Increase of Tryptophan Hydroxylase Enzyme Protein by Dexamethasone in Adrenalectomized Rat Midbrain

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Using two specific anti-peptide antibodies (WH-66 and WH-412) against tryptophan hydroxylase (TPH or WH), a single principle band from the midbrain raphe area was seen (approximately 49 kDa) in immunoblots. Densitometric comparison of the immunoreactivity of the 49 kDa band was greater (60-75%) in immunoblots of midbrain raphe samples from adrenalectomized (ADX) rats given dexamethasone (DEX) in their drinking water (10 mg/liter) for 12-96 hr. No difference from ADX brains was seen in the 49 kDa band after only 4 hr of exposure to DEX in the drinking water. Immunocytochemical staining with WH-66 of sections from rat brainstem showed specific cellular staining in all of the serotonergic raphe nuclei but not in substantia nigra or locus coeruleus. More cellular staining of WH-66-immunoreactive (WH-66-IR) cells was observed in the dorsal and median raphe nuclei in ADX rats given DEX for 72 hr, especially in the perikarya and in the primary dendrites. Quantification of staining per cell soma with an automatic image analyzer indicates that amount of WH-66-IR in neurons from both the lateral wing subdivision of the dorsal raphe nucleus and in the supraplum collateral nuclei, B-9, was 80% higher in the ADX+DEX compared to ADX animals. Interestingly, morphometric analysis of these same cells showed a corresponding increase of 37-60% in somal area. It is suggested that part of the increase in TPH/WH staining may be a consequence of cellular hypertrophy due to DEX treatment of the ADX rats.

[Key words: 5-HT, stress, dorsal raphe nucleus, Western blots, immunocytochemistry, anti-peptide antibodies]

Tryptophan hydroxylase (TPH or WH) is the enzyme that limits the synthesis of 5-HT in the brain and melatonin in the pineal. Antibodies raised against purified TPH/WH have been previously reported by several laboratories (Joh et al., 1975; Cash et al., 1985; Weissmann et al., 1987). These antibodies have the potential to show substantial cross reactivity with phenylalanine hydroxylase (PH) and tyrosine hydroxylase (TH) because of the potential to show substantial cross reactivity with phenylalanine hydroxylase (PH) and tyrosine hydroxylase (TH) because of the substantial sequence similarities among the three enzymes. For example, rat WH has 55.7% identity with rat PH and 50.1% identity with rat TH (Darmon et al., 1986). These authors noted that the greatest homology (73.3% with PH and 71.8% with TH) occurred in the central part of the molecule [amino acids (aa) 187-324], which contains five conserved cysteines. However, several sequences in WH were found to be unique for WH, and two of these regions were selected as immunogens for generating WH-specific anti-peptide antibodies.

Antibodies against synthetic peptides have been used to study many previously uncharacterized proteins for which only the cDNA sequence is known. Anti peptide antibodies can be directed against a specific short region of the molecule, which usually is bound to a larger carrier protein to increase its antigenicity (Hollow and Lane, 1988). We have previously reported success in producing anti-peptide antibodies specific for the 5-HT_{\alpha} receptor (Azmitia et al., 1992). In the present studies the amino acid sequences were first selected on the basis of non-overlap with sequences in TII and PII. Additional criteria based on the hydrophilicity and secondary structure were also used to arrive at aa 66-91 and aa 412-443, which corresponded to the N-terminal and C-terminal regions of the enzyme. We report that antibodies WH-66 and WH-412 react with a single major band in immunoblots from midbrain raphe tissue, where the serotonergic neurons were highly concentrated (Azmitia, 1978; Jacobs and Azmitia, 1992).

