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Abstract

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An Amino-terminal Extension Is Required for the Secretion of Chick Agrin and Its Binding to Extracellular Matrix

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Abstract. Agrin is an extracellular matrix (ECM) protein with a calculated relative molecular mass of more than 200 kD that induces the aggregation of acetylcholine receptors (AChRs) at the neuromuscular junction. This activity has been mapped to its COOH terminus. In an attempt to identify the functions of the NH2-terminal end, we have now characterized full-length chick agrin. We show that chick agrin encoded by a previously described cDNA is not secreted from transfected cells. Secretion is achieved with a construct that includes an additional 350 bp derived from the 5' end of chick agrin mRNA. Recombinant agrin is a heparan sulfate proteoglycan (HSPG) of more than 400 kD with glycosaminoglycan side chains attached only to the NH2-terminal half. Endogenous agrin in tissue homogenates also has an apparent molecular mass of >400 kD. While the amino acid sequence encoded by the 350-bp extension has no homology to published rat agrin, it includes a stretch of 15 amino acids that is 80% identical to a previously identified bovine HSPG. The extension is required for binding of agrin to ECM. AChR aggregates induced by recombinant agrin that includes the extension are considerably smaller than those induced by agrin fragments, suggesting that binding of agrin to ECM modulates the size of receptor clusters. In addition, we found a site encoding seven amino acids at the NH2-terminal end of agrin that is alternatively spliced. While motor neurons express the splice variant with the seven amino acid long insert, muscle cells mainly synthesize isoforms that lack this insert. In conclusion, the cDNAs described here code for chick agrin that has all the characteristics previously allocated to endogenous agrin.

Proper synaptic transmission requires a high degree of cell specialization at the site of contact between pre- and postsynaptic cells. The best studied chemical synapse is the neuromuscular junction (NMJ), where the nerve terminal of a motor neuron synapses on a muscle fiber. The ingrowing motor axon initiates in muscle fibers a local accumulation of several proteins of the cytosol, the cell membrane and the extracellular matrix (ECM; for a review see Hall and Sanes, 1993). The formation of these specializations in the muscle fiber is triggered by molecules released from the nerve (Frank and Fischbach, 1979; Dahm and Landmesser, 1991). The best-characterized molecule known to be involved in this process is agrin. When added to cultured myotubes, agrin induces the aggregation of AChRs and other molecules concentrated at the NMJ (Nitkin et al., 1987; Wallace, 1989). This process is believed to be mediated by binding of agrin to specific signaling receptors on the muscle cell surface (for reviews see Fallon and Hall, 1994; Bowe and Fallon, 1995). Active agrin is synthesized by motor neurons and is transported to the nerve terminal where it is released and becomes incorporated into the nascent muscle cell basal lamina (reviewed by McMahan, 1990). Nevertheless, agrin is also found in nonsynaptic basal laminae of muscle, in kidney and capillaries (Reist et al., 1987; Fallon and Gelfman, 1989; Godfrey, 1991; Rupp et al., 1991; Ruegg et al., 1992), suggesting that it may also have other functions.

The molecular mass of agrin deduced from cDNAs in rat, chick, and the marine ray is more than 200 kD (Rupp et al., 1991; Tsim et al., 1992; Smith et al., 1992). However a COOH-terminal, 45-kD fragment of agrin is sufficient to induce AChR clustering on cultured myotubes with high efficacy (Gesemann et al., 1995). Within this active fragment, two sites (called A and B in chick and ray, and y and z in rat) whose positions and amino acid sequences are conserved among the three species are subject to alternative mRNA splicing. When expressed in eukaryotic cells, only recombinant agrin isoforms with inserts at both sites are active in inducing AChR clusters, while those lacking in-
sers are much less or not at all active (Ruegg et al., 1992; Ferns et al., 1992, 1993; Gesemann et al., 1995). In vivo, transcripts encoding agrin isoforms are differentially distributed. The most active ones are expressed by cholinergic neurons, such as motor neurons of the spinal cord and ciliary ganglia neurons, and the inactive ones are highly expressed by nonneuronal cells like myotubes and Schwann cells (Ruegg et al., 1992; McMahan et al., 1992; Hoch et al., 1993; Thomas et al., 1993; Smith and O'Dowd, 1994; Ma et al., 1994, 1995). Thus, expression of specific isoforms is most likely the basis of the observed difference between muscle cell-derived and motor neuron-derived agrin in inducing AChR aggregates (Reist et al., 1992; Cohen and Godfrey, 1992).

As in other extracellular matrix proteins, agrin consists of several repeated regions. There are nine Kazal inhibitor-like domains that are also found in follistatin, six EGF-like domains, and three globular domains homologous to COOH-terminal domains of laminin α-chains. Homologous domains in other ECM molecules have been implicated in particular functions, such as cell–cell recognition and growth factor binding, which has led to many proposals for additional functions of agrin (e.g., Pathy and Nikolaos, 1993). The NH2-terminal half of agrin also contains several potential sites for carbohydrate attachment. There are five potential N-glycosylation sites, two serine/threonine-rich regions, and at least six potential glycosaminoglycan (GAG) chain attachment sites.

As a first step in searching for additional functions of agrin isoforms, we aimed to express full-length chick agrin and to characterize its biochemical properties in more detail. When we transfected COS-7 cells with the previously described agrin cDNA (Tsim et al., 1992) the recombinant protein was not targeted to the ER and hence, was not secreted from the cells. Here we present a chick agrin cDNA, whose coding sequence at the 5′ end is extended by ~350 bp. Agrin with this extension is secreted from transfected cells. The new sequence includes a stretch of 15 amino acids that is highly homologous to a peptide previously characterized in a heparan sulfate proteoglycan (HSPG) of bovine kidney (Hagen et al., 1993). Only recombinant agrin that includes this extension binds to Matrigel™, a solubilized mixture of ECM components. Recombinant chick agrin, depending on the inserts at site A and B, is active in inducing AChR aggregates on cultured myotubes. However, AChR clusters are considerably smaller than those induced by fragments of agrin. In addition, we have found a new 21-bp-long site at the 5′ end that is alternatively spliced. At early stages of synaptogenesis, motor neurons synthesize agrin mRNA with the 21-bp-long insert and the majority of agrin transcripts expressed in muscle lack this insert. In conclusion, these five extended constructs code for chick agrin isoforms that appear to fulfill all criteria to be full-length.

