Neural cell adhesion molecule promotes accumulation of TGN organelles at sites of neuron-to-neuron contacts

Vladimir Sytnyk,1,2 Iryna Leshchyns’ka,1 Markus Delling,1 Galina Dityateva,1 Alexander Dityatev,1 and Melitta Schachner1

1Zentrum für Molekulare Neurobiologie, Universität Hamburg, D-20246 Hamburg, Germany
2Laboratory for Biophysics and Bioelectronics, Dnepropetrovsk State University, 49050 Dnepropetrovsk, Ukraine

Transformation of a contact between axon and dendrite into a synapse is accompanied by accumulation of the synaptic machinery at this site, being delivered in intracellular organelles mainly of TGN origin. Here, we report that in cultured hippocampal neurons, TGN organelles are linked via spectrin to clusters of the neural cell adhesion molecule (NCAM) in the plasma membrane. These complexes are translocated along neurites and trapped at sites of initial neurite-to-neurite contacts within several minutes after initial contact formation. The accumulation of TGN organelles at contacts with NCAM-deficient neurons is reduced when compared with wild-type cells, suggesting that NCAM mediates the anchoring of intracellular organelles in nascent synapses.

Introduction

Synaptogenesis in the CNS is accompanied by accumulation of synaptic organelles and proteins at the sites of contact between axons and dendrites (Mammen et al., 1997; Rao et al., 1998; Friedman et al., 2000; Lee and Sheng, 2000; Zhai et al., 2001). The culmination of this process is the transformation of the initial contacts into functional synapses. In axons, synaptic proteins are transported within intracellular tubulovesicular membrane aggregates of TGN origin that probably represent synaptic vesicle precursors (Nakata et al., 1998). These precursors then concentrate at sites of contact in parallel with their transformation into synapses (Ahmari et al., 2000). In dendrites, TGN organelles undergo regulated exocytosis and are suggested to mediate the delivery of synaptic proteins to postsynaptic sites (Lledo et al., 1998; Maletic-Savatic and Malinow, 1998; Shi et al., 1999). The question is how these organelles are captured and stabilized at sites of contact that would initiate their transformation into synapses or mediate synaptic rearrangements.

Cell adhesion molecules are probably the best candidates to execute this task. In Drosophila, a deficit in the cell adhesion molecule fasciclin II leads to a loss of synapses (Schuster et al., 1996). In mammals, the neural cell adhesion molecule (NCAM),* the closest homologue of fasciclin II, accumulates at sites of intercellular contacts, where it has been proposed to stabilize the contact structure (Pollerberg et al., 1986, 1987). The role of NCAM in synaptogenesis was demonstrated in experiments showing a higher number of synapses on NCAM-expressing neurons compared with NCAM-deficient cells in a choice situation in heterogenotypic cocultures of NCAM-deficient and wild-type neurons (Dityatev et al., 2000). NCAM is expressed in three major splicing isoforms: GPI-linked NCAM120 (120 kD) and transmembrane NCAM140 (140 kD) and NCAM180 (180 kD). NCAM180, the isoform with the largest intracellular domain, interacts with spectrin (Pollerberg et al., 1986, 1987) and could provide a direct link to intracellular organelles, such as TGN organelles, which are lined by a spectrin cytoskeleton (De Matteis and Morrow, 1998; Holleran and Holzbaur, 1998; Lippincott-Schwartz, 1998).

Here, we used time-lapse confocal microscopic video recordings, markers of subcellular compartments, and subcellular fractionation to describe an association between NCAM clusters and intracellular aggregates of TGN organelles that are transported en ensemble to initial neuron–neuron contacts.

*Abbreviations used in this paper: DIC, differential interference contrast; NCAM, neural cell adhesion molecule.
We demonstrate that NCAM mediates the capture of organelles at initial sites of contact that is followed by the transformation of these contacts into functional synapses.

Results
NCAM clusters associate with a subpopulation of intracellular organelles
To study the relationship between localization of intracellular organelles and distribution of NCAM at the neuronal surface, we used differential interference contrast (DIC) microscopy combined with subsequent immunofluorescence analysis. In hippocampal neurons maintained for 1–3 d in vitro, intracellular organelles (observed as dark granules by DIC microscopy) were typically localized within neurite swellings of 1–2 μm in diameter. As observed by time-lapse video recording, intracellular organelles underwent rapid intermittent movement along neurites with a speed that reached ~0.5 μm/s. These intracellular aggregates often resembled transport packets as described previously (Ahmari et al., 2000; Washbourne et al., 2002) (Fig. 1 a).

After time-lapse imaging, neurons were fixed and stained with antibodies to NCAM, showing that a subpopulation of organelles that had moved during video recording were colocalized with intensely labeled clusters of NCAM (Fig. 1, a and b). NCAM clusters occupied areas of the plasma membrane of 0.4–2 μm in diameter that covered the plasma membrane over the intracellular organelles. The immunofluorescence intensity associated with NCAM clusters was more than two times higher than the basal level of immunofluorescence along the neurite. Because no detergents were used for immunofluorescence staining, the observed NCAM immunostaining pattern represented plasma membrane, and not intracellular, NCAM localization. In support of this argument, antibodies to tubulin applied in mixture with NCAM antibodies to neurons not treated with Triton X-100 did not give any staining (Fig. 1 b, tubulin, control), whereas antibodies to tubulin applied to cells treated after fixation with 0.25% Triton X-100 yielded a strong and uniform staining of microtubules in soma and neurites (unpublished data). Because intracellular organelles were usually located within varicosities, the question arose whether the apparent peaks of NCAM immunofluorescence intensity associated with organelles were due to the larger diameter of neurites at these sites. To resolve this, we stained neurons with the lipophilic dye DiI, which intercalates into the surface membrane by lateral diffusion. DiI showed a uniform distribution along neurites independently of neurite thickness and presence of varicosities (unpublished data), indicating that the peaks of NCAM immunofluorescence intensity at the cell surface corresponded to a higher density of NCAM at these sites.

