Development/Plasticity/Repair

**Complexity of BDNF and Neurotrophin-3 on the Firing Patterns and Synaptic Composition of Motoneurons**

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Neurotrophins, as target-derived factors, are essential for neuronal survival during development, but during adulthood, their scope of actions widens to become also mediators of synaptic and morphological plasticity. Target disconnection by axotomy produces an initial synaptic stripping ensued by synaptic rearrangement upon target reinnervation. Using abducent motoneurons of the oculomotor system as a model for axotomy, we report that trophic support by brain-derived neurotrophic factor (BDNF), neurotrophin-3 (NT-3) or a mixture of both, delivered to the stump of severed axons, results in either the prevention of synaptic stripping when administered immediately after lesion or in a promotion of reinnervation of afferents to abducent motoneurons once synaptic stripping had occurred, in concert with the recovery of synaptic potentials evoked from the vestibular nerve. Synaptotropic effects, however, were larger when both neurotrophins were applied together. The axotomy-induced reduction in firing sensitivities related to eye movements were also restored to normal values when BDNF and NT-3 were administered, but discharge characteristics recovered in a complementary manner when only one neurotrophin was used. This is the first report to show selective retrograde trophic dependence of circuit-driven firing properties *in vivo* indicating that NT-3 restored the phasic firing, whereas BDNF supported the tonic firing of motoneurons during eye movement performance. Therefore, our data report a link between the synaptotropic actions of neurotrophins, retrogradely delivered, and the alterations of neuronal firing patterns during motor behaviors. These trophic actions could be responsible, in part, for synaptic rearrangements that alter circuit stability and synaptic balance during plastic events of the brain.

**Key words:** abducent motoneurons; synaptic plasticity; BDNF; NT-3; Trk receptors; axotomy; oculomotor; vestibular

**Introduction**

Neurotrophins provide part of the molecular background that promotes both synaptogenesis and stabilization of existing synapses in the adult brain, and therefore are crucial for plastic events such as regeneration or learning and memory (Vicario-Abejón et al., 2002). These neurotrophin-mediated trophic effects, mostly studied *in vitro*, include the regulation of the number, composition and activity of synapses (McAllister et al., 1999; Mendell et al., 1999), as well as the regulation of firing patterns (Zhou et al., 2005; Youssoufian and Walmsley, 2007). Nonetheless, given the diversity of synaptic organization in the adult brain and the patterns of distribution of neurotrophins and receptors, an *in vivo* approach is required for ascertaining their specific roles under physiological conditions (Schinder and Poo, 2000; Caleo et al., 2003).

Lack of retrograde trophic support induced by axotomy in the adult CNS produces a profound synaptic stripping that decreases the motoneuronal responsiveness to afferent signals (Titmus and Faber, 1990). These alterations normally revert when target reinnervation is allowed to occur or when a new target is provided (Benítez-Temíño et al., 2005). The sequence of events for neurotrophins as target-derived factors is established. Neurotrophins are produced by muscle cells (Steljes et al., 1999; Omura et al., 2005) and then retrogradely and transsynaptically transported to motoneurons (Rind et al., 2005) that are endowed with neurotrophin receptors (Benítez-Temíño et al., 2004). Retrograde neurotrophin uptake and transport increases after nerve injury (Curtis et al., 1998), and neurotrophin signaling reaches the soma which influences cellular functions by different effectors and transcriptional pathways (Zweifel et al., 2005). The abducent motoneurons of the oculomotor system have firing patterns and afferent synaptic connectivity well correlated to the parameters of eye movement (Delgado-García et al., 1986), and, thus, can be used to test whether firing and synaptic properties can be modulated by extrinsic trophic support after target-disconnection by means of axotomy.

If neurotrophins play a causal role, as target-derived factors, in regulating the discharge characteristics via alterations of the synaptic composition, manipulations that either decrease or increase neurotrophin supply should result in altered firing and neuronal sensitivities. Here, to test this prediction, we administered brain-derived neurotrophic factor (BDNF) and/or...
neurotrophin-3 (NT-3) to the stump of the severed axons of abducens motoneurons. Thus, we sought to challenge the synaptic stripping by two ways. We either prevented it by neurotrophin treatment starting at the moment of the lesion or promoted an event-related neurotrophin exposure by performing a second lesion. The difference in the extent of synaptic stripping made the difference clear in the extent of synaptic connectivity. This firing restoration could be prevented by the blockade of the neurotrophin receptor signaling with K252a, which resembles the effects of axotomy on discharge properties of motoneurons. Our data demonstrate that during synaptic remodelling, NT-3 and BDNF play trophic effects on synaptic inputs to motoneurons being NT-3 specific for phasic inputs and BDNF selective for tonic inputs.

Materials and Methods

Animals and surgical procedures. Experiments were conducted on adult cats weighing 2.0–2.5 kg obtained from authorized breeders (Universidad de Córdoba, Córdoba, Spain). Animals were either prepared for the chronic recording of eye movements and neuronal extracellular recordings in the abducens nucleus (n = 12) or those for histology were simply instrumented for the chronic delivery of neurotrophins (n = 10). Surgical and handling procedures for chronic and acute experiments followed the guidelines of the National Institutes of Health (http://oacu.od.nih.gov) and the specific recommendations for the preparation and maintenance of higher mammals during neuroscience experiments (NIH publication #94-3207, 1994) and were in accordance with national legislation for the use and care of laboratory animals (R.D. 120/2005 BOE 252/ 34367-91, 2005).