**Materials and Methods**

**Polymerase Chain Reaction**

Conditions for PCR were as described elsewhere (Gesemann et al., 1995). Numbering of oligonucleotides is according to the sequences published by Tsim et al. (1992). Oligonucleotides of new 5′ sequences are negatively numbered. All PCR products used to generate expression constructs were cloned into PCR II cloning vector (Invitrogen, San Diego, CA) and sequenced to ensure that no mutations were introduced by DNA amplification.

**mRNA Isolation, Reverse Transcription and PCR**

Poly A+ RNA was isolated with the Micro-FastTrack mRNA Isolation Kit (Invitrogen) according to the manufacturer's advice. RACE experiments (Froehmann et al., 1988; Loh et al., 1989) were performed with the 5′ RACE System (GIBCO BRL, Gaithersburg, MD) according to the manufacturer's advice. The oligonucleotide used for the reverse transcription reaction was as366 (CGTTGGTCGTTAGGTTGGAATA) and the oligonucleotide as232 (ATCCCTGGCAGGCATCTCG) served as primer in the subsequent PCR reaction. Reverse transcriptase-polymerase chain reaction was carried out as described in Ruegg et al. (1992) except that Superscript™ II reverse transcriptase (GIBCO BRL) was used. To construct cDNAs encoding complete agrin isoforms, first strand synthesis was primed with as366 and subsequent PCR was conducted using EcoRI 5′-289 (GCATAGAATTCGCCGCGGCATGGG) as the sense primer and as232 as the antisense primer (see also below).

**Chicken Genomic Library Screening**

A chicken genomic library from adult chicken liver (Clontech, Palo Alto, CA) was screened according to the manufacturer's advice. Recombinant phage (9 × 105) from the amplified EMBL-3 library were screened with 32P-labeled DNA probe (nucleotides 134 to 547; for location see Fig. 3). Hybridization was at 42°C in 5× SSC, 0.1% SDS, 5× Denhardt's reagent, 30% formamide, and 0.1 mg/ml salmon sperm DNA. Filters were washed at a maximal stringency of 0.2× SSC, 0.1% SDS at 65°C. Two rounds of dilution were applied for cloning. DNA of positive recombinant phage was prepared as follows: 450 ml of 50 mM Tris-HCl, pH 7.5, 10 mM NaCl, 8 mM MgSO4, 0.1% gelatin were inoculated with a single positive plaque. 60 μl of maltose-induced NM 538 (Sambrook et al., 1989) in 10 mM MgSO4 (OD260 = 2) were added and incubated for 15 min at 37°C. After adding 30 ml of growth medium, 10 mM MgSO4 cells were incubated at 37°C until lysis. 300 μl chloroform were added and incubation was continued for another 45 min. The lysate was centrifuged for 15 min at 15,000 g to pellet cell debris. The supernatant was incubated with approximately 100 μg DNAse (Boehringer Mannheim, Mannheim, Germany) and 100 μg RNAse (Boehringer Mannheim) for 1 h at 37°C. 7 ml of 20% PEG (6,000) in 2.5 M NaCl were added and incubation was continued for 1 h on ice. After centrifugation for 15 min at 13,000 g, the pellet was dissolved in 450 μl 10 mM Tris-HCl, pH 8.0, 0.5% SDS, 10 mM EDTA, 30% formamide, 0.1 mg/ml proteinase K (Boehringer Mannheim). Digestion was performed for 30 min at 60°C. After phenol/chloroform extraction, DNA was precipitated with ethanol. Inserts were excised by Sall digestion, analyzed on agarose gel, and subcloned into Bluescript KS (Stratagene, La Jolla, CA).

**Alternative mRNA Splicing**

To examine the relative abundance of agrin mRNA, randomly primed cDNAs were subjected to PCR using either primers s86 (CCCACC-GCAAGGATTGATAG) and as232, which flank the novel splice site, or primers s5377 (TTTGATTGTTAGGCTATCTG) and as5511 (CCTCTTCTTGTAGGTGGAATG) as the sense primer and as232 as the antisense primer (see above).

**Expression Constructs**

To distinguish expression constructs from recombinant protein, cDNA constructs are named with the prefix p followed by the name of the protein they encode. All partial chick agrin cDNA constructs, with the exception of pcFull ps, pcNFull ps, pcNAgrin, and pcAgrin (see below), contain the cDNA encoding the NH2-terminal part of the recombinant protein, with the exception of pcFull ps, pcNFull ps, pcNAgrin, and pcAgrin (see below), contain the cDNA encoding the NH2-terminal part of the recombinant protein, with the exception of pcFull ps, pcNFull ps, pcNAgrin, and pcAgrin (see below), contain the cDNA encoding the NH2-terminal part of the recombinant protein, with the exception of pcFull ps, pcNFull ps, pcNAgrin, and pcAgrin (see below), contain the cDNA encoding the NH2-terminal part of the recombinant protein, with the exception of pcFull ps, pcNFull ps, pcNAgrin, and pcAgrin (see below), contain the cDNA encoding the NH2-terminal part of the recombinant protein.
using s5377 as the sense primer (TTTGATGGTAGGACGTACATG), TCAAGTCCTCCTCGCTGATTAACTITFGTTCTITGGCTA/GG/A-pcDNA I (Invitrogen) to obtain pMyc. To fuse these sequences to the CAGTGTAATA) and pC95AOB0 as template. The gel-purified PCR product, yielding pcNFullh. The analogous NH2-terminal construct (pcNFullh) containing the putative ER signal sequence as proposed by Tsim et al. (1992) was constructed by replacing the Sall/Xhol fragment of pcNFullh with that of pcFullh.

Construction of complete chick agrin cDNA was accomplished by three cloning steps: PCR products encoding the 5' extension were excised from the PCR II cloning vector with endonucleases EcoRI and Xhol and subcloned into a modified pcDNA I vector lacking the BamHI restriction site. The insert was then fused to the region encoding the NH2-terminal half of agrin by digestion with Xhol and BamHI and subsequent ligation to the corresponding CDNA fragment of pcNFull. The constructs were named pcNAgrinl (no 21-bp insert) and pcNAgrin2 (with 21-bp insert). In the last step, sequence for the COOH-terminal part of chick agrin was fused to pcNAgrin2, pcNAgrin3 using BamHI and Xhol restriction sites. To generate complete chick agrin CDNAs with different A and B sites, both recombinant pcNFullh constructs were used as parent constructs. Unless specifically stated, cDNA constructs used throughout this study contain inserts at the novel site and lack inserts at sites A and B (pcAgrinlABO).