NCAM clusters interact with TGN organelles via spectrin
To identify the composition of intracellular organelles associated with NCAM-immunoreactive clusters, neurons were stained with NCAM antibodies and labeled with antibodies to different organelle-specific markers. To label the TGN and TGN-derived organelles, we used antibodies to γ-adaptin (Robinson and Kreis, 1992; Girotti and Banting, 1996). This protein belongs to the AP-1 complex associated with the TGN and clathrin-coated vesicles that bud from the TGN (Robinson and Kreis, 1992; Schmid, 1997; Heimann et al., 1999) and that are distinct from clathrin-coated endocytic vesicles, which incorporate another adaptor complex, AP-2 (Clague, 1998). Also, we used antibodies to β-COP, a coat protein associated with the TGN and non-clathrin-coated vesicles that bud from the TGN (Robinson and Kreis, 1992). To label endosomal vesicles, we used anti-
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bodies to EEA1, an early endosome-associated protein (Mu et al., 1995), Rab4, characteristic of early and recycling endosomes (Sonnichsen et al., 2000), and lamp-1, a lysosomal membrane glycoprotein (Fukuda, 1991). These markers were highly concentrated in the soma and showed a patchy distribution along neurites. Thick tapering neurites were identified as dendrites, whereas thin neurites of a uniform diameter with multiple varicosities were identified as axons. This classification was verified in a separate set of experiments using the established markers, such as tau and synaptophysin for axons (Ahmari et al., 2000; Paglini et al., 2000) and MAP2b for dendrites (Shaﬁt-Zagardo and Kalcheva, 1998). Along dendrites and axons, NCAM-immunoreactive clusters were signiﬁcantly more associated with intracellular aggregates containing γ-adaptin– and β-COP–positive organelles than with the other markers, with ~70% of all NCAM clusters overlapping with γ-adaptin and β-COP immunopositive organelles (Fig. 2, a, b, and d). Approximately 70% of γ-adaptin– and β-COP–positive organelles overlapped with NCAM clusters both in dendrites and axons (Fig. 2 e). γ-Adaptin and β-COP accumulate in the TGN where they mediate budding of two different types of vesicles (Robinson and Kreis, 1992; Heimann et al., 1999). TGN and TGN-derived organelles could form large, up to several micrometers in diameter, vesicular–tubular structures (Nakata et al., 1998; Toomre et al., 1999, 2000; Polishchuk et al., 2000), which have been shown to transport synapse-speciﬁc proteins (Nakata et al., 1998). The synaptic vesicle protein synaptophysin was accumulated in varicosities of neurites (Fig. 2 c), identiﬁed as axons by this presynaptic marker. Double immunostaining for NCAM and synaptophysin showed that 80% of all NCAM clusters in axons coincided with synaptophysin clusters, and ~90% of all synaptophysin clusters overlapped with NCAM clusters (Fig. 2, c–e). Thus, NCAM clusters strongly overlap with TGN and synaptic vesicle markers.

We next studied whether the intracellular domains of NCAM140 or NCAM180 interact with TGN organelles. To this aim, TGN membranes were isolated from the brains of NCAM-deﬁcient mice. NCAM-deﬁcient mice were used instead of wild-type mice to ensure that binding sites for the intracellular domains of NCAM would not be saturated by endogenous NCAM. Immunoblotting conﬁrmed that the isolated membranes were positive for the NCAM markers γ-adaptin and β-COP and the synaptic vesicle marker synaptophysin, and negative for the endosomal markers EEA1, Rab4, and lamp-1 (Fig. 3 a). TGN membranes were incubated with the intracellular domains of NCAM140 (IC140) and NCAM180 (IC180), washed, and assayed for IC140 and IC180 binding after separation of the complexes by
SDS-PAGE and immunoblotting with antibodies to NCAM (Fig. 3 b). Only IC180 pelleted with TGN membranes, whereas IC140 did not.