Previous studies had shown that WH activity was higher in stressed animals than in control or stressed adrenalectomized (ADX) rats (Azmitia and McEwen, 1974; Boadle-Biber et al., 1989). Studies using a protein synthesis inhibitor had suggested that at least part of the increased activity was due to de novo protein synthesis (Azmitia et al., 1976; Boadle-Biber et al., 1989; Singh et al., 1990). Does the regulation of WH by adrenal steroids produce changes in the amount of WH protein present in the serotonergic neurons of the rat midbrain? We now present evidence from immunocytochemical and blot immunolabeling studies that WH protein levels are higher in ADX rats given dexamethasone (DEX) than in ADX rats without steroid replacement.

Abstracts have been previously published (Azmitia et al., 1991; Liao and Azmitia, 1992).

Materials and Methods

Experimental procedures. Thirty-six female Sprague-Dawley rats (Taconic Farms) weighing about 200 gm were used. Thirty-two rats were anesthetized with ketamine (40 mg/kg; Sigma) and xylazine (2.5 mg/kg; Sigma), and both adrenals were removed. The ADX rats were maintained with food and 0.9% NaCl solution on a 12 hr:12 hr light/dark cycle. After 3 or 14 d, the ADX rats were grouped two in each cage, exposed to strong unrythmical sound and 24 hr continuous light, with half of the rats (n = 16) receiving DEX (Sigma) at 10 mg/liter in the 0.9% NaCl drinking solution for various periods before death. Rats were randomly selected from the ADX and ADX+DEX groups for immunochemical versus blot immunolabeling studies. All animals were killed according to NIH guidelines by a protocol approved by the NYU Animal Welfare Committee.
Figure 1. Immunoblots for TPH/WH with antibody WH-412 (1:2000 dilution) using 4-20% SDS-PAGE run at 125 V and stained with avidin-biotin peroxidase. The lanes were loaded with 12.5 μg (lanes 1 and 2) or 25 μg (lanes 3 and 4) of proteins solubilized from midbrain of rats ADX for 8 d with or without DEX treatment on the last 3 d. The lanes of midbrain proteins treated with DEX show increased immunoreactivity at both loading concentrations of proteins for the principle band (MW = 49 kDa) and three heavier minor bands. Lane 5 contains low-molecular-weight biotinylated markers (Bio-Rad).

Immunocytochemistry. Fourteen rats (2 control, 6 ADX, and 6 ADX+DEX) were used for these studies. The control animals were used to establish the staining characteristics of the two antibodies. In the first series of experiments, four ADX rats were kept for 8 d and two of these given DEX in the drinking saline solution for the last 3 d. In the last experiment, eight animals were selected to compare the response of short- (1 week) and long- (2 weeks) term ADX on the effects of giving 3 d of DEX treatment before death.

All animals were perfused through the ascending aorta with one liter of 4% paraformaldehyde and 0.1% MgSO₄ in 0.1 M phosphate buffer (pH 7.4). The brains were postfixed at 5°C for at least 4 hr before processed for immunocytochemistry; 60 μm coronal sections of the brain were cut on a Vibratome (Oxford) and incubated in primary antiserum (WH-66, 1:500 to 1:50,000, 16-24 hr at room temperature) diluted in 0.1 M Tris-HCl-buffered (pH 7.6) saline (TBS) containing 1% normal sheep serum and 0.1% Triton X-100. Immunoreactivity was visualized with the elite Vectastain ABC kit. Lane 5 contains low-molecular-weight biotinylated markers (Bio-Rad).

Western blots for TPH/WH with antibody WH-412 (1:2000 dilution) using 10% SDS-PAGE. Lane 1 is the biotinylated heavy-molecular-weight marker. Lanes 2-9 represent the times of treatment of 0.9% saline or DEX to rats ADX for 5 d. Increase immunoreactivity was seen in the principle band at 49 kDa and the three heavier minor bands after 1 d (lanes 2 and 3). The differences in immunostaining between saline- and DEX-treated brains were maximal at 2-10 d.

