Aggregation of Sodium Channels Induced by a Postnatally Upregulated Isoform of Agrin

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Agrin is involved in signaling the formation of high concentrations of acetylcholine receptors (AChRs) at the neuromuscular junction (NMJ). There are multiple isoforms of agrin attributable to alternative splicing, and these isoforms are differentially expressed during development and between tissues. The ability to cluster AChRs varies among the agrin isoforms. Sodium channels (NaChs) are also concentrated at the NMJ. We have tested various agrin isoforms for their ability to induce formation of clusters of NaChs. We grew cocultures of dissociated adult rat muscle fibers with chinese hamster ovary (CHO) cells that had been transfected with different isoforms of agrin. Using immunocytochemical techniques, we determined that after 1 d in culture, CHO cells synthesizing the neuronally expressed isoform with an eight amino acid insert (Agrin8) were able to form NaCh clusters at sites of contact between the CHO cell and muscle cell. Clusters of NaChs could be formed anywhere along a muscle fiber, but more clusters were detected close to the endplate where the endogenous level of NaChs was higher. None of the other neuronal-specific agrin isoforms was able to cluster NaChs. Because Agrin8 is the only agrin isoform that is upregulated at birth when NaChs begin to cluster at the NMJ, we conclude that Agrin8 expression by motor neurons is a signal for NaCh clustering at the NMJ during normal development.

Key words: sodium channel; agrin; neuromuscular junction; synapse development; ion channel aggregation

Synapses are highly organized structures that require complex molecular interactions to be generated and maintained. Many molecules are highly enriched at the synapse [e.g., at the neuromuscular junction (NMJ); sodium channels (NaChs), acetylcholine receptors (AChRs), and acetylcholine esterase] (Froehner, 1991). Such specialization is required for proper interneuronal and neuromuscular signaling. Perhaps the best understood molecular pathway of synaptic organization is the involvement of agrin in the clustering of AChRs at the NMJ (for review, see McMahan, 1990; Nastuk and Fallon, 1993; Apel and Merlie, 1995).

Agrin is an extracellular matrix protein that is produced by neurons, muscle, and other tissues (McMahan et al., 1992). It is present in multiple isoforms as a result of alternative splicing at three sites designated x, y, and z in rat (Rupp et al., 1991, 1992). Inserts at the x and y sites have little or no effect on AChR distribution. There are three neuronal-specific isoforms of agrin (Ruegg et al., 1992; Hoch et al., 1993), which show varying degrees of activity in clustering AChRs (Ferns et al., 1992, 1993; Gesemann et al., 1995); these have 8 and/or 11 amino acid inserts at the z site (referred to here as Agrin8, Agrin11, or Agrin19). The isoforms are differentially expressed by different tissues and at different developmental stages (Hoch et al., 1993). The isoform with no insert (Agrin0) is widely expressed in the nervous system, muscle, and other non-neuronal cells (Hoch et al., 1993; Thomas et al., 1993).

Compared with the clustering of AChRs, little is known about the mechanism of NaCh accumulation at the NMJ. The density of NaChs is ~10-fold higher at the NMJ than in nonjunctional regions, and this increased density appears ~1 week after birth (Caldwell and Milton, 1988; Lupa et al., 1993). Lupa and Caldwell (1994) showed that some component of the basal lamina or Schwann cells was sufficient to induce NaCh clustering in the absence of innervation in regenerating muscle, as has been shown for AChRs in regenerating frog muscle (Burden et al., 1979). However, bath application of Torpedo agrin to cultures of dissociated adult rat muscle fibers did not induce clustering of NaChs (Lupa and Caldwell, 1991). This result suggests that either agrin alone is not sufficient to signal NaCh clustering or that a form of agrin other than that present in Torpedo extracts is required.

The goal of this study was to determine whether any of the rat neuronally expressed agrin isoforms are able to initiate NaCh clustering. We generated cocultures of adult rat muscle fibers and chinese hamster ovary (CHO) cells that had been transfected with cDNAs for various isoforms of rat agrin. Immunofluorescence techniques were used to assay for the presence of agrin-induced NaCh clusters. Only CHO cells expressing the Agrin8 construct were able to induce NaCh clustering. This suggests that the developmentally regulated neuronal expression of Agrin8 is a signal for the induction of NaCh clustering at the NMJ.

