pCREB in the Neonate Rat Olfactory Bulb Is Selectively and Transiently Increased by Odor Preference–Conditioned Training

John H. McLean,1,3 Carolyn W. Harley,2 Andrea Darby-King,1 and Qi Yuan1

1Division of Basic Medical Sciences and 2Department of Psychology, Memorial University of Newfoundland, St. John’s, Newfoundland, Canada A1B 3V6

Abstract

Early olfactory preference learning in rat pups occurs when novel odors are paired with tactile stimulation, for example stroking. cAMP-triggered phosphorylation of cAMP response element binding protein (pCREB) has been implicated as a mediator of learning and memory changes in various animals (Frank and Greenberg 1994). In the present study we investigate whether CREB is phosphorylated in response to conditioned olfactory training as might be predicted given the proposed role of the phosphorylated protein in learning.

On postnatal day 6, pups were trained for 10 min using a standard conditioned olfactory learning paradigm in which a conditioned stimulus, Odor, was either used alone or paired with an unconditioned stimulus, Stroking (using a fine brush to stroke the pup). In some instances stroking only was used. The pups were sacrificed at 0, 10, 30, or 60 min after the training. Using Western blot analysis, we observed that the majority of olfactory bulbs in conditioned pups (Odor + Stroking) had a greater increase in pCREB activation at 10 min after training than pups given nonlearning training (Odor only or Stroking only). The phosphorylated protein levels were low at 0 min and at 60 min after training. This is in keeping with the slightly delayed and short-lived activation period for this protein.

The localization of pCREB increases within the olfactory bulb as seen by immunocytochemistry. Naive pups were not exposed to odor or training. There was a significantly higher level of label in mitral cell nuclei within the dorsolateral quadrant of the bulb of pups undergoing odor-stroke pairing. No significant differences were observed among nonlearning groups (Naive, Odor only, or Stroking only) or among any training groups in the granule or periglomerular cells of the dorsolateral region. The localized changes in the nuclear protein are consistent with studies showing localized changes in the bulb in response to a learned familiar odor. The present study demonstrates that selective increases in pCREB occur as an early step following pairing procedures that normally lead to the development of long-term olfactory memories in rat pups. These results support the hypothesized link between pCREB and memory formation.

Introduction

The recently hypothesized role for cAMP response element binding protein (CREB) as a universal modulator of processes required for memory formation is supported by a set of empirical studies in which CREB levels are manipulated. Enhancement or attenuation of CREB levels, respectively, enhance or attenuate aversively motivated, and more recently, appetitively motivated, long-term memories (Silva et al. 1998). In particular, CREB levels have been shown to critically influence memories for olfactory avoidance conditioning in fruit flies (Yin et al. 1995; Tully 1996), defensive withdrawal conditioning in Aplysia (Abel and Kandel 1998; Bartsch et al. 1998; Michael et al. 1998) and avoidance conditioning, spatial escape learning, and social preference learning in rodents (Silva et al. 1998). CREB has also been shown to be phos-
phorylated in songbirds which hear a song from their own species (Sakaguchi et al. 1999).

The pathway by which CREB is hypothesized to be recruited during learning has been outlined by Frank and Greenberg (1994). Exposure to a neurotransmitter leads to the generation of increased amounts of cAMP, allowing the catalytic subunit of protein kinase A to translocate to the nucleus. In the nucleus, protein kinase A phosphorylates CREB at the Ser-133 site. Phosphorylated CREB (pCREB), in turn, can activate the transcription of immediate-early genes. The protein products of the immediate-early genes are transcription factors that go on to activate the transcription of late response genes such as brain-derived neurotrophic factor (Tao et al. 1998). It is these late response genes that are likely to be responsible for changes in neurotransmitters, neurotrophins/receptors, adhesion molecules, and cytoskeletal protein—the necessary components of long-term plasticity (learning) changes.