One of the known binding partners of the intracellular domain of NCAM180 is spectrin (Pollerberg et al., 1986, 1987). At least two spectrin isoforms, spectrin βI and βIII, form a meshwork lining Golgi membranes (De Matteis and Morrow, 1998, 2000; Holleran and Holzbaur, 1998; Lippincott-Schwartz, 1998). In neurons maintained for 1–5 d in vitro, βI spectrin (hereafter called spectrin) was detected in all neurites, whereas in neurons maintained for 20 d in vitro, αβI spectrin was detected only in dendrites. Spectrin accumulated in varicosities and colocalized with NCAM clusters (Fig. 3 c) that overlapped completely with NCAM180 labeling (unpublished data). Immunoblotting confirmed that the isolated TGN membranes contained βI spectrin, as detected with βI spectrin–specific antibodies (Fig. 3 a) and polyclonal antibodies against αβI spectrin (Fig. 3 d). We then determined whether the ability of IC180 to bind TGN membranes depended on the presence of noncovalently associated proteins, including the spectrin meshwork, by briefly incubating TGN membranes in alkaline sodium carbonate solution, thereby removing peripheral proteins. After this treatment, spectrin was no longer detectable on TGN membranes (Fig. 3 d). Also, IC180 no longer precipitated with TGN membranes (Fig. 3 b). Because alkaline treatment could have not only removed spectrin from TGN membranes but may also have denatured other TGN membrane-associated binding partners for IC180, TGN membranes were incubated with rabbit polyclonal antibodies to spectrin before assaying for IC180 binding. After this treatment, binding of IC180 was significantly inhibited, indicating that spectrin is a linker molecule that mediates binding of IC180 to TGN membranes (Fig. 3 e). As a control, the immunoglobulin fraction from nonimmune rabbits did not show any inhibition of binding (Fig. 3 e). It is therefore conceivable that NCAM180 at the cell surface interacts with TGN organelles via spectrin.

Movement of NCAM clusters and associated intracellular organelles

To visualize the distribution and movement of NCAM in association with TGN organelles, two strategies were pursued. Live hippocampal neurons were stained by indirect immunofluorescence using monoclonal antibodies to NCAM under conditions in which antibody-induced cross-linking of primary antibodies was minimized. Alternatively, neurons were transfected with NCAM180 tagged with GFP (NCAM180–GFP). NCAM180–GFP showed a distribution identical to that of immunocytochemically detectable endogenous NCAM180 and “total” cell surface NCAM in that it accumulated in clusters associated with varicosities along neurites and, in particular, also in growth cones (unpublished data). In both cases, intracellular organelles colocalizing with NCAM were observed using DIC optics. We also made use of the possibility to label TGN organelles with the vital styryl fluorescent dyes FM1-43 or FM4-64. Short-term application of these dyes is known to label presynaptic boutons via activity-dependent uptake of dyes into synaptic...
vesicles (Cochilla et al., 1999). However, after prolonged exposure of neurons to these dyes, they label multiple intracellular organelles, mostly TGN or Golgi-like structures (Maletic-Savatic and Malinow, 1998; Maletic-Savatic et al., 1998). Most of the organelles labeled for 24 h with the dyes (80%) coincided with NCAM-immunoreactive and NCAM180–GFP clusters in transfected neurons. Video recordings showed that NCAM-immunoreactive clusters and intracellular organelles moved bidirectionally along neurites (Fig. 4 a; see also Videos 1 and 2, available at http://www.jcb.org/cgi/content/full/jcb.200205098/DC1). Analyses of DIC images and FM4-64 fluorescence showed that NCAM clusters were consistently associated with intracellular organelles loaded with FM4-64 during their movement.

To show that NCAM-immunoreactive clusters that moved during the time-lapse recordings were located at the plasma membrane and not endocytosed during image acquisition, neurons were fixed and stained with polyclonal antibodies to NCAM using secondary antibodies carrying a different fluorochrome to distinguish live-labeled from fixed-labeled NCAM. All NCAM-immunoreactive clusters that moved during the time-lapse recording were stained with polyclonal antibodies to NCAM on nonpermeabilized cells, confirming that they had moved in the plasma membrane rather than in endosomes or other intracellular organelles (Fig. 4 b). Antibodies to tubulin applied in a mixture with NCAM polyclonal antibodies to control for membrane integrity did not give any staining. The same results were obtained when polyclonal antibodies were applied to live neurons at 4°C to block possible endocytosis (unpublished data), confirming surface location of NCAM-immunoreactive clusters. Also, staining of NCAM180–GFP-transfected nonpermeabilized neurons with polyclonal antibodies to NCAM showed that NCAM180–GFP clusters were located at the cell surface (unpublished data). To further confirm that organelles loaded with FM4-64 were TGN organelles, we acquired images of the hippocampal neurons loaded with FM4-64 and subsequently labeled the neurons with antibodies to γ-adaptin: the majority of FM4-64–loaded organelles (89%, n = 162) were γ-adaptin positive (Fig. 4 c). Approximately 10% of γ-adaptin–positive organelles were not loaded with FM4-64, probably due to the limited time of incubation with FM4-64.

Autophagic vacuoles constitute another class of large vesicles that can be observed in axons (Hollenbeck, 1993; Overly et al., 1996). To test whether NCAM clusters move in association with TGN organelles | Sytnyk et al. 653
with these organelles, we labeled them with RITC-dextran, which has been shown to accumulate in these organelles when applied to the culture medium for several hours (Hollenbeck, 1993). After loading with RITC-dextran, live hippocampal neurons were labeled with NCAM antibodies. Time-lapse analysis showed that NCAM clusters were very rarely associated with RITC-dextran–labeled organelles (<5%) and moved independently of them (Fig. 4, d and e).

**NCAM clusters and associated intracellular organelles accumulate at sites of neurite-to-neurite contacts**

Hippocampal neurons maintained for 2–3 d in culture formed multiple intercellular contacts. These contacts were intensely stained for NCAM and often contained intracellular organelles. To study the movement of NCAM clusters and associated intracellular organelles during contact formation, we visualized NCAM distribution by indirect immunofluorescence and recorded a time series of movements of NCAM-immunoreactive clusters during contact formation between hippocampal neurons.