Figure 2. Western blots for TPH/WH with antibody WH-412 (1:2000) using 10% SDS-PAGE. Lane 1 is the biotinylated heavy-molecular-weight marker. Lanes 2-9 represent the times of treatment of 0.9% saline or DEX to rats ADX for 5 d. Increase immunoreactivity was seen in the principle band at 49 kDa and the three heavier minor bands after 1 d (lanes 2 and 3). The differences in immunostaining between saline- and DEX-treated brains were maximal at 2-10 d.
ROSTRAL DRN

S A L I N E

D E X

Figure 3. Photographs of the rostral DRN at the level of the oculomotor motor nucleus in ADX rats (8 d) treated with or without DEX for the last 3 d. The neurons were stained with the WH-66 antibody (1:7500). The staining of the nucleus can be clearly seen in both animals. However, the intensity and density of staining appears greater in the animal given DEX. This was especially clear along the midline of the nucleus and just below the cerebral aqueduct. Scale bar, 200 μm.

Blot immunolabeling studies

Anti-peptide antibodies WH-66 and WH-412 were used at dilutions of 1/7500 and 1/2000, respectively, to immunostain nitrocellulose strips of electrophoretic transfers midbrain raphe subjected to SDS-PAGE on slab gels. The midbrain extract was run at two different protein amounts (25 and 12.5 μg) on the SDS gels. Antibodies WH-66 and WH-412 produced similar patterns of immunoreactivity although the bands appeared sharper and more easily resolved with the WH-412 than with the WH-66 antibody. The staining pattern obtained with the

to SDS-PAGE 10% and 4–20% slab gels at 125 V. Proteins were transferred electrophoretically (100 V, 1 hr) to nitrocellulose membrane (0.45 μ; Bio-Rad) at 5°C using a Bio-Rad Trans Blot Cell.

Nitrocellulose sheets were incubated overnight with antiserum WH 412 (1:2000) in 0.1 M TBS (pH 7.6) with 0.1% (v/v) Tween-20 (Sigma) (TTBS). The sheets were rinsed in TTBS for 10 min three times between the reactions with biotinylated goat anti-rabbit IgG (Boehringer-Mannheim) (60 min incubation), streptavidin–horseradish peroxidase conjugate (Boehringer-Mannheim) (60 min incubation), and 0.05% diaminobenzidine (DAB) in TBS, 0.2% nickel ammonium sulfate (Fisher), and 0.01% (v/v) H₂O₂, (5–10 min). The nitrocellulose sheets were photographed using a Leitz camera with TMax 100 film. The negatives were scanned using a soft laser scanning densitometer (model SLR-504-XL; Biomedical Instruments Inc., Fullerton, CA).

Results

Blot immunolabeling studies

Anti-peptide antibodies WH-66 and WH-412 were used at dilutions of 1/7500 and 1/2000, respectively, to immunostain nitrocellulose strips of electrophoretic transfers midbrain raphe subjected to SDS-PAGE on slab gels. The midbrain extract was run at two different protein amounts (25 and 12.5 μg) on the SDS gels. Antibodies WH-66 and WH-412 produced similar patterns of immunoreactivity although the bands appeared sharper and more easily resolved with the WH-412 than with the WH-66 antibody. The staining pattern obtained with the
DORSAL RAPHE NUCLEUS

Figure 4. Photographs of the DRN at its most developed extent. The four main subdivisions were clearly shown. The centromedian subdivision is directly below the cerebral aqueduct. The interfascicular subdivision is funnel shaped and between the medial longitudinal fasciculus. The bilateral lateral group of WH-66-IR neurons comprise the lateral wing subdivisions. Individual WH-66-IR neurons can be seen in the saline-treated animals throughout the three major subdivisions. However, in the DEX-treated animals, the density of WH-66-IR label obscures the individual cells in all subdivisions. This was especially obvious in the midline of the interfascicular subdivision. Scale bar, 200 μm.

WH-412 antibody is shown in Figure 1. A single major band of approximately 49 kDa was visible in lanes from saline- and DEX-treated tissues. Several minor bands were apparent at higher molecular weights (approximately 73, 100, and 116 kDa).