MATERIALS AND METHODS

Muscle fiber dissociation. Muscle fibers from the rat flexor digitorum brevis (FDB) were prepared as described previously (Bekoff and Betz, 1977; Lupa and Caldwell, 1991). Briefly, FDB muscles were dissected from the hind feet of adult rats. Individual intact muscle fibers were
dissociated from the muscle by first incubating the muscle in 2 mg/ml dispase and 10 mM aprotinin for 1 hr at room temperature. The muscle was then placed in 2 ml of sterile, rat saline-containing (in mM): 129 NaCl, 5 KCl, 2 CaCl₂, 1 MgCl₂, 11 glucose, 10 Na₂-PIPES, pH 7.3, and triturated with fire-polished Pasteur pipettes of decreasing size until single, intact fibers were obtained. The suspension was allowed to stand for 5 min for the muscle fibers to settle. Supernatant (1.5 ml) was then removed.

CHO cells. CHO cells were obtained from Drs. J. Campanelli and R. Scheller (Stanford University). In addition to wild-type CHO cells, cell lines transected with full-length cDNAs for four rat agrin isoforms (Agrin0, Agrin8, Agrin11, and Agrin19) were used. These cell lines have been characterized previously (Campanelli et al., 1991; Ferns et al., 1992). The agrin constructs included inserts of 12 amino acids at the N-terminus and 4 amino acids at the C-terminus. In addition to the cDNA for agrin, the plasmid transfected into CHO cells contained an insert coding for neomycin resistance (cells not containing the plasmid are killed by the neomycin analog, geneticin). Before coculturing with the FDB fibers, the transfected CHO cells were maintained in DMEM containing 200 µg/ml geneticin (Life Technologies), 10% FBS (Gemini Bioproducts, Calabasas, CA) and 1% pen/strep (Life Technologies) at 37°C and 5% CO₂. After three passages, the cells were discarded and fresh cultures were begun. The frozen frozen stocks (prepared from the original frozen stock provided by Drs. Campanelli and Scheller). This reduced the potential for the loss of the agrin construct.

Substrate for cultures. Glass coverslips were coated with Matrigel (Collaborative Biomedical Products, Bedford, MA) immediately before the addition of cells. A Corning (Corning, NY) coverslip (22 mm square, no. 1), dipped in ethanol to sterilize and allowed to dry, was placed into each well of a six-well culture plate. Matrigel (150 µl; 0.5 mg/ml in 0.1 M PBS) was then spread on the coverslips. Coverslips were allowed to stand for 1 hr at room temperature, and the excess Matrigel was then removed. Cells were added to the coverslips within 15 min.

Cocultures of FDB fibers and CHO cells. When the CHO cell cultures had reached confluence, they were rinsed with Dulbecco’s PBS and trypsinized for 3 min (0.25% trypsin, Ca²⁺- and Mg²⁺-free) (Life Technologies). DMEM containing 10% FBS was added. The cells were pelleted and resuspended in rat saline. Approximately 10⁵ cells in 50 µl of saline were added to the muscle fibers from one FDB muscle. The muscle fibers and CHO cells were mixed with gentle trituration and divided between six coated coverslips (~90 µl per coverslip). The coverslips were placed in a 37°C, 5% CO₂ incubator for 3–4 hr before culture medium (DMEM with 2.5% FBS, 5% horse serum (Life Technologies) and 5% pen/strep) was added. Cultures were then returned to the incubator for 1–2 days prior to processing for immunocytochemistry.

Antibodies and biotinylation. NaChs were detected with a rabbit polyclonal antibody [provided by Dr. S. R. Levinson, University of Colorado Health Sciences Center (UCHSC) and available from Upstate Biotechnology, Lake Placid, NY], which recognizes an intracellular epitope that is common to many NaCh subtypes. Characterization and properties of this antibody are described in Dugandzija-Novakovic et al. (1995). This was followed by a Cy3-conjugated donkey anti-rabbit IgG secondary antibody (Jackson ImmunoResearch Laboratories, West Grove, PA).

Agrin expression in the CHO cell cultures was detected using a mouse monoclonal antibody that recognizes a region near the second epidermal growth factor (EGF)-like domain and binds to all agrin isoforms (Hoch et al., 1994) (catalog #AGR-530, StressGen, Victoria, BC, Canada). In preliminary experiments, we used a mouse monoclonal that recognizes agrin with any insert at the z position, with similar results for Agrin8 (catalog #AGR-520) (StressGen). This was followed by a biotin-Sp-conjugated donkey anti-mouse IgG secondary antibody and finally fluorescein-conjugated streptavidin (both from Jackson). Detection of wild-type CHO cells was achieved by biotinylating them before trypsin treatment. Cultures were rinsed with Dulbecco’s PBS and then reacted for 1 hr at room temperature with 1 mg/ml biotinyl-ε-aminocaproic acid N-hydroxysuccinimide ester (biotin-LC-NHS) (Pierce, Rockford, IL) in Dulbecco’s PBS. The reaction was quenched with a 10-fold dilution in DMEM containing 10% FBS. Fluorescein-conjugated streptavidin was used for final visualization.

