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Permalink https://escholarship.org/uc/item/3vr1842m

Journal Journal of Neuroscience, 30(45)

ISSN 1529-2401

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Publication Date 2010-11-10

Peer reviewed



NIH Public Access

Author Manuscript

J Neurosci. Author manuscript; available in PMC 2011 May 10.

Published in final edited form as:

J Neurosci. 2010 November 10; 30(45): 15097–15101. doi:10.1523/JNEUROSCI.3549-10.2010.

BDNF Signaling During Learning Is Regionally Differentiated Within Hippocampus

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Abstract

Learning-induced neurotrophic signaling at synapses is widely held to be critical for neuronal viability in adult brain. A previous study provided evidence that unsupervised learning of a novel environment is accompanied by activation of the TrkB receptor for Brain-Derived Neurotrophic Factor (BDNF) in hippocampal field CA1b of adult rats. Here we report that this effect is regionally differentiated, in accord with 'engram' type memory encoding. A 30-min exposure to a novel, complex environment caused a marked, NMDA receptor-dependent increase in post-synaptic densities associated with activated (phosphorylated) Trk receptors in rostral hippocampus. Increases were pronounced in field CA3a, moderate in the dentate gyrus, and absent in field CA1a. Synapses with Trk activation were significantly larger than their neighbors. Surprisingly, unsupervised learning had no effect on Trk phosphorylation in more temporal sections of hippocampus. It thus appears that commonplace forms of learning interact with regional predispositions to produce spatially differentiated effects on BDNF signaling.

Keywords

spatial memory; NMDA receptor; phosphorylation; dentate gyrus; exploration; post-synaptic; neurotrophin; synaptic plasticity

Introduction

Identification of discrete brain sites associated with the encoding of particular types of memory has been a central theme in neurobiological studies of learning since the 'engram' concept was first advanced early in the last century (Schacter, 1982). While helping to organize work on the storage and retrieval of information, the engram concept also has fundamental implications for work on neurotrophic signaling in brain. A substantial literature has grown up around the broad idea that learning, acting through neurotrophic systems, promotes neuronal viability and thus slows the effects of aging (Valenzuela et al., 2007; Tapia-Arancibia et al., 2008). If so, then it follows from the engram hypothesis that a given form of learning will trigger growth chemistries in discrete portions of the cortical telencephalon; widespread activation would, by the same argument, require diverse forms of experience. The present studies tested the first of these propositions.

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Brain Derived Neurotrophic Factor (BDNF), which is synthesized in an activity-dependent manner by central neurons (Isackson et al., 1991; Castren et al., 1998; Greenberg et al., 2009), enhances neuronal growth and survival in a variety of experimental circumstances (Chao, 2003; Kalb, 2005). Recent work established that the learning-related theta pattern of afferent activity causes rapid activation (phosphorylation) of BDNF's TrkB receptor at synapses in apical field CA1 of adult hippocampal slices (Chen et al., 2010). The effect depended upon the presence of endogenous extracellular BDNF, presumably released during stimulation, and NMDA receptor-driven engagement of Src family kinases (Chen et al., 2010; Huang and McNamara, 2010).

Theta was considerably more effective than other stimulation regimens in the slice studies, suggesting that synaptic BDNF signaling is, as generally assumed, closely related to learning (Minichiello, 2009). Tests of this idea proved positive: A 30 minute session of unsupervised learning in a complex environment caused a robust increase in the number of synapses associated with dense concentrations of phosphorylated TrkB in the same CA1 field evaluated in the slice experiments (Chen et al., 2010). Here we describe results from studies comparing the magnitude of learning-related synaptic Trk activation in the primary subfields of hippocampus (dentate gyrus, CA3, CA1), and at different points along the structure's septo-temporal axis. The results demonstrate striking regional differences in the neurotrophic response of hippocampus to a brief learning episode.

Materials and Methods

All animal procedures were conducted according to the National Institutes of Health Guide for the Care and Use of Laboratory Animals and with protocols approved by the Institutional Animal Care and Use Committee of the University of California at Irvine, including efforts to minimize suffering and numbers of animals used.