In the first series of studies the animals were ADX for 8 d and half the rats were given DEX in the drinking solution (10 μg/ml saline) for the last 3 d. Comparison of the raphe tissue lanes from saline-treated and DEX-treated animals showed more dense staining of the 49 kDa band after DEX treatment (Fig. 1). This was apparent after loading with either 12.5 or 25 μg of protein. It also appeared that the higher-molecular-weight bands were similarly increased after DEX treatment (Fig. 1). Densitometric analysis of staining intensity measure in the 49 kDa bands showed a DEX-induced increase of 72% and 45% in the 12.5 and 25 μg lanes, respectively, compared to saline-ADX control animals.

The time course of the DEX-induced increase in staining of the gels was tested by making protein extracts after 4 hr and 1, 2, 3, or 10 d of DEX treatment to rats that were ADX 5 d before steroid treatment (Fig. 2). The results with loading 10 μg of protein showed that the difference in the major band between DEX and saline treatment was not seen after 4 hr, but was apparent after 1 d of treatment with a difference in the density of 32%. The difference after 2 d was 53%, and no further increase
CAUDAL DRN/MRN

Figure 5. Photographs of the caudal DRN and the MRN at the level of the ventral nucleus of Gudden ventrolateral to the medial longitudinal fasciculi. Notice at this level that the interfascicular subdivisions of the DRN and MRN are fused (this grouping was termed the nucleus centralis superior). Comparison of the two photographs shows that individual cells appear to be equally reactive in both saline- and DEX-treated ADX rats. However, as seen in the previous sections, the midline zone especially in the DRN was more dense after DEX treatment. Scale bar, 200 μm.

was noted after either 3 or 10 d (52% and 48%, respectively). The high-molecular-weight minor bands also showed an apparent increase after a single day of DEX treatment.

Immunocytochemistry of brainstem sections

Serial sections were made through the brainstem of saline- and DEX-treated ADX animals to include the entire superior groups of 5-HT-containing neurons. Antibodies against WH-66 showed specific DAB staining compared to preimmune staining at dilution from 1:1000 to 1:50,000, with the best labeling at a dilution of 1:7500. The preimmune serum for WH-66 showed no specific staining at any dilution. WH-412 showed no specific staining over a wide range of dilutions (1:500 to 1:50,000).

WH-66-IR cell bodies and dendrites were clearly seen in all nuclei previously reported to contain 5-HT. No specific cellular staining was seen in the substantia nigra or in the locus coeruleus. Detailed immunocytochemical analysis was performed on the main ascending 5-HT-containing nuclei in the midbrain and rostral pons. These included the dorsal raphe nucleus (DRN) from its most rostral to its caudal extent (Figs. 3-5, respectively), the median raphe nucleus (MRN) at its rostral (Fig. 6) and caudal end (Fig. 5), and the ventrolateral supramesencephalic nucleus [SLN (B9), Fig. 6]. The principle subdivisions of the DRN were all filled with WH-66-IR neurons; the lateral wings located dorsolateral to the medial longitudinal fasciculus (Fig. 7), the centromedian cluster located on the midline immediately below
Figure 6. Photograph of the rostral MRN just dorsal to the decussating fibers of the brachium conjunctivum and the ventrolaterally stretching cells of the SLN (B9) immediately dorsal to the medial lemniscal fibers. The cellular and nuclear pattern of WH-66-IR neurons appears similar in the MRN, but a markedly increase level of staining was seen in the SLN. Scale bar, 100 μm.
dra differences seen in the DRN cells. However, the WH-66-IR neurons in SLN (B-9), which extends ventrolaterally from the MRN, showed a much greater intensity of staining in the DEX-treated ADX rats (Fig. 6).