From this sequence it follows that if CREB has a critical role in initiating the genomic transcription underlying long-term memory, then CREB phosphorylation should be an early step in the development of long-term memories. Translocation of a protein kinase A subunit to the nucleus of Aplysia sensory neurons following serotonin application has been demonstrated as critical for serotonin facilitation (Abel and Kandel 1998), implying that an increase in CREB phosphorylation occurs in this form of plasticity. In mammalian studies, increased levels of pCREB have also been demonstrated to accompany increases in synaptic strength in several model systems (Bito et al. 1996; Deisseroth et al. 1996, 1998; Impey et al. 1996; Moore et al. 1996). However, no mammalian studies have demonstrated CREB phosphorylation that is linked specifically to associative learning procedures.

In rat neocortex, the level of pCREB is normally increased significantly during the waking phase of the daily cycle. This increase depends on intact locus coeruleus innervation (Cirelli et al. 1996). A critical role for noradrenergic locus coeruleus input in supporting CREB phosphorylation is consistent with the role of cAMP and the catalytic unit of protein kinase A in mediating phosphorylation. Calcium can also mediate CREB phosphorylation, but the failure to see normal pCREB elevation in the adult brain following noradrenergic lesions suggests calcium may not be sufficient.

In the present study we measure the occurrence of CREB phosphorylation in an olfactory learning paradigm in neonate rats in which noradrenergic input from the locus coeruleus has been critically implicated. Stroking of a pup is known to activate cells of the noradrenergic locus coeruleus (Nakamura et al. 1987) and an olfactory preference is conditioned in neonate rats by pairing stroking with a novel odor (Sullivan and Leon 1986b, 1987). The acquisition of a preference for the novel odor requires activation of β adrenoceptors in the rat olfactory bulb (Sullivan et al. 1989a). Thus, the odor conditioning is dependent on a noradrenergic unconditioned stimulus signal coupled to cAMP increases which could initiate CREB phosphorylation through translocation of the catalytic subunit of protein kinase A. The glutamatergic odor input (Berkowicz et al. 1994; Ennis et al. 1996) occurring simultaneously with β-adrenoceptor activation could further heighten CREB phosphorylation through the synergistic actions of calcium on Ca2+-dependent kinases, which also induce CREB phosphorylation (Deisseroth et al. 1996; Tao et al. 1998). CREB phosphorylation, if sufficiently sustained, should activate the genomic transcription necessary to support the long-term odor preference memory. To test the hypothesis that CREB phosphorylation is selectively triggered by odor preference acquisition, pups were either given Odor + Stroking pairings, which reliably lead to odor preferences in tests 24 hr later, or Odor only or Stroking only, which does not lead to odor preference in subsequent testing. Pups were sacrificed at several intervals following training and assessed for pCREB levels by Western blot. Neuroanatomical distribution of pCREB was also visualized using immunocytochemistry. We found that pCREB protein was increased transiently in the olfactory bulb after olfactory preference conditioning and that the pCREB increases occurred in a region associated previously with processing of the preferred odor. These results support a selective role for CREB phosphorylation as an initial step in mammalian learning.

Materials and Methods

ANIMALS

Sprague-Dawley rats were used in this study, which was approved by the Institutional Animal Care Committee of Memorial University of Newfoundland. Litters were culled to 12 pups on postnatal day (PND) 1, whereas the day of birth is con-
sidered PND0. A total of 27 pups from 10 litters (for Western blot analysis) and 24 pups from 6 litters (for immunocytochemistry) were used in this study. These numbers were the minimum possible to provide statistically significant results. A maximum of one male and female pup from each litter was used for each training group described below.