Unsupervised learning (USL) paradigm

Procedures were slightly modified from published work (Fedulov et al., 2007; Chen et al., 2010). The USL behavioral box (46 X 91 X 41 cm for length, width, and height respectively) included a large open field and a closed-top dark compartment with a small entrance. Six-week old male Long-Evans rats (Harlan laboratories) were handled and received daily intraperitoneal (i.p.) vehicle (0.9% saline) injections for 4 days. On the fifth day, the rats were injected i.p. with NMDA receptor antagonist 3-(2-carboxypiperazin-4-yl)propyl-1-phosphonic acid (CPP) (10 mg/kg in 0.9% saline) or vehicle 30 min before placement in the video monitored USL box or being returned to home cages. Thirty minutes later, the animals were anesthetized with fluorothane and decapitated. Brains were rapidly removed, frozen in 2-methylbutane (-55° C, \sim 1 min) and cryostat sectioned (20 µm, coronal). Sections were cold-mounted onto slides, gradually warmed to room temperature, fixed in methanol (-20° C, 10 min) and stored at -20° C.

Immunohistochemistry

Sections to be compared were processed simultaneously for double immunofluorescence as described (Chen et al., 2007) using a primary antisera cocktail containing mouse anti-PSD95 (1:1000; Affinity Bioreagents, #MA1-045) and rabbit antisera to phosphorylated (p) Trk^{Y490} (1:200; Cell Signaling, #9141), pTrk^{Y515} (1:200; Abcam, #ab51187), or TrkB (1:1000; Biosource, #611641) and a secondary antisera cocktail containing Alexa594 anti-rabbit IgG and Alexa488 anti-mouse IgG (Molecular Probes). Control sections were similarly processed but with individual primary antisera omitted.

Imaging and quantitative analysis

Immunolabeled sections were examined using an epifluorescence microscope (Leica DM6000) with a 63x PlanApo objective (numerical aperature 1.4) and a CCD camera (ORCA-ER, Hamamatsu). Digital image z-stacks ($0.2 \mu m z$ -steps through $3 \mu m$) were collected from sample fields ($136 \times 105 \times 3 \mu m$) within stratum (str.) radiatum of CA3a, CA1a and CA1b, CA3a str. lucidum and the dentate gyrus molecular layer (internal blade). Images were processed for restorative deconvolution (Volocity 5.0, Improvision) corrected by point-spread functions from a 3D reconstruction of tetraspeck fluorescent microspheres (Invitrogen). In-house built software (Matlab, C, Perl) was used to perform image intensity normalization and segmentation to identify immunolabeled elements and quantify the numbers and volumes of immunoreactive puncta as described (Rex et al., 2009). Element counts from each of 10–15 serial sections per zone (subfield and septo-temporal placement) were averaged to produce an animal mean for that region. For cross-section reconstructions, image montages were collected automatically across a pre-assigned contiguous grid and stitched using Leica Metamorph Acquisition software (Molecular Devices).

Statistics

Unless otherwise stated, values presented are group means \pm sem. Significance of treatment effects was determined using two-tailed t-tests or ANOVAs followed by Tukey's HSD posthoc test (SPSS Statistics, IBM). Frequency and cumulative frequency distributions represent pooled data across all animals within a group. Group frequency distribution differences were assessed using the Kolmolgorov-Smirnov (K-S) z-test followed by the more stringent one-way Kruskal-Wallis non-parametric test to account for differences between animals.