Antibodies
To raise polyclonal antisera against cFullhSA4B19, the procedures described by Gesemann et al. (1995) were used.

Tissue Culture and Transfections
COS-7 cells (Gluzman, 1981) were cultured in DME (GIBCO BRL) supplemented with 10% newborn calf serum, 10 mM sodium pyruvate, 100 i/ml penicillin, and 100 g/ml streptomycin (all from GIBCO BRL). COS-7 cells were transiently transfected with the DEAE-dextran–based method described by Cullen (1987). Chick myotube cultures were prepared as described (Gesemann et al., 1995).

Metabolic Labeling, Immunoprecipitation, and Deglycosylation
Metabolic labeling of intracellular proteins and immunoprecipitation followed by deglycosylation with endo-β-N-acetylglucosaminidase H (Boehringer Mannheim) was essentially done as described by Wessels et al. (1991) with some minor modifications: cell lysis and immunoprecipitation was performed at 0.5 M NaCl in the presence of 1 mM PMSF, 1 mg/ml pepstatin, 10 mg/ml aprotinin, 1 mg/ml leupeptin, and 1.75 mg/ml benzamidine.

Immunoblot Analysis of Chick Tissues
Tissues from E11-E15 White Leghorn chick embryos were harvested and homogenized on ice with a 15 ml Dounce-homogenizer in the appropriate volume of PBS containing 1 mM PMSF, 1 mg/ml pepstatin, 1 mg/ml antipain, 10 mg/ml aprotinin, 1 mg/ml leupeptin, and 1.75 mg/ml benzamidine. Homogenized tissues were subjected to SDS-PAGE on a 3-12% gradient gel. Proteins (~100 mg per lane) were transferred to nitrocellulose membranes (Millipore, Marlborough, MA) according to Towbin et al. (1979) in the absence of methanol. The quality of protein transfer was tested by staining the membrane with Ponceau S (Serva, Heidelberg, Germany). Agrin-like immunoreactivity was visualized by the ECL method (Amer sham, Buckinghamshire, UK).

Enrichment of Motor Neurons
Spinal cord segments were dissected from E5 White Leghorn chick embryos (developmental stages 26 and 27 according to the criteria of Hamburger and Hamilton, 1951). After digestion with 0.1% trypsin in PBS containing 1% glucose for 30 min at 37°C, the tissue was washed with 5 ml PBS supplemented with 300 U/ml DNase. A single cell suspension was obtained by gentle trituration. Cell debris were removed by centrifugation through a 3 ml cushion of 3.5% BSA in L-15, pH 7.4 for 15 min at 100 g. The cell pellet was resuspended in PBS, 0.5 mM EDTA. Motor neurons were enriched by density centrifugation on metrizamide (Dohrmann et al., 1982), by panning with mAb SCI (Tanaka and Ohtsuka, 1984; Bloch-Gallego et al., 1991), or by metrizamide centrifugation followed by SCI panning. The panning of spinal cord cells on SCI-coated petri dishes (Bloch-Gallego et al., 1991) was slightly modified in the following way: 100 mm polystyrene petri dishes (Greiner, Nürtin gen, Germany) were coated with 10 mg/ml purified IgG of mAb SCI in 50 mM sodium carbonate buffer, pH 9.6. Dishes were washed twice with PBS and incubated with 3% BSA in PBS for 1 h at room temperature. Cells were resuspended in MEM, 10% horse serum and incubated for 1 h at room temperature. The plates were washed four times with PBS with gentle swirling to remove nonadherent cells. Panned cells were lysed directly using the lysis buffer supplied by the Micro-FastTrack mRNA Isolation Kit.

Immunocytochemistry
Transiently transfected COS-7 cells were chilled to 4°C and anti-c55 antibodies (1:1,000) were incubated for 30 min at 4°C. After three washes with PBS, cells were fixed for 30 min at 4°C and 30 min at room temperature with 4% paraformaldehyde, 11% sucrose in 0.1 M potassium phosphate buffer, pH 7.2. After rinsing with PBS and 20 ml glycine in PBS, cells were permeabilized with 0.1% saponin in PBS (PBS/S). 10 mg/ml IgG of mAb SBI in PBS/S + 10% normal goat serum (NGS) was incubated for 2 h. After four washes with PBS/S, cells were incubated for 1 h with rhodamine-conjugated sheep anti-mouse IgG (1/200; Cappel, Durham, NC) and fluorescein-conjugated goat anti-rabbit IgG (1/200; Jackson Immunoresearch Laboratories, Inc., West Grove, PA) in PBS + 10% NGS. Cultures were washed with PBS/S, mounted on glass cover slips with Vectashield® (Vector Laboratories, Inc., Burlingame, CA), and examined with fluorescence microscopy.

Quantification of AChR Aggregation
Incubation of cultured chick muscle cells with agrin, labeling of AChRs with rhodamine-conjugated a-bungarotoxin, and quantification of AChR clusters using an imaging system (Image-i/AT; Universal Imaging, West Chester, PA) was done as described by Gesemann et al. (1995). Due to the small size of AChR clusters induced by cAgrin sAB; and cAgrin aAB; aggregates with the longer axis of ≥2 μm were included. Concentration of recombinant agrin in conditioned medium of transfected COS cells was determined with an ELISA as described (Gesemann et al., 1995).

Results
The Coding Sequence of Agrin Is Extended at the 5' End
To study properties of the entire chick agrin, the overlapping cDNAs isolated by Tsim et al. (1992) were fused. This cDNA construct is expected to encode full-length agrin.
which we called cFullp (for chick full-length with putative signal sequence; Fig. 1). COS-7 cells were transiently transfected with this cDNA construct to obtain recombinant protein. The presence of cFullp in the conditioned medium was tested by immunoprecipitation after labeling with [35S]methionine. No immunoreactive protein was detected in the supernatant with either mAb 5B1 (Reist et al., 1987; data not shown) or a polyclonal antiserum raised against the 95 kD, COOH-terminal fragment of chick agrin (Gesemann et al., 1995; Fig. 2), indicating that cFullp is not secreted from COS-7 cells. No agrin-like immunoreactivity, as assayed by immunohistochemistry, was found associated with the cell surface of transfected COS cells (data not shown).