CONDITIONING EXPERIMENTS AND SACRIFICE

Rat pups (PND6) were subjected to conditions that had been shown previously (McLean et al. 1993; Langdon et al. 1997; Price et al. 1998) to produce conditioned olfactory learning (Odor + Stroking) or not (Naive, Odor only, Stroking only). To this end, the pups were placed on normal (Stroking only) or peppermint-scented bedding (250 µl/300 ml of bedding) and stroked vigorously on the hind region using a paintbrush every other 30 sec for 30 sec (Odor + Stroking) or not (Odor only) over a 10 min period. The pups in the present experiment were subjected to training procedures but were not tested for recall because we wished to examine the role of pCREB specifically in the acquisition of olfactory learning. In previous studies, we (McLean et al. 1993, 1996; Langdon et al. 1997; Price et al. 1998), as well as others (Sullivan and Leon 1986a, 1987; Sullivan et al. 1989a, 1991), have shown that pups subjected to the different training procedures show predictable behavioral outcomes.

For Western blot analysis after the various training conditions (Odor + Stroking, Odor only, Stroking only, or Naive), the pups were sacrificed by decapitation at various intervals. The post-training time giving the strongest pCREB induction in Western blot analysis was used as the sacrifice time in subsequent experiments. For immunocytochemical analysis of pCREB, pups were sacrificed 10 min after training (Odor + Stroking, Odor only, Stroking only, or Naive) by overdose with sodium pentobarbital and perfused transcardially with fixative as described previously (McLean et al. 1993).

PROTEIN PREPARATION AND DETERMINATION FOR WESTERN BLOTS

After sacrifice, both olfactory bulbs were immediately frozen on dry ice and subsequently stored in microcentrifuge tubes at −70°C until use. Ice-cold lysis buffer containing 0.1% SDS, 1% NP-40, 20 mM Tris, 10% glycerine, and 1.37 mM sodium chloride with 1 µg/ml leupeptin, 2 mM PMSF, 8.9 U/ml aprotinin, and 1 mM sodium orthovanadate was added to each tube and each pair of bulbs was then ground using a homogenizer. The homogenate was left on ice for 30 min and then centrifuged at 13,000 rpm for 15 min at 4°C. The clear lysate containing the proteins of interest was stored in 50-µl aliquots at −70°C. The protein concentration of each pair of olfactory bulbs was determined separately using a Pierce BCA Protein Assay kit.

WESTERN BLOTTING

Samples were prepared in 4 µl of 5× sample buffer (0.25 M Tris-HCl, 10% SDS, 50% glycercer, 0.025% bromophenol blue, and 0.5 mM dithiothreitol added just prior to use) plus sufficient sample and water to constitute 20 µl of sample per well. Positive controls using forskolin-treated PC12 cells showed the appropriate band for pCREB. Samples and positive controls were boiled for 5 min and placed on ice until use. Equal weights of protein (100 µg) from each sample were loaded into a lane. Proteins were separated on the basis of gel mobility using SDS-PAGE. Color-coded molecular weight standards (Bio-Rad) were loaded into a separate lane for each blot. The gel-running buffer was a solution containing 25 mM Tris, 250 mM glycine, and 3.5 mM SDS at pH 8.3. Once proteins were loaded into one of 10 lanes, each gel apparatus was attached to a Bio-Rad power supply set to 100 mV. The protein was then transferred at 4°C from the gel to nitrocellulose paper (Hybond ECL, Amersham) using a Bio-Rad power supply set to 0.2 A for 45 min. The transfer solution contained 25 mM Tris, 192 mM glycine, and 20% methanol. After the transfer, the nitrocellulose blots were processed for detection of pCREB or placed in Ponceau S to visualize the total amount of protein transferred. Ponceau S was used on every blot (either after protein transfer or immunoblot) to determine if equal amounts of protein were loaded for comparisons.

The blots were rinsed 3 × 5 min in PBS with 0.05% Tween 20 (PBST) and then immersed in 5% milk in PBS for 1 hr. Following 3 × 5 min rinses in PBST, the blots were immersed in a polyclonal antibody to pCREB (1/1500 dilution, Upstate Biotechnology) with PBST overnight on a shaker at 4°C. Following further washes in PBST and incubation (1 hr) in anti-rabbit IgG-horseradish peroxi-
The blot was rinsed in PBST and visualized using enhanced chemiluminescence (Amersham).

