cite this article as Cold Spring Harb Perspect Biol 2012;4:a008045
Many of these aneural clusters are incorporated into neural AChRs clusters upon nerve arrival (Flanagan-Steet et al. 2005). However, the function of prepatterning has been a matter of debate, partly owing to the observations that some nerves can ignore preformed AChR clusters and initiate NMJ development at new areas (Anderson and Cohen 1977), raising the possibility that the nerve, or nerve-derived factors, are instructive for NMJ formation. Elegant studies by Jing et al. (2009) address this long-standing dilemma. Live imaging of wild-type and MusK−/− mutants (which in zebrafish is called “unplugged”) revealed that neuronal growth cones explored a wider muscle area and invaded muscle territories that they usually avoid. Because unplugged/MuSK is a muscle-specific gene, these data indicate that muscle plays a role in restricting the area that growth cones explore during innervation. Based on expression time and pattern (spinal cord and adaxial somites) in addition to its mutant phenotype, the investigators identified the non-conventional Wnt11r, as required for AChR prepatterning on the adaxial muscles. Live imaging of motor growth cones revealed that wnt11r morpholino-induced abnormalities in prepatterning were virtually identical to those observed in unplugged/MuSK−/− mutants, suggesting that Wnt11r might act through unplugged/MuSK. In contrast, neurally induced AChR clusters were formed and developed normally in wnt11r morpholino-treated animals, suggesting that Wnt11r function is restricted to prepatterning. That Wnt11r and Unplugged/MuSK were mutually required for prepatterning was confirmed by showing genetic interactions between the genes. In addition, by using rescue experiments with the short (containing the CRD and Kringle domains) or the long (containing three amino-terminal Ig domains) Unplugged, it was shown that the CRD domain of Unplugged/MuSK was required for physical interaction between Wnt11r and Unplugged. Expressing a Dvl isoform that interferes with non-canonical Wnt signaling also led to prepatterning defects, implicating it in the signaling pathway downstream from Wnt11r.

Interestingly, in the absence of prepatterning, neurally induced AChR clusters and functional NMJs were formed, albeit being slightly smaller. However, in the absence of prepatterning, axonal growth cones were not directed to the narrower mid-region of the prospective NMJs, and they explored much broader muscle areas. The investigators conclude that one of the functions of prepatterning is to guide incoming axons (perhaps via the Wnt11r or Unplugged signaling-induced modifications to the extra-cellular matrix) and to restrict muscle AChRs cluster formation to a common muscle area for efficient NMJ formation (Fig. 2A) (Jing et al. 2009).

NEGATIVE REGULATION OF NMJs BY WNT

Negative regulation of AChR clustering by Wnts has also been documented. Wnt3a expression in the muscle gradually increases during the phase of embryogenesis that coincides with NMJ development (Wang et al. 2008). However, pretreatment of cultured myotubes with Wnt3a-containing conditioned medium before Agrin application significantly reduced AChR aggregation. Moreover, transfection of C2C12 myotubes with Wnt3a, but not Wnt5, Wnt7a, or Wnt11, reduced Agrin-mediated AChR clustering. Transfection with Dkk1, which antagonizes the canonical Wnt pathway, increased AChR clustering, indicating that canonical Wnt signaling negatively regulates NMJ development. The stability of AChR clusters was also reduced when Wnt3a was applied after Agrin-induced clustering. The effects of Wnt3a were mediated via β-catenin and resulted in reduced Rapsyn expression. Reduction of β-catenin by shRNA resulted in decreased Agrin-dependent AChR clustering, likely because of the disrupted β-catenin–Rapsyn interaction (Zhang et al. 2008) or the increased Rapsyn expression (Yoshihara and Hall 1993; Wang et al. 2008). Downstream from Wnt3a, β-catenin is known to function as a cotranscription factor with TCF (Korinek et al. 1997). However, dominant negative TCF had no effect on AChR clustering, indicating that β-catenin function is independent of TCF-mediated...
transcription (Zhang et al. 2007; Li et al. 2008). These results indicate that muscle Wnt3a disperses AChR clusters by down-regulating Rapsyn expression in a β-catenin-dependent manner (Fig. 3). In a larger context, these observations highlight the importance of both positive and negative Wnt signals in sculpting the mature NMJ.