Animals were surgically prepared for chronic recordings of eye movements and the electrical activity of abducens nucleus neurons. After a protective injection of atropine sulfate (0.5 mg/kg, i.m.) to reduce vagal reflexes, the animal was anesthetized with sodium pentobarbital (35 mg/kg, i.p.) and placed in a stereotaxic frame. Surgery was then performed under sterile conditions to implant stimulating electrodes, acrylic rods, and the recording chamber. Two silver bipolar electrodes were implanted (supplemental Fig. 1, available at www.jneurosci.org as supplemental material) to stimulate the left VIth nerve at its exit from the brainstem and the right medial longitudinal fascicle (MLF) next to the oculomotor nucleus. Coils, made up of 2 turns of Teflon-insulated stainless-steel wire, were implanted in the sclera of both eyes. A square-shaped window (5 × 5 mm) was drilled in the occipital bone to allow transcranial access to the brainstem with glass micropipettes during the recording sessions. A recording chamber made of acrylic resin was constructed around the window, which was sealed between recording sessions. Finally, a pedastal made of dental acrylic with embedded bolts was constructed to immobilize the head during the recordings. Postoperative care was provided to ensure the healthy state of the animal.

Chronic extracellular recordings. After 10–15 d of postoperative recovery, recording sessions started. The animal was restrained with elastic bandages, and placed in a Plexiglas box inside the scleral coil frame. Both the micromanipulator and the head-restraining device were located inside the coil frame. Eye movements were recorded by means of the magnetic field search-coil technique (Püschl and Robinson, 1966). Eye coils were calibrated by rotating at known angles the magnetic field frame relative to the cat.

Extracellular recordings were performed with glass micropipettes bevelled to a resistance of 1–2 MΩ and filled with 2 M NaCl. The (left) abducens nucleus was approached stereotactically and located with the aid of the antidromic field potential produced by electrical stimulation of the VIth nerve. Abducens motoneurons were identified by their antidromic activation from the VIth nerve and by the collision test between the spontaneously occurring orthodromic and antidromically evoked action potentials. Abducens interneuron neurons were identified from the electrode implanted in the MLF. The extracellular neuronal activity was amplified and filtered at a bandwidth of 10 Hz–10 kHz for display and digitalization purposes.

Data storage and analysis. Horizontal and vertical eye position of both eyes and neuronal activity were digitally stored for off-line analysis using a digitizing card (Power 1401, Cambridge Electronics Design). Computer programs were written in Matlab 6.5 to display the histogram of instantaneous firing frequency of the neuronal discharge (i.e., the reciprocal of the interval between two adjacent spikes) and the position of both eyes, and to select data between cursors. Relationships between neuronal firing rate (FR) (in spikes/s) and horizontal eye position (EP) (in degrees) were obtained by linear regression analysis to calculate the slope, i.e., the neuronal sensitivity to eye position (k) (in spikes/s/degree), the intercept (EP) (in spikes/s), i.e., the neuronal firing rate at straight-ahead gaze. Firing rate during fixations responded to the equation FR = k·EP + Fp. Relationships between neuronal firing and eye velocity during spontaneous saccades were also obtained by linear regression analysis after subtraction of the position component (k·EP) calculated from the previously known sensitivity to eye position. Thus, the equation was used as FR = k·EP + r·EV + Fp, where r [in spikes/s/degree/s] is the neuronal sensitivity to eye velocity (EV) (in degrees/s). Due to the impairment of eye movements of the affected eye, we used for computations the eye movement of the contralateral eye.

Synaptic potentials and peristimulus time histograms. Four animals were additionally implanted with silver bipolar electrodes placed in the vestibular nerve bilaterally (supplemental Fig. 1, available at www.jneurosci.org as supplemental material) during either the main surgical procedure or in a second surgical session. Single pulse electrical stimulation to the vestibular nerve with currents ≤50 μA produced typical extracellular field potentials in the abducens nucleus (Pastor et al., 1997) that were positive or negative for the ipsilateral stimulation and mainly negative for stimulation applied to the contralateral vestibular nerve. One or more wavelets representing the afferent volley of activated presynaptic axons were typically seen at short latency after the stimulus artifact. To record extracellular field potentials, filters were set at DC–10 kHz. For constructing the peristimulus time histograms (PSTH), we generated binned histograms (bin width 0.5 ms) spanning 20 ms before and 50 ms after the stimulus for 200–400 cycles of stimulation applied to either the contralateral or the ipsilateral vestibular nerves. The Schmidt-trigger pulse obtained from action potentials was used to construct the PSTH and the stimulus artifact pooled at the time 0 bin, but excluded from computations. The intensity applied was 1.1× the current threshold necessary to evoke a detectable eye movement (Baker et al., 1981; Pastor et al., 1997). Neuronal responses were normalized to calculate the z-scores of a normal probability distribution to detect those bins that were statistically different from the mean of the response previous to the stimulus artifact. The contralateral and ipsilateral responses were considered to represent the brainstem response within the 4–5 ms that followed the electrical stimulus onset. The ipsilateral stimulation produced a desynchronization of the basal firing at the same latency. To quantify PSTH, the area above or below mean before the stimulus application was computed to compare responses among experimental groups.