ACHs were detected using biotinylated a-bungarotoxin prepared as follows: biotin-LC-NHS (0.5 mg) was dissolved in dimethylformamide (9 µl). Biotin-LC-NHS (1 µl), a-bungarotoxin (0.1 mg) (Sigma, St. Louis, MO), and NaHCO₃ (10 µl, 0.1 M, pH 8.3) were mixed and incubated at room temperature for 1 hr. The reaction was stopped by adding 1 µl glycine (0.1 M). Fluorescein-conjugated streptavidin was used to visualize the biotin-labeled bungarotoxin.

Immunocytochemistry. After 1–2 d in culture, coverslips were removed from the incubator, rinsed three times (5 min each) in PBS, and fixed in 4% paraformaldehyde for 10 min. Three PBS rinses were used between each subsequent treatment. Cells were permeabilized with 0.5% Triton X-100 (Sigma) in PBS. Non-specific binding was blocked using 15% goat serum (Sigma) in PBS for 30 min. Primary antibodies and/or α-bungarotoxin were then applied overnight at room temperature diluted in 15% goat serum as follows: anti-NaCh, 1:100; anti-Agrin, 1:1000; a-bungarotoxin, 1:10⁵. Secondary antibodies were also diluted in 15% goat serum and were applied for 1 hr at the following dilutions: Cy3-conjugated donkey anti-rabbit, 1:600; biotin-conjugated donkey anti-mouse, 1:200. Finally, fluorescein-conjugated streptavidin was added for 1 hr (1:600 in 15% goat serum). Coverslips were mounted in photoobleachable protectivetm media (Vectorshield, Vector Laboratories, Burlingame, CA) and sealed with fingernail polish.

Slides were observed on a Nikon Optiphot-2 fluorescence microscope. Digital images were collected using a Zeiss Axioskop fluorescence microscope (40X Planapoch objective, 1.3 numerical aperture) equipped with a cooled CCD camera (MCD 1000, SpectraSource Instruments, Westlake, CA). Fluorescence resulting from emission outside the plane of focus was usually removed using a no-neighbour deconvolution ( software written by Drs. C. Monks and A. Kupfer, National Jewish Center and UCHSC). Final image processing was performed using the Corel Graphics software package (Ottawa, Canada). The images were pseudocolored for illustration (see Fig. 3) with the colors reversed (NaCh, green; AChR, red). Linearity was maintained in all transformations.

Quantification of clustering. For contacts between CHO cells and muscle fibers to be scored for agrin-induced clusters of NaChs, several criteria had to be met. First, a CHO cell had to be clearly in contact with a healthy muscle fiber, and the muscle fiber had to be isolated (bundles of partially dissociated fibers were ignored). Second, the CHO cell had to express agrin (or be biotinylated in the case of wild-type cells). Third, the endogenous NaCh signal of the CHO cell (which was primarily intracellular) had to be separable from the muscle fiber membrane; this meant that it had to be possible by changing the focal plane or by having the cells side-by-side to attribute the signal to one cell or the other. For a contact to be scored as having agrin-induced clustering, the muscle fiber membrane adjacent to the CHO cell had to show greater fluorescence than the area adjacent to the contact. The signal at the majority of these clusters was scored by visual inspection; for six clusters the increased signal over background was quantitated by digitizing the image and comparing the intensity at the area of contact with adjacent regions of the cell using SigmaScan software (Jandel Scientific, San Rafael, CA), and the relative increase in intensity varied from two- to fivefold.

RESULTS

NaCh labeling of muscle fibers

In selecting our assay system, we wished to use a tissue that would maximize levels of endogenous NaChs available for clustering. We chose to culture adult muscle fibers dissociated from the rat FDB rather than embryonic myotubes, because adult muscle fibers have an extrajunctional sodium current density of ~10 nA/cm² (Caldwell and Milton, 1988; Lupa et al., 1993), which is ~10 times that in embryonic myotubes (R. Milton and J. Caldwell, unpublished observations). Unlike AChRs, NaCh density changes very little after denervation (Lupa et al., 1995); therefore, we did not denervate the muscles before dissociation.