Higher-magnification examination of the subdivisions of the DRN was performed. The lateral wing WH-66-IR neurons in the saline-treated brains showed poorly stained cell bodies and short processes extending from the neurons (Fig. 7). In contrast, the lateral wing WH-66-IR neurons in the DEX-treated brains appeared more intensely stained in the cytoplasm and in the proximal dendrites (Fig. 7). In addition, a number of long, fine fibers projected between and from these labeled neurons in the DEX-treated rats. A similar pattern was seen in the centromedian cluster (Fig. 8). The most striking change after DEX treatment to ADX rats could be seen in the fine fiber plexus above the labeled cells immediately below the cerebral aqueduct where the DEX-treated section showed intense labeling. Finally, in the interfascicular group of WH-66-IR neurons from the ADX rats, a large number of moderate- to heavy-stained cells were seen, a few with long processes (Fig. 9). In comparison, the WH-66-IR staining in the DEX-treated brain was more intense and in places individual cells were difficult to resolve. Note the very large number of processes between the cells in the DEX-treated compared to the saline-treated ADX brains.

Figure 7. Photograph taken at a higher magnification (250×) of the lateral wing subdivision of the DRN. The WH-66-IR neurons look equally stained, but the size seems larger and the branching more extensive after DEX treatment. Note the large number of dendrites branching from the soma of DEX-treated neurons compared to the saline control. Scale bar, 100 μm.
Figure 8. Photograph taken at 250× of the centromedian subdivision of the DRN. The WH-66-IR neurons lie below the cerebral aqueduct and individual neurons appear equally dark in both sections from the saline- and DEX-treated ADX animals. Noticeably, more dendrites and unidentified processes were seen immediately beneath the cerebral aqueduct in the DEX-treated brain. Scale bar, 100 μm.

Oil high-magnification (Leitz 63×/1.40 Planapo) examination of the lateral wing subdivision of the DRN (Fig. 10) and the SLN (B9) (Fig. 11) compares individual cells from the saline- and DEX-treated brains. In the saline-treated brains, the cells show heterogeneous label with some cells darkly and the others lightly labeled. The label was mainly in the perikaryon with short, thin processes emanating from the cell body. In contrast, the DEX-treated neurons in both areas (Figs. 10, 11) were more uniformly heavily labeled and thick, often bulbous, processes extending from the cytoplasm. The photograph of the lateral wing neurons of the DEX-treated brains (Fig. 10) illustrates many examples of WH-66-IR bulbous fibers. In the SLN (B9), several of the WH-66-IR processes also have enlarged varicosities that extend from the cell body. Finally, the size of cell body and primary dendrites of the WH-66-IR labeled neurons in the DEX-treated animals in both areas appeared to be larger.

In order to determine if long-term ADX rats responded differently to DEX treatment than short-term ADX rats, ADX rats were maintained on saline solution alone for either 1 (short) or 2 weeks before DEX was added to the drinking for 3 d. The increase in immunostaining described above was again seen in the short-term ADX rats when given DEX in the saline drinking water (Fig. 12). However, in the long-term ADX animals, the staining was weaker in both the ADX rats given saline alone and after DEX treatment (Fig. 13). Nevertheless, the difference between saline alone and saline with DEX in the drinking water...
was still apparent in the WH-66-IR staining pattern in raphe nuclei, especially in the lateral wings of the DRN of the short-term ADX + DEX.

**Morphometric analysis**

The increase staining observed at high magnification (see Figs. 10, 11) after ADX in the short-term ADX + DEX appeared to be accompanied by an increase in neuronal size. Image intensity measurements of the cellular WH-66-IR quantity and morphometric analysis of soma area were performed in selected neurons from the lateral wings of the DRN and the neurons in SLN (Table 1). The WH-66-IR level in individual neurons was increased after treatment with DEX by 80% in the DRN-lateral wing ($p < 0.05$) and by 76% in the SLN (B9). In conjunction, the soma area ($\mu m^2$) measurements of these same WH-66-IR neurons showed that the area in the ADX + DEX was increased by 79% in the DRN-lateral wing ($p < 0.05$) and by 37% in the SLN (B9) compared to ADX rats maintained on saline alone. If these results were expressed as a ratio of the amount of WH-66-IR divided by the area for each neuron, the significant differences seen in DRN-lateral wing was no longer present. In fact, the ratios were essentially the same between the two groups (ADX + DEX was 101% of ADX alone). A similar, but not as dramatic, decrease in the differences seen when the results were expressed as a ratio in the SLN (B9) neurons.