Abducens nerve section and preparation of the implantable intraocular device. For the delivery of neurotrophins, we implanted the stump of the abducens nerve in a custom-made chamber to control the milieu that bathed it and to prevent axonal reinnervation of any orbital structure or other extraocular muscles (Delgado-García et al., 1988). The intraorbital device consisted of a chamber that had two pieces of tubing inserted for fluid replacement and two electrodes as we recently described (Davila-López de Carriozza et al., 2008). The chamber was made with a standard polypropylene pipette tip shortened to ~15 mm and thus having conical shape. The wider side of the cone had inserted the tip of two 6-cm long pieces of thin wall polytetrafluoroethylene (PTFE) (26 gauge; Small parts). The electrodes for electroneurographic recordings at the stump of the severed nerve were two PTFE insulated pieces, 6 cm long, 1 mm bare tips, made of multistranded stainless steel (located inside the chamber) and a single stranded silver wire (330 μm outer diameter) located outside of the chamber that served as reference electrode (A-M Systems). The tubing and the inside electrode were glued in place to the wide side of the
chamber. The narrow part of the conical chamber was left open to ~1 mm wide to fit tightly the stump of the abducens nerve.

After a few control recording sessions to locate the abducens nucleus, the intraocular device was implanted under general anesthesia in a second surgical procedure. The orbit was approached temporarily through a skin incision in the eyelid canthus at the left side. The abducens nerve was dissected free and the lateral rectus muscle removed. The distal end of the abducens nerve was inserted in the chamber by applying suction through the tubing system. The electrodes and tubing were passed under the skin and were affixed with denatured acrylic to the head retainer. The injection socket was then cemented to the electrodes and a plastic chamber with a knurled cap to protect the tubing endings. The total volume of the system was ~10–15 μl including the dead space of the inlet and outlet tubing.

**Administration of neurotrophins and inhibitor K252a.** The protocol for neurotrophin delivery consisted of single injections of 15 μl in the nerve chamber on alternate days during 4–6 weeks of a solution at a concentration of 15 μg/μl in saline. During the recording session, the chamber was rinsed with sterile saline and followed by the injection in the chamber of a dose of NT-3, BDNF or a mixture or both neurotrophins. The concentrations we used were in accordance with the doses used in studies in vivo (Mendell et al., 1999; Novikov et al., 2000). The selective inhibitor of the tyrosine protein kinase activity, K252a, was delivered at the concentration of 100 nM dissolved in saline with 2% dimethylsulfoxide and the mixture of both neurotrophins (Tapley et al., 1992). Finally, vehicle treatment consisted in the administration alone of sterile saline. Administration of neurotrophins either started the same day of axotomy [immediate administration protocol (IAP)] or delayed 15 d after the axotomy day [delayed administration protocol (DAP)].

**Immunocytochemical procedures for fluorescent confocal microscopy.** Animals were perfused intracardially with physiological saline followed by 4% phosphate buffer paraformaldehyde. The brainstem at the abducens nucleus level was cut at 50-μm-thick coronal sections on a vibratome. The sections were washed in 0.1 M phosphate buffer, pH 7.4, with 0.9% saline and containing 0.1% Triton X-100 (PBS/TX) and then blocked with normal horse serum (1:10 in PBS/TX) and placed in different mixtures of primary antibodies overnight at room temperature. Dual or triple color immunofluorescence for choline acetyltransferase (ChAT) (for motoneurons), synaptophysin (SYN) (for synaptic boutons) and glial fibrillary acidic protein (GFAP) (for astrocytes) was performed by mixing the following primary antibodies and dilutions: ChAT (goat polyclonal antibody (pAb); 1:300; Millipore), SYN (rabbit pAb; 1:500; Zymed Laboratories) and GFAP [mouse monoclonal antibody (mAb) 7a; 1:800, Sigma]. To determine which fraction of the boutons was inhibitory, a series of experiments used the marker vesicular GABA/glycine transporter (VGAT) (Dumoulin et al., 1999; Gonzalez-Foreto et al., 2004), which is the vesicular γ-aminobutyric acid transporter also known as vesicular inhibitory amino acid transporter (VIAAT) that transports GABA and glycine into synaptic vesicles (rabbit pAb; 1:500, Millipore). Labeling of terminals with vesicular glutamate transporters 1 and 2 (VGluT1 and VGluT2) (guinea pig pAb; 1:300–1:1000; Millipore) yielded very low number of boutons on abducens motoneurons and no further attempts to characterize them were done. After several washes, the sections were incubated for 2–3 h in mixtures of secondary antibodies. Immunofluorescence was visualized with donkey anti-rabbit IgG coupled to fluorescein isothiocyanate (FITC) or Cy3, donkey anti-goat IgG coupled to FITC, tetramethylrhodamine isothiocyanate or Cy5 and donkey mouse IgG coupled Cy5 (The Jackson Laboratory). Secondary antibodies were diluted 1:50 in PBS/TX. Finally, sections were washed in PBS and mounted on glass slides and coverslipped with Vectashield (Vector Laboratories).