Soluble secretory molecules usually contain a signal sequence located at their NH₂ terminus that targets protein synthesis to the endoplasmatic reticulum (for review see Verner and Schatz, 1988). Such sequences are characterized by a stretch of 16–26 residues with a polar, basic NH₂ terminus and a central hydrophobic core of 7–15 residues (von Heijne, 1983; Perlman and Halvarson, 1983). Because the NH₂-terminal sequence of the cDNA of Tsim et al. (1992) does not meet these criteria, we replaced it with the signal sequence of hemagglutinin. The latter sequence is capable of targeting protein synthesis of agrin fragments to the secretory pathway (Tsim et al., 1992; Ruegg et al., 1992; Gesemann et al., 1995). As shown in Fig. 2, cFullh (for chick full-length with hemagglutinin signal sequence; Fig. 1) accumulates in the medium of transfected cells. As reported previously (Gesemann et al., 1995), it has an apparent molecular mass is more than 400 kD, indicating the high content of carbohydrates. Molecular weights in kD of standard proteins are indicated.
This result shows that the NH$_2$-terminal sequence of cFull ps does not serve as a signal sequence in COS cells.

In an attempt to isolate the endogenous signal sequence of chick agrin, we performed RACE experiments (rapid amplification of cDNA ends; Frohmann et al., 1988; Loh et al., 1989) on poly A$^+$ RNA from E5 to E6 chick spinal cord. Agrin-derived primers as366 used for reverse transcription, and as232 for the subsequent anchored PCR were both designed to anneal at the 5’ end of the agrin mRNA where chick and rat agrin are still highly homologous (see Fig. 3). We analyzed more than 40 individual PCR products. The longest PCR product comprised sequences from position 107 through 620 (Fig. 3). In all the clones analyzed, an additional thymidine residue was inserted at position 419 (V, Fig. 3). This insertion neutralizes the in-frame stop codon upstream of the initiator methionine postulated by Tsim et al. (1992) and extends the open reading frame. The 5’ end of the longest PCR product coded for a stretch of hydrophobic amino acids indicative of a signal sequence (von Heijne, 1983; Perlman and Halvorson, 1983), but no new initiator methionine was found. Several attempts with altered conditions did not yield longer RACE products (data not shown). One reason may be that the secondary structure of agrin mRNA induced a premature stop of reverse transcription (Sambrook et al., 1989).

We therefore screened a chick genomic library with a DNA probe spanning nucleotides 134 to 547 (for the location see Fig. 3). Out of $9 \times 10^6$ phages screened, two positive clones were analyzed. In both clones, an in-frame ATG was found 16 nucleotides upstream of the longest PCR product. This ATG is part of a consensus for initiation of translation in eukaryotes (Kozak, 1986) and is followed by 23 mainly hydrophobic residues and the proposed signal sequence cleavage site at Asn 25 (von Heijne, 1986). In addition, an in-frame stop codon is located 63 nucleotides upstream of the potential site of initiation. To prove that these novel sequences are within the transcribed portion of the agrin gene, we performed a reverse transcriptase–polymerase chain reaction (RT-PCR). Amplification was conducted using primers s-368 and as-273 (see Fig. 3). Although the yield was very low, the resulting PCR product had the expected size and could be re-amplified using the primer s-368 and a nested primer (data not shown).

**Recombinant Agrin with the Extension Is Secreted**

To generate a cDNA construct that covers the entire coding sequence of chick agrin, RT-PCR was performed with the sense oligonucleotide EcoRI_s-289, spanning the putative initiator methionine, and the antisense oligonucleotide as232 (see Fig. 3). The PCR product was then fused to the cDNA encoding cFullps. The recombinant protein encoded by this construct was called cAgrin (Fig. 1). To show that cAgrin was targeted to the secretory pathway of COS cells, we investigated the endo-1,3-N-acetylglucosaminidase (endo H) sensitivity of the newly synthesized protein. Endo H removes N-linked carbohydrates that become attached to the protein backbone in the ER. A decrease of the apparent molecular mass upon endo H treatment indicates that the protein is targeted to the ER. To see this shift more clearly, we transfected COS cells with cDNA constructs encoding only the NH$_2$-terminal half (termed cNFullps; see Fig. 1), which contains all the potential sites for N-linked glycosylation. After a pulse of [35S]methionine for 30 min, the cells were lysed and agrin-like proteins were immunoprecipitated. After incubation with endo H, agrin was analyzed on SDS-PAGE. As shown in Fig. 4, the construct with the signal sequence of hemagglutinin, cNFullps, is sensitive to endo H, indicating that the proteins had reached the lumen of the ER where they became N-glycosylated. In contrast, cNFullps is only weakly expressed and it is not sensitive to endo H (Fig. 4). The relative molecular mass of cNagrin is also shifted by endo H digestion (Fig. 4). These results show that the new NH$_2$-
Chick agrin containing the novel NH₂-terminal sequence is targeted to the endoplasmic reticulum. COS-7 cells that were transiently transfected with cDNA constructs encoding NH₂-terminal agrin fragments (see Fig. 1) were pulsed with [³⁵S]methionine for 30 min. Recombinant agrin was then isolated from the COS cell lysates by immunoprecipitation with anti-agrin antisera. Immunoprecipitates were incubated in buffer alone (−) or in the presence of endo H (+) to determine the presence of N-linked carbohydrates. Proteins were analyzed on a 4–10% SDS-PAGE. The apparent molecular mass of both cNFULLhs and cNAgrin is decreased upon Endo H treatment, indicating that the proteins had reached the lumen of the ER. In contrast, cNFULLp~ is only weakly expressed and is not sensitive to endo H. The higher molecular mass of cNAgrin compared to both, cNFULLhs and cNFULLp~ is consistent with its 123-amino acid-long extension.

Molecular weights in kDa of standard proteins are shown.

terminal extension of cAgrin targets biosynthesis to the ER. In addition, they demonstrate that agrin encoded by the previously proposed full-length chick agrin cDNA is not targeted to the ER confirming that its NHE-terminal sequence does not serve as signal sequence.

**Glycosaminoglycan Side Chain Attachment**

Several lines of evidence indicate that agrin is a HSPG: (a) Recombinant cFULLhs with a calculated molecular mass of 206 kDa, when synthesized by COS-7 and 293 cells, has an apparent molecular mass between 400 and 600 kDa (Geschmann et al., 1995; Fig. 2 and 7). (b) Cloning of a previously characterized major HSPG from chick brain (Halfter, 1993) has revealed that this HSPG is chick agrin (Tsen et al., 1995a). (c) Tryptic peptides derived from a HSPG isolated from bovine renal tubular basement membranes are highly homologous to chick agrin (Hagen et al., 1993; see below).

