INTRODUCTION

Tryptophan hydroxylase (TPH, EC 1.14.16.4) is the first and the rate-limiting enzyme in 5-hydroxytryptamine (5-HT, serotonin) biosynthesis (1, 2). It catalyzes the conversion of L-tryptophan into 5-hydroxytryptophan in brain, pineal gland (PG), and enterochromaffin cells of gut. In the central nervous system serotonergic cell bodies are mainly localized in the raphe system and its terminals are spread throughout the central nervous system (3-6). Studies of TPH characteristics and regulation have been hampered partly because of the enzyme instability and partly due to the low abundance in animal tissues resulting in the low availability of purified enzyme protein. End product regulation of 5-HT biosynthesis at the rate-limiting step appeared to exist in vivo (7, 8) but even high concentrations of 5-HT did not inhibit TPH activity in vitro (9). Meanwhile TPH mRNA levels are greater in PG than in the dorsal raphe nucleus (DRN), while TPH activity is higher in DRN than in PG (10, 11). It has been also reported that brain TPH was immunochemically different from the mouse intestinal TPH (peripheral enzyme) because antibodies against the mouse mastocytoma TPH immunoprecipitated TPH from intestine in a dose-dependent manner but did not significantly immunoprecipitate TPH from mouse brain (12). In addition, some biochemical characteristics of brain TPH have been reported to differ from pineal TPH in the previous reports (11, 13). Although TPH was extensively purified from rat brain and a mouse mastocytoma cell line, the quantity of purified enzyme protein was minimal (14-16). Antiserum against rat brain TPH has been produced, but it also contained some antibodies to catalase (16). Therefore, extensive purification to remove contaminant catalase antibodies was required (16). The production of anti-peptide antibodies to rat TPH has also been reported (17) but these antibodies detect not only a TPH-like protein band on immunoblot but also three additional non-defined higher molecular weight bands from extracts of the rat raphe area. Recently, recombinant mouse and rabbit TPH were successfully expressed in large quantities in bacteria, using bacterial expression vectors and was shown to be enzymatically highly active (18, 19). Specific polyclonal antiserum to the

Immunochemical Characterization of Brain and Pineal Tryptophan Hydroxylase

Recombinant mouse tryptophan hydroxylase (TPH) was expressed in Escherichia coli, using a bacterial expression vector and has been purified to homogeneity by sonication followed by Sepharose 4B column chromatography and native slab gel electrophoresis. This purified enzymatically active TPH protein was used for production of a specific antiserum. This antiserum identified the predicted TPH band (molecular weight, 54 kDa) on Western blot of crude extracts from the rat and mouse dorsal raphe, and the rat pineal gland. However, this antiserum recognized an additional protein band of lower molecular weight (48 kDa) in pineal extract. It is not clear whether the 48 kDa TPH band represents an isozyme or a protease cleavage product of TPH. Since the pineal gland contains higher TPH mRNA and lower TPH activity when it is compared with dorsal raphe nucleus enzyme, this lower molecular weight TPH may participate in the reduced TPH specific activity. In addition, there are no specific TPH inhibitors in the pineal gland and this lower molecular weight TPH is inactive or has a very low specific activity. This antiserum specifically immunostained serotonergic cell bodies in the dorsal raphe nuclei, some large caliber serotonergic processes in the dorsal raphe area as well as terminals in the olfactory bulb. It also immunolabeled the pineal gland and immunoprecipitated equally well TPH protein from the dorsal raphe nucleus and the pineal gland in a concentration-dependent manner.

Key Words : Tryptophan Hydroxylase; Recombinant; Raphe Nuclei; Pineal Body; Immunochemistry

Received : 1 December 2000
Accepted : 26 March 2001

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subunits of the recombinant mouse TPH (the holoenzyme consists of four identical subunits) has been produced in rabbit by injecting TPH band cut from SDS-polyacrylamide gels (18). However, this antiserum, although specific, had a major drawback for its use in immunocytochemical localization of serotoninergic neurons in brain due to some nonspecific immunostaining which perhaps resulted from usage of SDS-containing antigen for immunization.