**Confocal microscopy.** (Leica; TCS SP2) at higher magnification ([×63 oil immersion digitally zoomed at ×2] was used to sequentially capture images at the same focal plane by means of different filters and later merged in Sigma scan or Image J (NIH). Gray scales were adjusted to expand the maximum dynamic range of the image. Neurons included in the quantitative analysis were randomly sampled at high magnification ([×63] in 1–4 different focal planes separated by 2–3 μm in the z-axis around the nuclear plane. Synaptophysin or VGAT boutons were counted to calculate the linear density of boutons. The synaptic coverage was also measured as the percentage ratio between the perimeter occupied by synaptophysin-immunoreactive (IR), VGAT-IR terminals or GFAP-IR profiles over the ChAT-positive somata of motoneurons was also measured. Mean gray value as a measurement of the optical density of the section was also measured using square boxes of 30.4 μm side (80 pixels side) sampled through the neuropil of images captured at ×40. This measurement was considered as an index of the intensity of immunolabeling in the abducens nucleus and computed as the average gray value within the selection after background level subtraction. That is, the sum of the gray values of all the pixels (8-bit resolution) in the selection divided by the number of pixels. For comparisons between groups, one-way ANOVA at an overall level of significance of p < 0.05 followed by post hoc comparisons were performed in SigmaStat version 3.1. Results

Using the axotomy model of the well characterized abducens motoneurons, we tested the hypothesis that target-derived neurotrophic support modulates in adult motoneurons their synaptic composition and firing patterns. Axotomy leads to a profound reorganization of inputs to motoneurons that undergo stripping of synaptic boutons rendering these cells to a very plastic mode that helped us to test the synaptotrophic actions of neurotrophins.

**Axotomy leads to firing alterations in abducens motoneurons.** Control abducens neurons maintained a tonic and regular discharge during eye fixations. Their firing rate increased monotonically for ocular fixations at successive higher angular positions directed toward the ipsilateral side of recording (the on-direction; leftward or temporal-directed for a left-side recorded motoneuron), and decreased or eventually ceased as the eye adopted more eccentric eye positions in the off-direction (Fig. 1A). The slope of the linear regression line between the firing rate and the eye position allowed us to calculate the neuronal sensitivity to eye position (measured in spikes/s/deg) (Fig. 1D, circles). Motoneuronal activity was also related to eye velocity during saccades, thus motoneurons displayed high-frequency bursts during on-directed saccades or pause preceding off-directed saccades (Fig. 1A). The slope of the linear regression analysis between firing rate and eye velocity was the sensitivity to eye velocity, namely r (in spikes/s/deg/s).

The firing pattern of abducens motoneurons (Fig. 1B), but not interneurons neurons (Fig. 1C), was dramatically changed after lesion of the VTh nerve. Changes appeared gradually during the first week that followed lesion and were permanent and more noticeable throughout the second week and continued so during the entire course of the experiment (8–10 weeks). During the initial 15 d postlesion, axotomized motoneurons showed an overall reduction in firing rate. Motoneurons were accompanied also by a severe reduction of modulation related to eye position and velocity (Fig. 1B). Changes in the neuronal sensitivity to eye movement were noticed as reductions in the slopes between firing rate and both eye position and eye velocity (Fig. 1D,E, dots). Plotting the time course of changes in these two neuronal sensitivities (k and r) during axotomy revealed significant differences with control by 2 weeks after lesion (Fig. 1F,G). We used 15 d as a time border between two administration protocols of neurotrophins, because it is known that during these initial 15 d, motoneurons undergo a profound synaptic stripping (our data, see below) (Delgado-Garcia et al., 1988). Thus, if neurotrophins were administered immediately after the axotomy, we hypothesized that synaptic stripping could be, at least partly, prevented, and we termed this procedure the IAP. To the contrary, if synaptotrophic stripping was allowed to occur during the 15 d that ensued.
atotomy, the subsequent neurotrophin administration would in turn facilitate the afferent reinnervation during the so-called DAP.

**Complementary firing restoration after single-neurotrophin administration**

Neurotrophin administration through the IOD directly into the nerve stump of the VIth nerve led to firing restoration in abducens motoneurons that could be appreciated as early as 48 h after treatment. Treatment with BDNF and NT-3 produced distinct changes in the firing pattern of motoneurons. Thus, while BDNF led to the restoration of the tonic firing (Fig. 2B), NT-3, in turn, restored the bursting behavior of motoneurons (Fig. 2C) (see Fig. 2A for comparison with control), without altering in any case (during axotomy of the VIth nerve or during treatment with neurotrophins) the firing of abducens internuclear neurons (data not shown). Thus, BDNF and NT-3 produced complementary aspects of recovery in the firing of abducens motoneurons. BDNF treatment produced a significant recovery of the eye position sensitivity for both the immediate- and the delayed-administration protocols (Fig. 3A). To the contrary, BDNF did not produce any recovery in the eye velocity sensitivity (Fig. 3E) as bursts were of reduced amplitude (Fig. 2B). Sorting the firing rate-to-eye position plots in those fixations obtained after on-directed saccades versus those obtained after off-directed saccades yielded two eye position sensitivities, k-on and k-off. Comparison of the results obtained indicated that both recovered to control values after BDNF or BDNF+NT-3, but not after NT-3 treatment (not illustrated).

