Activation of Tyrosine Hydroxylase mRNA Translation by cAMP in Midbrain Dopaminergic Neurons

Xiqun Chen, Lu Xu, Pheona Radcliffe, Baoyong Sun, and A. William Tank

Department of Pharmacology and Physiology, University of Rochester Medical Center, Rochester, New York 14642

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ABSTRACT

During prolonged stress or chronic treatment with neurotoxins, robust compensatory mechanisms occur that maintain sufficient levels of catecholamine neurotransmitters in terminal regions. One of these mechanisms is the up-regulation of tyrosine hydroxylase (TH), the enzyme that controls catecholamine biosynthesis. In neurons of the periphery and locus coeruleus, this up-regulation is associated with an initial induction of TH mRNA. In contrast, this induction either does not occur or it is nominal in mesencephalic dopamine neurons. The reasons for this lack of compensatory TH mRNA induction remain obscure, because so little is known about the regulation of TH expression in these neurons. In this study, we test whether activation of the cAMP signaling pathway regulates TH gene expression in two rodent models of midbrain dopamine neurons, ventral midbrain organotypic slice cultures and MN9D cells. Our results demonstrate that elevation of cAMP leads to induction of TH protein and TH activity in both model systems; however, TH mRNA levels are not up-regulated by cAMP. The induction of TH protein is the result of a novel post-transcriptional mechanism that activates TH mRNA translation. This translational activation is mediated by sequences within the 3′ untranslated region (UTR) of TH mRNA. Our results support a model in which cAMP induces or activates trans-factors that interact with the TH mRNA 3′UTR to increase TH protein synthesis. An understanding of this novel regulatory mechanism may help to explain the control of TH gene expression and consequently dopamine biosynthesis in midbrain neurons under different physiological and pathological conditions.

Tyrosine hydroxylase (TH) is the enzyme that controls the biosynthesis of dopamine, norepinephrine, and epinephrine. It is one of the most meticulously regulated enzymes in the nervous system (Kumer and Vrana, 1996; Sabban and Kvetnansky, 2001). Its regulation is mediated by short-term and long-term mechanisms, both of which are adaptive responses to increased release of catecholamines during nerve stimulation, permitting neurons to maintain an appropriate supply of these vital neurotransmitters. Short-term mechanisms control the activity of pre-existing enzyme via phosphorylation of serine sites within its N-terminal domain. Long-term mechanisms involve induction of TH enzyme protein; these mechanisms are activated during chronic stimulation, such as long-term stress or repeated drug treatment. In adrenal medulla and locus coeruleus, long-term regulation is mediated by mechanisms that modulate TH gene transcription rate and TH mRNA stability (Sabban and Kvetnansky, 2001; Wong and Tank, 2007).

Much less is known about regulation of TH gene expression in mesencephalic dopaminergic neurons. Some workers have shown that severe stress, chronic nicotine treatment, or re-erpine treatment induces TH gene expression in midbrain, with those neurons having their cell bodies in the ventral tegmentum being more responsive than those with their cell bodies in the substantia nigra (Pasinetti et al., 1990; Ortiz et al., 1996; Serova and Sabban, 2002). However, this induction has not been observed in other studies and when observed, the effects are usually small or short-lived compared with those seen in adrenal medulla or locus coeruleus (Biguet et al., 1986; Smith et al., 1991; Melia et al., 1992).

In a previous collaborative study (Bowyer et al., 1998), we used high doses of amphetamine to destroy dopaminergic nerve terminals in striatum without producing significant loss of dopaminergic cell bodies in substantia nigra. Our expectation was that this loss of nerve terminals would lead to an adaptive induction of TH mRNA in midbrain cell bod-

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ABBREVIATIONS: TH, tyrosine hydroxylase; ACSF, artificial spinal fluid; DIV, days in vitro; SV40, simian virus 40; bp, base pair(s); UTR, untranslated region; 6MPH₄, 6-methyl-5,6,7,8-tetrahydropterin; Luc, luciferase; RT-PCR, reverse transcription-polymerase chain reaction; GAPDH, glyceraldehyde-3-phosphate dehydrogenase; 8-CPT-cAMP, 8-chlorothio-cyclic AMP; C₉, cycle threshold; nt, nucleotide(s); CRE, cAMP response element.
ies. To our surprise, midbrain TH mRNA levels were unaffected by this amphetamine-induced loss of striatal nerve terminals. One explanation for this lack of response in midbrain is that the signaling pathways mediating TH mRNA induction are not activated in this amphetamine neurotoxicity paradigm. However, to test this hypothesis is difficult, because so little is known about the regulation of TH in midbrain neurons.

Cyclic AMP is a powerful inducer of TH gene transcription rate and TH mRNA in many model systems (Kumer and Vrana, 1996; Sabban and Kvetnansky, 2001). Furthermore, it has been reported that forskolin induces TH mRNA, when it is injected into the rat substantia nigra under in vivo conditions (Leviel et al., 1991). In the present study, we test the effect of cAMP on TH gene expression in two rodent models of midbrain dopamine neurons under culture conditions. Our results unexpectedly demonstrate that cAMP does not induce TH mRNA in these two midbrain models. Instead, cAMP induces TH protein by activating a novel post-transcriptional mechanism that enhances the translation of pre-existing TH mRNA.

Materials and Methods

Ventral Midbrain Slice Cultures. Brains were removed from Sprague-Dawley rat pups on postnatal day 7 to 10 and immediately placed in oxygenated (95% O2, 5% CO2), ice-cold artificial spinal fluid (ACSF: 125 mM NaCl, 26 mM NaHCO3, 2.5 mM KCl, 1.25 mM NaH2PO4, 2 mM CaCl2, 1 mM MgSO4 (anhydrous), and 25 mM glucose, pH 7.4). The midbrain was dissected with the use of a brain block. The first cut was made at the side of the third ventricle, and a second cut was made below the pons-midbrain junction. The cortex was removed, and the ventral midbrain was isolated and placed in 5% low-melting agarose (dissolved in ACSF), which was then placed in an ice bath. Coronal slices (350–400 μm) were obtained using an Ingridslice 7550 PSDS Vibraslicer (Instrumments Ltd., Lafayette, IN), and they were perfused in ice-cold oxygenated ACSF for 1 to 2 h. The slices from each rat pup were placed on micellinc inserts (Millipore Corporation, Billerica, MA), and they were cultured at 35°C in an atmosphere of 95% air and 5% CO2 in medium containing 50% Earle’s balanced salt solution and 25% Hanks’ balanced salt solution (with phenol red), 2 mM l-glutamine, 0.01 μg/ml penicillin/streptomycin, 25% heat-inactivated horse serum (Invitrogen, Carlsbad, CA), and 6.4 g/l glucose. This procedure is a modification of the method of Stoppini et al. (1991). Unless otherwise stated, the slices were cultured for 7 days in vitro (DIV) before experimentation. Previous studies using anatomical and electrophysiological analyses indicated that neonatal rat brain slices cultured under these conditions survived well for many weeks in vitro (Stoppini et al., 1991; Thomas et al., 1998). In our initial studies using these midbrain cultures, TH-positive cells looked morphologically healthy for at least 14 DIV. They had intact cell bodies along with extended fibers; this was apparent at each time point examined. The slices possessed small and large unipolar, bipolar, and multipolar TH-immunoreactive cells. Nissl body staining also depicted healthy and viable neurons that remained viable for at least 14 DIV. Biochemical analyses of numerous slice cultures demonstrated initial decreases after 24 h in vitro in TH mRNA and TH activity that varied in extent compared with that observed in freshly dissected midbrain tissues. However, from 1 to 7 DIV, basal TH mRNA levels and TH activity in the cultures did not change significantly. These initial results suggested that after an initial variable loss of neurons that occurred rapidly after placing the slices into explant culture, neurons remained healthy and viable in the midbrain slices and basal TH expression remained relatively constant for at least 14 DIV.