In summary, Wnt proteins have been found to play multiple roles in all stages of vertebrate NMJ formation, both in the form of positive and negative signaling pathways. At least two Wnts, muscle Wnt3 and non-muscle Wnt11r, acting through non-canonical pathways, appear to function during prepatterning predominantly by modulating AChR precluster formation, which are subsequently stabilized into mature AChRs in an agrin-dependent manner. In contrast, Wnt3a induces the dispersal of AChR clusters in the presence of Agrin, suggesting that the balance of positive and negative signaling pathways is at work during NMJ development.

**WNT SIGNALING AT INVERTEBRATE NMJs**

Two invertebrate systems, *Drosophila melanogaster* (fruit fly) and *Caenorhabditis elegans* (roundworm), have been influential in uncovering the function of Wnt family molecules at the neuromuscular junction (NMJ). The ease of genetic manipulation in these relatively simple and highly accessible model organisms has provided a wealth of groundbreaking insight into the development and plasticity of NMJs.

**THE ROLE OF WNTs AT THE LARVAL NMJ OF DROSOPHILA**

*Drosophila* larvae offer an exceptionally accessible model system to study the development and plasticity of NMJs. The larval body wall muscles are innervated by excitatory glutamatergic motor neurons that terminate in stereotypical NMJs with easily discernible synaptic boutons. In response to the rapid growth of the larva and the consequent growth of its body wall muscle...
cells after hatching from the egg, NMJs show a 10-fold increase in synaptic bouton number and size during larval development (Gorczyca et al. 1993; Schuster et al. 1996). In addition, larval NMJs undergo structural and functional modifications in response to experience and electrical activity (Budnik et al. 1990; Sigrist et al. 2003; Steinert et al. 2006; Ataman et al. 2008; Koon et al. 2011), offering a powerful experimental system to study the molecular processes underlying synaptic growth and plasticity.

A role for *Drosophila* Wnt1, Wingless (Wg), in NMJ development was first discovered by Packard et al. (2002), when they found that Wg was present at synaptic boutons of the NMJ and that transient reduction in Wg secretion prevented NMJ expansion during muscle growth (Packard et al. 2002). NMJs from these temperature-sensitive *wg* mutants (wgts) shifted to restrictive temperature for 12–18 h during late larval development, also had NMJs containing unusually large and irregularly shaped boutons. These abnormal boutons also displayed abnormal distribution of glutamate receptor subunit, DGlutRIA, and the PDZ protein of the PSD95-family Discs-Large (Dlg). Most strikingly, several synaptic boutons, later called “ghost boutons” (Ataman et al. 2006), were filled with synaptic vesicles yet lacked mitochondria, presynaptic active zones, and the postsynaptic subsynaptic reticulum (SSR) (Packard et al. 2002). These ghost boutons were later found to be devoid of any postsynaptic protein and lacked most labeling for the active-zone protein Bruchpilot (Brp) (Ataman et al. 2006, 2008; Korkut et al. 2009). Live imaging of NMJs during acute elevation of synaptic activity, which resulted in rapid formation of ghost boutons, together with the observation that ghost boutons are also observed at wild-type NMJs, albeit at very low frequencies (Ataman et al. 2008), led to the conclusion that ghost boutons are a normal undifferentiated stage during the development of synaptic boutons. In Wnt mutants, the differentiation of ghost boutons is blocked, leading to a reduced number of boutons and the accumulation of ghost boutons.

The presynaptic microtubule cytoskeleton at the NMJ was also altered, because labeling NMJs with the microtubule-associated protein 1B (MAP-1B)–related protein Futsch, which is phosphorylated by Gsk3β/SHaggy (Sgg) (Gogel et al. 2006), revealed abnormal microtubule bundling (Packard et al. 2002). Wild-type NMJs display a characteristic proportion of microtubule loops (thought to be stable microtubule bundles), splayed microtubule bundles, and fragmented microtubules (both likely representing microtubules undergoing remodeling) throughout NMJ branches (Roos et al. 2000; Packard et al. 2002; Ruiz-Canada et al. 2004). However, the proportion of looped, splayed, and fragmented microtubules was altered in the *wg* mutant (Packard et al. 2002), again suggesting an abnormal development of the NMJ cytoskeleton in the absence of Wg signaling.