A mirror-image pattern occurred after NT-3 administration; in essence, the bursting behavior recovered but not the tonic firing of the motoneuron during spontaneous fixations (Fig. 2C). Quantification of the neuronal sensitivities for both the immediate and delayed administration protocols showed that the eye position sensitivity adopted an intermediate value, different from the axotomy value but different also from the control value (Fig. 3A). However, the eye velocity sensitivity showed a dramatic recovery reaching an average value higher than control (Fig. 3E) as motoneurons showed robust bursts during saccades (Fig. 2C). This supranormal recovery of the eye velocity sensitivity resembled that found in the Ia-spinal motoneuron synaptic efficacy described by Mendell et al. (1999) when using NT-3 in axotomized and also in control motoneurons.

Plotting of the eye velocity versus eye velocity

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**Figure 1.** Effects of VIth nerve axotomy on the discharge properties of abducens neurons with prevented regeneration and reinnervation of a target. **A.** Firing profile of a control abducens motoneuron during spontaneous eye movements. L and R are left and right directions of movement, respectively. **B.** Same as A, but for a motoneuron 20 d after axotomy. Note reduced firing during both saccades and spontaneous fixations. **C.** Abducens internuclear neurons always showed a normal firing pattern after lesion of the lateral rectus nerve. **D.** Calculation of eye position sensitivity for the motoneurons shown in **A** and **B**. The slopes of the regression lines are the neuronal sensitivity to spontaneous eye movements (in spikes/s/deg). Linear relationships were FR = 6.4 EP + 62.6 for the control (circles; r = 0.92; p < 0.001), and FR = 3.7 EP + 7.0 (dots; r = 0.9; p < 0.001) for the axotomized motoneuron. **E.** Same as **B**, but for eye velocity sensitivity. Linear relationships were FR = 0.74 EP (Figure legend continues.)
position sensitivity in control motoneurons demonstrated that a relationship existed and is described by the equation $r = 0.095k + 0.32$ ($r = 0.63; p < 0.001$). Thus, the larger the $k$, the larger was the $r$ value (Fig. 2D, blue) with $r$ being $-1/10th$ of the $k$ value. As a consequence of the treatment with individual neurotrophins (BDNF or NT-3), the population scatterplot was above the control group after NT-3 treatment (Fig. 2D, green), because they were dominated by higher $r$ for a given $k$ ($r = 0.164k + 0.36; r = 0.71; p < 0.001$). To the contrary, treatment with BDNF shifted the population scatterplot below the control group (Fig. 2D, red).

Joint administration of BDNF and NT-3 produces recovery of motoneuronal sensitivities to eye movement

A series of experiments consisted in the simultaneous administration of both neurotrophins. As expected from their individual actions, the recovery was significant for both the eye position and the eye velocity sensitivities and for both types of protocol, either immediate or delayed administration after axotomy (Fig. 3A, B). Firing of motoneurons appeared balanced in burst and tonic modulation, as shown in Figure 4A for an example of a motoneuron axotomized for 28 d and with the immediate trophic support. This result indicated that trophic restoration could be maintained for long-term after lesion and changes could also be seen in the electrophysiographic recordings (Fig. 4A, middle trace). The effects of the trophic support waned after administration ceased and motoneuron firing appeared again axotomy-like as well as the electrophysiographic recording (Fig. 4B). However, trophic support could be restored once the neurotrophin administration resumed and cells turned again into an apparently normal firing pattern with bursts and tonic discharges similar to controls (Fig. 4C). Thus, the joint administration of BDNF and NT-3 restored the normal firing by scaling the appropriate sensitivities to eye position and velocity of axotomized abducens motoneurons. Removal of the trophic support by stopping the neurotrophin administration led neuronal sensitivities to return to the axotomy levels.
We tested whether the actions of neurotrophins on abducens motoneurons were mediated via the signaling of Trk receptors by using the selective inhibitor of tyrosine kinase action K252a in the presence of a mixture of BDNF and NT-3. Recorded cells showed firing patterns that resembled axotomy when neurotrophins were applied together with the inhibitor from the distal end of sectioned axons (supplemental Fig. 2, available at www.jneurosci.org as supplemental material). The eye position and eye velocity sensitivities under the action of K252a decayed to axotomized values. Motoneurons returned to either control values once the inhibitor was eliminated from the treatment or stayed within the axotomized range of values when administration of neurotrophins stopped.

