Postnatal stabilization and maturation of the postsynaptic membrane are important for development and function of the neuromuscular junction (NMJ), but the underlying mechanisms remain poorly characterized. We examined the role of Src-family kinases (SFKs) \textit{in vivo}. Electroportation of kinase-inactive Src constructs into soleus muscles of adult mice caused NMJ disassembly: acetylcholine receptor (AChR)-rich areas became fragmented; the topology of nerve terminal, AChRs, and synaptic nuclei was disturbed; and occasionally nerves started to sprout. Electroportation of kinase-overactive Src produced similar but milder effects. We studied the mechanism of SFK action using cultured src$^{-/-}$/fyn$^{-/-}$ myotubes, focusing on clustering of postsynaptic proteins, their interaction with AChRs, and AChR phosphorylation. Rapsyn and the utrophin-glycoprotein complex were recruited normally into AChR-containing clusters by agrin in src$^{-/-}$/fyn$^{-/-}$ myotubes. But after agrin withdrawal, clusters of these proteins disappeared rapidly in parallel with AChRs, revealing that SFKs are of general importance in postsynaptic stability. At the same time, AChR interaction with rapsyn and dystrobrevin and AChR phosphorylation decreased after agrin withdrawal from mutant myotubes. Unexpectedly, levels of rapsyn protein were increased in src$^{-/-}$/fyn$^{-/-}$ myotubes, whereas rapsyn–cytoskeleton interactions were unaffected. The overall cytoskeletal link of AChRs was weak but still strengthened by agrin in mutant cells, consistent with the normal formation but decreased stability of AChR clusters. These data show that correctly balanced activity of SFKs is critical in maintaining adult NMJs \textit{in vivo}. SFKs hold the postsynaptic apparatus together through stabilization of AChR–rapsyn interaction and AChR phosphorylation. In addition, SFKs control rapsyn levels and AChR-cytoskeletal linkage.

\textit{Key words:} Src; acetylcholine receptor; neuromuscular synapse; agrin; tyrosine phosphorylation; postsynaptic membrane

\section*{Introduction}

Neuromuscular junctions (NMJs) develop in a series of steps in which the postsynaptic membrane first forms by concentrating acetylcholine receptors (AChRs) and associated proteins in a flat topology. Postnatally, NMJs mature and AChRs get arranged at the crests of postjunctional folds. Concomitantly, all but one axon withdrew, paralleled by destabilization of adjacent AChRs (Sanes and Lichtman, 2001). Maturation and stabilization of AChR clusters ensure proper synaptic development, which forms the basis for nerve-evoked muscle contractibility.

Much is known about the molecular pathways that first form NMJs. Neural agrin, by activating the muscle-specific kinase (MuSK), is crucial by triggering downstream cascades (for review, see Bezakova and Ruegg, 2003; Luo et al., 2003). Central in these is rapsyn, the main AChR-anchoring protein mediating clustering (Gautam et al., 2003), mediates agrin-induced phosphorylation of the AChR $\beta$ and $\delta$ subunits (Mittaud et al., 2001), and links the receptor to $\beta$-dystroglycan, a component of the postsynaptic utrophin-glycoprotein complex (UGC) (Cartaud et al., 1998; Bartoli et al., 2001). In clustering, AChRs become immobilized and less detergent extractable, both in agrin-treated myotubes (Prives et al., 1982; Stya and Axelrod, 1983; Podleski and Salpeter, 1988) and developing NMJs (Dennis, 1981; Slater, 1982). The players in this cytoskeletal link remain uncertain. Agrin-induced phosphorylation of AChR $\beta$ is involved (Borges and Ferns, 2001) and can occur through Abl- and Src-family kinases (SFKs) (Finn et al., 2003; Mittaud et al., 2004).

Much less is known about the mechanisms that mature NMJs and stabilize AChR clusters postnatally. Although MuSK is required (Kong et al., 2004), some of these pathways may not be essential in initial NMJ formation (Willmann and Fuhrer, 2002), as illustrated by mice lacking utrophin and dystrophin or the UGC components $\alpha$-dystrobrevin or dystroglycan (Grady et al.,...
1997; Grady et al., 2000; Jacobson et al., 2001). In these mice, NMJs form but fail to mature properly. In α-dystrobrevin−/− mice, AChR clusters are normal at birth but increasingly frag-
ment postnatally. Similarly, in cultured α-dystrobrevin−/− myo-
tubes, agrin induces normal AChR clustering, but these clusters are unstable and disperse rapidly when agrin is withdrawn from the myotubes (Grady et al., 2000). Thus, the UGC is a core player in postnatal NMJ stabilization. Additional candidates are SFKs. In src−/−/fyn−/− mice, NMJs appear normal around birth, when the animals die. In cultured src−/−/fyn−/− myotubes, agrin and laminin induce normal AChR aggregation, but the clusters dis-
perse rapidly after withdrawal of these factors (Smith et al., 2001; Marangi et al., 2002).

To elucidate the mechanisms of synaptic stabilization, we in-
vestigated the role of SFKs in vivo. In adult myofibers expressing dominant-negative Src, AChR-rich areas were severely frag-
mented. We addressed the mode of SFK action in AChR cluster stabilization using src−/−/fyn−/− myotubes. We found that Src and Fyn maintain clusters of rapsyn and UGC components, maintain AChR–rapsyn interactions and AChR β phosphoryla-
tion, mediate AChR-cytoskeletal linkage, and control rapsyn protein levels. Our data introduce SFKs as critical players in NMJ stabilization in vivo and reveal that complex signaling pathways underlie stabilization.