These defects were shown to be mediated by the Wg receptor *Drosophila* Frizzled-2 (DFz2) (Packard et al. 2002; Ataman et al. 2006) and its coreceptor Arrow/Lrp5/6 (low-density lipoprotein receptor-related protein) (Miech et al. 2008). These receptors are likely to function both presynaptically in the motor neurons and postsynaptically in the muscle. Efforts to delineate the exact signaling pathway triggered by Wg release led to the finding that Wg, likely released by synaptic boutons (Packard et al. 2002), activated alternative signaling pathways in the presynaptic compartment and the postsynaptic apparatus (Mathew et al. 2005; Ataman et al. 2006, 2008; Mosca and Schwarz 2010; Speese et al. 2012).

Using antibodies specifically recognizing the carboxyl terminus and amino terminus of DFz2, it was shown that in postsynaptic muscles, the DFz2 receptor was endocytosed and trafficked to the periphery of the muscle nucleus, likely in an
endocytic vesicle, where its cytoplasmic carboxy-terminal fragment was cleaved. Whereas the 8-kDa carboxy-terminal fragment of DFz2 (DFz2C) was imported into the nucleus, the amino-terminal fragment remained at the nuclear periphery. Within the nucleus, DFz2C formed submicrometer-sized foci (Mathew et al. 2005). Overexpressing Wg in motor neurons, which increased the levels of secreted Wg around synaptic boutons, led to an increase in the number of nuclear DFz2C foci in the muscles, and decreasing Wg secretion in wg mutants or interfering with DFz2 signaling in muscles led to the opposite effect (Mathew et al. 2005).

The cleavage of DFz2 was found to be required for Wg signaling in the muscle. Expression of a DFz2 transgene in the muscle, carrying a mutation that eliminated the cleavage site, in a dfz2 mutant background did not rescue the reduced bouton number observed in dfz2 mutants (Mathew et al. 2005). Importantly, attempts to circumvent the nuclear import pathway by expressing a tagged DFz2C fragment failed, because this did not rescue the reduced bouton number in dfz2 mutants (Mathew et al. 2005; but see Mosca and Schwarz 2010). These findings led the investigators to suggest that an interaction between the full-length DFz2 and Wg was likely required to make DFz2C competent for nuclear import and signaling.

The journey of DFz2 from the plasma membrane to the nucleus involved a direct interaction between the carboxyl terminus of DFz2C, which contains a PDZ binding sequence, and the 7 PDZ domain protein dGRIP (glutamate receptor interacting protein) (Ataman et al. 2006). In the vertebrate nervous system, GRIP mediates the synaptic trafficking and scaffolding of several proteins, including AMPA (Dong et al. 1997) and EphB (Hoogenraad et al. 2005) receptors. In Drosophila, mutations in dgrip or selective postsynaptic reduction in dGRIP levels led to phenotypes that highly resembled mutations in Wnt signaling (Ataman et al. 2006). These included a drastically reduced number of nuclear DFz2C foci, a reduced number of synaptic boutons, and an increase in the number of ghost boutons. Analysis of the trafficking of DFz2 receptors by using an antibody against the extracellular amino terminus of DFz2, which binds to the receptor in vivo, allowed the researchers to follow receptor trafficking through pulse-chase experiments. In wild-type larvae, DFz2 receptors localized at the postsynaptic region traffic toward the nucleus over time. However, when dGRIP levels were reduced, DFz2 accumulated at the postsynaptic region and failed to be transported toward the nucleus (Ataman et al. 2006). These observations suggested that dGRIP was required to transport DFz2C from the muscle plasma membrane to the nucleus.

Further insight into the FNI pathway was provided by Mosca and Schwarz (2010), who showed that the nuclear import of DFz2C requires the nucleoporin-associated proteins Importin-β11 (Imp-β11) and Importin-α2 (Imp-α2). In the absence of either of these two importins, synapses displayed phenotypes that resembled the alterations found at NMJs of wg, dfz2, and dgrip mutants, that is, a reduced number of synaptic boutons, an excessive number of ghost boutons, and a reduced thickness of the subsynaptic reticulum (SSR) (Mosca and Schwarz 2010). These results indicate that importins and therefore nuclear entry of DFz2C fragments through the canonical nuclear import pathway are key players in the FNI Wnt pathway.