Cell Cultures. MN9D cells were obtained from Dr. Lisa Opanashuk (Department of Environmental Medicine, University of Rochester Medical Center, Rochester, NY). These cells were created by Heller and coworkers (Choi et al., 1992), and they are a hybrid cell line derived from the fusion of mouse neuroblastoma N18TG2 cells with embryonic mouse mesencephalic neurons. The cells were cultured as described by Choi et al. (1992) in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum (Invitrogen) in an atmosphere of 95% air and 5% CO2. PC12 and cath.a cells were cultured as described previously (Pask et al., 1986; Ley et al., 1991).

Construction of TH-Luciferase Plasmids. The modified GL3 plasmid was constructed by first deleting the SV40 promoter in pGL3-Control (Promega, Madison, WI) using HindIII and BglII and inserting into this site the SV40 early enhancer/promoter from the pRL-SV40 vector (Promega), which was excised from pRL-SV40 using the same two enzymes. This modified GL3 was used as a control vector in some experiments, and it was also the starting backbone for the TH-luciferase (Luc) vectors used in this study. To construct TH(3’UTR)-Luc (see Fig. 6A), the 35-bp TH 5’UTR was inserted just upstream of the luciferase gene by cutting the modified GL3 vector with HindIII and Ncol and inserting an oligonucleotide encoding these sequences with HindIII and Ncol sites on the appropriate ends. The SV40 enhancer and SV40 late poly(A) signal sequences (downstream of the luciferase gene) in the modified GL3 vector were excised using XbaI and BamHI. TH cDNA sequences were then inserted into this XbaI and BamHI site; these TH cDNA sequences encoded 160 bp of coding sequence within the 5’ end of exon 13 of the TH gene, 280 bp of TH mRNA 3’UTR (encoding the 3’ end of exon 13), and 80 bp of genomic DNA downstream of the TH mRNA poly(A) addition site. TH(35’UTR)-Luc was created by excising the TH mRNA 5’UTR sequences using HindIII and Ncol, filling in the protruding ends with Klenow DNA polymerase I and religation. TH(Δex13)-Luc was created by excising the TH mRNA coding sequences in exon 13, using XbaI and Kpnl followed by blunt-end and religation. Both the 5’UTR and exon 13 coding sequences were excised to create TH(3’UTR)-Luc, TH(3’UTR)-Luc was generated by excising the TH mRNA 3’UTR sequences from Kpnl to SpfiHI from TH(3’UTR)-Luc, followed by blunt-end and religation.

Enzyme Assays and TH Protein Measurement. All procedures were performed on ice or in a cold room unless otherwise stated. For measurement of TH activity, midbrain slices or cells were homogenized in a buffer containing 30 mM potassium phosphate, pH 6.8, 50 mM NaF, and 10 mM EDTA. The resulting supernatant was assayed for TH activity at 30°C under Vmax conditions (unless otherwise stated) using the decarboxylase assay as described previously (Tangk et al., 1986), using 6-methyl-5,6,7,8-tetrahydropyridin-6-ol (MHPF) as cofactor. TH activity was expressed as picomoles of 14CO2 formed per minute divided by milligrams of protein. TH protein was measured using Western analysis as described previously (Yoshimura et al., 2004). The same slice or cell extracts used for assaying TH activity were used for measuring TH protein. Firefly and Renilla reniformis luciferase activities were measured using the Dual-Reporter Luciferase assay system (Promega). Protein determinations were made by the method of Bradford (1976), using bovine serum albumin as standard.

TH mRNA Measurements. TH mRNA levels were measured routinely using a semiquantitative RT-PCR assay as described previously (Sun et al., 2004; Yoshimura et al., 2004). In brief, 0.2 to 0.8 μg of total cellular RNA isolated from midbrain slices or MN9D cells was subjected to RT using random hexamer primers. Aliquots of the resulting single-stranded cDNA products were used along with the appropriate primers (see below) in the PCR to incorporate [32P]dATP (0.5 μCi/reaction) into double-stranded products encoding 537-bp TH cDNA or 294-bp 28S rDNA. The 5’ TH mRNA sense primer encoded rat or mouse cDNA sequences 390 to 410; the 3’ antisense primers were complementary to rat or mouse TH cDNA sequences 888 to 908. These primer sequences were based on the rat TH cDNA sequence; however, the mouse TH cDNA sequences in these primers diverged.
by only one (5′-primer) or two (3′-primer) bases; hence, these primers were successful for assaying TH mRNA in both rat and mouse RNA samples. The sequences for these primers were as follows: 5′ TH primer, 5′-ccc cac ctg gag tat ttt ggt-3′ and 3′ TH primer, 5′-act aag ggc gga cag tag acc-3′. The primer sequences used for PCR of 28S rRNA were identical in both rat and mouse cDNAs. The 5′ 28S rRNA primer encoded sequences 1 to 22 of rat or mouse 28S cDNA (5′-tgt aac agc gat tga aca ggt-3′); the 3′ 28S primer was complementary to rat or mouse 28S cDNA sequences 276 to 295 (5′-aac gcg gac gct ttc cau g-3′). For each experiment, the linearity of the RT-PCR was assessed with respect to both micrograms of RNA added to the RT reaction and the number of PCR cycles. Densitometric values of the bands corresponding to the TH mRNA and 28S rRNA PCR products observed on the dried-down electrophoretic gels were quantified using PhosphorImager analysis (GE Healthcare, Chalfont St. Giles, UK). TH mRNA values were normalized to 28S rRNA values for each sample. These normalized values (TH mRNA/28S rRNA) were then expressed as ∼fold increases over control values for each experiment, and the values are presented as such in the figures.