Materials and Methods

Src constructs, electroporation, whole-mount preparation, and immunohis-
tochemistry. Three different mutant Src expression constructs, each contain-
ing a cytomegalovirus promoter, and an empty pLNCX control vec-
tor were used for electroporation into the soleus muscle of mice. Src-AM (Kaplan et al., 1994) was kindly provided by Dr. Pam Schwartzberg (Na-
tional Institutes of Health, Bethesda, MD), Src-K295M (Mohamed et al., 2001) was provided by Dr. Sheridan Swope (Georgetown University, 
Washington, DC), and Src-Y527F was provided by Dr. Joan Brugge (Harvard Medical School, Boston, MA). Mutant Src constructs (8 μg/μl) and green fluorescent protein (GFP) containing a nuclear localization signal (NLS-GFP; 4 μg/μl) were mixed to yield final DNA concentrations of 2:1 (Src-GFP) and were first injected extrasynaptically into the soleus muscle of adult C57BL/6 mice (3–6 months of age). The wound was closed, electrodes were mounted against the leg, and electroporation was performed as described previously (Kong et al., 2004) using an ECM 830 electroporation system (BTX, Holliston, MA). Eight pulses of 20 ms were applied at a frequency of 1 Hz with voltage set to 200 V/cm. At the site of DNA injection, electroporation efficiency is highest, and once DNA con-
structs have entered muscle fibers, they diffuse within those regions, including the synapse (Kong et al., 2004). The electroporated muscles were analyzed after 6 weeks. Muscles were dissected and injected with 2% paraformaldehyde solution for fixation. This treatment is optimal for whole-mount analysis, because it swells the muscle, widening gaps be-
tween individual muscle fibers, facilitating additional fiber dissection. Fixed tissue was teased into thin fiber bundles of 5–10 myofibers. Whole-
mount preparations were triple-labeled as described previously (Kong et al., 2004). Briefly, AChRs were stained to visualize NMJs using rhodamine-coupled α-bungarotoxin. A mixture of rabbit polyclonal an-
tibodies against neurofilament (Sigma, St. Louis, MO) and synaptophy-
sin (Dako, Glostrup, Denmark), followed by cyanine 5 (Cy5)-conjugated goat anti-rabbit or Alexa 488-coupled goat anti-rabbit antibodies, was used to visualize motorneurons and nerve endings. The green channel was re-
served for GFP. For cytoskeletal staining, Alexa 350-coupled phalloidin (Invitrogen, Eugene, OR) was used to label F-actin. Monoclonal anti-α-
tubulin antibodies (clone DM1A; Sigma), followed by Cy3-conjugated goat anti-mouse antibodies, were used to visualize synaptic tubulin rings beneath the endplate. Conventional fluorescence imaging was done using a Zeiss (Feldbach, Switzerland) Axioskop 2 microscope equipped with a Hamamatsu (Shizuoka, Japan) OrcaCam digital camera.

Immunohistochemistry on cross sections. Soleus muscle tissue prepared
and fixed for whole-mount analysis (see above) was first placed overnight in 10% sucrose at 4°C and then embedded into O.C.T. compound (Tis-
tue-Tek; Sakura Finetek, Zoeterwoude, The Netherlands). The tissue was frozen in a cooling chamber (maintained at −4°C with surrounding liquid nitrogen) containing isopentane (Fluka, Buchs, Switzerland) for 5 min. Muscles were then cut into 14 μm cryosections (at −18°C). Sections were stained with rhodamine-coupled α-bungarotoxin (α-BT) to visualize the AChRs at the NMJ. GFP signals were enhanced with polyclonal anti-GFP antibody (Invitrogen) followed by Alexa 488-
coupled goat anti-rabbit antibodies. Nuclei were visualized with 4’,6’-
diamidino-2-phenylindole (DAPI) (Hoechst nuclear stain; Invitrogen).

The fixation procedure, optimized for whole-mount analysis, ex-
panded gaps between individual myofibers, rounding off some fibers and allowing entry of other cells (see Fig. 4C,D). Nonetheless, we could un-
equivocally identify muscle fibers (resulting from diffuse GFP signal) and the positionning of nuclei within those fibers (because most of the strong nuclear GFP signals overlapped with DAPI).

Confocal microscopy, imaging, and quantitation. Confocal laser-scanning microscopy was performed using an inverted Leica (Nussloch, Germany) confocal SP2 microscope coupled to a Silicon Graphics (Mountain View, CA) Workstation. A minimum of 40 stacks per image were taken, each section at −0.3 μm thickness. These confocal stacks were shown as a maximal projection and eventually imported into three-
dimensional (3D) image-processing software, Imaris 4.1.1, for image reconstruction and rotation (see Figs. 1–4). To quantify disassembly, synaptic contacts (as visualized by the postsynaptic AChRs and the pre-
synaptic nerve terminal) were scored based on size and integrity of the dis-
assembly. Intact endplates were large “pretzel”-shaped structures (25–40 μm) and continuous along their contours. A partial disassembly was scored when the endplate was broken up into more than two main frag-
ments and was porous. In such cases, it was still possible to identify the parts of a pretzel. In a complete disassembly, the structures completely dissolved into several fragments, which were <5 μm in length. These were scored as synaptic sites because of the nerve staining and surround-
ing intact endplates in GFP-negative fibers. Statistics were performed by averaging three sets of independent electroporation experiments, with a minimum of 30 pictures taken for each condition. In all cases, the NLS-
GFP signal colocalized with the AChR stain and lay beneath the end-
plates, suggesting that the electroporation was specific to the muscle nuclei and was not an indirect effect caused by electroporation of Schwann cells. Apart from synaptic muscle nuclei, extrasynaptic nuclei were also GFP positive in most cases.

Cell culture and agrin treatment. src−/−/fyn−/− cells (clones DM15 and 
and DM11) and their corresponding wild-type cells (clones SW10 and SW5) were grown as described previously (Smith et al., 2001) in DMEM sup-
plemented with 2% fetal bovine serum (FBS). To induce clustering, 10% fetal bovine serum, 2% chick embryo extract, penicillin–streptomycin, and 20 U/ml recombinant mouse interferon-γ. In all assays, we tested both mutant and both wild-type clones, also in pairwise comparison. There were no differences between clones of the same genotype, excluding the possibil-
ity of clonal variation.

For biochemical analysis, cells were plated on matrigel-coated tissue culture dishes (Nunc, Basel, Switzerland) at 0.18 million/10 cm dish (wild type) and 0.35 million/10 cm dish (mutants). For immunocyto-
chemistry, wild-type and mutant cells were plated on matrigel-coated 
chamber slides (Nunc) at 0.02 million/2.4 ml and 0.04 million/2.4 ml, respectively. Myoblasts were grown to 80% confluency at 33°C and 5% 
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ity of clonal variation.
AChR precipitation assays and immunoblotting. To examine the association of postsynaptic proteins with the AChR, myotubes were rinsed with ice-cold PBS containing 1 mM Na-orthovanadate and 50 mM NaF and extracted at 4°C in lysis buffer. The lysis buffer contained 1% NP-40 and an excess of protease and phosphatase inhibitors as described previously (Fuhrer et al., 1999). Lysates were processed and AChRs precipitated using biotinylated α-BT as detailed previously (Tox-P) (Mittaud et al., 2003). As a control, an excess of free α-BT (+T) was added to some lysates to judge the specificity of protein association with the AChR. Precipitates were subjected to SDS-PAGE and immunoblotting.