**IMMUNOCYTOCHEMISTRY**

Standard immunocytochemical methods were used as described previously (McLean et al. 1993). In brief, 30-µm frozen coronal sections through the bulb were cut and mounted on subbed slides (all sections were kept). The primary pCREB antibody was diluted 1/1500 in PBS containing 0.2% Triton X-100 and 2% normal goat serum. Following washes as described previously (McLean et al. 1993), the sections were incubated in biotinylated secondary antibody (Vectastain Elite) and then in a 0.05% solution of diaminobenzidine with 0.01% H₂O₂.

**ANALYSIS OF WESTERN BLOTS AND IMMUNOCYTOCHEMICAL SECTIONS**

Only those blots that showed equal protein levels for paired bulbs were analyzed subsequently. The average optical density (avg = integrated density value/area) of Western blots was determined using ChemiImager (Alpha Innotech Corp.). The integrated density value is the sum of all pixel values on a scale from 0 (white) to 255 (black) in the defined region and the area is the size in pixels of that defined region. The background integrated density was subtracted automatically from each blot area. The average optical densities were analyzed using a one-way repeated measures ANOVA. Tissues that were processed in the same Western blot run were grouped together. Using these criteria the analysis included seven triple sets (Odor + Stroking versus Odor only versus Stroking only). Additionally, three double sets (Odor + Stroking versus Odor only) were analyzed but could not be included in the repeated measures ANOVA because of the statistics criteria. Post hoc comparisons of the triple sets were made using the Newman–Keuls test.

For immunocytochemistry, the sections were analyzed systematically such that every third section through the bulb was examined. Image analysis (Bioquant, R&M Biometrics, Inc.) was performed by tracing the region/cell layer of interest. Labeled cells were then captured digitally using a background subtraction and threshold method to eliminate any uneven lighting conditions. The percent area reaching threshold in the region of interest was determined for each olfactory bulb. In addition, the number of cells reaching threshold were counted in the mitral cell layer. Threshold refers to pixels (which are converted to area) that fall within a given brightness level assigned to a gray level scale from the object of interest and provides a systematic and reliable means of identifying labeled cells. To assure objectivity, the slides were coded so that the person analyzing each section was unaware of the training procedure. The data were analyzed statistically by one-way ANOVA and the Newman–Keuls post hoc test.

**Results**

**WESTERN BLOT ANALYSIS**

**IDENTITY AND SENSITIVITY OF THE PCREB BAND**

In Western blots, PC12 cells treated with forskolin (known to increase pCREB levels) showed a band at 43 kD that was substantially more intense than that observed in untreated PC12 cells (Fig. 1). This observation demonstrates the specificity of the antibody for pCREB and confirms that the technique is sensitive to changes in pCREB. Furthermore, a band located at the same molecular mass (43 kD) was observed in the olfactory bulbs of PND6 pups (Fig. 1).

**TIME COURSE OF PCREB EXPRESSION**

Immediately after training or in naive pups, pCREB levels were very low (Fig. 2). Levels of pCREB were highest 10 min after training and began to subside at 30 min and 60 min (Fig. 2).

**PCREB PROTEIN LEVELS IN OLFACTORY BULBS OF CONDITIONED VS. NONCONDITIONED PUPS**

The olfactory bulbs of pups that were trained to learn (Odor + Stroking) generally showed in-
creased levels of pCREB 10 min after training (Figs. 3 and 4) compared to pups subjected to a non olfactory learning situation (i.e., Odor only or Stroking only). The one-way ANOVA revealed a significant effect of groups ($F_{2,12} = 4.6, P = 0.033$). In the Newman–Keuls test the Odor + Stroking pairing group (average density = 65.3) was significantly higher ($P < 0.05$) than either the Stroking only (average density = 44.0) or Odor only (average density = 44.9) groups, which did not differ from each other. In the three additional pairs of Odor + Stroking (average density = 67.3) and Odor only (average density = 49.9) pups, the results were similar to those for the larger group. Although the odor conditioned pups showed significantly greater pCREB (Fig. 4) 10 min after the training period compared to the nonlearning situations, not all bulbs exhibited the increase. When data from samples run on the same blots were compared, Odor + stroking levels were greater than Odor only in 9 out of 10 comparisons while the Odor + Stroking conditioned pups showed increased pCREB levels in 6 out of 7 comparisons with Stroking only pups.