When we tested the sensitivity of cFULLhs to chondroitinase ABC and keratanase, its molecular weight was not decreased (data not shown), while heparitinase treatment decreased the apparent molecular mass (Fig. 5). Agrin contains at least six potential GAG chain attachment sites with the consensus sequence E/DGSGE/D or SGXG (Bourdon et al., 1987; Zimmermann and Ruoslahti, 1989). Three sites are located in the 95-kDa, COOH-terminal part of agrin and three sites, whose positions are conserved across species, are within the NH₂-terminal half (see Fig. 1). To see which of these attachment sites in fact carry GAG side chains, COS cells were transiently transfected with cDNA constructs, encoding cAgrin, cFULLhs, and the NH₂-terminal half (cNFULLhs) or the COOH-terminal half (c95; see Fig. 1). As expected, incubation with heparitinase resulted in the reduction of the apparent molecular mass of both cFULLhs and cAgrin (Fig. 5). While cFULLhs, the NH₂-terminal portion of cFULLhs, was deglycosylated with heparitinase, the COOH-terminal fragment was not sensitive to the enzyme (Fig. 5). From this we conclude that HS-GAG chains are only attached to the NH₂-terminal portion of agrin.

**Agrin in Tissue Homogenates**

Independent evidence confirms that agrin is a HSPG in both mammals and birds. Hagen et al. (1993) isolated a HSPG from basement membranes of bovine kidney and subsequently determined amino acid sequences of three tryptic peptides. As shown in Fig. 6, two peptides are similar to both chick and rat agrin. Peptides 2 and 3 are homologous to sequences between the first and the second follistatin (PSTI)-like domain and to sequences preceding the first follistatin (PSTI)-like domain of agrin, respectively. Peptide 1 maps however to the extension presented here. These data provide additional evidence that the novel sequences described in this report are indeed contained in mature agrin and that they are conserved be-
Figure 6. Amino acids of the novel 5' sequence are conserved between chick and bovine agrin. Alignment of three tryptic peptides (single letter code) of a HSPG from bovine renal tubular basement membranes (BTBM; Hagen et al., 1993) with rat and chick agrin sequences. Conserved amino acids are shaded. Note that peptides 2 and 3 are homologous to both chick and rat agrin. Peptide 1 maps to a region encoded by the novel 5' extension (box) and hence is homologous only to chick. The position of the initiator methionine of Tsim et al. (1992) is indicated (A).

The calculated molecular mass of the protein backbone of chick agrin that includes the extension is 225 kD and agrin synthesized by transfected cells has an apparent molecular mass of more than 400 kD. However, agrin-like proteins purified from basal lamina extracts of different tissues and several species appear in two discrete doublets with molecular mass of 150/135 and 95/70 kD (Nitkin et al., 1987; Godfrey et al., 1988a,b; Godfrey, 1991). Similarly, Hagen et al. (1993) reported a calculated relative molecular mass of the core protein after heparitinase treatment of 145 and 125 kD. Hence, agrin seems to be degraded during purification. To examine whether the high molecular weight form is found in vivo and to determine whether proteolytic fragments of the reported size are also detected in freshly homogenized tissue, we performed Western blot analyses. To prevent proteolytic degradation, tissues were homogenized on ice in the presence of several protease inhibitors. Tissues examined were embryonic day 14 (E14) brain (Fig. 7, lanes B), spinal cord (Fig. 7, lanes S), hindlimb muscle (Fig. 7, lanes M), E10 vitreous humor (Fig. 7, lanes V), and E15 liver (Fig. 7, lanes L). Conditioned medium of 293 cells that had been stably transfected with cDNA encoding cFullhs provided a positive control (Fig. 7, lanes C). Agrin-like proteins were detected with antisera raised either against the 95 kD, COOH-terminal part of agrin (Fig. 7 A), or against cFullhs (Fig. 7 B). No immunoreactivity was observed with the corresponding preimmune sera (data not shown) except in the vitreous humor where the preimmune serum of the anti-c95 antiserum recognized two distinct bands of ~100 and 125 kD (* in Fig. 7 A, lane V). Agrin-like immunoreactivity was associated with a high molecular weight band in all tissues examined except liver (Fig. 7). The high molecular weight agrin was most abundant in the vitreous humor, followed by brain, muscle and spinal cord. It was seen more clearly with antibodies raised against cFullhs that carries HS-GAG chains (Fig. 7 B). Since this band was also detected with antibodies against c95 (Fig. 7 A), we conclude that the antigen recognized by anti-cFullhs antibo-

ies is agrin. With both antisera, several fragments were seen in most tissues. The detected fragments were sometimes of different size, which may be due to the divergent specificities of the antisera. In liver, only the anti-c95 antibodies recognized a single band with a relative molecular mass of ~125 kD (Fig. 7 A, lane L). We have not characterized this immunoreactive protein further. However, a similar band has been observed in adult rat liver using an antiserum raised against a fusion protein encoding sequences just upstream of the 95-kD fragment (Rupp et al., 1991). The distribution into doublets of 150/135 and 95/70 kD as observed by others (Nitkin et al., 1987; Godfrey et al., 1988a,b; Godfrey, 1991) was not seen consistently in all the tissues. Immunoreactive protein bands of ~150 and 90 kD were evident only in the vitreous humor (arrows in lanes V). At embryonic day 10, proteins secreted from retinal cells accumulate in this body fluid (e.g., Ruegg et al., 1989; Halfter, 1993). The presence of both high molecular weight band and agrin fragments therefore suggests that a fraction of agrin can be proteolytically processed in vivo.