TPH protein in rat brains is present in very small amounts (even in DRN) and very unstable due to the presence of small quantities of TPH protein during purification. The conventional many-step purification requires many rat brain tissues in addition to being difficult to maintain its enzyme activity. Also, the relative amount of TPH protein in brain tissue is minor compared to the other proteins present in the brain. Therefore, even if TPH preparation from one-step purification of Sepharose 4B column chromatography was applied to native gel electrophoresis, it is impossible to detect TPH activity with a maximum load of protein. On the other hand, the recombinant mouse TPH preparation after one-step purification of Sepharose 4B column chromatography was enzymatically more active (434 nmol/mg protein/min at 37°C, 2.2 mg from 100 mL of culture) in our previous study (20) and contained more abundant TPH protein (one of the major protein bands is TPH protein in native slab gel electrophoresis) than purified TPH (374 nmol/mg protein/min at 30°C, 19 μg from 70 rat brain stems) reported by others (14). A large quantity of enzymatically active recombinant mouse TPH preparation can be easily produced using an expression vector containing the full coding region of the TPH cDNA just by growing culture (20).

In the present study, we describe a simple two-step procedure for purification of recombinant mouse TPH and production of immunocytochemically specific TPH antisera to enzymatically active recombinant mouse TPH. Since we demonstrated that the coding sequence cDNA from DRN and PG was identical, obviously the next question raised is whether TPH protein from both sources are the same or not. In order to examine the identity of TPH protein from DRN and PG, we chose the commonly used techniques, i.e., immunocytochemical staining of central serotoninergic neurons as well as pinealocytes, determination of molecular weight by Western blotting, and immunoprecipitation of TPH protein by immunoirradiation with specific TPH antisera, to immunocytochemically characterize TPH protein from both sources.

MATERIALS AND METHODS
Expression of mouse TPH in Escherichia coli

We constructed a bacterial expression vector, pKSTPH, containing the full coding sequence of TPH (1.3 kb), as described elsewhere in detail (20). Briefly, the TPH coding region was ligated into the EcoRI site of the plasmid, pK S6.8. pK S6.8, a vector containing a strong tac promoter, which can be regulated in bacteria by lac repressor, and which is inducible by isopropyl-β-D-thiogalactoside (IPTG) (Gold Biotechnology, Inc., St. Louis, MO), was constructed by ligating a 1.5 kb DNA fragment from a PvuI, partial Sall digest of pK K233-3 (Pharmacia), with a PvuI/Sall digested 5.3 kb DNA fragment from PIN IIA3 (21) generously provided by Dr. M. Inouye (Department of Biochemistry, Rutgers University, N.J.). This plasmid contains an intact lacI gene, which permits induction of the foreign gene by IPTG regardless of the genotype of the host E. coli strain. pKSTPH was transformed into E. coli strain MC 1061, inoculated into LB broth containing 100 μg/mL ampicillin, and induced by IPTG. Cells were harvested by centrifugation at 4,000 g for 20 min. The sediment was suspended in sonic buffer (5 mM Tris-HCl, pH 7.5, 10% glycerol, 50 μM EDTA, 0.06% Tween 20, 2 × 10-3 M tryptophan, 1 mM phenylmethylsulfonyl fluoride, leupeptin (50 μg/mL), 1% v/v aprotinin (Sigma Chemical Co., St. Louis, MO) and lysed by sonication for 30 sec in ice-water with the use of an ultrasonic model set (Vibra Cell, Sonics and Materials, Inc., Danbury, CT) at 60% of maximum output, followed by freeze-thawing twice and centrifuged at 16,000 g for 10 min. The supernatant was used as a source of recombinant mouse TPH, unless otherwise specified. In case of antibody production the supernatant was partially purified by Sephrose 4B column (5 × 86 cm, equilibrated with 10 mM Tris-HCl buffer, pH 7.0, containing 2 × 10-3 M tryptophan and 0.5 mM dithiothreitol) chromatography prior to subjection of TPH preparation to gel electrophoresis.