**Neurotrophin-mediated recovery of vestibular synaptic potentials**

We tested synaptic potentials of vestibular origin in abducens motoneurons (Baker and Hightstein, 1975; Delgado-Garcia et al., 1988; de la Cruz et al., 1994; Pastor et al., 1997). Extracellular synaptic potentials elicited from the ipsilateral vestibular nerve (inhibitory) were a positive wave indicative of hyperpolarizing
action at the intracellular level. To the contrary, extracellular potentials elicited contralaterally (excitatory) were seen as a negative wave. As a result of axotomy, the vestibular potentials decreased in some cases almost completely although the presynaptic arrival of the afferent volleys were seen (Fig. 5B, dots). Treatment with NT-3 produced a significant recovery of the ipsilateral-inhibitory potentials (Fig. 5C,E), whereas treatment with BDNF produced mostly a recovery of contralateral-excitatory-synaptic potentials (Fig. 5D,E).

We used these stimuli while recording abducens motoneurons in the awake animal to elicit changes in firing probability immediately after application of the stimulus, thus testing synaptic efficacy on treated motoneurons (Broussard et al., 1995). Electrical stimulation at 1.1× the threshold for detecting eye movements directed toward the contralateral side resulted in transient changes of the discharge rate of abducens motoneurons, that once averaged, produced a characteristic pattern of facilitation and disfacilitation of the discharge rate produced by contralateral and ipsilateral stimuli, respectively (Fig. 6A). For the majority of cells, 200–300 stimuli were used to generate the PSTH. Figure 6B–E, shows the averaged response for each experimental condition using 13–17 cells in each group. Characterization of the bins that were significant using the z-score statistics demonstrated that for the bins within the initial 4–5 ms of the response there was a significant facilitation (for contralateral stimulation) or disfacilitation of the response (for ipsilateral stimulation). We quantified the responses through the computation of the area above or below the mean response previous to the stimulation (Fig. 6F, gray areas). Quantification of the results demonstrated that BDNF produced a significant recovery of the excitatory response (Fig. 6G). Contrary to the results found with the extracellular field potentials, NT-3 did not produce a significant recovery of the ipsilateral–inhibitory response, because the tonic resting level was low and the minimum response in a bin is zero, so the curving by inhibition of the PSTH was not very conspicuous.

The effects of neurotrophin administration to motoneurons studied during the vestibulo-ocular reflex demonstrated that (1) axotomy produced a decay of the neuronal sensitivities to eye position and velocity (supplemental Fig. 3B,C–F, available at www.jneurosci.org as supplemental material), (2) NT-3 produced the recovery of eye velocity sensitivity (supplemental Fig. 3E,F, available at www.jneurosci.org as supplemental material), (3) BDNF produced the complementary recovery of the eye position sensitivity (supplemental Fig. 3E,F, available at www.jneurosci.org as supplemental material), (4) the dual administration produced a full recovery of eye position and velocity sensitivities (supplemental Fig. 3A,C–F, available at www.jneurosci.org as supplemental material) that was (5) abolished by the inhibitor K252a (supplemental Fig. 3E,F, available at www.jneurosci.org as supplemental material).

BDNF and NT-3 prevent synaptic stripping
We found that the functional alterations of axotomy described here were directly related to the degree of synaptic stripping. To quantify these deficits of synaptic coverage, we performed immunohistochemistry for synaptophysin and GFAP which are markers for synaptic boutons and astrocytes, respectively. We also determined the fraction of inhibitory boutons by using immunostaining against VGAT. In the control abducens nucleus, the motoneurons were surrounded in a 48.1 ± 1.8% of the somatic perimeter by synaptophysin-IR boutons and 20.1 ± 1.1% of the perimeter was covered by astroglial filaments (Fig. 7, control; Fig.

Figure 5. Changes in synaptic potentials. A–D, Extracellular field potentials recorded at the center of the abducens nucleus upon single pulse electrical stimulation of the ipsilateral (left side recording) or the contralateral (right side recording) vestibular nerve. In control (A), the inhibitory input produces a positive extracellular wave, whereas the excitatory input produces a negative wave. During axotomy (B), both waves are extremely reduced, but note, however, the presynaptic wavelets (dots) indicative of the afferent volley of presynaptic fibers reaching the abducens nucleus. Treatment with NT-3 (C) restored the positive wave (inhibitory ipsilateral input), whereas BDNF treatment (D) produced the selective recovery of the negative wave (excitatory contralateral input). E, Summary of changes in mean amplitude of the field potentials for the different conditions. Mean and SEM are shown for 6–13 measurements taken between 6–12 d after treatment. Asterisks indicate significant differences with respect to control group (ANOVA test, Holm–Sidak method for post hoc multiple comparisons, p < 0.01).
Approximately 66% of the somatic boutons were VGAT-IR. The consequence of axotomy on surface coverage was that synaptophysin-IR bouton coverage dropped to 21.4 ± 2.1% being a large proportion of boutons VGAT-positive, whereas astrocytic filaments proliferate around the somata and doubled coverage to 40.1 ± 1.6% (Fig. 7, axotomy; Fig. 8A). Treatment with the mixture of BDNF and NT-3 in the immediate administration protocol prevented the drop of synaptophysin-IR and VGAT-IR boutons (Fig. 8A). An intermediate situation occurred by using a single neurotrophin, either BDNF (Fig. 7, BDNF; Fig. 8A) or NT-3 (Fig. 7, NT-3; Fig. 8A). In these cases the prevention of synaptic stripping produced intermediate results since coverage values were both different from control (thus, certain synaptic stripping still occurred) but also different from the axotomy values (Fig. 8A). Neurotrophin treatment with either single neurotrophins or the mixture of two neurotrophins diminished in all cases the astrocytic reaction around the somata of motoneurons (Fig. 8B).