In previous studies, NaCh density and distribution was determined primarily with the loose patch voltage clamp technique. In the experiments reported here, we expected there to be changes in channel density at the sites of contact between the agrin-expressing CHO cells and the muscle fibers, a region difficult or inaccessible for patch clamping. Consequently, we chose to use an anti-NaCh antibody to determine NaCh distribution. It was possible to visualize NaChs at the endplate of dissociated muscle fibers as well as in the perijunctional membrane of the muscle fiber (Fig. 1). Figure 1A1 shows staining for
NaChs on a muscle fiber that was fixed 4 hr after dissociation. NaCh labeling was highest at the endplate (left edge of the fiber), which was identified by AChR labeling with α-bungarotoxin (Fig. 1A2), and labeling of the perijunctional membrane on both edges of the fiber was also evident. The highest NaCh signal on the fiber always coincided with the AChR labeling; this allowed us to identify the endplate by NaCh labeling alone. Preincubation of the anti-NaCh antibody with the peptide to which it was generated (0.1 mg/ml) blocked labeling of the endplate region (Fig. 1B, Anti-NaCh antibody control (B1–B3 are identical fields of view)). Before addition to the muscle fibers, the antibody was incubated for 8 hr at 4°C with the peptide against which it was generated. B1, Blocked-antibody labeling. Preincubation abolished labeling of NaChs in muscle. B2, AChR labeling with α-bungarotoxin. B3, Nomarski image. Scale bar, 10 μm.

Figure 1. A, NaCh and AChR expression on a dissociated FDB muscle fiber (A1–A3 are identical fields of view). The fiber was fixed 4 hr after dissociation. A1, NaCh labeling. Notice the bright staining of the endplate on the left edge of the fiber and the perijunctional staining on both sides of the muscle fiber. A myonucleus forms a bulge on the right side of the fiber. A2, AChRs labeled with α-bungarotoxin. Unlike the NaChs in A1, the AChRs are restricted to the endplate. A3, Nomarski image. B, Anti-NaCh antibody control (B1–B3 are identical fields of view). Before addition to the muscle fibers, the antibody was incubated for 8 hr at 4°C with the peptide against which it was generated. B1, Blocked-antibody labeling. Preincubation abolished labeling of NaChs in muscle. B2, AChR labeling with α-bungarotoxin. B3, Nomarski image. Scale bar, 10 μm.

NaCh labeling fell to background levels within 20–80 μm of the endplate. This corresponds to a decrease in sodium current density from 100 mA/cm² to 20 mA/cm², as measured with the loose-patch technique (Caldwell and Milton, 1988; Lupa et al., 1993). If we assume that 1 mA/cm² corresponds to 20 NaChs/μm² (see Caldwell et al., 1986), this comparison implies that we can detect NaChs with the antibody at a density of ~400 NaChs/μm².

NaCh labeling of the endplate region remained high for 2 d in culture (the longest time used in these experiments). After 3 d in culture, labeling of the endplate started to decrease significantly (data not shown). This has also been observed with the loose-patch technique (Lupa and Caldwell, 1991). Identification of the endplate allowed us to determine whether the frequency of NaCh clusters varied with distance from the endplate.

Agrin and NaCh expression in CHO cells

We used CHO cell lines that had been transfected previously with various isoforms of agrin (Campanelli et al., 1991; Ferns et al., 1992). These cells are known to express agrin both on the cell surface and intracellularly. Figure 2C2 shows labeling of Agrin8-transfected CHO cells with an antibody that recognizes all isoforms of agrin. By changing the focal plane, labeling could be seen both on the surface of the cells as well as in the cytoplasm. Agrin labeling was not uniform either within a given cell or between different cells. Exclusion of the primary antibody resulted in almost no labeling by the secondary antibody (Fig. 2A2, B2). Agrin labeling of CHO cells transfected with the other isoforms (Agrin11, Agrin11, and Agrin19) yielded similar results. Wild-type cells showed no labeling by the anti-agrin antibody (data not shown).

CHO cells express NaChs (Lalik et al., 1993) at a level that is detected with anti-NaCh antibodies (Fig. 2C1). Treatment with blocked antibody (preincubated with a large molar excess of the antigenic peptide) also revealed a detectable but lower level of staining (Fig. 2B1); this weak staining was completely abolished if secondary antibody was used alone (Fig. 2A1). Thus, anti-NaCh antibodies bind both specifically and nonspecifically to CHO cells. NaCh labeling of the CHO cells was similar for wild-type and for each of the transfected cell lines.
Clustering of NaChs by transfected CHO cells

Dissociated FDB muscle fibers and CHO cells were plated at densities that resulted in only a few contacts between CHO cells and each muscle fiber. The cultures were fixed after either 1 or 2 d in culture and double-labeled for NaChs and agrin. The results were the same for the 1 and 2 d cultures. It was necessary to label the cultures with the anti-agrin antibody, not only to determine whether CHO cells contacting muscle fibers expressed agrin (see above) but also to distinguish CHO cells from other cells that originated from the muscle dissociation. To distinguish wild-type CHO cells from other cell types, CHO cells were biotinylated before mixing them with muscle fibers; these wild-type cells were visualized with FITC-conjugated streptavidin.