Fig. 5 shows the formation of three contacts between filopodia of a neurite approaching the central part of the target neurite. We also visualized NCAM clusters and associated intracellular organelles during contact formation (see also Video 3, available at http://www.jcb.org/cgi/content/full/jcb.200205098/DC1). The clusters were loaded with FM1-43 for 24 h before application of NCAM monoclonal antibody. Prior to contact formation, NCAM-immunoreactive clusters moved bidirectionally along the neurite in association with FM1-43–positive intracellular organelles (Fig. 5). Growth cones approaching neurites were often not labeled by NCAM antibodies, probably due to the fact that the growth cones were formed after the antibody was applied (usually >20 min after antibody labeling). After establishment of physical contacts between the growth cone and the target neurite (Fig. 5 a, 40 s), NCAM-immunoreactive clusters and associated organelles that had moved along the target neurite started to accumulate at the site of contact (Fig. 5 a, 90 s, cluster No. 1). The clusters and associated intracellular aggregates often passed several times through the actual site of contact, but finally one or several clusters and associated organelles were “trapped” at the contact site and remained there until the end of the recordings (10–100 min). After the initial contact, neurites formed a “varicosity” or “bouton,” probably representing the site where apposing membranes approach each other. Organelles were trapped within this thickening that was in close proximity to the intersection point between neurites, but did not always exactly coincide with the intersection point between neurites (Fig. 5, organelles 1 and 2). Sometimes, the growth cone contacted the target neurite at the site of an NCAM-immunoreactive cluster. In this case, the NCAM-immunoreactive cluster and associated intracellular aggregates remained at the site of contact from the moment of its formation. Out of 14 contacts recorded, 4 contacts were formed at the sites where NCAM-immunoreactive clusters and intracellular organelles were already present, whereas 10 of the 14 contacts were initially formed at NCAM-immunonegative sites in the organelle-free areas of neurites.

**FM4-64–loaded TGN organelles are capable of exocytosis**

FM4-64–loaded TGN organelles are the major carriers between the TGN and plasma membranes (Nakata et al., 1998; Toomre...
et al., 1999, 2000; Polishchuk et al., 2000), being capable of fusing with the plasma membrane. To show that TGN organelles loaded with FM4-64 undergo exocytosis in response to external stimuli, we stimulated neurons with 90 mM K⁺. In agreement with previous observations on exocytosis of axonal synaptic precursor organelles (Matteoli et al., 1992; Kraszewski et al., 1995; Dai and Peng, 1996; Zakharenko et al. 1999) and dendritic and somatic TGN organelles (Maletic-Savatic and Malinow, 1998; Maletic-Savatic et al., 1998), this stimulation induced destaining of the FM4-64–labeled organelles in neurites and soma, demonstrating their fusion with the plasma membrane (Fig. 5 b). Exocytosis was observed not only for axonal but also for dendritic organelles, as shown by the absence of synaptophysin immunostaining on most of the processes that display FM dye–labeled organelles (Fig. 5 b). Evoked exocytosis in somatic and dendritic compartments was observed in our experiments starting already from day four in culture, which is earlier than observed by Maletic-Savatic and Malinow (1998). The reason for this could be that hippocampal neurons in our experiments were obtained from postnatal mice and maintained in serum-containing medium, whereas neurons used by Maletic-Savatic and Malinow (1998) were derived from embryonic rat brains and maintained in serum-free medium.

Accumulation of NCAM at sites of contact accompanies synaptic differentiation of initial contacts
To show that accumulation of NCAM accompanies synaptic differentiation of the initial contact, cultures were stained by short-term application of FM4-64 in a solution containing 90 mM K⁺ according to a protocol that is accepted to stain functional synapses (Ryan et al., 1993; Liu and Tsien, 1995). The loading of synapses with FM4-64 was performed three times with a 1-h interval between loadings to allow for the detection of contacts de novo undergoing activity-stimulated endocytosis during this time interval. Because recycling synaptic vesicles and synapse-specific proteins can be found distributed along neurites without any evident contacts (Matteoli et al., 1992; Kraszewski et al., 1995; Dai and Peng, 1996; Rao et al., 1998; Zakharenko et al., 1999; Ahmari et al., 2000), we identified differentiating contacts as contacts between neurites visualized by DIC optics and labeled by acute application of an FM4-64 dye in response to K⁺ stimulation (Fig. 6). Fig. 6 a shows an NCAM180–GFP-transfected neuron and a few synapses distributed along neurites that had been loaded with FM4-64. The second and third loading with FM4-64 (Fig. 6, b and c) at 60 and 120 min afterwards revealed several new endocytosis–competent contacts that had formed during this time period. These events occurred at sites of NCAM180–GFP accumulation (Fig. 6 d). To further show that these contacts are exocytosis competent, we stimulated synaptic activity by application of 90 mM K⁺ for 1 min (Fig. 6 e). This treatment led to complete destaining of the newly labeled contacts, confirming that they were functionally active. Neurons were then fixed and stained with antibodies to synaptophysin (Fig. 6 f), confirming that the observed FM4-64–labeled contacts had accumulated synaptic vesicles and thus likely represent nascent synapses. FM4-64–negative and synaptophysin-positive puncta probably represent functionally immature synapses or transport packages moving along axons (Ahmari et al., 2000). Altogether, we recorded 29 cases of synaptic differentiation. Only three cases were detected at contact sites without accumulation of
NCAM180–GFP, whereas 26 nascent synapses coincided with NCAM180–GFP clusters. The average profile of NCAM180–GFP distribution along neurites showed a peak that was apposed to FM4-64–loaded synapses (Fig. 6 h). Interestingly, two of the three nascent synapses that were formed at contact sites without NCAM180–GFP accumulation were not stained with FM4-64 within 1 h after their appearance, suggesting that they were not stabilized. In contrast, only one of the recorded 26 nascent synapses coinciding with NCAM180–GFP clusters was not stabilized. These results indicate that accumulation and stabilization of intracellular organelles coupled to clusters of NCAM180–GFP accompany synaptic differentiation and suggest that NCAM is important for the stabilization of nascent synapses.