To quantify proteins in total cell extracts, parts of lysates were processed, without precipitation, in parallel by SDS-PAGE and immunoblotting. To ensure proper loading, protein concentrations of total lysates were done before loading using a standard BCA protein assay kit (Pierce, Rockford, IL), and amounts of each protein were normalized to the β-subunit of the AChR on the immunoblot.

In Western blots, α-dystrobrevin-2 was visualized using rabbit polyclonal B1CT-FP antibodies (gift from Dr. Derek J. Blake, University of Oxford, Oxford, UK). Conditions and antibodies to detect rapsyn, phosphorysine, AChR α and β subunit, utrophin, and MuSK were as detailed previously (Marangi et al., 2001; Moransard et al., 2003). Quantitations of the immunoblots were done by scanning exposed films containing gray, non-saturated signals with a computerized densitometer (Scantouch 210; Nikon, Tokyo, Japan) and using the NIH Image J 1.29X software. Experiments were repeated at least five times to obtain consistent results.

AChR extractability assay. We used a modified version of a sequential extraction procedure detailed previously (Borges and Ferns, 2001; Moransard et al., 2003). All steps were performed on ice if not specified otherwise. Two- to 3-d-old myotubes grown in 10 cm dishes were washed briefly with ice-cold PBS (+ 1 mM Na-orthovanadate) before adding 1 ml of lysis buffer (30 mM triethanolamine, pH 7.5, 50 mM NaCl, 5 mM EDTA, 5 mM EGTA, 50 mM NaF, 1 mM Na-orthovanadate, 1 mM benzamidine, 1 mM N-ethylmaleimide, 1 mM Na-tetradethionate, 50 μM phenylarsine oxide, 10 mM p-nitrophenylphosphate, 25 μg/ml aprotinin and leupeptin, 1 mM PMSF) containing a final concentration of Triton X-100 ranging from 0.03 to 0.09%, according to the conditions tested. The exact time for the first extraction was 21 min; a timer was started just after the addition of lysis buffer to the first plate. Cells were scraped from each plate and the extracts were homogenized by pipetting up and down 10 times and then transferred to tubes that were rotated for the rest of the time. Lysates were centrifuged at 14,000 rpm and 4°C in a table eppendorf centrifuge, and supernatants were transferred to fresh tubes and called “first extraction.” pellets were resuspended in 1 ml of lysis buffer containing 1% Triton X-100 for a total extraction time of exactly 15 min. Again, the timer was started after the first resuspension, and tubes were rotated together at 4°C for the rest of the time and finally centrifuged under the same conditions. Supernatants were transferred to fresh tubes and called “second extraction.” Both series of tubes were then subjected to AChR-precipitation with biotinylated α-BT, followed by immunoblotting for AChR α as described above.

Immunochemical staining procedures and quantitation of clusters. Cultured myotubes were grown in matril-coated multiwell chamber slides. Stainings for AChR, α-dystrobrevin-1, utrophin, rapsyn, α-dystroglycan, syntrophin isoforms, and phosphorysine were all done exactly as described previously (Marangi et al., 2001; Moransard et al., 2003). Myotubes were examined at 400X magnification in both rhodamine and fluorescein channels with a fluorescence microscope (Axioskop II; Zeiss). Representative pictures were taken and processed with a cooled digital camera (Orcacam; Hamamatsu).

To quantitate clusters, a total of 20 representative pictures per condition were taken from several chamber slides. Clusters of AChR and the respective postsynaptic proteins were counted as done previously (Marangi et al., 2001), based on signal intensity (clearly distinguishable from the diffuse background) and the length of the cluster being at least 10 μm. Numbers of clusters per myotubes were calculated independently for the AChR and the postsynaptic marker and averaged. Co-localization of clusters were calculated as the percentage of number of AChR clusters containing the postsynaptic marker protein. Staining experiments were repeated several times to ensure reproducibility.

Results

Role of SFKs: correctly balanced kinase activity is required to maintain adult NMJs in vivo

We addressed the role of SFKs in NMJ stabilization using an in vivo electroporation paradigm in soleus muscle of adult mice. This method allows one to import plasmids into synaptic and extrasynaptic regions of individual myofibers (Kong et al., 2004). Successfully electroporated fibers are identified by nuclear GFP staining, because a GFP construct containing a nuclear localization signal is coelectroporated (Kong et al., 2004).

We first used a kinase-inactive Src expression construct, Src-AM, in this approach. Src-AM harbors two mutations, K295M and Y527F, which inactivate the kinase activity and the inhibitory C-terminal phosphorylation site, respectively. The resulting kinase-dead molecule preferentially adopts an open conformation and acts strongly in a dominant-negative way (Kaplan et al., 1994; Thomas and Brugge, 1997). Such dominant-negative Src constructs interfere with many members of the Src family (e.g., Src, Fyn, and Yes), not just Src itself. The constructs therefore lead to reduction of cellular SFK function (Tsamale-Stein et al., 1993; Roche et al., 1995).

After electroporation of plasmids expressing short-interfering RNA against MuSK, postsynaptic AChR clusters are normal after 2 weeks but disassembled after 6 weeks (Kong et al., 2004). Apparently, adult NMJs are stabilized, presumably through multiple protein interactions and the cytoskeleton, such that it takes 6 weeks to see disassembly when a critical kinase (MuSK) is inactivated. Because SFKs are also tyrosine kinases, can interact with MuSK, and can act within the MuSK signaling pathway (Mohamed et al., 2001; Mittaud et al., 2004), we concentrated our analysis to 6 weeks after electroporation of Src-AM. Whole-mount preparations of myofibers were subjected to α-BT- rhodamine, neurofilament, and synaptophysin staining (Fig. 1, blue) and fluorescence microscopy. In GFP-positive fibers (expressing Src-AM), AChR clusters often appeared partially disassembled: the typical pretzel shapes were disturbed in that they did not form one single continuous structure but contained substan- tial holes, fragmenting pretzels often into two or more subregions (Fig. 1A, middle). More dramatically, in some GFP-positive fibers, NMJs were completely disassembled, so that no pretzels were observed at all but only small fragments of AChR-clusters (Fig. 1A, right). Confocal imaging confirmed these results and allowed, in three-dimensional reconstruction, to better visualize the disassembly of AChR clusters in fibers expressing GFP and Src-AM (Fig. 1B). We again observed several degrees of fragmentation, ranging from partial (Fig. 1B, middle two columns) to complete disassembly (Fig. 1B, right). As controls, we used GFP-negative fibers in Src-AM experiments (Fig. 1A) or an empty expression vector lacking the Src-AM insert (Fig. 1B). In both cases, NMJs appeared mostly intact.