In select experiments (see Fig. 2), TH mRNA was quantified using a quantitative RT-PCR assay using a Prism 7000 real-time PCR cycler (Applied Biosystems, Foster City, CA). Initial experiments were performed with RNA extracts from each cell line to determine the linearity of the TH mRNA and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) mRNA signals with respect to total cellular RNA concentrations added to the RT reactions. For MN9D, PC18, and cath.a cells, linearity was achieved using total cellular RNA in each assay. The RT reaction was performed as described for the semiquantitative assay (see above). Real-time PCR was performed as recommended by Applied Biosystems using a 2-μl aliquot of the cDNA produced in the 20-μl RT reaction and SYBR Green (Bio-Rad Laboratories, Hercules, CA) indicator to measure the amount of cDNA product formed in each PCR cycle. Forward and reverse primers were present at 1 μM in the PCR reactions. GAPDH mRNA was measured in each sample for normalization purposes, using sample cDNA transcribed from the same RT reaction as that used for measuring TH mRNA. The TH primers were as follows: rat forward primer encoded sequences 620 to 639 of rat TH cDNA (5′-tgg gaa ggt gtg tag aga-3′) and rat reverse primer was complementary to sequences 695 to 705 (5′-tcc cgc tga ttc cac atg-3′); mouse forward primer encoded sequences 1016 to 1033 of mouse TH cDNA (5′-tgg tgt gtc acc gca cat-3′) and mouse reverse primer was complementary to sequences 1058 to 1076 (5′-gcc ccc gag aga gca cag c-3′). GAPDH mRNA primers were the same for both rat and mouse samples: forward primer encoded sequences 507 to 517 of rodent GAPDH cDNA (5′-ggg agg gct cat gac gag atg-3′) and reverse primer was complementary to sequences 570 to 589 (5′-acc tta cag ccc act acc-3′). Standard curves for TH mRNA and GAPDH mRNA were created for each experiment. Cloned TH cDNA and GAPDH cDNA sequences encompassing the amplified regions were transcribed in vitro using a MAXIScript transcription kit (Ambion, Austin, TX). Known amounts (picomoles) of the in vitro-transcribed mRNAs were used to construct these standard curves using the same RT-PCR conditions used for the sample mRNAs. The C_{T} values derived from the real-time PCRs for different concentrations of standard TH mRNA or GAPDH mRNA were plotted against the known concentrations of the standard mRNAs used in the RT reaction to construct these curves. The C_{T} values for the sample TH mRNAs and GAPDH mRNAs were then compared with the C_{T} values on the standard curves to obtain picomoles of TH mRNA and GAPDH mRNA. The TH mRNA values were then normalized to the GAPDH mRNA values for each sample.

Luciferase mRNA Measurements. Firefly and R. reniformis luciferase mRNA levels were measured using semiquantitative RT-PCR essentially as described above, with minor modifications. Because of significant contamination of the RNA samples with plasmid DNA, each sample was treated with 2 units of TURBO-DNase (Ambion) before the RT step. Furthermore, all samples were assayed with or without the RT step, to check for DNA contamination; only those assays that showed negligible or very low levels of signal in the absence of the RT step (relative to that observed in assays run in the presence of the RT step) were used for quantitation of luciferase mRNA. As for the TH mRNA assays, all RT-PCR reactions were run under conditions of linearity with respect to both RNA input and PCR cycle number. The primers used for firefly luciferase mRNA were as follows: 5′ primer encoded sequences 92 to 112 of the luciferase mRNA coding region (5′-aga gat gcc ccc tgt tgc ctg-3′) and the 3′ primer was complementary to sequences 469 to 492 of the coding region (5′-cat cga tgt aaa ttc ctc atc-3′). The primers used for R. reniformis luciferase mRNA were as follows: 5′ primer encoded sequences 672 to 676 of the coding region (5′-ggc tga aat ccc ggt ag-3′) and 3′ primer was complementary to sequences 853 to 858 of the coding region (5′-cca ttt cat cag gat ctc ct-3′).

Measurement of TH Rate of Synthesis. The rate of TH protein synthesis in midbrain slices or MN9D cells was estimated using pulse-labeling procedures essentially as described by Tank et al. (1986). Slices or cells were incubated in the presence or absence of 1 mM 8-CPT-cAMP for 24 h, at which time the medium was changed to leucine-free Dulbecco’s modified Eagle’s medium (supplemented with 10% fetal bovine serum) containing 25 μCi/ml [4,5-3H]leucine (60 Ci/mmol). The cells were incubated for an additional 1.5 h in the labeling medium in the continued presence of the cAMP analog, when appropriate. The cells were then harvested and washed twice in phosphate-buffered saline containing 1 mM leucine. Radiolabeled TH protein was isolated by immunoprecipitation, and radioactivity was measured in the immunoprecipitated protein as described previously (Tank et al., 1986). The [3H]TH cpm values denoted in the table in Fig. 3 represent the amount of radioactivity incorporated into TH protein normalized to the micrograms of total protein present in the cell or slice extract. This value was calculated by dividing the cpm of [3H]leucine incorporated into 50 units of TH by the micrograms of total protein present in that 50 units of TH activity. Incorporation of radiolabeled leucine into total soluble protein was measured as described previously (Tank et al., 1986). Rates of synthesis were calculated as cpm incorporated into TH protein per microgram of protein divided by cpm incorporated into total soluble protein per microgram of protein. In select experiments the gel was incubated in EN3HANCE (PerkinElmer Life and Analytical Sciences, Boston, MA) for 1 h as described by the manufacturer, dried down onto 3MM paper (Clifton, Trenton, NJ), and exposed to X-ray film for 7 days in a −80°C freezer. The radiolabeled band corresponding to TH was then identified on the autoradiogram.

Polysome Profile Analysis of MN9D TH mRNA. Polysome profile analysis was performed as described by Xu et al. (2007). In brief, MN9D cells were homogenized in ice-cold low-salt lysis buffer (20 mM Tris-HCl, pH 7.5, 10 mM NaCl, 3 mM MgCl₂, 1 mM RNasin, 1 mM dithiothreitol, 0.3% Triton X-100, and 0.05 M sucrose). Nuclei and the majority of mitochondria were sedimented by centrifugation for 10 min at 10,000 g at 4°C. The supernatants (1 ml) were adjusted to 170 mM NaCl and 13 mM MgCl₂, layered onto 9-ml linear sucrose gradients (15–45% w/w), and centrifuged at 36,000 rpm for 2 h at 4°C in a SW40 rotor. Fractions (800 μl) were collected after centrifugation, and RNA in each fraction was precipitated and resuspended in 15 μl of diethyl pyrocarbonate-treated water. TH mRNA levels in each fraction were assayed using the semiquantitative RT-PCR assay described above. To control for differences in recovery of RNA from each fraction during the isolation procedure, a known amount (100 ng) of a standard control RNA was added to each fraction before RNA isolation. This standard RNA was obtained by in vitro transcription (using a kit from Ambion) of TH genomic DNA promoter sequences and detected using specific primers (for details, see Xu et al., 2007). The amount of TH mRNA in each fraction was first normalized to the recovery of this standard RNA, and then it was expressed as a percentage of the total mRNA isolated from the


**Results**

**TH Protein and TH Activity Were Increased in Midbrain Slice Cultures in Response to Treatment with 8-CPT-cAMP.** When midbrain slices were treated for 24 h with the cAMP analog, 8-CPT-cAMP, TH activity measured under saturating cofactor conditions (4 mM 6MPH4) increased in a dose-dependent manner (Fig. 1A). A 1.5-fold increase in TH activity was observed using 0.1 mM 8-CPT-cAMP, and a maximal increase was observed using 0.3 mM cAMP analog. When assayed under these apparent V_{max} conditions, an increase in TH enzyme activity is usually indicative of an increase in TH protein. To test this hypothesis directly, Western analysis was performed using the same supernatants assayed for TH activity. A representative Western blot is presented above the bar graph in Fig. 1A. TH protein levels increased to the same extent as TH activity at each concentration of 8-CPT-cAMP.