Agrin8-expressing CHO cells were able to induce clusters of NaChs on muscle fibers (Fig. 3) after 1 d in culture (the shortest time we tested). Figure 3A shows NaCh labeling of two muscle

Figure 2. NaCh and agrin labeling of Agrin8-transfected CHO cells after 1 d of culture on Matrigel-coated coverslips. A, CHO cell labeling with secondary antibodies alone. Fluorescence when anti-NaCh (A1) and anti-agrin (A2) antibodies were omitted. A3, Nomarski image. B1, CHO cell labeling with “blocked” anti-NaCh antibody. Notice that there is some non-NaCh labeling of the CHO cells. B2, Weak, diffuse labeling when primary antibody to agrin (same protocol as A2) was omitted. B3, Nomarski image. C1, NaCh labeling of CHO cells. This signal is attributable both to a low level of endogenous NaCh expression in some CHO cells and to nonspecific labeling by the anti-NaCh antibody (see B1). C2, Agrin labeling of CHO cells. Notice that expression of agrin is heterogeneous between and within cells. C3, Nomarski image. Scale bars: A, 20 μm; B, C, 10 μm.
fibers lying next to each other. Each endplate is indicated by an asterisk. The arrows indicate four clusters of NaChs induced by two Agrin8-expressing CHO cells lying between and contacting the fibers. Notice that the CHO cell membrane not in contact with the muscle fiber has a NaCh signal below that of the nonjunctional muscle fiber membrane. Figure 3A shows agrin labeling of the same field. Endogenous expression of agrin by the muscle fiber does not generate an appreciable signal; this is probably attributable to the removal of the basal lamina by the enzymatic dissociation (Betz and Sakmann, 1973). Only one of the two CHO cells in the figure appeared heavily labeled for agrin in this focal plane. The other CHO cell was clearly labeled by the agrin antibody in another focal plane (data not shown). Figure 3B shows another example of an Agrin8-transfected cell that has induced NaCh clusters on two muscle fibers. Notice in the composite images of Figure 3, A3 and B3, that the increase in NaCh labeling occurs at the contact with the CHO cells.

Although CHO cells labeled only weakly with the anti-NaCh antibody (and NaCh labeling of CHO cells was primarily intracellular rather than membrane-associated) (Fig. 2C), we were concerned that the superposition of the weak signal from the CHO cell with that of the muscle fiber membrane could make it appear that there was an increase in NaCh concentration at that point. For that reason, only contacts where muscle fiber labeling and CHO cell labeling could be distinguished were considered as candidates for clustering. To exclude any contribution to the NaCh signal from labeling of the CHO cells, most contacts that could be analyzed occurred at the side of muscle fibers. All of the contacts illustrated in Figure 3 were on the side of fibers except Figure 3C, which shows an example in which an Agrin8-expressing CHO cell had contacted the upper surface of a muscle fiber (see Fig. 3C4). In most cases of this latter type, it was not possible to determine whether there was NaCh clustering in the muscle membrane at the contact site, because the intracellular labeling of the CHO cell contributed to the signal. However, in this case, the labeling occurred at thin, veil-like CHO cell membrane contacts, and the intensity of labeling exceeded that in any isolated CHO cell. We found that approximately one-third (39%) of the contacts that could be analyzed between Agrin8-expressing CHO cells and muscle fibers resulted in NaCh labeling (70 of 181 contacts); 154 of 335 contacts (46%) were not scorable. This is similar to the percentage of contacts between transfected CHO cells and rat myotubes that produced AChR clusters (10–50%) (Campanelli et al., 1991). Because the muscles used in the present experiments were not predenervated, AChR clusters were not expected and, thus, we did not test whether NaCh clusters coincided with AChR clusters. No spontaneous clustering of NaChs was observed under any of our culture conditions (with or without CHO cells). Because clusters were only observed at contact sites, we also conclude that CHO cells do not induce stable clusters and subsequently migrate away during the time course of these experiments.

One unexpected finding was that NaCh clustering did not always occur where Agrin8 expression was highest. For example, in Figure 3C3 the intense NaCh labeling was not co-localized with the maximal agrin labeling. The NaCh labeling was associated with the contact of the veil-like area of the CHO cell (see Fig. 3C4). Explanations for this incongruence include the possibility that much of the agrin signal is intracellular, that the density of agrin on the CHO cells exceeds that required for NaCh clustering, and that NaCh clustering may occur, followed by a redistribution of agrin within the CHO cell.

Agrin8-induced clusters of NaChs could occur anywhere on a muscle fiber. Figure 3D shows an example of Agrin8-induced NaCh clustering near the end of a muscle fiber (bottom of the figure, 400 µm from the endplate). We had originally predicted that we might be able to observe clustering only near the perijunctional region where NaCh density is greatest. This is clearly not the case, because we have observed clusters at distances >600 µm from the endplate (Fig. 4).