Rotation of 3D reconstructions using Imaris software revealed the typical architecture of intact NMJs in muscles electroporated with empty control vector: GFP-labeled synaptic nuclei were properly clustered underneath the AChR pretzel and AChRs mostly underneath the nerve (Figs. 1B, 2A–E). This topology was disturbed in fragmented NMJs in fibers expressing Src-AM. Here, remnants of AChR pretzels were in the same focal plane as
the nerve with no preferential labeling underneath it (confirmed by analysis of single confocal stacks; data not shown), such that the nerve was equally visible from a top view and a view from inside the muscle (Figs. 1 B, 2 F–J). In addition, clusters of synaptic nuclei became dispersed after the fragmentation of the AChR pretzels. In fragmented NMJs, we occasionally observed nerve labeling that resembled sprouting (Fig. 1 B, middle Src-AM column). We quantitated the proportion of normal, partially, and completely disassembled AChR clusters for Src-AM-expressing and control fibers. In the control, most endplates contained intact AChR pretzels, some scored as partially disassembled clusters, but we observed almost no completely disassembled cases (Fig. 2 K). In Src-AM-expressing fibers, the proportion of partially and completely disassembled endplates was much higher (Fig. 2 K).

We repeated these experiments using another Src mutant, a construct that carried only a single mutation, K295M. This is the classic kinase-inactive, dominant-negative Src form used most often to investigate the involvement of SFKs in signaling pathways (Kaplan et al., 1994; Roche et al., 1995; Thomas and Brugge, 1997). We observed results very similar to those with Src-AM (Fig. 2 L). Together, these data demonstrate that normal SFK activity is a requirement to maintain the proper structure of adult NMJs in vivo.

In most cells, SFK activation is under tight regulation by a variety of extracellular signals and intracellular protein interactions (Thomas and Brugge, 1997). Experimentally, reducing or increasing SFK activity can produce changes in downstream signaling pathways (Thomas et al., 1995; Brandt et al., 2002; Kilarski et al., 2003). We therefore addressed whether increased SFK activity, leading to gain of SFK function, would disturb the postsynaptic organization of AChRs. For this purpose, we electroporated an Src construct, Src-Y527F, in which the inhibitory C-terminal phosphorylation site is replaced by phenylalanine. The resulting Src kinase is disinhibited and constitutively activated (Kaplan et al., 1994; Kilarski et al., 2003). In GFP-positive fibers, many AChR structures resembled perforated pretzels, containing holes or broken-up regions, very similar to partially disassembled endplates observed with Src-AM (Fig. 3 A). Confocal three-dimensional imaging illustrated the process of postsynaptic disassembly, revealing that synaptic nuclei were more dispersed, as the AChRs

Figure 1. Electroporation of kinase-inactive Src-AM into soleus muscle leads to disassembly of NMJs. Muscles were electroporated in vivo with a mixture of Src-AM and NLS-GFP or empty control vector and NLS-GFP. Six weeks later, muscles were dissected and whole mounts of fibers were stained with α-BT-rhodamine and a mixture of neurofilament (NF) and synaptophysin (Syn) antibodies (blue). A, Conventional microscopy shows that in GFP-positive fibers, NMJs are partially (middle, arrow) or completely (right) disassembled, whereas GFP-negative fibers show intact NMJs (left, arrowhead in middle). B, Confocal microscopy with 3D reconstruction to visualize the degree of NMJ disassembly and the topology of nerve, AChRs, and synaptic myonuclei. Control vector electroporation leaves NMJs intact (left), whereas Src-AM electroporation disassembles NMJs in GFP-positive fibers (arrows, middle columns) but not GFP-negative fibers (arrowheads, middle columns). Disassembly ranges from partial (middle columns) to complete (right). Occasionally, nerves of disassembled NMJs show sprouting as indicated by the small arrowheads. Clusters of synaptic nuclei are less dense in disassembled NMJs, and the topology of nerve versus AChRs is disturbed.
Interestingly, in fibers expressing Src-Y527F, we observed no completely disassembled AChR clusters, unlike for Src-AM. Quantitation showed that Src-Y527F expression strongly increased the proportion of partially disassembled AChR pretzels in comparison with control situations (Fig. 3).

Together, the data demonstrate that proper regulation of kinase activity of SFKs in myofibers is a critical aspect in maintaining a normal morphology of the NMJ. If SFK activity is decreased, resulting from dominant-negative kinase-dead Src expression, postsynaptic organization is strongly affected, ranging from partial to complete disassembly of AChR pretzels and alterations in the relative topology of nerve terminals, AChRs, and synaptic nuclei. If SFK activity is increased by expressing overactive Src, the morphology is also disrupted, albeit to a lesser degree.
Expression of dominant-negative Src alters the subsynaptic but not extrasynaptic cytoskeleton and does not cause muscle degeneration

It was important to ascertain that Src-AM electroporation directly affects postsynaptic processes and not global muscle substrates that may indirectly lead to synaptic changes. We focused on two cytoskeletal elements and known substrates of SFK-activated signaling cascades, F-actin and α-tubulin. In other cells, kinase-dead as well as kinase-overactive Src expression affects their organization (Cox and Maness, 1993; Thomas et al., 1995; Brandt et al., 2002; Kilarski et al., 2003). We stained whole-mount preparations with fluorescent phalloidin to visualize actin filaments and found a transverse pattern throughout myofibers. This is indicative of costameric structures (Rybakova et al., 2000), and there was no difference between Src-AM-expressing and control fibers that had disassembled and intact NMJs, respectively (Fig. 4A). α-Tubulin stains revealed typical ring-like structures in the subsynaptic zone of the NMJ, as reported previously (Ralston et al., 1999) (Fig. 4B, arrows). Analysis of single confocal sections located the rings subsynaptically underneath AChR clusters (data not shown). These rings often surrounded synaptic nuclei as shown previously (Ralston et al., 1999) and as verified in our control experiments using empty vector and GFP (data not shown). Importantly, the rings and the overall intensity of synaptic tubulin staining were much less pronounced at disassembled endplates in fibers expressing Src-AM (Fig. 4B). The irregular extrasynaptic α-tubulin signal showed no difference between Src-AM-expressing and control fibers. These data show cytoskeletal changes at dispersed NMJs but not throughout the muscle fiber, strongly suggesting that Src-AM acts via a specific postsynaptic mechanism to disassemble NMJs.