Results

Localization of phospho-Trk in rostral hippocampus

A sizable literature indicates that the hippocampus is critical for unsupervised learning (USL) of novel environments (Roberts et al., 1962; Cohen, 1970; O'Keefe, 1978). We therefore used a variant of this paradigm for the present studies. Rats were handled and acclimated to i.p. vehicle injections for four days. On the following test day, home cage (HC) control and experimental/USL rats were transported to the test room but only the latter were introduced into the complex environment for the 30 min test session. The two groups (HC and USL) were subdivided into subsets pretreated with vehicle or the centrally active NMDA receptor antagonist CPP, at concentrations (10 mg/kg) that block LTP (Abraham and Mason, 1988) and long-term memory in multiple tasks (Ward et al., 1990; Mele et al., 1996), including USL (Fedulov et al., 2007; Chen et al., 2010). In agreement with previous reports (Fedulov et al., 2007; Chen et al., 2010), an analysis of motion tracking data showed that the distance traveled by vehicle-treated animals decreased steadily over the course of the unsupervised learning session (p<0.05 initial 5 min vs. final 5 min; Tukey's HSD). CPP did not measurably influence these time-dependent effects or the mean velocity of movement (p>0.05, vehicle vs. CPP groups, repeated measures ANOVA).

Phospho-specific antisera directed against the conserved Trk Tyr490 residue (Trk^{Y490}), which is phosphorylated with Trk activation, were used to study the effects of learning on BDNF signaling. Most immunostaining for pTrk was localized to small dense structures that were evenly distributed across str. radiatum of hippocampal fields CA3 and CA1 (Fig. 1A; Supplemental Fig. 1). Deconvolved immunofluorescence images showed that pTrk-immunopositive (+) elements were commonly, though not ubiquitously, double-labeled with antisera to the post-synaptic density (PSD) scaffold protein PSD95 (Fig. 1B-C). As described below, only a small percentage of the PSDs were pTrk-immunoreactive.

Blind, automated counting indicated that $3.7 \pm 1.0\%$ (mean \pm sem) of the total PSD population was immunolabeled for pTrk in CA3a str. radiatum of home cage control rats; values for the dentate gyrus molecular layer (DG) and field CA1a were also low $(3.5 \pm 0.4\%)$ and $5.6 \pm 1.8\%$, respectively). Similar results were obtained for home cage rats injected with CPP (CA3a: $4.4 \pm 1.3\%$; DG: $4.6 \pm 1.0\%$; CA1a: $4.3 \pm 1.0\%$), indicating that the NMDAR antagonist by itself had little if any effect on synaptic pTrk levels. Accordingly, we combined the results for the two home cage groups into a single control group. To facilitate comparisons of experimental effects across regions and groups, we expressed all counts from USL rats as a fraction of the mean for each region of the appropriate voked, home cage control cohort. The experimental predictions were that i) 30 minutes of unsupervised learning would increase the number of pTrk+ synapses above that found in home cage controls, and ii) this effect would be reduced by pretreatment with CPP. Automated analysis of pTrk+ PSDs confirmed both points: Mean counts across the three sampling regions were approximately 70% greater in USL-vehicle rats than in home cage controls (p=0.014, t-test, 2-tailed) or USL-CPP groups (p=0.04) (Fig. 2A). As noted, the CPP dose used here did not affect the distance traveled or movement velocity, although it is known from past work to reduce long-term (24 h) retention of USL (Fedulov et al., 2007). Accordingly, the marked increase in double-labeling found in the present USL-vehicle rats is not likely due to locomotor activity.

Learning-related phosphorylation of synaptic Trk varies across subfields

We next tested if the increases in double-labeled PSDs were uniformly distributed across sampling zones in the USL-vehicle group. While values were clearly elevated above control for the DG (1.7 ± 0.3 -fold; p=0.001) and CA3a (2.5 ± 1.1 ; p=0.004), those for CA1a were not detectably different from the control groups (1.0 ± 0.4) (Fig. 2B). The regional differences (among DG, CA3a, and CA1a) in the USL-vehicle group were highly significant (p=0.003; repeated measures ANOVA) and CA3a had more pTrk+ synapses than the DG (p<0.05) or CA1a (p<0.01). The numerical results for CA3a are comparable to those described previously for field CA1b (2.3 ± 0.2 -fold control) (Chen et al., 2010). No differences were observed among subfields for numbers of PSDs double-labeled for total (phosphorylated and non-phosphorylated) TrkB (Supplemental Fig. 2), confirming that the above effects were not due to regional variations in the expression of the receptor itself. These results provide the first evidence that USL-associated increases in Trk signaling occur in a regionally differentiated manner.