Figure 4 shows the distribution of clusters as a function of distance from the endplate. This is a summary of 70 positive contacts of 181 scorable contacts measured from four cultures. The data have been plotted as the percentage of scorable contacts that showed clusters. The number of contacts scored is shown above each bar. Notice that the percentage of contacts that caused clustering decreased beyond 80 µm. Using intensity measurements of the images, we determined that the average NaCh signal in the first 80 µm from the endplate (excluding the signal from the endplate itself) was 2.5 times that of the background intensity measured at distances farther from the endplate. This is in agreement with loose patch-clamp measurements of sodium current density (Caldwell and Milton, 1988; Lupa et al., 1993). If the number of clusters at a given distance from the endplate is proportional to the density of NaChs, there should be 2.5 times as many clusters in the first 80 µm than in subsequent 80 µm sections, and the sections >80 µm from the endplate should contain an equal number of clusters. We observed that the first 80 µm contained 1.7 times as many clusters as the average contained in the 80 µm segments between 80 and 400 µm from the endplate. This suggests that the probability of NaChs being clustered increases with the concentration of NaChs in the membrane but not in a linear manner. The number of contacts beyond 400 µm decreased markedly, most likely because FDB fibers are ~1 mm in length and the endplate usually is not in the exact middle of the fiber. This reduced the probability of finding contacts at distances >400 µm, and the clusters observed at these greater distances, therefore, were not included in the average of the extrajunctional segments.

We did not observe clusters of NaChs in cocultures with wild-type CHO cells or the Agrin0-, Agrin11-, or Agrin19-transfected CHO cell lines. At least three separate experiments, each including twelve coverslips, were examined for the wild-type and for each transfected cell line; in each of these, >100 scorable contacts were observed with no NaCh clustering. Figure 3E shows an example of an Agrin0-transfected CHO cell in contact with a muscle fiber. Even though this CHO cell was covering the endplate and the perijunctional area, there was no CHO cell-induced NaCh clustering.

**DISCUSSION**

During normal development, proper organization of the postsynaptic membrane at the NMJ is initiated by contact from the presynaptic terminal of a motor neuron. Neuronally expressed and secreted isoforms of agrin are known to be involved in the concentration of a variety of postsynaptic molecules at the NMJ (Wallace, 1989; McMahan, 1990). The purpose of this study was to determine whether any of the neuronally expressed isoforms of agrin could induce clustering of NaChs in dissociated adult muscle fibers and, therefore, perhaps be involved in the *in vivo* increased density of NaChs at the NMJ.

**Agrin-induced NaCh clustering**

We used a culture system that combined dissociated adult rat FDB muscle fibers and CHO cells that had been transfected with
various isoforms of rat agrin. This system provided us with several advantages. The adult muscle fibers expressed a higher level of extrajunctional NaChs than embryonic myotubes. This allowed us to use an antibody against NaChs to assay for clustering rather than the more sensitive, yet spatially constrained, loose patch-clamp technique. Using CHO cells that express agrin as a surface molecule may present a more physiological situation than bath application of soluble agrin, because it provides a focal presentation of agrin, as does a presynaptic nerve terminal.

We found that the Agrin8-transfected CHO cell line was able to cluster NaChs in cultured adult FDB fibers and that it was the only CHO cell line of the ones tested (one wild type and four cluster NaChs in cultured adult FDB fibers and that it was the most effective. Limitations on measuring very low levels of NaCh density may have caused us to miss weak clustering produced by the other agrin isoforms. It is also possible that longer exposure to the agrin-transfected CHO cells would have revealed clustering by other agrin isoforms. Additionally, our in vitro assay system may not be an adequate model of the in vivo maturation of the NMJ. For example, differentiated adult muscle fibers may not respond to the same isoforms as muscle fibers from embryonic and neonatal animals. Ferns et al. (1993) found that membrane glycoproteins were necessary for the clustering of AChRs by some isoforms of agrin; adult FDB muscle fibers may lack receptors or glycoproteins specific for the other agrin isoforms.

A previous study (Lupa and Caldwell, 1991) showed that *Torpedo* agrin applied to the bath was unable to generate clusters of NaChs in adult FDB muscle fibers. Together with the results reported here, this suggests that unlike AChR aggregation, NaCh clustering may require agrin from the correct species, that *Torpedo* extract may not contain Agrin8, or that NaCh clustering may require focal application of agrin.

All of our analysis of NaCh clustering was done with light microscopy. Because light microscopy does not allow one to distinguish the CHO cell membrane from the muscle membrane at the site of contact, we cannot exclude the possibility that the NaCh signal originates in the CHO cell rather than in the muscle. We believe this explanation is unlikely, because the density of NaChs in CHO cell membranes is much lower than that in muscle cells and because, of the five CHO cell lines tested, only one was able to induce clustering. Moreover, we never observed NaCh clustering between pairs of Agrin8-expressing CHO cells. Thus, we interpret the increased NaCh signal at contact sites as coming from muscle membrane.