**IMMUNOCYTOCHEMISTRY**

There were many cells labelled in the olfactory bulb of all animals examined, but the mitral cells were significant in label intensity. The dorsolateral quadrant, in particular, appeared to have increased labeled cells and was, thus, analyzed in some detail (Fig. 5). In addition, the ventromedial quadrant, which appeared to have few mitral cells labeled, was analyzed in detail. Quantitative analysis of nuclei areas reaching threshold revealed a significantly higher level of pCREB label within the mitral cell layer in the dorsolateral quadrant of pups subjected to the learning (Odor + Stroking) paradigm compared to pups which received no training (Naive), Odor only, or Stroking only. The ventromedial quadrant showed less density of mitral cell label and no significant difference in label among learning (Odor + Stroking) and nonlearning (Naive, Stroking only, Odor only) conditions (Fig. 6). Neither the periglomerular nor granule cell populations within those same regions showed any significant differences among training groups (Fig. 6). The number of mitral cells reaching threshold in the dorsolateral quadrant was also analyzed and, similar to the analysis of threshold areas, one way ANOVA showed significance ($F_{3,20} = 3.20, P = 0.045$). The post hoc Newman–Keuls test revealed that the density of cells in the Odor + Stroking

---

**Figure 2:** Western blot showing the change in pCREB expression (arrow) over time following conditioned olfactory training. (O/S) Odor + Stroking (learning); (O/O) Odor only; (Naive) not exposed to peppermint odor or stroking. Note that the most intense activation is ~10 min after training.

**Figure 3:** Representative Western blot showing pCREB levels in the olfactory bulbs of pups subjected to the conditioned learning of odor paired with stroking (O/S) or to non learning training (Odor only (O/O) or Stroking only (S)). The arrow indicates the pCREB protein band from both olfactory bulbs of one animal for each training condition.

**Figure 4:** The average optical density (mean ± S.E.M.) of pCREB Western blot analysis in the olfactory bulbs of pups trained to Odor only, Stroking only, or Odor + Stroking (conditioned learning). (*) $P < 0.05$. Each treatment included seven pups.
group (avg = 2.0 cells/10,000 µm²) was significantly different from the Stroking only group (avg = 1.2 cells/10,000 µm², P < 0.05) and Naive group (avg = 1.2 cells/10,000 µm², P < 0.05) while the Odor only group (avg = 1.1 cells/10,000 µm²) was not significant.

Discussion

The significantly greater increase in pCREB in the olfactory bulb of rat pups undergoing a training procedure known to produce a conditioned preference for peppermint odor 24 hr later (McLean et al. 1993; Langdon et al. 1997; Price et al. 1998) supports the hypothesis that CREB is a universal memory molecule implicated in the initiation of long-term memories. Neither Odor alone, which can increase c-fos expression in similar regions of the olfactory bulb (Guthrie and Gall 1995a,b; Johnson et al. 1995), nor Stroking alone, which is known to activate noradrenergic neurons involved in modulating pCREB levels (Cirelli et al. 1996), was able to produce a comparable increase in CREB phosphorylation. In initial experiments we also observed that the olfactory bulbs of Naive pups exhibited relatively low levels of pCREB (Fig. 2). Low levels of pCREB were quantitated by immunocytochemistry in the bulbs of Naive pups, an observation which was in keeping with the qualitative observations in Western blots. Thus, baseline levels of pCREB were observed in Naive pups as well as pups given nonlearning training. The qualitative and quantitative data suggest that pCREB is increased only when pups are trained by conditioning. This is consistent with a model in which both a calcium signal initiated by the odor input (Bozza and Kauer 1998) and a cAMP signal initiated by norepinephrine are required synergistically to elevate significantly pCREB levels. While the selectivity of the pCREB increase for the pairing condition supports a special role of the CREB promoter in associative learning, future investigations should compare forward (Odor + Stroking) and backward (Stroking followed by Odor, nonlearning) pairing. A demonstration of an effect of pairing order, which is known to affect memory formation in this paradigm (McLean et al. 1996; Sullivan and Hall 1988; Sullivan et al. 1991; Sullivan et al. 1989b; Wilson and Sullivan 1991), would provide the strongest test of the claim for a unique association between pCREB increases and associative memory formation.