Phosphorylated Trk is associated with expanded PSDs in dorsal hippocampus

Certain actin signaling events found at synapses during the production of LTP in hippocampal slices (Chen et al., 2007; Lynch et al., 2008) also occur during unsupervised learning (Fedulov et al., 2007). In both cases, PSD95+ synapses double-labeled for the LTP marker pCofilin were larger than neighboring contacts. The present results provide a second label (pTrk) for testing the idea that LTP-related signaling is associated with synapse expansion in vivo. We therefore measured the volumes of the PSD95+ elements co-localized with dense pTrk^{Y490} immunoreactivity and compared these values with those for surrounding PSDs that were not double-labeled.

Single- and double-labeled PSD volume frequency distributions were constructed from the 1.3 million measurements collected for the DG and CA3a of USL-vehicle rats. The distributions were clearly different for pTrk^{Y490}-negative vs. pTrk^{Y490}-positive PSDs in both sampling zones (Fig. 3A, p<0.000001 each region, K-S test). The pTrk^{Y490} -positive contacts generated a near Gaussian distribution while the much larger population of single-labeled synapses formed a Poisson (Fig. 3A). As expected from these differences, the individual animal median volumes for PSD95+ puncta were markedly different for single-

vs. double-labeled PSDs in each field (Fig. 3B, DG: p=0.004: CA3a: p=0.0004; paired t-test, 2-tails). These effects were also evident in cumulative volume distributions for the three hippocampal subfields in both USL-vehicle and control-vehicle animals (p<0.000001 each region, two-sample K-S test; Supplemental Fig. 3).

Given that previous work established that measures of synapse size are not affected by double labeling *per se* (Chen et al., 2007), the above results support the conclusion that synapses associated with LTP-related markers are significantly larger than surrounding contacts in vivo.

TrkB changes are absent in the temporal hippocampus

Neurobehavioral studies indicate that the rostral hippocampus is critical for spatial learning while more temporal portions are not required for place coding in paradigms lacking an explicit reward (Moser and Moser, 1998; Bast et al., 2009; Fanselow and Dong, 2010). We therefore tested if increases in pTrk found in rostral hippocampus are equally well developed in more temporal regions. Serial coronal sections were evaluated from two segments along the septo-temporal axis, beginning 3 mm (mid-temporal) and 5 mm (temporal) posterior to bregma (Paxinos, 1986).

In marked contrast to the more than two-fold increases in pTrk+ PSDs obtained in rostral hippocampus (Chen et al., 2010), there was no evident effect of learning on Trk phosphorylation in field CA1b in temporal hippocampus (fold of home cage controls: HC-vehicle vs. USL-vehicle: 1.0 ± 0.1 vs. 1.1 ± 0.3 ; n=12 animals/group). Indeed, the number of double-labeled synapses in CA1b was not detectably different across the two home cage (vehicle, CPP) and two USL groups (Fig. 4A).

Given that CA1a did not show a learning effect in rostral hippocampus, and the possibility that boundaries between CA1a and CA1b might differ across the septo-temporal axis, we repeated the experiment by counting double-labeled synapses in the clearly defined temporal CA3a. We again found no evidence for a USL-induced increase in pTrk+ contacts (p>0.3) (Fig. 4B).

Combined, the results indicate that the robust effects of USL on BDNF signaling in rostral hippocampus do not occur in more temporal regions.

Discussion

The present results indicate that a form of learning common to mammals, and prominent in humans, is associated with the activation of synaptic BDNF receptors in hippocampus and that synapses with active Trk signaling are larger than their neighbors. The increases in Trk activation were likely related to encoding, rather than to locomotor activity, because they were blocked by an amnestic dose of an NMDA receptor antagonist that does not cause changes in the distance traveled in the complex environment. The same drug treatment had no measurable influence on basal pTrk immunolabeling in home cage control rats, so its actions in the learning paradigm can be attributed to a selective suppression of behaviorally induced increases. The larger volume of pTrk+ synapses relative to surrounding contacts lacking Trk activation is novel but accords with previous evidence for an enlargement of synapses exhibiting LTP-related signaling in vitro and in vivo (Chen et al., 2007; Fedulov et al., 2007).