**Distribution of NaCh clustering**

Milton and Caldwell (1990) found that approximately half of the NaChs in extrajunctional regions of adult rodent FDB muscle fibers were free to diffuse within the membrane, and they proposed a model in which NaChs can attach and detach from binding sites during their lifetime in the membrane. We found that Agrin8-transfected CHO cells were more likely to induce NaCh clusters when they were close to the muscle fiber endplate.

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**Figure 3.** CHO cells expressing Agrin8-induced NaCh clusters on muscle fibers after 1 d in culture (*A–D*). Note that the images have been pseudocolored with the natural colors reversed. *A1*, NaCh labeling of two muscle fibers. The endplate on each fiber is indicated with an asterisk (endplate on top fiber is out of focus). Notice the NaCh clusters on the edge of both fibers (arrows). The endplate of the right fiber protruded into the space between the two fibers and produced an apparent NaCh signal on the left fiber that was considered to be an artifact. *A2*, Agrin labeling. Fluorescence in the top left corner is the result of detritus out of the focal plane. *A3*, Composite image of *A1* and *A2*. Notice that the NaCh clusters are coincident with the CHO cells (the top CHO cell contained agrin that was not in this plane of focus). *A4*, Nomarski image. The cluster of cells in the bottom left corner did not label for agrin (see *A2*) and was not scored for clustering. *B1*, NaCh labeling of two muscle fibers with a CHO cell nestled at their juncture. *Arrows* show NaCh clustering at the CHO cell contact. *Asterisk* indicates out-of-focus endplate. Notice the myonucleus bulging from the bottom fiber in the perijunctional region. *B2*, Agrin labeling. *B3*, Composite image of *B1* and *B2*. *B4*, Nomarski image. *C*, NaCh clustering induced by an Agrin8-expressing CHO cell on top of a muscle fiber (110 μm from endplate). *C1*, NaCh labeling. *C2*, Agrin labeling. *C3*, Composite image of *C1* and *C2*. *C4*, Nomarski image. The NaCh labeling was coincident with the thin, flat contact between CHO cell and muscle fiber. *D*, NaCh clustering induced by two Agrin8-expressing CHO cells near the end of a muscle fiber (400 μm from endplate). *D1*, NaCh labeling. The NaCh signal from the CHO cell was one of the strongest seen; a more typical example is illustrated in *E1*. *D2*, Agrin labeling. *D3*, Composite image of *D1* and *D2*. Both CHO cells contained agrin, but the agrin signal in one cell was not in this focal plane. Notice that there is NaCh labeling on the fiber adjacent to both cells. *D4*, Nomarski image. *E*, Agrin0-transfected CHO cells do not induce NaCh clustering. This example was chosen because the contact is close to the endplate where channel density is highest and where the highest incidence of clusters occurred with Agrin8 cells (Fig. 4). *E1*, NaCh labeling. Notice the labeling of the endplate (asterisk) and the perijunctional region directly across from the endplate. *E2*, Agrin labeling. The morphous signals in the top portion of the panel are from CHO cells out of the plane of focus. *E3*, Composite of *E1* and *E2*. The CHO cell next to the endplate is not able to cluster NaChs, even though it is directly adjacent to the endplate. *E4*, Nomarski image. Scale bars: *A–D*, 20 μm; *C, D*, 10 μm.

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**Figure 4.** Distribution of Agrin8-induced NaCh clusters relative to the endplate of a muscle fiber. Data are the summary from four cultures. The vertical axis represents the percentage of contacts that induced clustering out of the scorable contacts observed (the number above each bar). The binwidth is 80 μm. Distance from the endplate was measured from the center of the endplate using an eyepiece reticle.
Because the concentration of NaChs increases with proximity to the endplate, it raises the possibility that the Agrin8-transfected CHO cells have access to more NaChs closer to the endplate and that this increases the likelihood of clustering close to the endplate. This is based on the assumption that new clusters of NaChs are formed from NaChs that are not permanently anchored to the cytoskeleton, as has been suggested for AChRs (Wallace, 1992).

Only approximately one-third of the Agrin8 CHO cell–muscle contacts produced clusters of NaChs, which is similar to the percentage of Agrin8 CHO cell contacts that induced AChRs (Campanelli et al., 1991). This low fraction could be attributable to a variable amount of time that the CHO cells contacted the fiber; however, when we observed identified CHO cells in the same culture over a 4–6 hr period, there were small changes in position but little evidence for significant migration. Alternatively, the density of the membrane receptor for agrin may be limiting. α-Dystroglycan is a receptor for agrin (Bowé et al., 1994; Campanelli et al., 1994; Gee et al., 1994; Sugiyama et al., 1994), but there is evidence that another receptor exists that mediates agrin-induced AChR clustering (Gesemann et al., 1996; Meier et al., 1996); the identity and membrane density of this receptor are unknown.