More recently, Speese et al. (2012) have begun to unravel the function of DFz2C foci within the nucleus. The foci are composed of large ribonucleoprotein (RNP) particles bearing synaptic protein transcripts. These granules localize to the space between the inner and outer nuclear membrane and require Lamin C, a nuclear protein linked to muscular dystrophies (Worman and Bonne 2007). lamC mutations mimicked NMJ phenotypes of mutations interfering with the FNI pathway. Notably, it was shown that the granules exit the nucleus via a budding mechanism akin to the nuclear egress of Herpes-type viruses (Mettenleiter et al. 2006), a process previously thought to be exclusive for these viruses. Like HSV nuclear egress, this process required PKC, which is known to disrupt the lamin through phosphorylation. It was suggested that nuclear envelope budding is an endogenous nuclear export mechanism for large
RNPs (Speese et al. 2012). Interestingly, the nuclear DFz2C foci contained transcripts encoding synaptic proteins, such as Par6 and CaMKII, and some of the transcripts were also found at the NMJ. These observations suggested that the large RNPs were required for local translation of the postsynaptic apparatus during new synaptic bouton formation (Speese et al. 2012). Although the exact function of DFz2C fragments within the large RNP granules is not known, the authors speculate that the PDZ binding domain of DFz2C may serve as a zipcode to properly target large RNPs to synaptic sites. Several PDZ-containing scaffolding proteins are localized at postsynaptic sites and are required for synaptic development at the NMJ (Lahey et al. 1994; Thomas et al. 1997; Mathew et al. 2002). Furthermore, proteins containing PDZ-binding motifs are known to be targeted to postsynaptic sites (Zito et al. 1999). Although the generality of the FNI signaling pathway for other systems and tissues is presently unknown, nuclear DFz2C foci are also observed in Schneider-2 cells and at Drosophila larval salivary glands (Speese et al. 2012), where the nuclear envelope granules were first observed along with epithelial cells of the midgut (Gay 1956; Hochstrasser and Sedat 1987). In addition, these foci are observed in peripheral glial cells in the larva (SD Speese and V Budnik, unpubl.). Notably, granules accumulating between the outer and inner nuclear membrane resembling the DFz2-encoded large RNP granules have been observed in diverse contexts, from plants (Dickinson 1971) to mammals, where they are particularly prevalent at early embryonic stages (Hadek and Swift 1962; Szollosi 1965). Indeed, that such granules might reflect an alternate mode of nucleocytoplasmic export has been previously proposed (Gay 1956; Szollosi and Szollosi 1988), but it had not been experimentally validated until now. However, whether these granules are also associated with Frizzled receptor fragments is currently unknown.

As discussed above, Wg also turns on a presynaptic signaling pathway with resemblance to the canonical Wnt signaling pathway (Ataman et al. 2008; Miech et al. 2008). Several components of the canonical Wnt signaling pathway can be detected in presynaptic boutons (Packard et al. 2002; Miech et al. 2008), including Dishevelled (Dvl), Gsk3β/Shaggy, DFz2 and its coreceptor Arrow/Lrp5/6, but not Armadillo/β-catenin (Miech et al. 2008). Mutations in arrow/Lrp5/6, a receptor thought to be involved in canonical Wnt signaling (He et al. 2004), showed phenotypes similar to those observed in wg mutants such as reduced bouton number, increase in large (>5 μm), irregularly shaped boutons, and decrease in the number of boutons with presynaptic Futsch loops (Miech et al. 2008). However, the presence of ghost boutons was not examined in those studies. All of these phenotypes were rescued by the presynaptic expression of Arrow. Notably, expressing an arrow transgene in muscles of arr mutants also rescued the reduced bouton number while not rescuing the presynaptic microtubule phenotypes. In addition, Arrow overexpression in neurons resulted in an increase in the number of bouton and satellite boutons, similarly to what was shown for presynaptic Wg overexpression (Packard et al. 2002; Miech et al. 2008). Subsequent experiments indicated that Wg binding to presynaptic Arrow leads to the activation of a divergent canonical pathway. Indeed, overexpression of a dominant negative Dvl in neurons, but not in muscle, led to the prototypical wg mutant phenotypes, whereas overexpression of dominant negative Gsk3β/Sgg, as expected, resulted in the opposite phenotype, mimicking Wg overexpression (Franco et al. 2004; Miech et al. 2008). No phenotypic changes were observed upon dominant negative TCF/pangolin or Armadillo/β-catenin overexpression in neurons (Miech et al. 2008), suggesting that the dominant negative Dvl and Gsk3β overexpression phenotypes did not result from interfering with canonical Wnt signaling. Thus, by activating alternative signaling pathways, a single Wnt simultaneously coordinates presynaptic and postsynaptic development.