Alternative Splicing of Agrin mRNA at a Novel Site
In our RACE experiments on mRNA isolated from E5 to E6 chick spinal cord, we obtained two populations of PCR
Figure 8. Tissue distribution of the new splice variant. (A) Nucleotide and deduced amino acid sequence of cDNAs around the novel splice site. Numbering is according to Fig. 3. (B) Autoradiogram of PCR products from randomly reverse transcribed mRNA of E5-E6 chick spinal cord (spinal cord) and of cultured chick myotubes (muscle). PCR was performed using oligonucleotides flanking the novel splice site. Control PCR using no template (1), cDNA with 21 bp (2), cDNA without 21 bp (3), or a mixture of both cDNAs (4). Motor neurons of total spinal cord (T) were enriched by panning spinal cord cells with mAb SC1, by fractionating them on a metrizamide gradient (M), or by the combination of both methods (S+M). Finally, both methods were applied sequentially to eliminate small, SC1-positive cells (Tanaka and Obata, 1984). With each enrichment the relative content of agrin mRNA containing the 21-bp insert increased (Fig. 8, B and C, spinal cord, lane T), while conversely, muscle cells synthesize mainly mRNA lacking this insert (Fig. 8 B and C, muscle). Since we were interested in the distribution of the splice variants in motor neurons, we either panned freshly dissociated spinal cord cells on petri dishes that had been coated with mAb SC1, which recognizes an antigen expressed by motor neurons and floor plate cells (Tanaka and Obata, 1984; Bloch-Gallego et al., 1991), or separated them from other cells according to their size on a metrizamide gradient (Dohrmann et al., 1986). Finally, the last column depicts PCR products from chick myotubes grown in culture. (C) Quantification of the relative abundance of agrin mRNA including or lacking the 21-bp insert. For each lane S3P counts were measured with a PhosphorImager and the relative abundance of each PCR product was determined. Values are means ± SEM (n = 3). In E5/E6 spinal cord 82% of agrin mRNA includes the 21-bp insert (gray) and 18% lacks it. In muscle cells (lane muscle) the ratio between the two transcripts is reversed (19% + 21 bp; 81% – 21 bp). After enrichment for motor neurons by SC1 panning (lane S), metrizamide gradient centrifugation (lane M) and a combination of both procedures (lane S+M) the amount of mRNA including the 21 bp is increased to 97%, suggesting that motor neurons express specifically this agrin transcript.

Biding of Recombinant Agrin to Extracellular Matrix Requires the NH2-terminal Extension

Agrin was originally purified from basal lamina extracts of the electric organ of *Torpedo californica* (Nitkin et al., 1987). Moreover, agrin-like immunoreactivity remains associated with synaptic basal lamina for at least three weeks after degeneration of the cells at the NMJ (Reist et al., 1987).
Figure 9. Binding of recombinant agrin to extracellular matrix requires the NH₂-terminal extension. Double immunofluorescence staining of transiently transfected COS-7 cells grown on Matrigel™. The constructs and agrin isoforms used are indicated on the left of each row (see Fig. 1 for structure of the constructs). Extracellular agrin (nonpermeabilized) was visualized by incubating COS cells with antibodies directed against the 95 kD, COOH-terminal fragment at 4°C for 30 min and an appropriate, fluorescein-conjugated secondary antibody. To identify transfected cells, cells were permeabilized and agrin was subsequently visualized with mAb 5B1 and a rhodamine-conjugated secondary antibody (permeabilized). No deposition of agrin in Matrigel™ is observed with c95A4B8 and cFullhsA4B8 (A and B), while agrin is deposited in a gradient with both constructs that include the extension, cAgrin_0A4B8 and cAgrinTA4B8 (C and D). Bar, 40 μm.
Hence, in addition to inducing AChR aggregates, agrin also must bind to basal lamina (McMahen, 1990). To localize the region in agrin that confers its binding to ECM and to test whether alternative mRNA splicing affects the binding, transiently transfected COS cells were grown overnight on Matrigel™, a solubilized basement membrane extracted from the Engelbreth-Holm-Swarm (EHS) mouse sarcoma (Kleinman et al., 1982). Deposition of agrin isoforms was assayed by extracellular staining with antibodies directed against the 95-kD COOH-terminal fragment of agrin. COS cells were then permeabilized and stained with mAb 5B1 (Reist et al., 1987) to identify transfected cells. No deposits in Matrigel™ are detected with cells that express the 95-kD COOH-terminal fragment of agrin (Fig. 9 A) although intracellular staining shows that c95AABB is synthesized (Fig. 9 A'). The cDNA construct encoding recombinant agrin, cFullhSAABB, that does not include the 5' extension shows the same staining pattern as C95A4B8 (Fig. 9 A'). In contrast, cAgrinTAABB, a splice variant that includes the 7-amino acid insert at the new site, is efficiently deposited in the Matrigel™ predominantly in the vicinity of transfected COS cells (Fig. 9 C). Exclusion of the 7-amino acid insert at the new site did not change the staining pattern on Matrigel™ (Fig. 9 D, but see below). Similarly, no effect of splicing at sites A and B was observed (data not shown). No staining was observed on the cell surface of COS cells. This contrasts results by Campanelli et al. (1991) where full-length rat agrin remained associated with the cell surface. The close association of the COS cells with Matrigel™ also prevented anti-agrin antibodies from staining extracellular agrin at the basal site of the transfected cell. After permeabilization, the staining with mAb 5B1 was much stronger when COS cells had been transfected with constructs including the extension than with cDNAs lacking it (compare Fig. 9, A' and B', with C' and D'), suggesting that agrin was also deposited on the basal site of the COS cells. The deposition of recombinant agrin was not seen on uncoated or gelatin-coated tissue culture dishes (data not shown), indicating that the binding occurs to specific components of the Matrigel™.

To quantify the binding of recombinant agrin to Matrigel™ and to see whether alternative splicing at the novel site may influence this binding, we next compared the concentration of recombinant protein in the conditioned medium of transiently transfected COS cells that were either grown on normal tissue culture dishes or on Matrigel™-coated dishes. As shown in Table I, no significant difference in the concentration of recombinant agrin on either substrate was seen with c95AABB and cFullhAABB. In contrast, the concentration of recombinant agrin that remained soluble on Matrigel™-coated dishes was less than 50% with both constructs that include the 5' extension. Moreover, in two independent experiments, considerably more cAgrin0AABB was bound to Matrigel™ than cAgrinTAABB. Hence, alternative splicing at the novel site may influence binding of agrin to extracellular matrix.

The NH₂-terminal Extension Influences Size But Not Extent of Agrin-induced AChR Clusters