A time course of this response to 8-CPT-cAMP is presented in Fig. 1B. TH activity assayed using saturating pterin cofactor concentration (4 mM) increased significantly at each time point between 12 and 48 h of treatment with the cAMP analog. Similar increases in TH activity were observed when the enzyme was assayed using a subsaturating pterin cofactor concentration. TH protein levels increased 2- to 3-fold after 24, 36, or 48 h of 8-CPT-cAMP treatment. However, in contrast to the observed increase in TH activity, TH protein levels were not elevated at the 12-h time point. Taken together, these results indicate that TH is induced 2- to 3-fold by 8-CPT-cAMP in midbrain slice cultures. Similar increases in TH activity and TH protein were observed when midbrain slice cultures were treated with 10 μM forskolin for 24 h (data not shown).

**TH mRNA Levels Were Not Affected by Treatment of Midbrain Slices with 8-CPT-cAMP.** Midbrain slices were treated with 1 mM 8-CPT-cAMP, and total cellular RNA was isolated at different time points. TH mRNA levels were measured using semiquantitative RT-PCR and normalized to the levels of 28S rRNA measured in the same samples (Fig. 1C). This assay was used in several of our previous studies to detect induction of TH mRNA in adrenal medulla, locus coeruleus, and pheochromocytoma cell lines (Sun et al., 2003; Sun et al., 2004; Yoshimura et al., 2004; Osterhout et al., 2005). An example of an autoradiogram showing results of this assay using midbrain slice RNA samples is presented in Fig. 1C; a more extensive time course is shown in the table below the autoradiogram. It is evident from the autoradiogram and the more quantitative analysis in the table that TH mRNA levels did not increase significantly in response to 8-CPT-cAMP at any time point tested. In contrast, as shown in the autoradiogram and table in Fig. 1C, dexamethasone (used as a positive control) induced TH mRNA 2- to 3-fold in midbrain slice cultures.

The induction of TH protein without an induction of TH mRNA in response to cAMP suggested that a post-transcriptional mechanism mediated this response in midbrain dopamine neurons. However, it was possible that this unexpected response was due to changes in the neurons that occurred during culture of the slices for 7 days. To minimize this possibility, midbrain slices were prepared as usual, but only cultured for 1 day (DIV 1) before treatment with 8-CPT-cAMP (see table in Fig. 1C). Similar to the results obtained using the DIV 7 cultures, cAMP treatment did not induce TH mRNA at any time point tested in the DIV 1 cultures. In contrast, TH activity and TH protein increased 2- to 3-fold after 24 h of treatment with 8-CPT-cAMP in the DIV 1 cultures (TH activity: controls = 24 ± 3 pmol/min/mg protein, 8-CPT-cAMP-treated = 52 ± 4, n = 10, p < 0.01; TH protein: controls = 1.0 ± 0.1, 8-CPT-cAMP-treated = 2.4 ± 0.4, n = 8, p < 0.01).

**TH Protein and TH Activity Were Increased in MN9D Cells in Response to Treatment with 8-CPT-cAMP.** To study the mechanisms for this apparent post-transcriptional response in more detail, we tested whether cultured MN9D cells could be used as a model system, in which TH expression responded to cAMP in a similar manner as that observed in the midbrain slice cultures. These results are summarized in Fig. 2. MN9D cells were treated for 48 h with different concentrations of either 8-CPT-cAMP or forskolin (Fig. 2A). TH activity was measured under apparent V_{max} conditions, using a saturating concentration of pterin cofactor (4 mM 6MPH4). Both drugs produced concentration-dependent increases in TH activity. Maximal or near-maximal effects were observed using 300 μM 8-CPT-cAMP or 10 μM forskolin; the concentration-response curve for 8-CPT-cAMP was very similar to that observed in midbrain slice cultures. To test whether these increases in TH activity were due to induction of TH protein, Western analysis was performed; a representative immunoblot is presented in Fig. 2B, top. As in midbrain slices, it is evident from this blot that TH protein levels increased significantly after 12 to 24 h of treatment with 8-CPT-cAMP in MN9D cells. In a more detailed time course study, TH activity and TH protein levels were measured in MN9D cells treated with 1 mM 8-CPT-cAMP for different durations of time (Fig. 2B, bar graph). Significant increases in TH activity (measured under apparent V_{max} conditions with respect to pterin cofactor) were observed after 3, 6, and 12 h of treatment with the cAMP analog. These early increases in enzyme activity did not correlate with
changes in TH protein levels, which were measured in the same MN9D supernatants. TH activity increased more dramatically (1.9–2.5 fold) after 24, 36, and 48 h of treatment; these increases correlated closely with changes in TH protein levels. In many experiments performed in MN9D cells over the past 2 to 3 years, the changes in TH activity and TH protein levels in response to 24 h of treatment with 8-CPT-cAMP varied from 1.5- to 2.5-fold; however, in every experiment significant increases in TH activity and TH protein were measured in response to cAMP.

The results of these MN9D time courses as well as those performed in the midbrain slice cultures suggested that the increases in TH activity observed at the later time points (24, 36, and 48 h) were due to induction of TH protein, whereas those at the earlier time points (3 and 6 h) were due to enzyme activation. To test further this latter hypothesis, we assayed TH activity in samples from the early time points using low pterin cofactor concentrations in the assays (Fig. 2C). Under these assay conditions, TH activity increased by 4-fold at both the 3- and 6-h time points, when TH protein was not induced. In contrast, TH activity measured in the same samples under saturating cofactor concentrations elicited smaller increases (~1.7-fold) in TH activity (similar to that seen in the other experiments presented in Fig. 2B). These data are suggestive of a cAMP-mediated activation of TH at these early time points, leading to a decrease in the $K_m$ value for pterin cofactor, as observed in many other studies in other cell types (Kumer and Vrana, 1996); however, a more thorough enzyme kinetic analysis is necessary to confirm this conclusion.

**Fig. 1.** TH activity and TH protein, but not TH mRNA were induced in cultured rat midbrain slices by treatment with 8-CPT-cAMP. A, midbrain slices were treated for 24 h with 8-CPT-cAMP at the concentrations designated in the figure. Control slices were left untreated. B, midbrain slices were treated with 1 mM 8-CPT-cAMP for the durations designated in the figure. Control slices were left untreated. All slices were harvested at the same time, quick-frozen, and stored at $-80^\circ$C. TH activity was assayed using 4 mM 6MPH$_4$ (A) and either 0.2 or 4 mM 6MPH$_4$ (B). Control TH activity in A was $27 \pm 3$ pmol/min/mg protein. TH protein was measured using Western analysis. An autoradiogram depicting a representative western blot is shown above the bar graphs in A. The results in A and B represent the means ± S.E. from three to six slice cultures. C, midbrain slices were treated with $1 \times 10^{-3}$ M 8-CPT-cAMP or $0.1 \times 10^{-3}$ M dexamethasone for the durations of time designated in the figure. Control cultures were left untreated or treated with vehicle (dimethyl sulfoxide). All cultures were harvested at the same time, quick-frozen, and stored at $-80^\circ$C. Total cellular RNA was isolated, and TH mRNA was assayed using semiquantitative RT-PCR. Autoradiograms depicting representative RT-PCR assays are shown for midbrain slice cultures (DIV 7) treated for 24 h with either drug. The data in the table represent values for TH mRNA divided by those for 28S rRNA measured in the same samples and expressed as $\pm$-fold increase over controls. The data represent the means ± S.E. from 6 to 12 midbrain slices for the 8-CPT-cAMP experiments and from four midbrain slices for the dexamethasone experiments. $a, p < 0.05$ compared with controls.
Treatment of MN9D Cells with 8-CPT-cAMP Was Not Associated with Induction of TH mRNA. We tested whether the cAMP-mediated induction of TH protein in MN9D cells was associated with an initial induction of TH mRNA. A representative autoradiogram depicting TH mRNA and 28S rRNA levels in RNA samples isolated from MN9D cells is shown in Fig. 2B; data collected from a number of time course experiments are shown in the bar graph of Fig. 2B. TH mRNA levels did not increase in MN9D cells treated with the cAMP analog at any time point tested between 3 and 36 h; similar results were observed using forskolin as a stimulus (data not shown).