An important avenue of inquiry examined the involvement of Wnt signaling during synaptic plasticity. Ataman et al. (2008) showed that larval NMJs respond to repeated or patterned stimulation administered via high-potassium depolarization, motor nerve stimulation,
or blue light stimulation of channelrhodopsin-2 (Nagel et al. 2002) expressing motor neurons, by extending new “synaptopods,” dynamic filopodia-like structures that are observed, albeit at lower frequency, in unstimulated animals (Ataman et al. 2008). In addition, NMJs respond to these stimuli by forming new ghost boutons. Analysis by time-lapse microscopy revealed that a subset of these ghost boutons could subsequently mature by acquiring active-zone and postsynaptic proteins, suggesting that ghost boutons are an undifferentiated state of mature boutons. In addition to the above morphological changes, the frequency of spontaneous miniature synaptic potentials (mEJPs) was potentiated by these stimulation paradigms. The above changes required at least five stimulation bouts separated by a resting period, and intact transcriptional and translational machinery. Most importantly, Wg signaling was necessary for these changes downstream from motor neuron stimulation, because removing just one copy of the wg gene was sufficient to eliminate the structural and functional changes elicited by patterned stimulation. Furthermore, neuronal overexpression of Wg, which increased Wg secretion by synaptic boutons, reduced the number of cycles required to initiate the structural and functional changes (Ataman et al. 2008). Thus, Wnts are not readily diffusible in the extracellular space. Several mechanisms have been documented to be involved in the diffusion of Wnt signals, including the association of Wnt proteins with lipoprotein particles (Panakova et al. 2005), the transmission of argosomes (Greco et al. 2001), cytonemes (Ramirez-Weber and Kornberg 1999), and very recently, Secreted Wg-interacting molecule (Swim), a member of the Lipocalin family that is suggested to maintain Wg in a soluble form for long-range signaling (Mulligan et al. 2012). The studies at the NMJ suggest yet another mechanism, the shuttling of Wnts associated with Evi, which is released in vesicles (Fig. 4A). Most recently, analysis of the Evi vesicles showed that they indeed correspond to exosomes (K Koles and V Budnik, unpubl.). Exosomes are released by the fusion of multivesicular bodies with the plasma membrane, thus releasing their vesicular content. Our initial studies indicate that Evi-exosome release from presynaptic terminals requires Rab11, Myosin 5, and Syntaxin-1A, suggesting that exosome release is an integral component of the synaptic bouton release machinery.

Most recently, Wg signaling has been implicated in negatively regulating homeostatic compensation (Marie et al. 2010). Neurotransmitter release and postsynaptic muscle depolarization are tightly coupled processes, and their...
Figure 4. Evi-mediated synaptic release of Drosophila Wg and regulation of ACR-16 receptor abundance in postsynaptic muscle membranes by CWN-2 in C. elegans. (A) Schematic representation of Evi-mediated Wg release at Drosophila larval NMJs. Wg carried by Evi-exosomes is sorted into multivesicular bodies in synaptic terminals that fuse with the synaptic membrane, releasing their content to the cisternae of underlying subsynaptic reticulum. Potential fates of postsynaptic exosomes are shown, such as fusion with the postsynaptic membrane and subsequent endocytosis. The endocytosed DFz2/Wg/Evi-containing vesicles can be either sorted and targeted to lysosomes or, via DFz2 binding to Grip, transported near the muscle nucleus (also see Fig. 1E). (B) In C. elegans, motor neurons release CWN-2, which binds to the heteromeric CAM-1/Lin-17 receptor and activates Dishevelled-1. Activated Dvl-1, in turn, leads to the mobilization and membrane delivery of subsynaptic ACR-16 receptor pools.
coordination is required to maintain synaptic efficacy. The process of synaptic homeostasis involves the presynaptic (increase in neurotransmitter release) or postsynaptic (abundance of glutamate receptors) compensatory adjustment to changes in postsynaptic or presynaptic parameters, resulting in the restoration of wild-type synaptic efficacy. Synaptic homeostasis is thought to occur in two phases, an induction phase that is transcription and protein synthesis independent, and a maintenance phase that requires transcription and the presynaptic BMP-signaling pathway (Goold and Davis 2007). Marie et al. (2010) have recently shown that Goosberry (Gsb) is required for the maintenance phase of this homeostatic compensation process. Using the GluRIIA^{Asp16-null} mutant background (Petersen et al. 1997), which leads to a compensatory increase in presynaptic neurotransmitter release due to a decrease in the amplitude of spontaneous mEPSPs, the investigators discovered that this compensation was significantly reduced in heterozygous gsb01155 mutants (Marie et al. 2010). Interestingly, Gsb also functions early during embryonic development to inhibit Wg signaling in the Wg-secreting cell (Duman-Scheel et al. 1997). Indeed, when the investigators examined the possibility of whether Gsb may function in homeostatic compensation in motor neurons by inhibiting presynaptic Wg signaling, they found that replacing one copy of the wild-type wg gene with a temperature-sensitive wgts allele in the gsb01155/1 background restored homeostatic compensation (Marie et al. 2010). Therefore, in addition to its other structural and functional roles at the NMJ, these results raise the interesting possibility that Wg, along with BMP signaling, regulates synaptic homeostasis (Marie et al. 2010).