Enzyme and protein assay

TPH activity was determined by a minor modification of the method of Park et al. (18, 22), in which the conversion rate of L-[1-14C]tryptophan to 14CO2 was measured. Briefly, the assay mixture containing 25 μmole of Tris-acetate buffer, pH 7.5, 112.5 nmol of DL-6-methyltetrahydropteridine (Calbiochem., La Jolla, CA, U.S.A.), 600 nM of DL-dithiothreitol, 5.1 nM of Fe (NH4)2 (SO4)2 6H2O, 43 units of catalase (Sigma C-10), and 37.5 nM of L-[1-14C]tryptophan (0.1 μCi) American Radiolabeled Chemicals, Inc., St. Louis, MO in a total volume of 50 μL was added to 100 μL of enzyme solution. The mixture was incubated at 37°C for 15 min and then 50 μL of solution containing partially purified hog kidney aromatic L-amino acid decarboxylase (30-50% ammonium sulfate fraction, the activity in 18 nM product formed/min at 37°C) and 50 nM of pyridoxal phosphate in H2O was added. The assay tube was then covered by a rubber cap in which a plastic well containing a piece of filter paper soaked in 100 μL of NCS II (Amersham)
Characterization of Tryptophan Hydroxylase

491

was suspended. The mixture was further incubated at 37°C for 30 min. The reaction was then stopped by injecting 100 µL of 30% HClO4 and 3H2O was trapped at 37°C for 30 min and counted with 15 mL of Econofluor (DuPont-NEN). Activity was expressed as nmol of CO2 formed per 15 min at 37°C. Protein concentration was measured according to Lowry et al. (23) using bovine serum albumin as standard.

Electrophoresis

Polyacrylamide gel electrophoresis was carried out by a minor modification to the standard procedure (24, 25) using 0.4 M glycine-Tris buffer, pH 8.2 in a 10% slab gel including 4% stacking gel (0.15 cm in thickness × 14 cm in width × 16 cm in length). Using partially purified recombinant mouse TPH (2.78 mg protein loaded on each gel), electrophoresis was performed at 25 watts for 3 hr at 12°C, following 1 hr prerun. The gels were also stained for proteins with Coomassie blue R-250. Following electrophoresis, the gels were cut consecutively (0.2 cm in length × 0.15 cm in thickness × 1 cm in width) and TPH activity measured as described above.

Production of Antiserum

Antibodies to recombinant mouse TPH were raised in rabbits by injecting the sections of the gels containing peak enzymatic activity. Active band containing about 200 µg protein were cut and homogenized in 1 mL of 0.9% NaCl and an equal volume of complete Freud's adjuvant. The resultant emulsion was injected subcutaneously at multiple sites on the back of rabbits. Booster injections of the antigen, using incomplete Freund's adjuvant were given on days 43 and 64 and the animals were bled 1 week later. Handling of rabbits, as well as rats and mice (see below) followed the guidelines of the Institutional Animal Care and Use Committee of the Cornell University Medical College.

Immunocytochemistry

For immunocytochemistry, rats and mice were anesthetized with sodium pentobarbital (50 mg/kg, i.p.) and perfused transcardially with heparinized saline containing 0.5% sodium nitrite, followed by 0.1 M sodium phosphate buffer, pH 7.2, containing 4% formaldehyde generated from paraformaldehyde. The brain and pineal gland were post-fixed in 4% buffered formaldehyde for 1 hr, then rinsed in the buffer and placed in 30% sucrose at 4°C overnight. Tissue sections of 30-40 µm thickness were cut on a sliding microtome and collected in standard wells filled with 0.1 M sodium phosphate buffer. Tissue sections were incubated in 0.1 M sodium phosphate-buffered saline pH 7.4, containing 0.2% Triton X-100 and 1% bovine serum albumin. Tissues were washed in phosphate-buffered saline containing 0.5% bovine serum albumin and incubated overnight with a specific rabbit polyclonal antiserum to recombinant full-length TPH described above. The antiserum was used at a dilution of 1:10,000. A diluted 5-HT antiserum (1:30,000) was used for the purpose of comparison. Tissue was incubated with biotinylated goat anti-rabbit IgG (Vector Laboratories, Burlingame, CA) for 1 hr and then treated with a Vecta Stain Elite Kit as per manufacturer's direction. The antigen was visualized with 3,3'-diaminobenzidine-HCl (50 mg/100 mL) and 0.005% hydrogen peroxide as a chromogen. Sections were mounted onto gelatin-coated slides, dehydrated through graded ethanols and coverslipped with Permount (Fisher Scientific).