TEMPORAL AND SPATIAL CHARACTERISTICS OF PCREB INCREASES

As predicted by the Frank and Greenberg
model (1994) changes in pCREB were transient with increases peaking at 10 min after the session and returning near the control range at 60 min after the session. This pattern is similar to previously reported temporal patterns of pCREB activation (Bito et al. 1996; Impey et al. 1996; Moore et al. 1996; Matthies et al. 1997; Deisseroth et al. 1998) and is consistent with the rapid and transient time course required for the catalytic C subunit of protein kinase A to translocate to the nucleus after adenylate cyclase/cAMP activation (Nigg et al. 1985).

Given the overall increase in pCREB protein (Western blots) in the bulb associated with learning (Odor + Stroking), one might have predicted substantial increases in pCREB observed by immunocytochemistry. While we did observe regional and cell specific increases, the overall number of pCREB cells localized in all populations in the bulb did not appear to be dramatically increased in the learning (Odor + Stroking) situation. A possible explanation for this apparent discrepancy may be that large mitral cells produce a large amount of pCREB, which may not be reflected by cellular localization with immunocytochemistry but is detectable by Western blot analysis.

In the present study, the localization of the pairing-associated pCREB changes in the dorsolateral olfactory bulb overlaps with the midlateral portion of the bulb shown to be associated with increased 2-deoxyglucose activity in rats that have been conditioned to the same peppermint odor (Coopersmith and Leon 1986; Sullivan and Leon 1986a, 1987; Sullivan et al. 1988; Woo et al. 1987). The localization of pCREB in the dorsolateral region correlates very well to c-fos mRNA expression in response to peppermint odor (Guthrie et al. 1993), which was shown to activate periglomerular, mitral, and granule cells in adult rats and to the general region (midlateral) of c-fos activated by retrieval of a conditioned peppermint odor in older pups (Johnson et al. 1995). However, in retrieval studies, increases in fos-reactive cells have been associated selectively with the periglomerular region in response to peppermint odors (Guthrie et al. 1993; Johnson et al. 1995; Schwab 1996). Cell
numbers in the periglomerular region (Cooper-Smith and Leon 1986; Woo and Leon 1987, 1991; Sullivan et al. 1990, 1991; Matsutani and Leon 1993; Johnson et al. 1995), the density of glial processes in the glomeruli (Matsutani and Leon 1993) and the area of the glomerular neuropil (Woo et al. 1987) are all increased in pups that have undergone 18 days of conditioning procedures. The results from the present study, however, highlight a role of mitral cell activation in the initial acquisition of conditioned preference to a novel odor. Whether such activation leads to alterations in mitral cell dendrites which might contribute to the later observations of increases in glomerular neuropil or periglomerular cells remains to be investigated. It will be important to assess more directly the relationship between odor input and pCREB activation during acquisition by measuring metabolic activation simultaneously using 2-deoxyglucose techniques and pCREB activation using immunocytochemical techniques on sections from the same subjects.

The pCREB changes in the present study are widespread within the dorsolateral quadrant, a pattern consistent with encoding changes occurring throughout a neural network rather than at a few hot spots within the olfactory bulb. Both excitatory and inhibitory connections are likely to be altered during learning (Wilson and Sullivan 1990) so it may not be surprising to have widespread regional activation of CREB.