The lack of cAMP-mediated induction of TH mRNA in both midbrain slice cultures and MN9D cells was very unexpected. To further verify that this result was not an artifact of the method used to assay TH mRNA, we developed a quantitative RT-PCR assay for TH mRNA using a real-time PCR cycler. The results of experiments using this quantitati-

Fig. 2. TH activity and TH protein, but not TH mRNA were induced by 8-CPT-cAMP in MN9D cells. A, MN9D cells were treated for 48 h with either 8-CPT-cAMP or forskolin at the concentrations designated in the figure. Control cells for the 8-CPT-cAMP experiments were left untreated, whereas controls for the forskolin experiments were treated with vehicle (dimethyl sulfoxide). Cells were harvested, quick-frozen, and stored at −80°C until assayed for TH activity. The results represent the means ± S.E. from three dishes. B, MN9D cells were treated with 1 mM 8-CPT-cAMP for the times designated in the figure; control cells were left untreated. Top, representative autoradiogram of a Western blot measuring TH protein and tubulin, which was used to control for variability in sample loading or protein transfer to the membrane. Middle, representative autoradiogram of semiquantitative RT-PCR assays measuring TH mRNA and 28S rRNA, which was used for normalization purposes. Bottom, bar graph depicts a time course for TH activity (measured under apparent V\textsubscript{max} conditions), TH protein (normalized to tubulin), and TH mRNA (normalized to 28S rRNA), each expressed as fold increase over controls. The results represent the means ± S.E. from three to eight dishes. C, MN9D cells were treated with 1 mM 8-CPT-cAMP for 3 or 6 h, and then they were harvested and stored at −80°C until assayed. Control cells were left untreated. TH activity was assayed using either 0.2 or 4 mM 6MPH\textsubscript{4} as cofactor. The results represent the means ± S.E. from three dishes. D, MN9D, PC18, and cath.a cells were treated with 0.5 mM 8-CPT-cAMP for 6 or 24 h; controls were left untreated. TH mRNA was measured using quantitative RT-PCR and normalized to GAPDH mRNA measured in the same samples. The results represent the means ± S.E. from five to six dishes. a, \( p < 0.01 \) compared with controls. b, \( p < 0.05 \) compared with controls.
tive assay (Fig. 2D) agreed closely with those obtained using the semiquantitative assay, in that treatment of MN9D cells with 8-CPT-cAMP for 6 or 24 h did not induce TH mRNA. In contrast, as expected, cAMP treatment induced TH mRNA significantly in both adrenal medulla-derived rat pheochromocytoma PC18 cells and locus coeruleus-derived cath.a cells (Fig. 2D).

Treatment with 8-CPT-cAMP Was Associated with an Increased Rate of Synthesis of TH Protein in Midbrain Slices and MN9D Cells. To test whether the cAMP-mediated induction of TH protein was due to an increase in the rate of synthesis of the enzyme, pulse-labeling experiments were performed in both model systems (Fig. 3). Midbrain slices or MN9D cells were incubated in the presence or absence of 1 mM 8-CPT-cAMP for 24 h, at which time the slices or cells were treated for an additional 1.5 h with [3H]leucine in the continued presence of the cAMP analog, when appropriate. In preliminary experiments using pulse-chase procedures, we found that the apparent half-life of TH protein in MN9D cells was ~40 h (data not shown). Hence, the degradation of TH protein during the 1.5-h pulse period would be predicted to be negligible. The slices or cells were harvested after the 1.5-h pulse and the incorporation of [3H]leucine into TH protein was determined using immunoprecipitation procedures. Cell extracts were incubated overnight with TH antiserum; the immunoprecipitates were collected by centrifugation and subjected to SDS-polyacrylamide gel electrophoresis. The gels were stained for protein and the ~60-kDa TH protein band was visualized. A representative gel stained for protein is shown in Fig. 3. A major band that comigrated with purified TH was easily identified, and it was excised from the gel and the radioactivity was counted by liquid scintillation spectrometry. In an initial experiment, the specificity of the antisera was tested by exposing a dried-down gel to X-ray film for 7 days. A single major radioactive band that comigrated with purified TH was observed on the autoradiogram (Fig. 3). Using this assay, we measured an ~2-fold increase in the incorporation of radiolabel into TH protein in both model systems (see table in Fig. 3). There was no change in the incorporation of [3H]leucine into total cellular protein in either model. The relative rate of synthesis of TH protein, measured as the incorporation of radiolabeled leucine into TH divided by that into total protein increased by 2- to 3-fold in either midbrain slices or MN9D cells treated for 24 h with 8-CPT-cAMP. These increases in rate of synthesis correlated reasonably well with the increases in TH activity measured in the same cell extracts. Although we have not yet ruled out an effect on TH protein degradation, these results suggest that the induction of TH elicited by the cAMP analog was primarily due to an increase in TH protein synthesis.

Treatment with 8-CPT-cAMP Increased the Association of TH mRNA with Polysomes in MN9D Cells. If cAMP treatment was associated with increased translation of pre-existing TH mRNA, then there should be an increase in the association of TH mRNA with polyribosomes in cAMP-treated cells. To test this hypothesis, MN9D cell extracts were subjected to sucrose gradient centrifugation to separate polysomal and nonpolysomal RNAs. After centrifugation, fractions were collected from the gradients and TH mRNA levels in each fraction were measured using the semiquantitative RT-PCR assay. Autoradiograms from a representative sucrose gradient fractionation are shown at the top of Fig. 4; TH mRNA levels in each fraction collected from three separate experiments are presented in the bar graphs in Fig. 4. TH mRNA levels were isolated in fractions throughout the gradient, representing both polysomal-bound RNA (heavier fractions) and nonpolysomal-bound RNA (lighter fractions). In cells treated with 8-CPT-cAMP for 24 h, there was a significant shift in TH mRNA levels to the heavier fractions (Fig. 4). No such shift in 28S rRNA was observed. The percentage of TH mRNA isolated in the heaviest six fractions (which are those enriched in 28S rRNA, as might be expected if they represent polysomes) is 66 ± 3% of the total TH mRNA isolated from the gradient in control samples and 84 ± 1% in 8-CPT-cAMP-treated samples (p < 0.01). A sec-

![Fig. 3. Rate of synthesis of TH protein was increased in cultured rat midbrain slices and MN9D cells treated with 8-CPT-cAMP. Midbrain slices or MN9D cells were treated with 1 mM 8-CPT-cAMP for 24 h; control cells were untreated for the same period. The cultures were then incubated for an additional 1.5 h in the presence of leucine-free medium supplemented with serum, 25 μCi/ml [3H]leucine and 1 mM 8-CPT-cAMP, when appropriate. Cultures were harvested and radiolabeled TH protein was isolated using immunoprecipitation procedures. The final immunoprecipitates were subjected SDS-polyacrylamide gel electrophoresis. After electrophoresis, the gels were stained with Coomassie Blue; left, a representative Coomassie Blue-stained gel. In a select experiment, a gel was treated with EN3HANCE, dried down onto Whatman 3MM paper and exposed to X-ray film for 7 days. Right, autoradiographic signal. In most experiments, the TH band was excised from the wet gel, along with a region of the gel that did not contain measurable protein staining (used to determine background radioactivity). These gel pieces were solubilized and radioactivity counted by liquid scintillation spectrometry. The cpm incorporated into TH protein, along with the cpm incorporated into total cell protein, are presented in the table. TH activity was assayed in an aliquot of the same supernatants used for immunoprecipitation. The results represent the means ± S.E. from three experiments. a, p < 0.01 compared with controls.]