AChR-aggregating activity of agrin is strongly regulated by the alternatively spliced inserts A and B (Ruegg et al., 1992; Ferns et al., 1992, 1993; Gesemann et al., 1995). The dependence of activity on these inserts was observed with 95-kD COOH-terminal fragments and those that included the NH₂-terminal part of agrin (cFullhA; Gesemann et al., 1995). To test whether recombinant agrin isoforms with the extension were also active in AChR aggregation in an isoform-specific manner we incubated cultured chick myotubes for 16 h with an excess of recombinant protein. AChRs were visualized using rhodamine-labeled α-bungarotoxin. No AChR clustering was detected after incubation with conditioned medium of mock-transfected cells (Fig. 10 A). Agrin isoforms that lacked the A and the B inserts also did not induce receptor clustering (cFullhA0B0, Fig. 10 C; cAgrinTA0B0, Fig. 10 E; cAgrin0A0B0, Fig. 10 G). As shown previously, clustering of AChRs on Matrigel™ was clearly induced by 500 pM of C95AABB (Fig. 10 B) and cFullhAABB (Fig. 10 D). Agrin isoforms that included the extension were also active in inducing AChR clusters (cAgrinTAABB, Fig. 10 F; cAgrin0AABB, Fig. 10 H). AChR clusters induced by these constructs were however smaller and more frequent (Fig. 10, F and H) than those induced by agrin fragments (Fig. 10, B and D). The average size of AChR clusters induced by cFullhAABB in 20 myotube segments was 26.2 ± 2.3 μm² (mean ± SEM), while induction by cAgrinTAABB yielded clusters with an average size of 10.8 ± 0.6 μm² (mean ± SEM). To determine whether the change in size was based on decreased AChR-aggregating activity, we measured the area of AChR clusters with a computerized image analysis system. As shown in Fig. 10 I, there was no significant difference between active agrin isoforms that have the extension and their corresponding fragments. The distribution of AChR clusters (aligned versus dispersed; compare Fig. 10, F and H) sometimes varied between cAgrinTAABB and cAgrin0AABB. However, we could not reproduce this phenomenon consistently. In summary, these results show that AChR-aggregating activity of complete chick agrin depends on splice inserts at sites A and B and that the most NH₂-terminal sequence of agrin alters the size of the induced clusters but not the overall extent of the induction.

Table I. Relative Concentration of Recombinant Agrin in the Medium of Transiently Transfected COS Cells Plated on Matrigel™-coated Dishes

<table>
<thead>
<tr>
<th>Protein</th>
<th>Plastic</th>
<th>Matrigel</th>
</tr>
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<tbody>
<tr>
<td>c95AABB</td>
<td>100.0 ± 3.4</td>
<td>99.2 ± 3.2</td>
</tr>
<tr>
<td>cFullhAABB</td>
<td>100.0 ± 4.8</td>
<td>100.7 ± 2.4</td>
</tr>
<tr>
<td>cAgrinTAABB</td>
<td>100.0 ± 1.7</td>
<td>45.2 ± 1.2</td>
</tr>
<tr>
<td>cAgrin0AABB</td>
<td>100.0 ± 3.4</td>
<td>22.3 ± 1.8</td>
</tr>
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</table>

COS cells were transfected with cDNAs encoding the indicated agrin constructs. One day after transfection, cells were replated in the same density on regular tissue culture dishes (plastic) or on Matrigel™-coated dishes (Matrigel). After 18 h, conditioned medium was collected and the concentration of recombinant agrin was determined by ELISA as described in Materials and Methods. Concentration of agrin with cells grown on regular tissue culture dishes was set as 100%. Values are means ± SEM of two independent experiments with triplicate cultures for each condition. The concentration of c95AABB and cFullhAABB does not change, indicating that these constructs do not bind to Matrigel™. In contrast, both cAgrinTAABB and cAgrin0AABB bind to Matrigel™. Note also the difference in binding to Matrigel™ between agrin constructs with different inserts at the novel splice site.
Fig. 10. Agrin with the NH$_2$-terminal extension induces smaller AChR clusters. Fluorescence micrographs of cultured chick myotubes labeled with rhodamine-α-bungarotoxin. Myotubes were incubated for 16 h with conditioned medium of COS cells, that had been mock-transfected (A) or with cDNAs encoding C95A4BS (B), cFullhSA0B0 (C), cFullhSA4B8 (D), cAgrin7n0Bo (E), cAgrin7A4B8 (F), cAgrin0g0B0 (G) and cAgrin0A4BS (H). The concentration of recombinant agrin was always 500 pM. Only agrin$_{7A4BS}$ isoforms induced AChR aggregates (B, D, F, and H), whereas agrin$_{0G0B0}$ isoforms were inactive (C, E, and G). Active agrin including the extension (cAgrin$_{7A4BS}$, F; cAgrin$_{0A4BS}$, H) induced smaller clusters. Bar, 50 μm. (I) Quantification of AChR aggregation. Due to the smaller size of AChR aggregates with the longer axis ≥2 μm are included. Activity is expressed as percentage area of AChR clusters. Each data point represents mean ± SEM of duplicate cultures, where 10 myotube segments were counted in each. The control corresponds to myotubes incubated with conditioned medium of mock-transfected COS cells. Results of one representative experiment are shown.

Discussion

Differences between Chick and Rat Agrin

We report here the heterologous expression of chick agrin comprising all the functional properties that have been postulated for endogenous agrin (McMahan, 1990). This was achieved by ligating the novel 5' end sequences to the cDNAs described by Tsim et al. (1992). When grown on noncoated tissue culture dishes, two different cell lines transfected with this construct secrete the recombinant protein into the medium (Figs. 5 and 7). This contrasts the results of Campanelli et al. (1991) where recombinant full-length rat agrin remained associated with the cell surface of transfected COS and CHO cells. Comparison of the protein sequences of chick and rat agrin suggests that the difference resides in the NH$_2$ terminus. Although chick and rat agrin are highly conserved (60% identity), no homology is found in the previously postulated putative signal peptides (Rupp et al., 1991; Tsim et al., 1992). The 80% identity of the 15-amino acid peptide of bovine kidney HSPG with a corresponding stretch in the 5' extended region in chick (Fig. 6) shows that the formerly postulated signal sequence of chick agrin is in fact part of the mature protein. This, together with our finding that the extension is important for the binding of agrin to ECM, makes it very likely that rat agrin also contains a homologous stretch. The published NH$_2$-terminal sequence of rat agrin (Rupp et al., 1991) is however completely different to our extension and to the bovine peptide and may therefore be derived from a rat agrin mRNA that encodes an alternative 5' sequence.

The calculated relative molecular mass of chick agrin is 225 kD. When expressed in COS and 293 cells, recombinant chick agrin has a relative molecular mass of 400 to 600 kD (Fig. 5) due to the attachment of carbohydrates. Similarly, agrin-like immunoreactivity in tissue homogenates has the same apparent molecular mass (Tsen et al., 1995a; Fig. 7). Full-length rat agrin expressed by COS cells and agrin-like protein in rat tissue homogenates has however an apparent molecular mass of ~200 kD (Rupp et al., 1991; Campanelli et al., 1991). In addition, ray and chick agrin-like proteins in earlier studies appear as distinct protein bands between 150 and 70 kD (Nitkin et al., 1987; Godfrey et al., 1988a,b; Godfrey, 1991). The difference in the molecular mass observed earlier and our results suggests that the previously described immunoreactive proteins result from degradation. Although we observed several smaller protein bands in our Western blots, the presence of agrin-like protein at a high molecular mass indicates that agrin in vivo can be released from cells without proteolytic processing. One possible explanation for the absence of the high molecular mass bands in earlier studies may be that the antibodies used were directed either against the nonglycosylated, COOH-terminal part of agrin or against fusion proteins expressed in Escherichia coli. As shown in Fig. 7, such antibodies have a lower avidity for the highly glycosylated agrin.