It was also important to verify that Src-AM does not induce muscle degeneration followed by regeneration, a process that could indirectly contribute to postsynaptic disassembly at affected NMJs. A hallmark of muscle degeneration/regeneration is the appearance of myotubes with smaller diameter and centrally positioned nuclei (Paoni et al., 2002). We analyzed cross sections of electroporated muscle by triple-labeling with anti-GFP antibodies, rhodamine-α-BT, and DAPI. This allowed to identify muscle fibers, resulting from diffuse GFP-signal, and to observe the positioning of nuclei within those fibers, because most of the strong GFP signals overlapped with the nuclear DAPI signal (Fig. 4C,D). In myofibers expressing Src-AM or empty control vector, all GFP signals were always at the fiber periphery, never in the center (Fig. 4C). Also, in fibers expressing Src-AM and having disassembled NMJs, nuclei were peripheral (Fig. 4D). Finally, we did not observe small-diameter myotubes. Thus, electroporation with empty vector or Src-AM does not lead to detectable degeneration and regeneration of muscle, implying that Src-AM expression disrupts the postsynaptic NMJ apparatus by a more direct subsynaptic mechanism.

Mechanism of SFK action: they do not act in recruitment but in stabilization of postsynaptic proteins in clusters

We examined the postsynaptic mode of action of SFKs in stabilizing AChR clusters by using cultured src−/−/fyn−/− myotubes for detailed cell biological and biochemical analysis. These cells are a useful model for postsynaptic stabilization. Synapse-
promoting factors such as neural agrin or laminin induce normal AChR clustering, but most clusters disassemble when these factors are withdrawn from the cell medium for a few hours, whereas clusters in wild-type cells remain stable (Smith et al., 2001; Marangi et al., 2002).

We first addressed whether SFKs act in stabilization of agrin-induced AChR clusters by recruiting other proteins into AChR-containing aggregates. We analyzed many UGC components, because this complex plays a central role in postsynaptic stabilization and maturation (Grady et al., 1997, 2000; Jacobson et al., 2001). We treated cultured src−/−;fyn−/− myotubes with agrin and examined them by immunocytochemical staining and fluorescence microscopy. Utrophin, α-dystrobrevin, α-dystroglycan, syntrophin, rapsyn, and phosphotyrosine-containing proteins were all clustered normally by agrin treatment, and all proteins efficiently colocalized with AChRs (Fig. 5). Recruitment of these proteins into AChR-containing clusters thus occurs independently of Src and Fyn.

We next asked whether SFKs act in stabilizing clusters of postsynaptic proteins by treating src−/−;fyn−/− myotubes with agrin to induce aggregates. Agrin was then withdrawn for 4–5 h to assess the stability of these clusters. Utrophin clusters disappeared rapidly in parallel with AChR aggregates; the number of remaining utrophin clusters was as low as the number of AChR aggregates after 4 h of withdrawal, and remaining AChR clusters colocalized efficiently with utrophin (Fig. 6A, B) (supplemental Fig. 1, available at www.jneurosci.org as supplemental material). We observed the same for α-dystrobrevin (Fig. 6C) and rapsyn (Fig. 6D) (see supplemental Fig. 1, available at www.jneurosci.org as supplemental material). These data demonstrate that SFKs are not essential in the formation of clusters of postsynaptic proteins. Rather, SFKs are required for stabilization of AChRs and all other proteins examined in coextensive clusters, illustrating a general importance.
of SFKs in postsynaptic maintenance by holding together the postsynaptic apparatus.

**SFKs maintain agrin-induced AChR–rapsyn interaction**

To unravel the mechanisms by which SFKs maintain clusters of AChRs and other postsynaptic proteins, we analyzed AChR-protein interactions biochemically in src−/−;fyn−/− myotubes, focusing first on rapsyn. Rapsyn is a key anchor for the AChR in clusters (Gautam et al., 1995), and its interaction with the receptor is highly regulated and increased by agrin (Moransard et al., 2003). Rapsyn exists in free and AChR-bound forms (Marangi et al., 2001; Moransard et al., 2003). Although intracellular and unclustered surface AChRs interact with some rapsyn, agrin induces increased rapsyn–AChR binding at the plasma membrane, and this increase tightly correlates with clustering and cytoskeletal anchoring. Likewise, synaptic AChRs contain more bound rapsyn than extrasynaptic AChRs in denervated muscle (Moransard et al., 2003). Because increased rapsyn–AChR interaction thus underlies clustering, we analyzed this interaction in cluster formation and stabilization in the absence of Src and Fyn. We treated src−/−;fyn−/− myotubes with agrin and then withdrew agrin for 3 or 5 h (Fig. 7A). AChRs were precipitated from lysates using biotinylated α-BT and streptavidin-agarose (Tox-P), and AChR-bound rapsyn was visualized by immunoblotting. Interestingly, in untreated src−/−;fyn−/− myotubes, α-BT-precipitated AChRs contained, on average, more associated rapsyn than in wild-type cells (Fig. 7A, B). This interaction was further increased approximately twofold by agrin treatment but returned to basal levels within 5 h of agrin withdrawal. In contrast, rapsyn binding to AChR in wild-type cells, increased by agrin to the same relative degree (approximately twofold) as in the mutant, remained increased even after 5 h of withdrawal (Fig. 7A, B). Thus, although AChRs contain, on average, more associated rapsyn in src−/−;fyn−/− myotubes, the agrin-induced increase in the interaction is very unstable in the absence of Src and Fyn.

We previously quantitated the degree of rapsyn coprecipitation with AChRs and found that not every precipitated receptor molecule is associated with one rapsyn molecule (Fuhrer et al., 1999). Other observations suggested that rapsyn exists in an equilibrium between free and AChR-associated state (Marangi et al., 2001; Moransard et al., 2003) and that the average ratio of rapsyn to AChR expression is ∼1:1 (LaRochelle and Froehner, 1987). The extraction step of our α-BT precipitation of AChRs is likely to break up some AChR–rapsyn interaction (Fuhrer et al., 1999). For these reasons, the higher basal degree of rapsyn coprecipitation with AChRs in src−/−;fyn−/− myotubes may originate from a weakened cytoskeletal linkage of rapsyn, making its extraction and coprecipitation with the receptor more efficient. Alternatively, rapsyn protein may be present in higher amount in the mutant cells, so that more rapsyn
Cells were labeled with rhodamine-labeled cluster maintenance, agrin was then withdrawn from cultures for 4–5 h (wd 4 h, wd 5 h). Double labeled with rhodamine-tic protein, in the case of agrin withdrawal for remaining AChR clusters. Values are mean quantitation (right) indicates the percentage of AChR clusters that contain the respective postsynaptic proteins disappear in parallel with AChRs after agrin withdrawal. Colocalization (left) is available at www.jneurosci.org as supplemental material. Quantitation (left) shows that clusters of these postsynaptic proteins are larger in the wild type. Error bars represent SEM. ag, Agrin treated; no ag, untreated.