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<th>Midbrain Slices Cultures</th>
<th>MN9D Cells</th>
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<tr>
<td></td>
<td>Controls</td>
<td>8-CPT-cAMP</td>
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<tr>
<td>cpm in TH protein/μg protein</td>
<td>9.4 ± 1.9</td>
<td>17.7 ± 2.0</td>
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<td>cpm in total protein/μg protein (10^6)</td>
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<td>cpm in TH/cpm in total protein (10^6)</td>
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<td>29 ± 3.0</td>
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<tr>
<td>TH activity (pmol/min/mg protein)</td>
<td>30 ± 3</td>
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OND method by which analyze these data is to use a weighted mean average approach as described by Gore et al. (1995). This is a measure of the fraction number in the gradient that contains the weighted average amount of TH mRNA relative to that measured throughout the entire gradient, and it is a method to quantitate a shift in the polysome profile. An increase in the weighted mean average indicates a shift of TH mRNA to heavier fractions containing polysomal-bound TH mRNA. The weighted mean averages for controls and 8-CPT-cAMP-treated cells were 7.8 ± 0.13 and 8.6 ± 0.15 (p < 0.02), respectively. Both of these analyses indicate a significant shift of TH mRNA levels to polysomes in cAMP-treated MN9D cells.

TH mRNA 3’UTR Sequences Conferred Activation of Reporter Gene Expression by 8-CPT-cAMP. Regulation of mRNA translation is often determined by sequences within the 5’ or 3’ UTR of the specific mRNA (Kwon and Hecht, 1991; Ostareck-Lederer et al., 1994). To test which sequences participated in the cAMP-mediated translational activation of TH mRNA, we constructed a reporter gene (designated TH(5’3’U)-luciferase) that encoded the 35-nt 5’UTR and 280-nt 3’UTR of TH mRNA, along with 160 nt of TH mRNA coding sequences that are present in exon 13 of the TH gene and 80 bp of TH 3’ genomic sequence downstream of the polyA+ addition site. These TH mRNA and genomic sequences were fused to the coding region of the firefly Luc gene as shown in Fig. 5. This hybrid gene was driven by the SV40 promoter/enhancer. Deletion mutations of this transgene were also constructed as shown in Fig. 5. A modified version of the GL3-control construct (Promega), which was used as the vector backbone for these TH-luciferase constructs, was used as a control for these experiments.

MN9D cells were transiently transfected with one of these vectors, and cells were incubated in the presence or absence of 0.5 mM 8-CPT-cAMP for 24 h. The cells were cotransfected with a vector expressing R. reniformis luciferase and the expression of this gene was used to control for transfection efficiency. The expression of firefly luciferase activity in cells transfected with the wild-type vector, TH(5’3’U)-luciferase, was increased by ~2-fold in response to cAMP analog treatment (Fig. 5). Deletion of either the 5’UTR [TH(Δ5’)U] or the TH mRNA coding sequences at the 5’ end of exon 13 [TH(ΔEx13)] did not diminish the response to 8-CPT-cAMP. Likewise, deletion of both the 5’UTR and exon 13 coding sequences [TH(3’U)] had no effect on the cAMP response.

Furthermore, deletion of the 5’UTR and/or exon 13 coding sequences had no significant effect on the basal expression of luciferase in the context of this hybrid gene. In contrast, deletion of the 3’UTR sequences between the KpnI and Sph1 sites (TH(3’UΔKS)) decreased the basal expression of luciferase by ~50% (Fig. 5). More importantly, the loss of these 3’UTR sequences ablated the response of the hybrid gene to cAMP. As expected, cAMP treatment did not elicit an effect on luciferase expression in cells transfected with the control modified GL3-control vector; however, basal luciferase expression in these cells was 3- to 4-fold greater than that observed in cells transfected with the wild-type TH-luciferase construct. This disparity may be due to differences in expression determined by the TH mRNA 3’UTR versus the SV40 3’UTR present in the modified GL3 vector. Alternatively, TH mRNA 3’UTR sequences may confer regulatory effects on luciferase mRNA translation or stability, as observed in analogous studies using 3’UTR sequences from other mRNAs (Kwon and Hecht, 1991; Ostareck-Lederer et al., 1994). To test whether this response to cAMP was specific for MN9D cells, mouse locus coeruleus-derived cath.a cells and rat pheochromocytoma PC12 cells were transfected with TH(3’U)-luciferase and treated for 24 h with either 8-CPT-cAMP or forskolin. Expression from TH(3’U)-luciferase was not significantly affected by either treatment in the cath.a cells or PC12 cells (data not shown).

Luciferase mRNA Expression from TH(3’U)-Luciferase Was Not Affected by 8-CPT-cAMP in MN9D Cells. The observed increase in TH(3’U)-luciferase activity elicited by cAMP in MN9D cells might be due to an increase in translation and/or stability of TH-luciferase mRNA molecules. Two approaches were used to differentiate between these different possibilities. In the first approach, MN9D cells were transiently transfected with TH(3’U)-luciferase and the R. reniformis luciferase expression vector. These transfected cells were then treated for 24 h with 0.5 mM 8-CPT-cAMP or vehicle, and total cellular RNA was isolated. Initial studies indicated that the levels of both firefly and R. reniformis luciferase mRNAs were very low in these transfected cells. Furthermore, measurement of these mRNAs was confounded by contamination of the RNA samples with transfected plasmid DNAs. To overcome this latter problem, the RNA samples were treated extensively with DNase (TURBO-DNase; Ambion) before the RT-PCR assay. Furthermore, controls were run for each sample in which the
RT step was omitted, to detect any residual plasmid DNA. Examples of this type of analysis are shown in Fig. 6A. These autoradiograms depict firefly and *R. reniformis* luciferase mRNAs measured using the radioactive semiquantitative RT-PCR assay. Both mRNAs were detected in RNA samples treated first with RT to produce corresponding cDNAs. In contrast, no significant signals were detected for *R. reniformis* luciferase mRNA and only minor signals were detected for firefly luciferase mRNA in samples that were not treated first with RT. As is evident from the autoradiogram, treatment of the MN9D cells with 8-CPT-cAMP for 24 h did not significantly increase firefly luciferase mRNA levels (Fig. 6A). To verify that the DNase treatment did not alter mRNA levels, TH mRNA and 28S rRNA levels were measured in MN9D RNA samples that were treated with or without TURBO-DNase (Fig. 6B). As seen in the first two lanes of the autoradiograms in Fig. 6B, DNase treatment totally abolished the signals representing firefly and *R. reniformis* mRNAs, when the samples were not first incubated with RT; these results indicated that the contaminating plasmid DNAs were degraded by this DNase treatment. In contrast, both TH mRNA and 28S rRNA signals were not affected by the DNase treatment, suggesting that TURBO-DNase did not significantly degrade RNA in the samples. Quantitative measurements of firefly and *R. reniformis* mRNAs using this assay are presented in the bar graph in Fig. 6C. Treatment with 8-CPT-cAMP for 24 h did not affect the levels of firefly luciferase mRNA expressed from the TH(3′U)-luciferase construct. In contrast, firefly luciferase enzyme activity increased ~1.9-fold in these experiments. These results argue against a stabilization of firefly luciferase mRNA, but were consistent with a translational activation of firefly luciferase mRNA conferred by the TH 3′UTR sequences.