Possible Role of Agrin as a Heparan Sulfate Proteoglycan

We show here that sites within the NH$_2$-terminal half of agrin carry N-linked carbohydrates and HS-GAG side chains (Figs. 4 and 5), and that the COOH-terminal end lacks these glycosylations. The attachment of GAG side chains is not necessary for agrin's activity to induce AChR clusters (Gesemann et al., 1995). Several lines of evidence suggest however that HSPGs expressed by the muscle cells...
do play a role (Wallace, 1991; Ferns et al., 1992, 1993; Gordon et al., 1993). We have therefore recently postulated that agrin may bind to HSPGs expressed in muscle cells and that these HSPGs may help to present agrin to its postulated signal-transducing receptor (Gesemann et al., 1995). Because agrin expressed by muscle cells is a HSPG (Fig. 7) and because it accumulates at AChR clusters (Lieth et al., 1992), muscle agrin itself may be such a helper protein. Indeed, the 95-kD fragment of the active isoform agrin_AABB binds weakly to the longer, inactive recombinant cFullba08 that carries HS-GAG chains (Gesemann, M., A. J. Denzer, and M. A. Ruegg, unpublished observation).

Agrin's role as a HSPG may be similar to those described for other HSPGs. They are believed to play a role in the scaffold for the attachment of various ECM components and to modulate interactions with cells during morphogenesis (for a review see Timpl, 1993). The presence of HS-GAG chains together with a similar arrangement of structural motifs makes agrin a homologue to perlecan (Noonan et al., 1991). Perlecan is a low affinity, auxiliary receptor for basic fibroblast growth factor, a potent growth factor involved in angiogenesis, and this interaction is dependent on its HS-GAG chains (Aviezer et al., 1995). Since agrin is highly expressed during the development of the nervous system (McMahan et al., 1992; O'Connor et al., 1994; Ma et al., 1994), it may have functions similar to those described for perlecan (reviewed in Iozzo et al., 1994). Recent evidence suggests that agrin, which is also expressed in axonal tracts of the developing chick visual system (Halter, 1993), modulates the homophilic binding of the cell adhesion molecule N-CAM. This function depends on the presence of HS-GAG chains in agrin and hence is constricted to its NH2-terminal half (Burg et al., 1995).

**Binding of Agrin to Extracellular Matrix and AChR-aggregating Activity**

We show that binding of agrin to ECM, detected as a deposition in Matrigel™, requires the novel NH2-terminal sequence (Fig. 9). This preparation of solubilized basement membrane contains mainly laminin-1 (EHS laminin), collagen IV, HSPGs, and nidogen (Kleinman et al., 1982). Hence, any of these proteins may be a binding partner for agrin. Deposition of agrin-like immunoreactivity on a similar substrate has also been observed with spinal cord neurons of Xenopus laevis (Cohen et al., 1994), supporting our conclusion that the construct we describe here codes for the entire chick agrin.

AChR aggregates induced by this construct are smaller than those induced by agrin fragments. Small AChR aggregates (<4 μm²) are formed within two hours when agrin isolated from Torpedo californica is added to medium bathing chick myotubes. Their size increases with time and reaches a maximum after 16 to 20 h (Wallace, 1988). This increase in size is, at least in part, based on the fusion of the small aggregates (Wallace, 1994). It is conceivable that Torpedo agrin bound to its postulated signal-transducing receptor is still mobile in the muscle cell membrane. Therefore, small clusters can merge to form bigger clusters. Because the partially purified agrin from Torpedo californica used in these studies contains mainly COOH-terminal fragments (Nitkin et al., 1987), it does not bind to ECM. We propose that recombinant agrin including the extension, which additionally binds to ECM, immobilizes the small clusters and prevents them from merging, or, alternatively, prolongs the time needed for the fusion.

**Alternative mRNA Splicing at the Novel Site Is Regulated**

We also describe a site of 21-bp length that undergoes alternative mRNA splicing. As reported for sites A and B, the splice variants are expressed differentially by presynaptic motor neurons and postsynaptic muscle cells (Fig. 8). As embryonic day 5 to 6 motor neurons (HH stage 26 and 27; Hamburger and Hamilton, 1951) extend their axons to muscle, AChRs begin to cluster in response to these ingrowing axons (Dahm and Landmesser, 1991). This is also the time when motor neurons express high levels of SCI (also known as DM-GRASP or BEN; Pourquie et al., 1990; Burns et al., 1991). In the purest preparation of motor neurons, 97% of agrin mRNA contains the 21-bp insert. In this cell fraction more than 95% are motor neurons (Bloch-Gallego et al., 1991). Hence, the 3% of agrin mRNA lacking the 21-bp insert most likely represents contamination by nonmotor neurons and we conclude that, early in synaptogenesis, chick motor neurons synthesize agrin mRNA that contains the 21-bp insert. On the other hand, in postsynaptic cells, such as E12 (HH stage 38; Hamburger and Hamilton, 1951) hindlimb muscle and myotubes grown in culture, the majority of agrin mRNA lacks the 21-bp insert. E10 dorsal root ganglia also synthesize predominantly agrin mRNA containing the 21-bp insert and Schwann cells of the sciatic nerve have a pattern similar to muscle cells (Denzer, A. J., and M. A. Ruegg, unpublished data). This suggests that neurons other than motor neurons synthesize agrin mRNA containing the 21 bp while nonneuronal cells show a preference for the splice variant lacking this insert. Indeed, Tsen et al. (1995b) have obtained similar results in the central nervous system, suggesting that nonneuronal cells express mainly agrin isoforms lacking the 21-bp insert and neurons synthesize agrin mRNA that contains the 21-bp insert. We have not closely looked at functional differences between the splice variants. Our finding that the isoforms seem to bind to Matrigel™ with different strength (Table I) suggests that this insert may regulate the interaction of agrin with ECM. We are currently aiming at identifying the binding partners of agrin in the ECM and studying in detail whether splicing at the novel site affects binding.

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