The learning-related increases in pTrk, while pronounced, were unexpectedly restricted in their spatial extent. With regard to subdivisions of hippocampus, increases in pTrk+ PSDs were evident in str. radiatum of CA3b and CA1b, present but less pronounced in the dentate

molecular layer, and absent from apical CA1a. More striking still, the effects obtained in fields CA1b and CA3a of rostral hippocampus were absent in more temporal regions of the structure. These results provide the first evidence that USL-induced, synaptic BDNF signaling is restricted to a relatively small portion of hippocampus.

What mechanisms could account for marked regional differences in neurotrophic responses within the same forebrain structure? Recent work has shown that LTP is less robust in temporal than in dorsal hippocampus and suggests that this is due to differential distributions of mineralocorticoid receptors (Maggio and Segal, 2009). There is also evidence that the LTP threshold is higher in temporal regions, possibly because of differences in adenosine 'tone' (Colgin et al., 2004). Given that the conditions needed for synaptic TrkB responses and LTP induction overlap extensively (Chen et al., 2010), these observations raise the possibility that the threshold for TrkB activation increases along the septo-temporal axis of hippocampus; if so, then the same USL-elicited firing patterns that pass threshold for TrkB activation in septal fields may fail to do so more temporally.

Pertinent to the above, we recently found short trains of theta burst stimulation to be the most effective pattern for activating synaptic TrkB among a set of afferent stimulation regimens, including a high frequency train, using the same number of electrical pulses. The mechanisms underlying the differential efficacy involved an NMDA receptor-initiated cascade targeted at BDNF receptors (Chen et al., 2010). Given these points, it is of interest that theta activity power and the incidence of spatially tuned neurons are both reduced in the temporal portion of the hippocampus relative to more rostral areas (Buzsaki, 2005). Possibly then, a relative weakness of theta activity, and the burst firing that accompanies it during behavior (Otto et al., 1991), results in a lower probability of engaging synaptic TrkB signaling in temporal hippocampus.

There have been no studies identifying optimal stimulation parameters for eliciting TrkB phosphorylation, or stable LTP, in temporal hippocampus. If regional differences do exist, then a synthetic hypothesis would be possible in which different types of learning trigger characteristic patterns of input to the hippocampus, and thereby elicit TrkB activation in portions of the structure 'tuned' to that pattern but not in areas responsive to other signals.

Functionally, the spatially restricted distribution of synaptic effects suggests that the hippocampus is economical with regard to neuronal resources used to store new information. The learning episode lasted for only 30 minutes and it is conceivable that different kinds of material acquired in the complex environment over additional days of testing would engage CA1a or regions further down the septo-temporal axis. The hippocampus could, for example, use space to segregate the types of data acquired through USL (e.g., spatial cues on day 1 vs. objects on day 2) or to maintain a record of when particular memories were encoded. Along this line, there is evidence that ventral hippocampus is involved in encoding the sequence of odor cue presentation (Hunsaker et al., 2008; Kesner et al., 2010). More generally, the striking absence of Trk phosphorylation in temporal areas is consonant with past arguments that different portions of hippocampus are involved in different types of memory. Results from lesion and chronic recording experiments suggest that the rostral, but not temporal, hippocampus is required for acquisition of spatial information in the absence of explicit reward (Morris et al., 1982; Packard and McGaugh, 1996), while temporal hippocampus plays a critical role in reward-based learning (Bast et al., 2009), something that might be expected given its close relationship to the amygdala (Monfils et al., 2007).