In a second approach, MN9D cells were stably transfected with TH(3′U)-luciferase, and the effects of treatment with 8-CPT-cAMP on luciferase mRNA and enzymatic activity were analyzed. This approach was used to avoid the plasmid DNA contamination problems that we encountered when trying to measure luciferase mRNA levels, as well as the very low expression of the luciferase gene using the transient transfection approach. Two stably transfected MN9D cell lines were isolated that expressed reasonable concentrations of firefly luciferase mRNA; these cell lines were designated U1 and U17. As shown in the autoradiogram in Fig. 7, the semiquantitative RT-PCR assay easily detected firefly luciferase mRNA in RNA samples from these cells. Only minor luciferase mRNA bands were observed in samples that were not first treated with RT; these minor bands were presumably due to contaminating genomic DNA. When the stably transfected cells were treated for 24 h with 8-CPT-cAMP, luciferase mRNA levels were the same or slightly less than that observed in control cells. In contrast, luciferase activity increased significantly (1.5- to 2-fold) in response to the cAMP analog in both cell lines. These results also support the

**Fig. 5.** TH mRNA 3′UTR conferred cAMP-mediated inducibility on the luciferase gene in MN9D cells. A, diagrams of the TH-luciferase constructs used in these experiments are shown in the figure. B, MN9D cells were transfected with one of the TH-luciferase constructs along with the *R. reniformis* luciferase expression vector used for normalization purposes. Twenty-four hours after transfection, the cells were treated with or without 0.5 mM 8-CPT-cAMP for another 24 h. The cells were then harvested, and firefly and *R. reniformis* luciferase activities were measured in cell extracts. The results represent the means ± S.E. from 11 to 12 dishes, except for the modified GL3 data, which were from four dishes. a, p < 0.01 compared with controls.
hypothesis that cAMP activates translation, but does not stabilize TH(3′U)-luciferase mRNA in MN9D cells.

**Discussion**

Very little information is available concerning the molecular mechanisms that control TH gene expression in mesencephalic dopamine neurons. This lack of information is especially important when trying to understand the compensatory mechanisms that regulate dopamine biosynthesis in nigrostriatal neurons in neurodegenerative paradigms. When mesencephalic dopamine neurons are completely destroyed in animal models with the use of neurotoxins, several cognitive, motor, and sensory deficits are observed, from which the animals never recover (Robinson et al., 1990; Zigmond et al., 1990). However, when these neurons are only partially destroyed (<80%), there is considerable recovery of function from these deficits over time. This recovery is due to compensatory mechanisms that occur in the surviving nerve terminals, as well as neuronal sprouting in the nerve terminal regions (Robinson et al., 1990; Zigmond et al., 1990; Bezard et al., 2000; Rothblat et al., 2001; Jakowec et al., 2004). These mechanisms maintain appropriate concentrations

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**Fig. 6.** TH-luciferase mRNA levels were not increased by treatment of transiently transfected MN9D cells with 8-CPT-cAMP. MN9D cells were transfected with TH(3′/U)-luciferase and the *R. reniformis* luciferase expression vector. Twenty-four hours after transfection, the cells were treated for another 24 h with or without 0.5 mM 8-CPT-cAMP. The cells were then harvested, and total cellular RNA was isolated. The RNA samples were treated with 2 units of TURBO-DNase (unless otherwise stated) before RT-PCR. A, autoradiograms showing both firefly and *R. reniformis* luciferase mRNA levels detected by semiquantitative RT-PCR with or without RT are shown in the figure. B, autoradiograms are shown depicting semiquantitative RT-PCR assays run under different conditions as noted in the figure measuring firefly and *R. reniformis* luciferase mRNAs, along with TH mRNA and 28S rRNA in the same samples. C, results are shown of four experiments in which both luciferase activity and luciferase mRNA levels expressed in MN9D cells transiently transfected with the TH(3′U)-luciferase construct were measured in the same samples. The results represent the means ± S.E. from 17 to 18 dishes. a, p < 0.01 compared with controls.

**Fig. 7.** TH-luciferase activity, but not TH-luciferase mRNA levels were increased by treatment of stably transfected MN9D cells with 8-CPT-cAMP. MN9D cells were stably transfected with TH(3′/U)-luciferase, and two clonal colonies expressing luciferase activity and mRNA were isolated; these colonies were designated U1 and U17. These cells were treated with or without 0.5 mM 8-CPT-cAMP for 24 h, and then both luciferase activity and luciferase mRNA levels were measured. Representative autoradiograms depicting the luciferase mRNA levels in each cell line are shown in the top panel. The results represent the means ± S.E. from six dishes. a, p < 0.01 compared with controls.
of dopamine at striatal synapses, even when large numbers of nerve terminals are destroyed.

In contrast to the vigorous compensatory mechanisms that occur at nerve terminals, only modest or insignificant increases and even decreases in TH mRNA levels are observed in surviving midbrain dopamine cell bodies after lesioning with neurotoxins (Pasinetti et al., 1992; Kastner et al., 1994; Blanchard et al., 1995; Bowyer et al., 1998; Rothblat et al., 2001). Furthermore, TH gene transcription rate does not increase in surviving midbrain cell bodies after lesioning with 6-hydroxydopamine (Sherman and Moody, 1995). In human Parkinsonian patients, decreases in midbrain TH mRNA levels have been observed (Javoy-Agid et al., 1990; Kastner et al., 1993). This lack of compensatory induction of TH gene expression in the midbrain is puzzling. One would anticipate the existence of robust homestatic mechanisms that elevate TH gene expression in dopamine cell bodies, to increase dopamine biosynthesis in spared striatal nerve terminals and to maintain high levels of TH protein in newly sprouted striatal nerve terminals. This idea is supported by studies in the peripheral nervous system; it is well documented that when norepinephrine is depleted from sympathetic nerve terminals by reserpine or 6-hydroxydopamine, TH is induced in ganglionic cell bodies (Mueller et al., 1969; Black et al., 1985; Trocmé et al., 2001).

An explanation for this lack of compensatory response is that receptors and/or signaling pathways that mediate the induction of the TH gene may be down-regulated, may not be activated or their signals may be overcome by increased inhibitory input during Parkinson’s disease or after chemical lesioning of striatal terminals. These stimulatory receptors and signaling pathways could be those activated by neurotransmitters released from afferent nerve terminals, those recognizing humoral factors such as glucocorticoids or retinooids, and/or those recognizing neurotrrophic factors released from glial cells or target tissues. However, there is almost no information concerning which receptors and signaling pathways are responsible for regulating TH gene expression in adult dopaminergic neurons in response to extracellular physiological signals. Leviel et al. (1991) have reported that intranigral microinjection of forskolin elicits the induction of TH mRNA, but not TH protein in substantia nigra. This early report suggests that activation of the cAMP signaling pathway may induce midbrain TH gene expression, presumably as observed in most other model systems. However, this finding has never been confirmed or expanded upon.