MONOAMINE INTERACTIONS IN CONDITIONED OLFATORY LEARNING: POSSIBLE PCREB SYNERGY

Although noradrenergic input appears to act as the critical unconditioned stimulus for olfactory conditioned preference learning, we have found that serotonergic depletion of the olfactory bulb can prevent such learning. Our earlier work suggests that 5-HT acting through 5-HT2 receptors normally promotes noradrenergic-induced plasticity. Although sufficient β-adrenoceptor activation can overcome the requirement for 5-HT2 input during olfactory preference learning (Langdon et al. 1997), an increase of 5-HT2 stimulation can produce learning with subthreshold noradrenergic stimulus (Price et al. 1998). 5-HT2 activation alone does not seem to produce learning. These results fit with the hypothesis that these two inputs may act synergistically on the same cells with respect to increasing cAMP. The β1 adrenoceptor is coupled to a G protein which, upon activation, stimulates adenyl cyclase induced cAMP production (De Blasi 1989). If cAMP-mediated long-term plasticity is the substrate for olfactory preference learning, as suggested by the present study, this would parallel the role of β-adrenoceptors and/or cAMP in the plasticity of the developing visual system (Kasamatsu 1983; Kasamatsu and Shirokawa 1985) and in learning-related plasticity in a variety of biological systems (Abrams et al. 1991; Levin et al. 1992; Frank and Greenberg 1994; Tully et al. 1994). Although 5-HT receptor activation does not by itself increase cAMP expression (Morin et al. 1992), which is consistent with our failure to induce learning with such activation alone, 5-HT2 activation can act through the phosphotidyl inositol second messenger system to enhance the generation of intracellular cAMP by isoproterenol or adenylyl cyclase (Rovescalli et al. 1993). It follows that serotonin in the olfactory bulb might act to promote or intensify the norepinephrine-induced cAMP unconditioned stimulus signal.

The present results suggest mitral cells may be the critical cellular substrate for such interaction. Other studies have shown that 5-HT2A receptor mRNA (McLean et al. 1995) and protein (Hamada et al. 1998) are localized in mitral/tufted cells and, to a much lesser extent, in periglomerular and granule cells (Hamada et al. 1998). Less is known concerning the cellular localization of the β-adrenoceptors, but one study has shown that mitral cells and periglomerular cells express β2 adrenoceptor mRNA in the neonate bulb (K.J. Ivins, E. Ibrahim, and M. Leon, unpubl.). Thus, it is possible that both serotonergic and noradrenergic axons act on mitral cells and/or interneurons to induce learning by activating pCREB. It is less likely that both act on periglomerular cells because very few noradrenergic axons reach the periglomerular region (McLean and Shipley 1991; McLean et al. 1989). However, it should be noted that there is relatively dense β-adrenoceptor binding in the glomerular layer (Woo and Leon 1995a) and lesions of the locus coeruleus increase β2 adrenoceptor density (Woo et al. 1996), whereas conditioned learning decreases β2 adrenoceptor density in glomerular foci (Woo and Leon 1995b). It will be of interest to observe changes in mitral cell pCREB following olfactory conditioning procedure in pups with various levels of 5-HT2 and β-adrenoceptor activation.

Interestingly, in one of the first reports of the necessary and sufficient role of 5-HT in producing long-term change in the Aplysia 5-HT2 nervous sys-
tem, Brunelli et al. (1976) suggested that the likely mammalian analog to *Aplysia* serotonin input would be norepinephrine. The involvement of CREB in long-term plasticity was first reported in cultured sensory and motor neurons of *Aplysia* exposed to a serotonergic signal. It now appears, as Brunelli et al. (1976) hypothesized, that norepinephrine functions in an analogous manner via CREB activation to induce a long-term memory in neonate rats.

**Acknowledgments**

This work was supported by the Medical Research Council of Canada (MT-14148).

The publication costs of this article were defrayed in part by payment of page charges. This article must therefore be hereby marked “advertisement” in accordance with 18 USC section 1734 solely to indicate this fact.

**References**


Received September 18, 1999; accepted in revised form October 15, 1999.