In addition to Wingless, two additional Wnt family members—Wnt5, its postsynaptic receptor Derailed (Drl) (Liebl et al. 2008), and Wnt2 (Liebl et al. 2010)—were also shown to regulate NMJ development. Mutations in Wnt5 led to structural as well as functional defects at the NMJ. Bouton numbers and NMJ areas were significantly decreased in wnt5 mutants, and the number and density of active zones, as assessed by Bruchpilot (Brp) immunolabeling, was also significantly lower. Interestingly, only neuronal Wnt5 expression was able to rescue bouton numbers to wild type, but either neuronal or muscle-specific rescue restored the number and density of active zones, suggesting a potential retrograde function for Wnt5 in regulating active-zone formation. Electrophysiological examination of wnt5 mutant synapses revealed a 40% decrease in evoked excitatory junctional currents (EJCs) and a similar decrease in the frequency of mEJCs (Liebl et al. 2008). Because the amplitude of mEJCs (which reflects postsynaptic glutamate receptor function) was not affected in wnt5 mutants, these data point toward the presynaptic function of Wnt5. Earlier studies have shown that Wnt5 can bind Derailed (Drl) (Yoshikawa et al. 2003). Indeed, drl mutant NMJs manifest similar structural defects to wnt5 mutants, such as a significant decrease in bouton number (Liebl et al. 2008). This phenotype could be rescued by postsynaptic expression of Drl. However, there were also differences in the phenotypes of drl and wnt5 mutant larvae. drl mutations, unlike wnt5 mutations, did not result in a decrease in active-zone density, nor in functional changes in synaptic transmission, suggesting that some aspects of Wnt5 function are Drl independent. It is interesting to note, however, that presynaptic expression of Wnt5 in the wild-type background led to a significant increase in bouton number, and that this increase was dependent on Drl, because in the drl mutant background no change in bouton number was observed upon neuronal Wnt5 overexpression. In summary, Wnt5 is poised to play a complex structural and functional role at the larval NMJ, in part by signaling via the Drl receptor, while also showing Drl-independent effects.

Subsequently, the same researchers have shown that Drosophila Wnt2, the ortholog of mammalian Wnt7b, is required for proper NMJ development and for the maintenance of proper Brp density and intensity at synapses (Liebl et al. 2010). In the absence of Wnt2, presynaptic arbors branch more frequently, although the number of boutons is unchanged, indicating that unlike other Wnts, Wnt2 negatively regulates
NMJ branching. The total number of Brp puncta did not change in Wnt2 mutants. However, the density of Brp puncta decreased significantly, as expected from the increased NMJ arbor size in these mutants. A combination of rescue experiments with Wnt2 overexpression and tissue-specific knockdown of Wnt2 suggests that it functions in the muscle, most likely in a retrograde manner to negatively regulate presynaptic growth. In agreement with these observations, the amplitude of evoked excitatory junctional potentials was also reduced, whereas the frequency and amplitude of spontaneous vesicle release were unaffected. It is as yet unclear what receptor Wnt2 signals through at the NMJ. Previous studies implicated Fz, DFz2, and Fz3 as potential receptors for Wnt2 (Wu and Nusse 2002), but genetic studies did not affirm these possibilities at the larval NMJ (Liebl et al. 2010).