In the present study, we test directly in two different cultured model systems whether the signaling pathways activated by cAMP induce TH expression in midbrain dopamine neurons. We have chosen to study the cAMP signaling pathway, because cAMP is a powerful inducer of TH gene transcription rate, TH mRNA, and TH protein in most other model systems (Kumer and Vrana, 1996) and because cAMP has been implicated in the induction of midbrain TH in a previous report (Leviel et al., 1991). As expected, our results demonstrate that TH protein is induced after treatment of either midbrain slices or MN9D cells for 24 to 48 h with either forskolin or the cAMP analog 8-CPT-cAMP. A cAMP-mediated activation of pre-existing TH enzyme molecules is also suggested at earlier time points in the MN9D cells and possibly in midbrain slice cultures. However, in the latter model, the increase in TH activity observed at 12 h (Fig. 1) without a concomitant increase in TH protein seems to be due to an increased apparent V max. This result is unexpected, because activation of TH by cAMP is usually associated with a decreased K m for pterin cofactor (Kumer and Vrana, 1996). However, Daubner et al. (1992) have shown that protein kinase A-mediated phosphorylation of TH in the presence of inhibitory concentrations of dopamine produces an activation of the enzyme that is characterized by an apparent increase in V max. More studies are needed to confirm the precise mechanism(s) mediating this enzyme activation process in this model system.

Our studies show that even though treatment with 8-CPT-cAMP induces TH protein, it does not induce TH mRNA in either DIV 1 or DIV 7 midbrain slice cultures or in MN9D cells. These results are in direct contrast to those reported by Leviel et al. (1991), who observed induction of TH mRNA, but not TH protein after microinjection of forskolin into the substantia nigra under in vivo conditions. The reason for this discrepancy is not clear; however, it is probably due to inherent differences using in vivo versus cell culture approaches. For example, microinjected forskolin may be raising cAMP levels in nondopaminergic afferent nerves or neighboring glial cells, leading to stimulation of TH gene transcription in dopaminergic neurons via indirect mechanisms not related to cAMP generation in the dopamine neuron itself. The use of anesthetic in the microinjection studies may be having an unexpected effect. Alternatively, neuronal damage produced by the microinjection may be causing the release of chemicals that stimulate TH mRNA expression. These confounding issues inherent to the in vivo approach are less likely to be observed in the cultured model systems used in the present study. However, it is important to note that the TH gene promoter has a perfect consensus CRE just 35 bp upstream of the transcriptional start site and many studies have shown that this CRE is responsive to cAMP in pheochromocytoma, neuroblastoma, and cath.a cells (Kumer and Vrana, 1996; Lewis-Tuffin et al., 2004). Hence, the lack of an apparent transcriptional response in the cultured dopamine cell model systems is very surprising and how these results relate to the response observed under in vivo conditions requires further investigation. Nevertheless, a significant finding of the present study is that the TH gene in cultured midbrain dopamine neurons apparently responds to cAMP in a different manner than in cultured cells derived from other lineages.

A second major finding in this report is that TH protein is induced by a cAMP-mediated mechanism that regulates TH mRNA translation in midbrain neurons. This conclusion is based on our findings that cAMP increases the rate of synthesis of TH protein without a concomitant increase in TH mRNA levels in both midbrain slices and MN9D cells and increases the percentage of TH mRNA localized with polysomes in MN9D cells. We also show that this translational response is dependent on sequences within the 3′UTR of TH mRNA. These sequences reside in a region that encodes the polyuridyline-rich binding site described by Czyzyk-Krzeska and coworkers (Paulding and Czyzyk-Krzeska, 1999). These workers have shown that this site plays an important role in mediating the stabilization of TH mRNA that occurs in response to hypoxia in PC12 cells. They have also shown that hypoxia induces the levels of a trans-acting protein, designated polyC-binding protein, that apparently binds to this...
site, leading presumably to a decreased degradation of TH mRNA. It is conceivable that proteins binding to this site (or an adjacent site) in MN9D cells may regulate the translation of TH mRNA in addition to its degradation. According to this model, cAMP may induce a protein (or set of proteins) that binds to the 3′UTR of TH mRNA, leading to its increased association with polyosomes and consequently, enhanced translation and synthesis of TH protein. More work is needed to identify the specific sequences within the TH mRNA 3′UTR and the trans-acting factors that mediate this translational response.

This cAMP-mediated translational activation of TH mRNA may be unique to midbrain dopamine neurons. As discussed above, in all other model systems studied so far, cAMP-stimulated increases in TH protein are associated with increases in TH mRNA and TH gene transcription rate. In these models, cAMP is thought to induce TH via signaling pathways that phosphorylate cAMP response element-binding protein or other CRE-binding proteins, which bind to the TH CRE and transactivate the TH gene promoter, leading to the observed transcriptional stimulation. Even though it has not been directly assessed in these other systems, there is no evidence that cAMP regulates TH mRNA translation, because the increases in TH mRNA and TH protein are usually very similar in size. The evidence in this article supports that for this reason, we have not been able to detect any cAMP-mediated increases in luciferase reporter expression directed by the TH mRNA 3′UTR in either PC12 cells or cat.h.a cells. However, in this regard, we have rather shown that under in vivo conditions TH mRNA translation is regulated by stress in rat adrenal medulla (Xu et al., 2007), but there is no evidence yet that this regulation is cAMP-mediated.

The reason for this possibly unique and novel cAMP response in midbrain dopaminergic neurons remains obscure. This mechanism may have developed in these cells, because the more common transcriptional response of the TH gene to cAMP is not functional. Alternatively, this mechanism may be necessary to induce TH in response to nerve stimulation in a particular cellular locale in midbrain neurons. Several synaptic proteins are known to be regulated by translational mechanisms at local dendritic synapses (Krichevsky and Kosik, 2001). Perhaps, TH mRNA is also localized in dendrites and local elevation of cAMP leads to local synthesis of TH protein and increased dopamine biosynthesis in these dendrites. From a pharmacological standpoint, drugs that block receptors linked to inhibition of adenyl cyclase, such as antagonists of D2 dopamine receptors, may raise cAMP in dopamine neurons for long periods, activating this mechanism. More work is needed to test these hypotheses. Nevertheless, our results clearly demonstrate a novel post-transcriptional mechanism that regulates TH mRNA translation. This mechanism may play an important role in maintaining dopamine levels in midbrain dopamine neurons. Further elucidation of the molecular and cellular mechanisms that mediate this translational response may lead to new therapeutic strategies to up-regulate dopamine biosynthesis in nigrostriatal neurons during Parkinson’s disease. It may also provide new avenues for engineering viral vectors designed to overexpress TH and replace dopamine in different disease states.

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**Address correspondence to:** Dr. William Tank, Department of Pharmacology and Physiology, Box 711, University of Rochester Medical Center, 601 Elmwood Ave., Rochester, NY 14642. E-mail: awilliam_tank@urmc.rochester.edu