Mean gray value, as a measure of optical density, measured in the neuropil produced a similar pattern of synaptic coverage with synaptophysin-IR and VGAT-IR profiles. The somatic coverage of motoneurons with GFAP was between the control and the axotomy values for the NT-3 treatment and showed a full recovery for the BDNF or the mixture of neurotrophins in the immediate administration protocol (Fig. 8B). The GFAP-IR profiles, however, demonstrated that reactive astrocytes were present in the neuropil for all treatments; therefore, the recovery was partial for treatment with a single neurotrophin but recovery was more profound for treatment with the mix of both neurotrophins (Fig. 8B). Thus, astrocytes were reactive as a consequence of the axotomy in the neuropil but locally withdrew from the surface of the motoneurons concomitantly with the lack of synaptic stripping.

**Neurotrophin treatment promotes afferent bouton reinnervation**

The delayed administration protocol of the mixture of neurotrophins allowed us to test the hypothesis that neurotrophins would have a reinnervation effect once the synaptic stripping had taken place. For doing so, abducens motoneurons were axotomized and 15 d later, once the stripping has taken place, treatment with neurotrophins started until the animals were killed after 30 d. The delayed administration protocol produced the largest recovery of bouton coverage (51.8 ± 1.57%) around motoneuron soma with synaptophysin-IR profiles (Fig. 7, BDNF + NT-3; Fig. 8A). Compatible with these results, the coverage with GFAP-IR profiles was the lowest of all treatments (15.9 ± 0.9%). In the neuropil, the mix of BDNF and NT-3 in the delayed administration protocol produced also the
largest recovery in the optical density (Fig. 8E). These results indicate that neurotrophins used here have a profound synaptic and synaptotropic effect on axotomized motoneurons.

Discussion

Present data show that BDNF and NT-3 administered to the severed axons of motoneurons results in both the prevention of stripping of synaptic afferents when immediately administered after lesion and the recovery of afferent innervation after synaptic stripping due to lesion. Axotomy also induced the decay in neuronal sensitivities to eye movements and a reduction in the amplitude of synaptic potentials of vestibular origin that were also restored to normal values when both neurotrophins were applied. Moreover, the discharge characteristics recovered in a complementary manner when only one neurotrophin was administered. Thus, it is shown the selective trophic dependence of bursting firing on NT-3 whereas BDNF supported the tonic firing of motoneurons. The present data support a link between the synaptotrophic actions of neurotrophins retrogradely delivered and the alter-
Figure 8. Prevention of synaptic stripping and promotion of reinnervation by neurotrophins. A, Changes in synaptophysin-IR (green) and VGAR-IR (red) boutons in the abducens nucleus after axotomy and different treatments. Up-directed bars represent the percentage of covered perimeter of the motoneurons in control, axotomy, single neurotrophin treatment (BDNF or NT-3) and double neurotrophin treatment for either IAP or DAP. Down-directed bars correspond to measurements in the neuropil of the same animals. Bars represent mean ± SEM for 15–45 motoneuronal profiles in each group and 25–50 measurements in the neuropil. B, Same as A, but for measurements of GFAP-IR profiles around somata (up-directed bars) or in the neuropil (down-directed bars). Asterisks indicate significant differences with control, whereas dots indicate significant differences with axotomy. Two-way ANOVA, LInm-Sidak method for pairwise multiple comparisons (p < 0.01).

Neurotrophin complementary action on specific inputs to motoneurons

Our findings show that neurotrophic support starting from the day of lesion prevents the withdrawal of synaptic inputs and that protection against synaptic stripping is more effective when BDNF and NT-3 were applied together rather than when only one neurotrophin alone was used. These results could indicate either an additive effect over all the different afferents to the motoneuron or a complementary effect over distinct afferents. Our neuronal recordings demonstrate the second alternative. Thus, NT-3 supported the phasic firing of motoneurons whereas BDNF supported the tonic firing. BDNF and NT-3 have a strong synaptotropic action regulating the synaptic composition and the growth of presynaptic and postsynaptic elements in the adult brain (Novikov et al., 2000; Cotrufo et al., 2003) and in vitro (McAllister et al., 1997; Je et al., 2006). It is noteworthy that BDNF and NT-3 frequently have opposing roles that account for the specification of electrophysiological properties in the spiral ganglion neurons (Adamson et al., 2002; Zhou et al., 2005) to the growth of dendrites in layer 4 pyramidal neurons (McAllister et al., 1997).