NCAM promotes stabilization of TGN organelles at sites of contact

To investigate whether NCAM plays a role in the stabilization of organelles at sites of contact, we used heterogenotypic cocultures of hippocampal neurons derived from wild-type (+/+) and NCAM-deficient (−/−) mice maintained in coculture for 4 d. Multiple γ-adaptin–positive organelles (red) distributed along neurites are seen (see overlay of DIC image and γ-adaptin staining). NCAM+/+ neurites were identified by NCAM staining. Contacts identified in the DIC image were classified as γ-adaptin positive (circles, solid lines) or γ-adaptin negative (circles, dashed line). (b) A histogram showing the number of γ-adaptin–positive contacts formed by NCAM+/+ axons on NCAM−/− dendrites, NCAM+/+ axons on NCAM−/− dendrites, NCAM−/− axons on NCAM+/+ dendrites, and NCAM−/− axons on NCAM−/− dendrites as percentage of the total number of all contacts analyzed. Data present mean ± SEM (six cultures, two independent experiments). *P < 0.01, paired t test shows a significant difference in the percentage of γ-adaptin–positive contacts between NCAM-negative and other types of contacts. Bar, 10 μm.

NCAM+/+ dendrites, and NCAM+/+ axons on NCAM+/+ dendrites as percentage of the total number of all contacts analyzed. Data present mean ± SEM (six cultures, two independent experiments). *P < 0.01, paired t test shows a significant difference in the percentage of γ-adaptin–positive contacts between NCAM-negative and other types of contacts. Bar, 10 μm.

To further investigate the mechanism of TGN organelle stabilization by NCAM, we analyzed contact formation and organelle accumulation at contact sites in homogenotypic cultures of hippocampal neurons from wild-type or NCAM-deficient mice (Fig. 8). We identified the accumulation of organelles as a persistent association of organelles with the contact until the end of recordings. In cultures of neurons from NCAM-deficient mice, the number of newly formed contacts with accumulated organelles was reduced when compared with cultures from wild-type mice. Whereas 20 of 24 contacts between wild-type neurons accumulated organelles 1–15 min after initial contact formation, NCAM-deficient neurons accumulated organelles at contact sites in only 3 of 18 cases (P < 0.001, chi-square test). The decreased capability to accumulate organelles at sites of contact in NCAM-deficient neurons could be due to the reduced time of contact.
Contacts between NCAM-deficient neurons were more often disrupted due to retraction of neurites when compared with wild-type neurons (10 of 18 newly formed contacts in NCAM-deficient neurons underwent retractions within 5–20 min after contact formation, but only 1 of 24 in wild-type neurons, $P < 0.001$, chi-square test) (Fig. 8 a). These retractions were not accompanied by any signs of damage to neurites and were often followed by regrowth of the retracted neurites in the previous direction. To determine more precisely the role of NCAM in the anchoring of organelles at sites of contact, we estimated the time that organelles spent at contact sites and the persistence of their association with the contact. The time that contacts contained organelles in NCAM-deficient neurons was $\sim 60\%$ of the total recording time, being $\sim 30\%$ less when compared with wild-type neurons ($90\%$ of the total recording time). Moreover, organelles moved away from the contacts approximately four times more often in NCAM-deficient neurons when compared with wild-type neurons (Fig. 8). We conclude that NCAM is important both for stabilization of initial contacts and for stabilization of organelles at sites of contact.

**Discussion**

We show in this study that NCAM mediates accumulation of TGN organelles at sites of cell-to-cell contact. TGN organelles that can form large pleiomorphic structures up to several microns in diameter mediate the majority of cargo transport from the Golgi network to the plasma membrane (Nakata et al., 1998; Toomre et al., 1999, 2000; Polishchuk et al., 2000; Stephens and Pepperkok, 2001). To label TGN organelles in live cells we used the styryl dyes FM1-43 or FM4-64. These dyes have been shown to accumulate in the Golgi or Golgi-like structures after prolonged ($>8$ h) incubation time (Maletic-Savatic and Malinow, 1998; Tarabal et al., 2001). To prove this, Maletic-Savatic and Malinow (1998) correlated the distribution of FM1-43-labeled organelles with NBD C6-ceramide, which localizes to the TGN. Moreover, the combined fluorescence and electron
microscopic analyses show that FM1-43–labeled organelles correspond to the TGN. In agreement with our data, only a small proportion of the FM1-43–loaded organelles were identified as endosomes or lysosomes (Maletic-Savatic and Malinow, 1998). The reason for this is that FM dyes reside in endosomal compartments only very transiently before being transferred to TGN-like structures. This view is supported by data showing that the trans-Golgi network interacts with the recycling endosomal system (Mallard et al., 1998) and that TGN-associated proteins, such as TGN-38, could be found in endosomes (Nakata et al., 1998). Recycling of the mannose-6-phosphate receptor from late endosomes to the TGN is also well characterized (Clague, 1998).