THE ROLE OF WNTs AT C. elegans NMJs

The muscles of C. elegans are innervated by excitatory cholinergic or inhibitory GABAergic neurons. Unlike Drosophila or vertebrate NMJs, muscles extend “muscle arms” toward the nerve cord to form en passant synapses (Dixon and Roy 2005). Two types of acetylcholine receptors (AChRs) have been identified in C. elegans, the N-type, likely a homomer of ACR-16 subunits (Francis et al. 2005) and activated by nicotine, and the L-type channels, which are heteropentamers and sensitive to levamisole. Two distinct mechanisms have evolved to localize these two types of receptors. N-type AChRs require CAM-1, a MuSK-related orphan receptor tyrosine kinase (Francis et al. 2005), and N-type AChRs require an extracellular scaffold in the synaptic cleft, composed of secreted Lev-9 (Gendrel et al. 2009), the extracellular domain of Lev-10 (Gally et al. 2004), and a small, secreted single immunoglobulin containing OIG-4 (Rapti et al. 2011).

Although in vertebrates and Drosophila, initial discoveries revealed pro-synaptogenic functions for Wnts (even though subsequent experiments identified negatively acting Wnt signals as well), the first C. elegans Wnt signaling pathway involved in regulating synapse formation served to restrict the position of synapses on axons. For a while these findings led to the notion that Wnts were inhibitory in C. elegans; however, some of the latest discoveries seem to dispel this notion (see below).

The first Wnt discovered to function in C. elegans neuromuscular junction formation was Lin-44 (Klassen and Shen 2007). Lin-44/Wnt is secreted from four hypodermal cells during embryonic and larval development. In lin-44/wnt mutants, ectopic synapses are formed, indicating that lin-44/wnt inhibits synapse formation. Lin-44/Wnt functions in DA9 motor neurons, where it recruits the Wnt receptor Lin-17/Frizzled to a subdomain of the DA9 axon via binding to the cysteine-rich domain (CRD) of Lin-17/Frizzled. Lin-44/Wnt cooperates with another C. elegans Wnt, EGL-20/Wnt, and by binding to and signaling through Lin-17/Frizzled, it prevents synapse formation at this region of the DA9 axon (Klassen and Shen 2007). Interestingly, except for Dvl, none of the known canonical, planar cell polarity, or calcium-dependent Wnt signaling pathway effectors were found to be involved in this process, suggesting that novel effectors mediate the local inhibition of synaptogenesis on DA9 motor neuron axons.

An additional aspect of Wnt signaling regulation at the C. elegans NMJ was revealed by the findings of Babu et al. (2011). In an RNAi screen for cell surface Ig-domain-containing (IgSF) proteins that are involved in modulating sensitivity to aldicarb (a cholinesterase inhibitor), Babu and colleagues identified RIG-3, a GPI-anchored IgSF-domain protein. Decreased levels of this protein, which is normally expressed in cholinergic neurons, led to heightened sensitivity to aldicarb. When the investigators examined the underlying mechanisms for this increased sensitivity, they found that in the absence of RIG-3, the delivery of ACR-16 (the homomer-forming subunit of nicotinic AChRs) from muscle mobile stores to the muscle membrane was significantly increased following aldicarb treatment. Furthermore, this increase was the result of increased levels of
CAM-1, which is an unconventional Wnt receptor of the Ror-type receptor tyrosine kinase family (Green et al. 2008) involved in ACR-16 delivery to synaptic sites (Francis et al. 2005). Interestingly, CAM-1 binds Wnt ligands, and mutations that reduce secreted Wnt levels, such as mig-14, led to the elimination of aldicarb-induced hypersensitivity in rig-3, mig-14 double mutants, indicating that CAM-1 was mediating this phenotype in a Wnt-dependent manner. These data show that RIG-3, a presynaptic molecule, negatively regulates CAM-1-mediated Wnt signaling at ACR-16-containing synapses following aldicarb treatment and thus may play a role in stabilizing synapses and synaptic circuits.

As mentioned above, the first discovery of *C. elegans* Wnt signaling in NMJ formation revealed the negative regulation of synapse formation, suggesting that Wnts may be inhibitory in *C. elegans*. However, a new study from Jensen et al. (2012) seems to dispel this notion. In a series of elegant and groundbreaking studies, the researchers showed that neuronally released CWN-2 (but not CWN-1 or EGL-20), by binding to the CAM-1/Lin-17 heteromeric receptor (but not those containing MIG-1 or CFZ-2), resulted in the increased postsynaptic muscle membrane delivery of ARC-16 (Fig. 4B). This signaling required ongoing CWN-2 release, because transient CWN-2 pulses led to only transient increases in postsynaptic ACR-16. For the first time in *C. elegans*, the investigators also showed that this Wnt signaling pathway was activity dependent. Channelrhodopsin2 stimulation of neurons led to an increased translocation of ACR-16 to the muscle membrane and a corresponding increase in ACR-16-mediated current, and these effects required the neuronal expression of CWN-2. In summary, an ongoing, activity-dependent CWN-2 release and signaling through CAM-1/Lin-17 receptors is required for activity-dependent synaptic plasticity and the maintenance of proper synaptic transmission in adult *C. elegans* neuromuscular junctions. These observations also point out the similarities of Wnt function at the NMJ across species. The activity-dependent release and anterograde function of CWN-2 are strikingly similar to what has been described for Wg at *Drosophila* larval NMJs (Ataman et al. 2008). In a broader context, these new results highlight that Wnts do not only function during development, but are also required throughout the life span of organisms for proper synaptic functioning.