The present findings also raise a series of novel questions about the function of learningrelated neurotrophic signaling at synapses. Increases in the number of synapses labeled for activated Trk were large relative to the low values found in home cage rats but nonetheless involved a very small percentage of the total synaptic population. Is this sufficient to trigger genomic responses in the neurons on which the Trk response occurred? Answering this experimentally tractable question would be a significant step in evaluating the argument that learning plays a critical role in maintaining the viability of neurons.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

Work was supported by NIMH grant MH082042 to C.G., NINDS grant NS45260 to C.G. and G.L., ONR MURI award N00014-10-1-0072 to G.L., NIMH Fellowship MH083396 to L.C., and NINDS fellowship NS045540 to C.R. We thank Rishi Mehta, Ignacio Rivera and Katherine M. Wong for expert technical assistance.

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Figure 1.

Phosphorylated Trk is localized to postsynaptic densities. **A**) Montages of photomicrographs collected from dorsal hippocampus show DAPI labeling of nuclei (top; image intensities inverted), immunofluorescence labeling for PSD95 (green) and pTrkB^{Y515} (red), and a merger of PSD and TrkB^{Y515} immunolabeling (sr: str. radiatum. Calibration bar: 500 μ m). The pTrkB^{Y515} and pTrk^{Y490} immunoreactivities have similar distributions in hippocampus (Supplemental Figure 1; also, Chen et al, 2010). **B**) Representative deconvolution micrographs show immunofluorescence for pTrk^{Y490} (red) and PSD95 (green) in CA3a str. radiatum. Arrows indicate double-labeled elements (yellow in 'merge'). Calibration bar: 10

 $\mu m.$ C) High magnification of a single synapse double-labeled for $pTrk^{Y490}$ (red) and PSD95 (green). Calibration bar: 1 $\mu m.$

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Figure 2.

Learning-related increases in numbers of pTrk^{Y490}+ synapses are regionally differentiated in rostral hippocampus. The mean (\pm sem) number of double-labeled synapses (pTrk(+)PSDs) in the sample field is expressed as a ratio of the home cage (HC) control mean (indicated by dashed line). **A**) Overall numbers of double-labeled synapses (cumulative across three sampling zones) were significantly greater in vehicle (veh)-treated rats that explored the complex environment for 30 min (USL veh) than in HC controls. Treatment with CPP prior to exploration (USL CPP) eliminated this increase (n=12–14/group; two-tailed t-tests). **B**) Plot shows that increases in pTrk+ PSDs in the USL-veh group were not equally distributed across hippocampal subfields (ANOVA: p=0.003): Values from USL-veh rats were greater than from HC controls for the dentate gyrus molecular layer (DG) and CA3a (p<0.01) but not for CA1a (p>0.4).



Figure 3.

Synapses associated with pTrk^{Y490} are larger than neighboring, single-labeled synapses. **A**) Volume frequency distributions for PSD95-immunoreactive elements (PSDs) that were double-labeled (black), or not double-labeled (grey) for dense pTrk^{Y490} immunoreactivity in the dentate gyrus molecular layer (DG) and CA3a str. radiatum. The pTrk+ PSD volumes were normally distributed, whereas pTrk-negative PSD volumes followed a Poisson distribution. (For each region: n>3,400 and >197,000 for double- and single-labeled PSDs, respectively; p<0.0000001, K-S test) **B**) Bar graph shows the mean (\pm sem) of the median volumes (per individual) of double-labeled (pTrk(+)PSD) and single-labeled (pTrk(-)PSD)

PSDs for the HC and USL-veh groups. (n>40 sections from 4 rats each; *p=0.004, **p=0.0004; paired two tailed t-test).

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Figure 4.

Unsupervised learning (USL) does not increase numbers of $pTrk^{Y490}$ + synapses in temporal hippocampus. Summary of double-labeled synapses (pTrk(+)PSDs) in CA1b (A) and CA3a (B) str. radiatum of temporal hippocampus expressed as a function of the home cage control mean (HC con). A) Plot shows that 30 minutes of USL, with or without CPP treatment, had no effect on numbers of pTrk+PSDs in temporal CA1b. B) There were no significant differences in numbers of pTrk+PSDs in temporal CA3a among the four groups (p>0.3 for HC-CPP rats and the two USL groups vs. HC controls).