TGN organelles are responsible for the trafficking of a variety of newly synthesized proteins, including synaptic-specific proteins such as synaptophysin and SNAP-25 (Nakata et al., 1998). They are involved in the constitutive and regulated trafficking pathways both in dendrites and axons (Maletic-Savatic and Malinow, 1998; Nakata et al., 1998). Being stabilized in the vicinity of a contact between an axon and dendrite, these organelles provide the proteins necessary for further synaptic differentiation.

The time course of accumulation of synaptic precursor organelles and different synaptic proteins at sites of contact and the molecular mechanisms underlying post- and presynaptic differentiation have been well studied in recent years (Burry, 1986, 1991; Dai and Peng, 1996; Mammen et al., 1997; Rao et al., 1998; Ahmari et al., 2000; Friedman et al., 2000; Lee and Sheng, 2000; Scheiffele et al., 2000; Zhai et al., 2001). We show that NCAM is one of the initial proteins that accumulates at sites of contact within several minutes after contact formation. Our study is the first, to our knowledge, showing that recognition molecules, such as NCAM, provide a direct link between extracellular cues and intracellular organelles to stabilize them at nascent synapses. This finding fits well with recent data on the involvement of NCAM in the mobilization and cycling of synaptic vesicles at the neuromuscular junction (Polo-Parada et al., 2001). The first step in this process is the formation of a complex between clusters of NCAM and TGN organelles before neurites contact each other (Fig. 9). It is therefore noteworthy that organelle-rich varicosities have also been described in the intact tissue (Fiala et al., 1998; Shepherd and Harris, 1998). The complexity between NCAM and intracellular organelles is generally formed outside of the initial contact. Because organelles are driven by intracellular motors, we infer that the mobility of NCAM clusters depends on these.

The link between organelles and cell surface–integrated NCAM180 requires the involvement of peripheral proteins associated with the membrane of the TGN organelle. Here, we show that spectrin is one of these molecules, underscoring a putative mechanism for recent findings in Drosophila on the importance of spectrin in accumulation of synaptic proteins and synaptic transmission (Featherstone et al., 2001). Spectrin has been shown to bind to NCAM180, the largest major isoform of NCAM, but does not bind to NCAM140 (Pollerberg et al., 1986, 1987). Spectrin is highly enriched in neurite varicosities containing vesicular and tubular membranous compartments (Koenig et al., 1985), and the cytoplasmic surface of a variety of intracellular organelles is lined with the spectrin–actin cytoskeleton (De Matteis and Morrow, 1998; Holleran and Holzbaur, 1998; Lippincott-Schwartz, 1998). Spectrin is also tightly colocalized both with TGN organelles and NCAM clusters and mediates binding of the intracellular domain of NCAM180, but not NCAM140, with TGN organelles, in agreement with the experiments by Pollerberg et al. (1986, 1987), which were performed under completely different conditions. It is conceivable that membrane phospholipids are also involved in this interaction (Haest et al., 1978; Sikorski and Kuczek, 1985; McKiernan et al., 1997).

That NCAM indeed mediates the stabilization of initially formed contacts between neurites is underscored by the observation that in heterogenotypic cocultures, contacts on NCAM-deficient neurons resulted in a reduction of synaptic coverage (Dityatev et al., 2000). Data presented in this paper provide further insights into the mechanisms of contact formation, showing that trapping of NCAM-associated TGN organelles represents an early event in synaptogenesis that entails stabilization of contacts. This stabilization could be shown in our study to result in the transformation of the initial contacts into functional contacts undergoing exo- and endocytosis of synaptic vesicles. Analysis of TGN organelle accumulations in heterogenotypic cultures suggests that heterophilic and homophilic interactions align the TGN at sites of contact. Interestingly, in our previous study (Dityatev et al., 2000), heterophilic interactions also appeared to be im-

![Figure 9. A model of NCAM-mediated accumulation of TGN organelles at sites of contact followed by synaptic differentiation.](image322x403to563x732)
important for neurons maintained for 7 d in vitro. In the present study, neurons maintained for 3–4 d in vitro were analyzed, i.e., at stages when the first contacts are formed. The interesting difference between the two studies is that at later developmental stages, only postsynaptically localized NCAM appears to be important for stabilization. These observations imply that in addition to the primary stabilization in the presence of NCAM either pre- or postsynaptically, there may be secondary selection processes, which destabilize synapses expressing NCAM presynaptically, but not postsynaptically. Interestingly, NCAM does not appear to be the only player in the secondary stabilization process, as in homogenotypic cultures comprising only NCAM-deficient neurons, synaptic coverage is as efficient as in cultures containing wild-type neurons (Dityaté et al., 2000). Thus, in a choice situation, NCAM is most likely a preferred candidate for the stabilization of contacts between cells. However, the present study also shows that NCAM may not be the only molecule involved in the stabilization of initial contacts and accumulation of TGN organelles at these contacts, which are transformed into functional synaptic machinery. It is therefore likely that other molecular mechanisms could contribute to the anchoring of intracellular organelles to sites of initial cell contact and thus contribute to the transformation of the nascent synapse to the mature, functionally active synapse.