NT-3 produced the recovery of the phasic firing during spontaneous saccades and increased the synaptic coverage of axotomized abducens motoneurons. The phasic component is provided by burst reticular neurons (excitatory and inhibitory; EBN, IBN) located in the paramedian pontine reticular formation (Hightstein et al., 1976; Hikosaka et al., 1978; Igusa et al., 1980) that convey a strong burst during on-directed saccades and a pause during movements in the opposite direction (Büttner and Bütter-Ennever, 2006). It is likely that NT-3 exerts a trophic effect over reticular synaptic contacts on motoneurons that explains the robust bursts during saccades (Fig. 9). Similarly the presence of strong pauses associated to off-directed saccades in-
axotomized spinal motoneurons in vitro and in vivo (Munson et al., 1997; Arvanian et al., 2003). NT-3 derived from muscle spindles or after NT-3 treatment increases the amplitude of this synapse over the control value, but results impaired for sustained high frequency firing (Mendell et al., 1999; Chen et al., 2002).

BDNF induced the recovery of the tonic firing of axotomized abducens motoneurons as well as an increase in their synaptic coverage. The tonic component arises from the output of a velocity-to-position integrator (Aksay et al., 2001) whose main output cells are the burst-tonic neurons of the prepositus hypoglossi nucleus (Escudero et al., 1992) that provide a reciprocal tonic excitatory/inhibitory drive during on- and off-directed eye fixations, respectively (Fig. 9). Chronic treatment with BDNF has been shown to produce a long-term enhancement of synaptic transmission with no changes in voltage-gated sodium and potassium currents (Sherwood and Lo, 1999). Nonetheless, it has been shown that BDNF increases the excitability to depolarizing and hyperpolarizing currents through modulation of potassium currents rendering tonic and rebounding cells in response to current steps (Youssoufian and Walsmey, 2007). BDNF, also in vitro, decreases membrane potential fluctuations promoting rhythmical firing through a K252a-sensitive mechanism that increases a persistent sodium current (Fujisawa et al., 2004) that others have identified as a tetrodotoxin-insensitive sodium current (Blum et al., 2002). BDNF has also been shown to regulate the synaptic composition of axotomized motoneurons (Novikov et al., 2000). From the signal analysis of firing rate we found that both the k-on and the k-off sensitivities were both restored to control values after BDNF treatment alone or in combination with NT-3. This result indicates a trophic action of BDNF over the tonic excitatory and inhibitory input to the motoneurons. If that were not the case, the rate-position plots used to obtain the eye position sensitivity would have a smaller slope than that found (Fig. 9).

A third major source of afferents to the abducens nucleus arises from the medial vestibular nucleus that provides reciprocal, excitatory-contralateral/inhibitory-ipsilateral input to the abducens motoneurons (Fig. 9). The firing of abducens neurons during the VOR can be expressed as the sum of a vestibular input and the output of a velocity-to-position integrator that provides abducens neurons a signal related to eye position. The vestibular signal is channeled through the position-vestibular-pause neurons (Ishizuka et al.,

dicates that also the inhibitory burst neurons are trophically maintained by NT-3, otherwise the rate-velocity plots found would have a smaller slope (Fig. 9). Interestingly, eye velocity sensitivity resulted after NT-3 treatment higher than in control motoneurons. A similar result occurs in the IA synapses over
1980, McCrea et al., 1980) and the tonic signal is also mainly channeled through prepositus hypoglossi and these vestibular neurons (Ramachandran and Lisberger, 2006). We found excitatory vestibular synaptic potentials recovered with BDNF as well as the amplitude of modulation in the PSTH after contralateral vestibular stimulation, indicating that the input from excitatory vestibular neurons contralateral to the abducens would be dependent on the trophic action of BDNF. However, inhibitory vestibular synaptic potentials recovered with NT-3 treatment indicating that the ipsilateral vestibular neurons would be dependent on trophic support from NT-3 (Fig. 9).

The present work opens the possibility for the existence of different motoneuronal types (tonic, phasic) according to their responsiveness to different mixtures of neurotrophins coming from the target. Our previous study (Bényez-Temíno et al., 2004) demonstrated the coexpression of TrkB and TrkC receptors in 72–82% of abducens motoneurons which also implies that 18–28% of the motoneurons might depend solely on either BDNF or NT-3. This would indicate that some control motoneurons might be purely tonic, others purely phasic and the vast majority tonic-phasic. This likely heterogeneity among motoneurons is in contrast, although, with the homogeneous firing patterns found in abducens motoneurons since every cell demonstrates a tonic and a phasic firing (Delgado-García et al., 1986) (present data). Nonetheless, at least two types of motoneurons have been proposed recently on the basis of retrograde and transneuronal tracing (i.e., twitch large cells and small, non-twitch, peripherally located in the nucleus) (Bütter-Ennever et al. 2001; Ugolini et al., 2006). Despite the fact that smaller, non-twitch motoneurons, having thus higher input resistance, could have a similar sensitivity to saccades arising from a reduced number of reticular boutons, our data suggest that twitch and non-twitch motoneurons might express a different number of TrkB and TrkC receptors and/or receive different amounts of BDNF and NT-3 neurotrophins from their innervated muscle fibers, which in turn would modulate the synaptic coverage and firing patterns of abducens motoneurons.

References
Ihizuka N, Mannen H, Sasaki S, Shimazu H (1980) Axonal branches and


Sumner BEH (1976) Quantitative ultrastructural observations of the inhibited recovery of the hypoglossal nucleus from the axotomy response when regeneration of the hypoglossal nerve is prevented. Exp Brain Res 26:141–150.