Figure 9. Photograph taken at 250 x of the interfascicular subdivision of the DRN. Many WH-66-IR neurons can be seen in the section from the ADX animals maintained with saline solution. The soma labeling does not stain the nucleus, which was clearly seen as a clear circle in a number of the neurons. There was substantially more WH-66-IR after DEX in this area and individual cells were difficult to identify. WH-66-IR processes within this subdivision appeared more numerous after DEX treatment. Scale bar, 100 $\mu m$. 
Figure 10. Photograph at highest magnification taken at 630x of WH-66-IR neurons in the lateral wing subdivision of DRN. Individual cellular labeling was easily seen in ADX rats given saline alone or with DEX. The WH-66-IR neurons in the saline-treated brains were fusiform or oval with few thin processes emanating from the soma. The cell nucleus can be seen to be unlabeled. The neurons of the DEX brain appear more pyramidal in shape and with more pronounced bulbous dendrites extending from the soma. The nucleus was also unlabeled in these neurons. Scale bar, 30 μm.

**Discussion**

The availability of a rodent TPH/WH-specific antibody with no cross-reactivity for TH has made this study possible. The peptide sequences selected for raising the two anti-peptide antibodies against WH showed no homology with the sequences of TH and PH (Darmon et al., 1986), and no specific staining was seen in brain nuclei that contain TH, the substantia nigra and locus coeruleus. Thus, the results presented in this report deal only with the changes in WH, the rate-limiting enzyme for 5-HT biosynthesis. Polyclonal antibodies raised against the whole WH protein have shown considerable cross-reactivity with both TH and PH (see Cash et al., 1985). A monoclonal antibody raised against PH from monkey liver (PH8) stains WH but not TH in human brain fixed with 4% paraformaldehyde and paraffin embedded, and in rat brain sections the PH8 antibody labels TH as well as WH (Haan et al., 1987; Törk et al., 1992). Among the two anti-peptide antibodies that we made against WH, WH-412 was more sensitive for immunoblotting analysis and only WH-66 was useful for immunocytochemistry of rat brain sections.

Rat WH has a predicted molecular weight of 51,010 Da (Darmon et al., 1988). Running SDS-PAGE gels with midbrain raphe tissue under reducing and denaturing conditions produced a single principle band with an apparent molecular weight of 49 kDa. A series of higher-molecular-weight bands, heavily labeled...
in nonreducing conditions (data not shown), were lightly stained in our gel (see Figs. 1, 2). There is evidence that WH exists as a protein complex with a molecular weight of 288 kDa and 300 kDa composed of proteins of 59 kDa (Nakata and Fujisawa, 1982). These results indicate that WH exists as a tetramer of four identical subunits, and in a cross-linking experiment with purified WH, SDS-PAGE showed bands of 45 kDa, 90 kDa, 135 kDa, 180 kDa, 225 kDa, and 270 kDa. In our studies, midbrain raphe brain tissue produced a single principle band of 49 kDa in normal and ADX rats with minor bands at 73 kDa, 90 kDa, and 117 kDa. The intensity of both the major and minor bands increased after administration of DEX in the drinking water (Figs. 1, 2).