Materials and methods

Antibodies and toxins

Rabbit polyclonal antibodies (Martin and Schachner, 1986) and rat monoclonal antibody H28 against mouse NCAM recognizing extracellular epitopes of the protein (Gennarini et al., 1984) were generated as previously described. We also used mouse monoclonal antibodies against γ-adaptin and EEA1 (Transduction Laboratories), rabbit polyclonal antibodies against Rab4 and goat polyclonal antibodies against βI spectrin (Santa Cruz Biotechnology, Inc.), rat monoclonal antibody against lamp-1, mouse monoclonal antibody against tubulin and mouse monoclonal antibody 5B8 against NCAM (Developmental Studies Hybridoma Bank, Iowa City, IA), rabbit polyclonal antibodies against erythropoietin, spectrin, mouse monoclonal antibodies against β-COP, rabbit nonimmune immunoglobulins (Sigma-Aldrich), and secondary antibodies against goat, rabbit, rat, and mouse Ig coupled to HRP, Cy2, Cy3, or Cy5 (Santa Cruz Biotechnology, Inc.) and mouse monoclonal antibody against tubulin and mouse monoclonal antibody 5B8 against NCAM (Developmental Studies Hybridoma Bank, Iowa City, IA). Indirect immunofluorescence staining of fixed nonpermeabilized cells was performed as previously described (Dityaté et al., 2000). Antibodies to NCAM were applied to fixed nonpermeabilized cells for 30 min at room temperature and visualized using Super Signal West Pico reagents (Pierce Chemical Co.) on nitrocellulose transfer membrane (PROTRAN; Schleicher & Schuell). Immunoblotting was performed using standard procedures (Venken et al., 1995). For immunofluorescence staining, a monoclonal NCAM antibody was applied in culture medium to live cultures, monoclonal NCAM antibody was applied in culture medium to live cultures for 15 min and detected with fluorescein isothiocyanate-conjugated secondary antibodies applied for 5 min, all in a CO2 incubator. To minimize antibody-induced cross-linking of NCAM in live cells, different concentrations of primary and secondary antibodies were tested to select the concentrations resulting in a staining pattern similar to that obtained after fixation of cells with paraformaldehyde.

Fluorescence labeling of intracellular organelles

The fluorescent dyes FM1-43 or FM4-64 (Molecular Probes) were used to label intracellular organelles mainly of TGN origin (Maletic-Savatic and Malinow, 1998). To label autophagic vacuoles, RITC-dextran (Sigma-Aldrich) was used (Hollenbeck, 1993). Cultures were incubated for 24 h in a CO2 incubator in culture medium containing 10 μM FM1-43 or FM4-64 and 1 mg/ml RITC-dextran and briefly washed four times in culture medium preheated to 37°C. In double labeling experiments, images of FM-stained structures were acquired before the permeabilization step that resulted in washout of the dye.

Colocalization analysis

A membrane area of 0.4–2 μm in diameter with a level of NCAM immunofluorescence intensity that was at least more than two times higher than that of adjacent membranes was identified as an NCAM cluster. An NCAM cluster was considered as overlapping with an organelle when >30% of the cluster area overlapped with the corresponding organelle marker.

Cell surface staining with Dil

Cells were fixed for 15 min at room temperature with PBS containing 4% paraformaldehyde and washed four times, 5 min each, in PBS. The lipophilic dye Dil (Molecular Probes) was dissolved in sesamine oil (Sigma-Aldrich) to saturation (Papa et al., 1995). A drop of this Dil solution (5–10 μm in diameter) was applied through the glass pipette onto the soma of the neuron to be labeled. The dye was allowed to spread over the surface of the neuron for 30 min at room temperature before confocal microscopic imaging.

Labeling of synapses by short-term FM4-64 loading

Cultures were exposed to 15 μM FM4-64 in a stimulating solution containing 31.5 mM NaCl, 90 mM KCl, 5 mM Hepes, 1 mM MgCl2, 2 mM CaCl2, and 30 mM glucose for 1 min in a CO2 incubator (Ryan et al., 1993; Liu and Tsien, 1995). Washing and imaging of neurons was performed in culture medium containing 10 μM 6-cyano-7-nitroquinolxaline-2,3-dione (CNQX) and 50 μM DL-AP5 to block spontaneous synaptic transmission and thus avoid desensitization of the synapses. To show that synapses are capable of active exocytosis, they were desensitized by incubation of cultures in stimulating solution without FM4-64 for 1 min in a CO2 incubator.

Production of intracellular domains of NCAM140 and NCAM180

The BamHI sites were introduced at the 5'-ends of the coding sequence of the EGFP cDNA (CLONTECH Laboratories, Inc.) using standard PCR technique. EGFP was cloned into the unique EheI site (position 2133) of an EcoRI-XhoI subfragment of NCAM180. The BamHI sites were introduced at the 5'-ends of the coding sequence of the EGFP cDNA (CLONTECH Laboratories, Inc.) using standard PCR technique. EGFP was cloned into the unique EheI site (position 2133) of an EcoRI-XhoI subfragment of NCAM180. PCR amplification of EGFP and direction of the EGFP insertion were verified by sequencing. The EcoRI-XhoI subfragment was cloned into the pE Dex vector, which contained the 5' HindIII-EcoRI fragment of NCAM180.