CONCLUDING REMARKS

In the last decade, we have witnessed a large number of new discoveries regarding the function of Wnt signaling at neuromuscular synapse formation and function. From these findings, it is apparent that Wnts are intricately involved in almost every aspect of NMJ development, maintenance, and function, including activity-dependent plasticity. They can play both essential and modulatory roles and promote or inhibit synapse formation. As surmised by Speese and Budnik (2007) and van Amerongen and Nusse (2009), the likely scenario of Wnt action is that of a parallel, combinatorial network of signals emanating from the neuron, muscle, and surrounding non-neuronal cells that work in anterograde, retrograde, and autocrine manner to effect and coordinate the developmental and synaptic plasticity-induced processes.

One of the key challenges of the Wnt signaling field will be to reconcile data from in vivo and in vitro experiments. For example, Dvl1 was identified via its MuSK interaction and was shown to be required for Agrin-induced AChR clustering, yet the *Dvl1* mutant mouse phenotypes are surprisingly mild compared with the Dvl1 knockout in C2C12 cell cultures (Luo et al. 2002). Furthermore, in muscle-specific β-catenin−/− knockout muscle cultures, Agrin-mediated AChR clustering was not affected (Li et al. 2008), unlike in the C2C12 cultured cell line, where β-catenin levels were reduced via shRNA treatment (Zhang et al. 2007; Wang et al. 2008). These data highlight the difficulties involved in translating the cell culture findings to in vivo situations in a living/developing organism. These data illustrate that multiple pathways may converge at the level of β-catenin and that the preceding developmental history of the cell could lead to diverse compensatory
mechanisms or adaptations that may lead to conflicting results if these factors are not controlled for.

Among the exciting future challenges will be the identification of retrograde signaling mechanisms that were revealed through examining the muscle specific β-catenin−/− NMJs (Li et al. 2008). Likewise, because many aspects of Wnt signaling are conserved across species, it will be exciting to test whether the continuous, ongoing requirement for C. elegans Wnt, CWN-2, in synaptic plasticity (Jensen et al. 2012) is also at play in other model organisms, such as the fly, zebrafish, or other vertebrates. Furthermore, the contribution to NMJ development and function of terminal Schwann cells in vertebrates (Feng and Ko 2008; Sugiura and Lin 2011) is poised to lead to novel insights.

In addition, future research is likely to lead to an integrated model of multiple Wnt signaling pathways (van Amerongen and Nusse 2009) that are acting simultaneously and synergistically from multiple tissue sources (nerve, muscle, somites, glia) to sculpt the synapse. Current efforts naturally focus on delineating single, rather than multiple, linear signaling pathway effects (e.g., as described above, Wnt1, Wnt5, and Wnt2 are all involved in Drosophila larval NMJ development); however, their integration into a dynamic, spatio-temporal map of signaling events will transform our understanding of NMJ development and function. Given how far the field has come in the last few decades and the continuous technical advances allowing more nuanced experiments to be performed, future developments in the field of Wnt signaling at the neuromuscular synapse are bound to be exciting.

ACKNOWLEDGMENTS

We thank Dr. Cassandra Brewer for critical review of the manuscript. V.B. is supported by NIH grants RO1 MH070000 and R01 NS063228.

REFERENCES


Cite this article as Cold Spring Harb Perspect Biol 2012;4:a008045
Wnt Signaling in NMJ Development


Jing L, Lefebvre JL, Gordon LR, Granato M. 2009. Wnt signals organize synaptic prepattern and axon guidance...


Valenzuela DM, Stitt TN, DiStefano PS, Rojas E, Mattsson K, Compton DL, Nunez L, Park JS, Stark JL, Gies DR,