A time course of WH-412 immunoblot staining after addition of DEX to the drinking water showed that an increase of 32% was seen after only 24 hr of treatment with DEX (Fig. 2) and a maximal increase of 50% was reached by 2 d. The rate of increase suggests an enzyme turnover rate of approximately 1.7 d that was consistent with the measure of 1.43 d measured after p-chlorophenylalanine (Weissmann et al., 1990). A 50% increase over the ADX value was seen at 2, 3, and 10 d. This indicates that the rate of translation rose to a new steady-state level (synthesis/degradation + transport) and was not permanently activated by a de novo supply of enzyme. Unpublished observations (E. C. Azmitia, D. M. Stone, T. H. Joh, and D. H. Park) suggest that short-term ADX rats treated for 3 d with...
Figure 12. Photograph shows the main extent of the DRN 7 d after ADX given saline alone or with DEX added for the last 3 d. The saline-treated animals show good WH-66-IR with the WH-66 antibody with individual cells seen in all subdivisions. Individual cells were not seen because of the density of labeling in the DEX-treated ADX animals. This was most apparent in the centromedian and interfascicular subdivisions where the intraneuronal space was filled with heavily labeled processes. This was best appreciated in the lateral wing subdivision of the DRN. Scale bar, 200 µm.

DEX do not show elevated WH mRNA levels as measured by in situ hybridization with a full-length WH probe (see Kim et al., 1991; Kim, 1992). Thus, the increase in protein amount may reflect a posttranslational mechanism that was coupled to the increase in soma size (see below).

In our preparation, the increase in WH-66-IR was accompanied by an increase in the size of neuronal dendrites and perikaryon. Most of the raphe nuclei showed increased WH-66-IR staining due to an increase in the amount of WH-66IR in the perikarya and primary dendrites of the neurons. Morphometric analysis of the perikarya area revealed an increase of 80% in the lateral wings of the DRN that was sufficient to account for the increase in both the image intensity readings of the immunocytochemically stained neurons and in the densitometry measures from 49 kDa bands in the immunoblots. In the WH-66-IR neurons of the SLN, the increase in somal area only partially compensated for the increased in the intensity of WH-66-IR.

It may be proposed that the increase in WH-66-IR was secondary to the changes in cytoplasmic area in the soma and dendrites. Similar effects of adrenalectomy on somal area in ADX rats have been reported in the hippocampus. Adrenalecto-
LONG-TERM ADX DORSAL RAPHE NUCLEUS

SALINE

DEX

Figure 13. Photograph shows the main extent of the DRN 2 weeks after ADX given saline alone or with DEX added for the last 3 d. The saline-treated brains show weak WH-66-IR with the WH-66 antibody, especially in the lateral wings and the centromedian subdivisions. The WH-66-IR neurons in the DEX-treated animals were well stained, but the processes were not as clearly visible as seen in the short-term ADX rats. This can best be appreciated in the neurons of the lateral wings of the DRN, where only the occasional process was visible. Scale bar, 200 μm.

The hippocampal neurons contain high levels both of corticosterone type I (aldosterone) and type II (dexamethasone, RU28362) receptors. In the hippocampus, aldosterone is more effective than RU28362 in reducing the number of pyknotic cells in granule layer of the dentate gyrus seen in short-term ADX rats, suggesting that type I receptors mediate this effect (Woolley et al., 1991). However, the loss of 5-HT₆ mRNA, the reduced molecular layer, and Nissl labeling of the granular layer seen in long-term ADX dentate gyrus are reversed by short-term exposure to a type II receptor agonist (Liao et al., 1993). The midbrain raphe neurons contain only type II corticosterone re-
Previous studies have shown that the activity of tryptophan hydroxylase (WH) in adrenalectomized (ADX) rats could be increased by replacement with adrenal steroids. In rats ADX for a week, a 107% increase in the activity of WH in the midbrain was seen after daily injections of 1 mg of corticosterone for 5 d before death (Azmitia and McEwen, 1969). Similarly, rats ADX for 8 d and treated with DEX (500 μg/d) for the last 5 d also showed an increase in WH activity in midbrain (47%) and cortex (60%) (Singh et al., 1990). In this latter report, the increase in WH activity was resistant to alkaline phosphatase treatment and was not attributed to an increase in phosphorylation of the enzyme (Izumoto et al., 1978). Furthermore, aldosterone was without effect and the effect of chronic treatment was proposed to occur through the type II corticosterone receptor, which has an extensive distribution throughout the brain (Reul and de Kloet, 1985; McEwen et al., 1986; Funder and Shephard, 1987). These results were consistent with the changes seen in our immunoblot analysis and the densitometry measurements reported here. The previous observations that WH activity was increased after chronic stressors in normal but not ADX rats (Azmitia and McEwen, 1976; Boadle-Biber et al., 1989) indicate that these changes in WH activity are physiologically relevant. Changes in 5-HT turnover after stress and ADX have been reported (Thierry et al., 1968; Azmitia et al., 1970; De Kloet et al., 1982).