Developmental regulation of NaCh clustering
During development, neuronal expression of Agrin8 increases markedly starting 1 week after birth; expression of other isoforms either changes very little or decreases (Hoch et al., 1993, their Figs. 6, 7). This increase in Agrin8 expression corresponds closely with the expression and concentration of the adult subtype of NaCh at the NMJ (Lupa et al., 1993, their Fig. 2). Figure 5 shows a plot of the contribution of various z site splice variants to the total agrin in the spinal cord at different developmental stages (RNA levels replotted from Hoch et al., 1993). Notice that the relative expression of Agrin8 corresponds temporally to the increase in junctional NaCh concentration (solid line with no symbols, redrawn from Lupa et al., 1993). This temporal relationship, in conjunction with our findings here, suggests that the onset of Agrin8 expression is responsible for clustering of NaChs at the NMJ during normal development.

The expression and concentration of the adult NaCh subtype at the NMJ begin shortly after birth as the muscle shifts expression of NaCh subtype from the embryonic form (identical to the cardiac subtype rH1 that is also designated SkM2 or μ2) to the adult form (SkM1). Although this switch coincides with the development of the high NaCh density at the endplate, both subtypes can be concentrated at the endplate (Caldwell and Milton, 1988; Lupa et al., 1995). The aggregation at the developing endplate coincides temporally with other late synaptic formation events such as infolding of the postsynaptic membrane and the appearance of a specific isoform of protein kinase C (PKCθ) (Hilgenberg and Miles, 1995). It is possible that Agrin8 is also responsible for induction of folds and the localization of PKCθ.

Related mechanisms of molecular aggregation
Because NaChs and AChRs are not precisely co-localized and because they become concentrated at different developmental stages, it follows that some of the mechanisms and molecules controlling expression and localization of these proteins are different. In addition to agrin, other molecules released by the nerve terminal might be involved in NaCh clustering. Neuregulin is a signaling molecule that is synthesized by motor neurons and deposited in the synaptic basal lamina (Jo et al., 1995). Multiple transcripts for neuregulin are generated by alternative splicing, and some of these isoforms can increase AChR synthesis and the density of NaChs on the surface of cultured muscle cells (for review, see Mudge, 1993). Receptors for neuregulin are members of the EGF receptor family and are protein tyrosine kinases. Three members of this family (erbB2, erbB3, and erbB4) are concentrated at the NMJ of adult skeletal muscle. Expression of neuregulin and its receptors increases during early postnatal development (Zhu et al., 1995). The similarity in developmental appearance of NaChs and neuregulin/erbBs suggests that in addition to agrin, some isoforms of neuregulin may regulate NaCh density at the endplate.

Intracellular molecules establish or maintain the specialized membrane at the endplate. Because NaChs and AChRs are in separate domains in the subsynaptic membrane (Flucher and Daniels, 1989), the molecules that link NaChs to the cytoskeleton are likely to be different from those that tether AChRs. Ankyrin, which co-purifies with NaChs (Sriniwasan et al., 1988), is a candidate for immobilizing NaChs. Three genes for ankyrin have been identified, and multiple isoforms are generated by alternative splicing; the specific isoform(s) present at the NMJ has not been determined.

Other membrane-associated, intracellular molecules that could tether NaChs are those that contain PDZ binding domains. The PDZ domain consists of ~90 amino acids and is present in several homologous proteins (Cho et al., 1992). Candidate molecules with a PDZ domain are the syntrophins, a family of proteins that bind to dystrophin and are part of the dystrophin–glycoprotein complex (Adams et al., 1995; Yang et al., 1995). One of the syntro-
phins is concentrated at the NMJ (Peters et al., 1994) and could participate in tethering NaChs.

**Conclusion**

The coculture experiments presented here show that a specific neural isoform of agrin (Agrin8) is capable of rapidly clustering NaChs. This is the one isoform for which temporal expression during development coincides with the postnatal aggregation of NaChs at the endplate and other late processes of synapse maturation. These results support the hypothesis that Agrin8 is a common signal that initiates not only AChR and NaCh clustering but also additional postsynaptic specializations required for synapse formation and maturation.

**REFERENCES**


Flucher BE, Daniels MP (1989) Distribution of Na+ channels and ankyrin in neuromuscular junctions is complementary to that of acetylcholine receptors and the 43 kDa protein. Neuron 3:163–175.