Figure 6. Src and Fyn are required to stabilize postsynaptic protein clusters along with AChR clusters. Cells were treated overnight with 0.5 nM agrin followed by withdrawal (wd) as indicated. AChRs were precipitated from lysates using α-BT-biotin and streptavidin-agarose (Tox-P), and associated rapsyn was detected by Western blotting. As control, we quantitated the cellular amounts of other postsynaptic proteins, such as utrophin, MuSK, and α-dystrobrevin-1 (DB) (C), or rapsyn (Rap) (D), followed by FITC-coupled secondary antibodies. Illustrative pictures of clusters and their disassembly are shown in supplemental Figure 1 (available at www.jneurosci.org as supplemental material). Quantitation (left) shows that clusters of these postsynaptic proteins disappear in parallel with AChRs after agrin withdrawal. Colocalization (right) indicates the percentage of AChR clusters that contain the respective postsynaptic protein, in the case of agrin withdrawal for remaining AChR clusters. Values are mean ± SEM, from 20 pictures for each marker and condition. Error bars represent SEM. ag, Agrin treated; no ag, untreated.

is available to bind the receptor in the absence of Src and Fyn. To distinguish between these two possibilities, we analyzed two aspects of rapsyn in src−/− fyn−/− myotubes.

First, we determined the amount of rapsyn protein in cellular lysates. Using immunoblotting, we observed higher amounts of rapsyn protein per milligram of cellular protein in standard lysates, made with 1% detergent, of src−/− fyn−/− cells compared with wild-type cells (Fig. 7C). In mutant cells, levels of rapsyn protein (normalized for AChR β subunits (mAb 124)) were 2.0 ± 0.1-fold increased (mean ± SEM; n = 5). In many further extractions using milder and harsher conditions (decreasing or increasing detergent and salt concentrations), we always found similarly elevated rapsyn protein levels in src−/− fyn−/− cells (data not shown). As control, we quantitated the cellular amounts of other postsynaptic proteins, such as utrophin, MuSK, and α-dystrobrevin. We observed no significant difference in these proteins between mutant and wild-type cells (data not shown), suggesting that the effect of Src and Fyn on rapsyn levels is specific.

Second, we measured the relative strength of cytoskeletal rapsyn interaction. We performed extraction experiments simi-
lar to those described in detail for AChRs in Figure 9, using low- and high-detergent concentrations. By immunoblotting, we determined the proportion of rapsyn in each of these extractions. The proportion of rapsyn in low- versus high-detergent extractions was the same for src$^{-/-}$;fyn$^{-/-}$ and wild-type myotubes (data not shown), showing that the cytoskeletal link of rapsyn is unchanged in the mutant.

Together, these data, combined with previous observations (Marangi et al., 2001; Moransard et al., 2003), strongly imply that SFK activity is required to maintain phosphorylation of the AChR, and that the overall link of rapsyn is a linker between AChR and the UGC (Apel et al., 1995; Cartaud et al., 1998; Bartoli et al., 2001).

Phosphorylation of the AChR β subunit is a key event in agrin signaling and important for efficient clustering and cytoskeletal linkage of the receptor (Borges and Ferns, 2001). Agrin-induced AChR β phosphorylation is an early step in agrin signaling and precedes receptor clustering, similar to agrin-triggered rapsyn–AChR interaction (Ferns et al., 1996; Moransard et al., 2003). We examined whether SFKs are required to maintain this phosphorylation in the process of cluster stabilization. Precipitation with biotin–α-BT and immunoblotting with anti-phosphotyrosine showed that agrin treatment caused normal β phosphorylation in src$^{-/-}$;fyn$^{-/-}$ myotubes, as described previously (Smith et al., 2001). But after agrin withdrawal, this phosphorylation was unstable and disappeared much faster than in wild-type cells (Fig. 8B). Although in the wild type significant β phosphorylation persisted 5 h after agrin withdrawal, β phosphorylation was already down at basal levels after 3 h in src$^{-/-}$;fyn$^{-/-}$ myotubes (Fig. 8B). We thus estimate the half-life time ($t_{1/2}$) of β phosphorylation to be ~4 h in the wild type but <90 min in the mutant. Because AChRs can be direct substrates for SFKs (Swope and Huganir, 1993; Fuhrer and Hall, 1996; Mohamed et al., 2001; Mittaud et al., 2004), these data strongly imply that SFK activity is required to maintain phosphorylation of the AChR, and that the receptor is a direct substrate for SFKs at that stage.

The correlation between loss of AChR phosphorylation and AChR–rapsyn binding after agrin removal raises the possibility that β phosphorylation may regulate rapsyn interaction. Collectively, the sum of our data on src$^{-/-}$;fyn$^{-/-}$ myotubes up to this point suggests that SFK-mediated stabilization of AChR phosphorylation and of AChR association with rapsyn and dystrobrevin form a core mechanism by which SFKs hold together proteins of the postsynaptic apparatus.

The overall link of AChRs to the cytoskeleton is weak in src$^{-/-}$;fyn$^{-/-}$ myotubes but still strengthened by agrin in addition to changes at the level of AChR clusters, the alterations in postsynaptic architecture in Src-AM-expressing myotubes in vivo suggest that the postsynaptic cytoskeleton (as shown...
for α-tubulin in Fig. 4B) and its protein interactions may be altered after loss of SFK function. To further test this hypothesis, we analyzed the role of SFKs in interactions of the AChR with the cytoskeleton. We quantitated the AChR extractability in src⁻/⁻/fyn⁻/⁻ myotubes by applying a sequential detergent extractability protocol in which a first extraction in low detergent is followed by a second extraction in higher-detergent concentration. From each extraction, we precipitated AChRs using biotin-BT, visualized them in Western blots, and calculated the receptor distribution between first and second extraction, using increasing detergent concentrations in the first extraction (Fig. 9A, B). Such methods are established measures for the relative strength of the interaction of the AChR to the cytoskeleton (Borges and Ferns, 2001; Moransard et al., 2003). We found that in wild-type myotubes, AChR started to become efficiently solubilized in the first extraction when the concentration of first detergent was 0.06% Triton X-100 (Fig. 9A). At lower Triton X-100 concentrations in the first extraction, receptors appeared in the second extraction, whereas at higher concentrations, AChRs participated into the first extraction (Fig. 9B). In contrast, in src⁻/⁻/fyn⁻/⁻ myotubes, a concentration of 0.04% Triton X-100 was sufficient to efficiently solubilize AChRs in the first extraction (Fig. 9B). These data reveal that the overall basal cytoskeletal link of the AChR is weakened in the absence of Src and Fyn.