Glucocorticoids have been shown to change the transcriptional rate of only a few proteins by acting through the steroid promotor sequence (Yamamoto, 1985). There is a decrease of 60% in glial fibrillary acidic protein (GFAP) mRNA in rat brain after 8 h (Nichols et al., 1990). The decrease was due to activation of the type II corticosterone receptor since RU 28362 was the most effective steroid in decreasing GFAP mRNA. This drop in mRNA levels was accompanied by a corresponding decrease in GFAP tissue levels (O’Callaghan et al., 1991). These reports again implicate the type II corticosterone receptor in the regulation of a brain protein.

An increase in the mRNA for TH in locus coeruleus have been reported after isolation stress (Angulo et al., 1991). The changes were 18%, 42%, or 68% above control after 7, 14, or 28 d of isolation stress, respectively. The long delay in this increase argues against a direct regulation by adrenal steroids on the transcriptional levels of TH in the locus coeruleus. This was in contrast to the rapid increase in TH activity seen after a variety of treatments (Zigmond, 1988). These results with TH activation provide evidence for increase protein amounts before an increase in mRNA.

The fact that adrenal steroids can influence the expression of a structural protein in hippocampal astrocytes (Nichols et al., 1990) raises the possibility that structural proteins in neurons may be regulated by adrenal steroids, and secondarily produce an increase in the levels of transmitter-related proteins without a change in transcription. It fact, nuclear runoff assays of WH gene transcription rates revealed similar levels of gene expression for pineal and midbrain although the RNase protection assay showed much higher steady-state mRNA levels in pineal than in midbrain (Hart et al., 1991). These authors concluded that the control of WH mRNA levels was posttranscriptional, which agrees with the findings showing a greater translational efficiency in brain (Dunas et al., 1989). These authors found that although the level of WH mRNA was at least 150 times lower in the raphe nuclei than in the pineal, the WH antigen was three times more abundant in raphe than pineal. The mechanism of posttranslational modification remains to be determined, but may involve structural proteins since our results indicate that the greater level of WH immunoreactivity seen after DEX treatment was correlated to an increase in soma area.

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### Table 1. The computer measurements of the intensity of WH-66-IR per cell, the soma cell area, and the ratio of intensity:area of 10 WH-66-IR neurons from either the lateral wing (LW) of the DRN or the neurons of the suprapulmonary nucleus (B9) are shown in ADX rats with or without DEX treatment

| Treatment               | Intensity   | Cell area (μm²) | I/CA            |
|-------------------------|-------------|----------------|-----------------|
| ADX, DRN-LW             | 20 ± 4      | 167.5 ± 26     | 0.1194 ± 0.015  |
| ADX + DEX, DRN-LW       | 36 ± 4*     | 300.0 ± 48*    | 0.1183 ± 0.026  |
| (% of ADX)              | (180%)      | (179.1%)       | (101%)          |
| ADX, B9                 | 22 ± 8      | 171.3 ± 40     | 0.119 ± 0.0257  |
| ADX + DEX, B9           | 39 ± 7      | 234.6 ± 40     | 0.1530 ± 0.012  |
| (% of ADX)              | (176%)      | (137%)         | (129%)          |

The results are presented as the average ± SD. I/CA is intensity over cell area.

* p < 0.05, two-tailed Student’s t test.
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