Indirect immunochemistry

Indirect immunofluorescence staining of fixed cells was performed as previously described (Dityaté et al., 2000). Antibodies to NCAM were applied to fixed nonpermeabilized cells for 30 min at room temperature and detected with fluorescein isothiocyanate-coupled secondary antibodies applied for 30 min at room temperature. Antibodies against γ-adaptin, β-COP, synaptophysin, EEA1, Rab4, lamp-1, spectrin, or tubulin were applied after permeabilization of cells with 0.25% Triton X-100 for 60 min at room temperature. The fluorescent dyes FM1-43 or FM4-64 (Molecular Probes) were used to label intracellular organelles mainly of TGN origin (Maletic-Savatic and Malinow, 1998). To label autophagic vacuoles, RITC-dextran (Sigma-Aldrich) was used (Hollenbeck, 1993). Cultures were incubated for 24 h in a CO2 incubator in culture medium containing 10 μM FM1-43 or FM4-64 and 1 mg/ml RITC-dextran and briefly washed four times in culture medium preheated to 37°C. In double labeling experiments, images of FM-stained structures were acquired before the permeabilization step that resulted in washout of the dye.

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Production of intracellular domains of NCAM140 and NCAM180

The BamHI sites were introduced at the 5’ and 3’ ends of the cDNAs encoding the NCAM140 or NCAM180 intracellular domains and were cloned in frame into the BamHI site of pQE30 (QIAGEN). Proteins were expressed in Escherichia coli strain M15 and purified on Ni-NTA-agarose (QIAGEN) according to the manufacturer's instructions.

Construction of the NCAM180–GFP chimera

The NCAM–EGFP chimera was constructed by introducing Ehel sites at the 5’ and 3’ ends of the coding sequence of the EGFP cDNA (CLONTECH Laboratories, Inc.) using standard PCR technique. EGFP was cloned into the unique Ehel site (position 2133) of an EcoRI-XhoI subfragment of NCAM180. PCR amplification of EGFP and direction of the EGFP insertion were verified by sequencing. The EcoRI-XhoI subfragment was cloned into the pE Dex vector, which contained the 5’ HindIII-EcoRI fragment of NCAM180.

Transfection of hippocampal neurons

Cells were transfected 24 h after seeding by the calcium phosphate method (Ethell and Yamaguchi, 1999) using the mammalian transfection kit (Stratagene).

Gel electrophoresis and immunoblotting

Proteins were separated by 8% SDS-PAGE and electroblotted to nitrocellulose transfer membrane (PROTRAN; Schleicher & Schuell) for 3 h at 250 mA. Immunoblots were incubated with appropriate primary antibodies followed by incubation with peroxidase-labeled secondary antibodies and visualized using Super Signal West Pico reagents (Pierce Chemical Co.) on
BIOMAX film (Sigma-Aldrich). Molecular weight markers were prestained protein standards from Bio-Rad Laboratories or Life Technologies.

**Isolation of TGN membranes**

Trans-Golgi membranes were isolated from the brains of 3-mo-old NCAM-deficient mice (Cremer et al., 1994) as previously described (Fath et al., 1997). TGN membranes were resuspended in PEMS (10 mM Pipes, pH 7.0, 1 mM EGTA, 2 mM MgCl₂, and 0.25 mM sucrose) containing complete EDTA-free protease inhibitor cocktail (Roche).

**Binding of NCAM140 and NCAM180 intracellular domains to TGN membranes**

TGN membranes or TGN membranes extracted on ice for 30 min in 0.1 M Na₂CO₃, pH 11.5, were used. The extracted TGN membranes were collected by centrifugation through a 0.5 M sucrose-PKM (100 mM potassium phosphate, 5 mM MgCl₂, 3 mM KCl, pH 6.5) cushion at 25,000 gₑₑₑ for 30 min at 4°C. Pellets were resuspended in PEMS and washed three times. Intracellular domains of NCAM180 and NCAM140 were added to TGN membranes at a concentration of 40 μg/ml and incubated for 30 min at room temperature. To monitor whether TGN membranes bind to the intracellular domain of NCAM180 via spectrin, TGN membranes were incubated with antibodies to erythrocyte spectrin (50 μg/ml) or nonimmune immunoglobulins (100 μg/ml) as a control for 60 min at room temperature. After incubation, the TGN membranes were collected by centrifugation through a 0.5 M sucrose-PKM cushion at 259,000 gₑₑₑ for 30 min at 4°C. Pellets were resuspended in PEMS buffer, washed three times, and analyzed by immunoblotting.

**Online supplemental material**

The supplementary videos (available online at http://www.jcb.org/cgi/content/full/jcb.200205098/DC1) show movement of NCAM-immunoreactive clusters associated with FM4-64–labeled organelles (Video 1), content/full/jcb.200205098/DC1) show movement of NCAM-immunoreactive clusters associated with FM4-64–labeled organelles (Video 1), and the supplementary videos (available online at http://www.jcb.org/cgi/content/full/jcb.200205098/DC1) show movement of NCAM-immunoreactive clusters associated with FM4-64–labeled organelles (Video 1).

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