Agrin treatment is known to strengthen the AChR-cytoskeletal link in the process of cluster formation (Wallace, 1992, 1995; Borges and Ferns, 2001; Moransard et al., 2003). This is manifested as a decrease of solubilized and precipitated AChR in the first extraction and an increase of receptor in the second extraction (Borges and Ferns, 2001; Moransard et al., 2003). Like-
wise, AChRs increasingly become cytoskeletally anchored at developing NMJs in vivo (Dennis, 1981; Slater, 1982). We analyzed whether agrin affects the weak overall cytoskeletal interaction of the AChR in src−/−;fyn−/− myotubes by quantitating the relative amounts of AChRs in the first and second extraction in agrin-treated and untreated cells. Agrin caused a shift of receptors from the first (0.05% Triton X-100) to the second (1% Triton X-100) extraction in src−/−;fyn−/− myotubes (Fig. 9C,D), similar to results previously seen in wild-type cells (Borges and Ferns, 2001; Moransard et al., 2003). The percentage of agrin-induced decrease of AChR precipitated from the first extraction was identical between C2 myotubes, wild-type myotubes, and src−/−;fyn−/− myotubes (Fig. 9E).

Together, these results show that although the overall basal cytoskeletal linkage of AChRs is weaker in the absence of Src and Fyn, agrin treatment still strengthens this link. This is consistent with the normal phosphorylation and cluster formation of AChRs in agrin-treated src−/−;fyn−/− myotubes and the decreased stability of such clusters after removal of agrin (Smith et al., 2001). Most likely, the agrin-induced strengthening of the AChR cytoskeletal link in mutant cells originates from linkage of the receptor to the UGC, because AChR–dystrobrevin interactions are normally induced by agrin in these cells (Fig. 8A) and because the UGC interacts with F-actin (Winder et al., 1995).

Discussion
Our data reveal that SFKs are key players in the pathways that stabilize the postsynaptic apparatus of the NMJ in vivo. They act by holding postsynaptic proteins together in clusters through stabilization of rapsyn–AChR interaction and AChR phosphorylation. In addition, they control rapsyn protein levels and AChR-cytoskeletal linkage.

SFKs hold together the postsynaptic apparatus
Interference with SFK function causes complex alterations at adult NMJs. After Src-AM expression, AChR pretzels fragment and attach to the nerve in the same focal plane (not underneath it), subsynaptic α-tubulin organization is disturbed, synaptic nuclei become more dispersed, and nerves occasionally sprout. These changes originate from Src-AM expression in muscle, acting specifically on postsynaptic mechanisms; GFP-positive nuclei were only seen in myofibers and never in other cells (e.g., Schwann cells), confirmed by 3D reconstruction (G. Sadasivam and C. Fuhrer, unpublished observations), and costameric F-actin organization along myofibers was not affected. Nerve sprouting is in accordance with studies showing that postsynaptic disturbance affects the nerve, leading to sprouting or, as is the case in rapsyn- or MuSK-deficient mice, extensive nerve growth (Gautam et al., 1995; DeChiara et al., 1996; Kong et al., 2004).

The effects of Src-Y527F expression are similar, illustrating that correctly balanced SFK activity is important to maintain the postsynaptic apparatus in vivo. Consistent with this, reducing or increasing SFK activity experimentally leads to changes in downstream pathways and affects cytoskeletal organization (for example, actin fibers) in other cell types (Thomas et al., 1995; Brandt et al., 2002; Kilarski et al., 2003).

We investigated the consequences of reduced SFK function using src−/−;fyn−/− myotubes, where agrin normally recruited postsynaptic proteins into AChR-containing clusters. This parallels the normal development, until birth, of endplates in src−/−;fyn−/− mice (Smith et al., 2001). But after agrin removal from src−/−;fyn−/− myotubes, clusters of UGC and rapsyn disintegrated in parallel with AChRs. Thus Src and Fyn hold together the postsynaptic apparatus, consistent with AChR pretzel disassembly in Src-AM-expressing myofibers.

SFKs maintain AChR–rapsyn interaction and AChR phosphorylation
In parallel with unstable clusters, AChR-protein interactions are unstable in src−/−;fyn−/− myotubes, and the key compromised interaction is that between AChRs and rapsyn. Agrin-induced increases in AChR–rapsyn interaction correlate highly with clustering (Moransard et al., 2003), and increased rapsyn binding slows metabolic AChR turnover (Gervasio and Phillips, 2005). AChRs and rapsyn are the most abundant postsynaptic components. Occasionally, AChR–rapsyn complexes are linked to the UGC through dystroglycan, giving rise to a corral model in which postsynaptic proteins such as the UGC are held together through dystroglycan, utrophin, utro, Utrophin; DB, α-dystrobrevin.

Figure 10. Model of SFK action in stabilization of the postsynaptic apparatus. Unclustered proteins such as rapsyn (rap) are in equilibrium between free and complexed form, indicated by double arrows. Agrin increases AChR-protein interactions in the process of clustering. SFKs act in postsynaptic stabilization by maintaining the AChR–rapsyn interaction (pathway 1). Because rapsyn and AChRs are the most abundant postsynaptic proteins, their interaction plays a core role and holds the postsynaptic apparatus together. This may occur through AChR–dystroglycan (p), which is maintained by SFKs (pathway 1). SFKs also negatively control the overall amount of rapsyn protein (pathway 2). In the absence of Src and Fyn, rapsyn amounts are high and may start saturating binding sites on β-dystroglycan (β-DG) and the AChR, although AChR–dystrobrevin interactions appear normal. SFKs may control another cytoskeletal link (cyto) of the AChR, independent of rapsyn as a linker (hypothesis pathway 3). This would explain the observed weak overall AChR-cytoskeletal linkage in src−/−;fyn−/− myotubes, which is still strengthened after agrin treatment resulting from AChR–UGC association. utro, Utrophin; DB, α-dystrobrevin.
1996). After agrin stimulation of myotubes, however, SFKs only act in the initial phase, are later compensated by Abl kinases and not necessary for cluster formation (Mittaud et al., 2004). We now find that Src and Fyn are required to maintain β phosphorylation, and this most likely reflects direct phosphorylation. Loss of β phosphorylation may lead to weaker AChR-cytoskeletal linkage, but we were not able to reliably quantify AChR extractability after agrin withdrawal caused by variation between experiments (R. Willmann and C. Fuhrer, unpublished observations).

In src−/−;fyn−/− myotubes, loss of β phosphorylation after agrin withdrawal is paralleled by loss of AChR–rapsyn interaction. Additional experiments have further corroborated a tight correlation. Whereas agrin induces pronounced β phosphorylation and rapsyn binding in C2 myotubes, pervanadate treatment causes stronger β phosphorylation and stronger rapsyn binding, revealing a linear relationship between the two events (M. Moransard and C. Fuhrer, unpublished observations). Furthermore, the time course of agrin-induced β phosphorylation exactly parallels that of AChR–rapsyn binding (Moransard and Fuhrer, unpublished observations). Thus, increased AChR–rapsyn binding may occur through β phosphorylation via direct protein interaction or through an intermediate linker (Fig. 10, pathway 1). Consequently, loss of β phosphorylation may diminish AChR–rapsyn interaction, causing postsynaptic disassembly. The proposal that rapsyn binds AChRs in several ways, in a basal state independent of AChR phosphorylation but after agrin addition through increased receptor phosphorylation (Fig. 10), is consistent with findings from heterologous cells. Here, rapsyn directly or indirectly interacts with AChRs in multiple ways through association with all receptor subunits (Maimone and Merlie, 1993; Maimone and Enigk, 1999; Bartoli et al., 2001; Huebsch and Maimone, 2003).

More surprisingly, SFKs repress the amount of rapsyn protein (Fig. 10, pathway 2). Correct expression level of rapsyn is important, because its overexpression reduces AChR clustering in myotubes (Yoshihara and Hall, 1993; Han et al., 1999). We did not investigate whether alterations in rapsyn synthesis, degradation, or turnover cause the increase in overall rapsyn protein. More importantly, we found that the cytoskeletal link of rapsyn is unaffected by Src and Fyn. In heterologous cells, SFKs form a complex with rapsyn, and rapsyn triggers their kinase activity, leading to AChR phosphorylation (Mohamed and Swope, 1999). In myotubes, rapsyn is required for agrin-induced activity of SFKs, implying an interaction between rapsyn and SFKs (Mittaud et al., 2001). Thus, SFK and rapsyn seem to be engaged in mutual control, leading to correct rapsyn protein levels and SFK activity. Fine tuning of such interactions and activities may be important for correct protein interactions in building up and stabilizing the postsynaptic apparatus and for appropriate linkage of associated signaling pathways.

SFKs control AChR-cytoskeletal interactions
Unstable of AChR-protein interactions is sufficient to explain the disintegration of postsynaptic protein clusters but may not account for all postsynaptic changes observed in vivo after Src-AM expression. Changes in nerve-AChR topology and synaptic nuclei position are likely to reflect additional changes in the postsynaptic cytoskeleton as illustrated by loss of synaptic α-tubulin rings. Indeed, cytoskeletal linkage of the AChR is weaker in src−/−;fyn−/− myotubes yet strengthened by agrin treatment. One candidate mechanism to explain this observation is the UGC and its interaction with F-actin (Winder et al., 1995). We do not think that the higher rapsyn amount in src−/−;fyn−/− cells saturates rapsyn-binding sites on AChRs or β-dystroglycan, thereby disturbing AChR-F-actin linkage, because the UGC component dystrobrevin associates normally with AChRs in the mutant cells (Fig. 8A). This leads to the conclusion that another mechanism, potentially involving microtubular organization, accounts for the weaker basal cytoskeletal link of the AChR in src−/−;fyn−/− myotubes. Such a mechanism may not involve rapsyn as a linker, because the rapsyn cytoskeletal interaction is normal in the mutant cells. More likely, the AChR interacts directly or indirectly with other elements of the cytoskeleton through a novel pathway, before and after agrin treatment, and such linkage depends on Src and Fyn (Fig. 10, putative pathway 3). Agrin-triggered strengthening of the overall AChR-cytoskeletal link may stem from agrin-induced AChR–UGC interaction, because agrin induces normal AChR-dystrobrevin association in mutant and wild-type myotubes. In such a manner, normal AChR-UGC association in combination with defects in a putative additional cytoskeletal pathway provides an explanation for the observed alteration in overall AChR extractability in SFK-defective cells.

The intermediate elements in the putative SFK-mediated cytoskeletal pathway remain to be identified. Candidates are known SFK substrates involved in cytoskeleton dynamics, such as contactin and WASP (Wiskott–Aldrich syndrome protein) (both regulating the Arp2/3 complex) (Daly, 2004; Martinez-Quiles et al., 2004) or p190RhoGAP (influencing Rho GTPase activity) (Chang et al., 1995), which all ultimately regulate F-actin assembly. Reduction or increase in SFK activity affects the organization of actin fibers in other cell types (Thomas et al., 1995; Brandt et al., 2002; Kilarski et al., 2003), consistent with our finding that both Src-AM and Src-Y527F expression disassembles AChR pretzels in vivo. Rho, along with Rac and Cdc42, is already known to play a role in AChR cluster formation in cultured myotubes (Weston et al., 2000, 2003). SFKs also influence the tubulin network (Cox and Maness, 1993), and we have observed changes in synaptic tubulin organization after Src-AM expression.

It remains to be investigated what other connections between SFKs and the cytoskeleton and its regulators exist at the NMJ, affecting postsynaptic stability. Because balanced SFK activity is important (Fig. 1–3), SFKs may be counteracted by tyrosine phosphatases. Phosphatase activity dissolves AChR hot spots in cultured Xenopus myocytes, and some of this activity is triggered by agrin application (Madhavan et al., 2005). The phosphatase Shp-2 is a possible candidate, because blocking Shp-2 increases spontaneous AChR clustering (Madhavan et al., 2005). The balance between SFKs and phosphatases offers a fine-tuning system to shape the postsynapse. It will be interesting to assess the role of such a system in vivo, also in the first weeks of postnatal NMJ development, when synapse elimination occurs. In this process, AChR regions can be selectively destabilized paralleled by nerve withdrawal (Lichtman and Colman, 2000), and a tyrosine kinase-phosphatase regulation is an attractive candidate mechanism.

References
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Src Action in Postsynaptic Stabilization