Western blotting

Enzyme sources are 16,000 g supernatant of rat and mouse DRN and rat PG homogenate in 50 mM Tris-acetate buffer, pH 7.5, containing 0.5 mM dithiothreitol and recombinant mouse TPH preparation mentioned above. Western blotting was performed according to our minor modification (26) of the standard methods described by Towbin et al. (27) and Burnette (28). Briefly, proteins were subjected to gel electrophoresis using 10% SDS-polyacrylamide slab gels containing 4% stacking gel as described by Laemmli (29). Following electrophoresis, the resolved proteins were transferred at room temperature (200 mA for 3 hr) to Nitro-Screen West membrane (Dupont-NEN) in 192 mM glycine/25 mM Tris buffer, pH 8.3 containing 20% methanol. The membrane was then incubated overnight in blocking buffer (150 mM NaCl, 50 mM Tris-HCl, pH 7.4 and 1% gelatin) to avoid nonspecific binding. The blot was rinsed with 150 mM NaCl, 50 mM Tris-HCl, pH 7.4, 5 mM EDTA, 0.25% gelatin and 5% normal goat serum and incubated for 2 hr with TPH antiserum described above, diluted (1:30,000 dilution for recombinant mouse TPH and 1:10,000 dilution for other sources of TPH) in rinsing buffer. The blot was rinsed in the above buffer and then incubated for 2 hr with a secondary antibody (peroxidase-conjugated goat anti-rabbit IgG (Kirkegaard and Perry Laboratories, Inc., Gaithersburg, MD) diluted (1:200) in rinsing buffer. The antigen-antibody complexes were visualized with 0.05% 3,3'-diaminobenzidine containing 0.005% hydrogen peroxide in 50 mM Tris-HCl buffer, pH 7.4.

Immunotitration

Increasing amounts of antiserum to recombinant mouse full-length TPH were added to the fixed quantity of enzyme to be titrated and the total volume was adjusted with premune serum. The mixture was incubated for 1 hr at room temperature with occasional shaking and then for 16 hr at 4°C. The antigen-antibody complex was removed by cen-
trifugation for 10 min at 16,000 g, and a 100-μL aliquot of
the supernatant was assayed for residual TPH activity (25).

RESULTS
Electrophoretic profile of recombinant mouse TPH
Crude recombinant mouse TPH (16,000 g supernatant,
103 mg protein) was subjected to Sepharose 4B column (5
× 86 cm) chromatography, proteins were eluted with 10
mM Tris-HCl buffer, pH 7.0, containing 2 × 10⁻⁵ M trypto-
phan and 0.5 mM dithiothreitol, and the most enzymatically
active fractions were combined. Subsequently, the partially
purified recombinant mouse TPH subjected to slab gel
electrophoresis with 10% gel at pH 8.2 showed a single
enzymatically active peak located at 2.3 cm from the top of
the separating gel (Fig. 1). This active peak overlapped with
one major protein band stained with Coomassie blue. The
distribution of enzyme activity (nM/15 min at 37 °C/section)
and staining pattern of proteins on gel are shown in Fig. 1.

Western immunoblotting
Western blot analysis was performed to determine whether
the antiserum raised is specific and TPH proteins from the
extracts of the rat and mouse DRN, the rat PG and recom-
binant mouse TPH preparation are immunochemically
identical. A specific rabbit polyclonal antiserum to recom-
binant mouse TPH was used for this analysis. The TPH
antiserum recognizes an identical band with the same
molecular weight (54 kDa) in all four TPH sources (Fig. 2).
This antiserum also detects an additional and equally abund-
ant immunoreactive band with lower molecular weight in
the extract of the rat PG. However, no immunoreactive
band was detected with the preimmune serum (data not
shown). Molecular weight was estimated by comparison to
Coomassie blue stained marker proteins as indicated in the
margin of Fig. 2. The main TPH band (MW 54 kDa)
among these four sources is indicated by an arrow. Possibly
because of the higher enzyme concentration of TPH in the
recombinant enzyme preparations as compared to the ani-
mal tissue extracts, a higher dilution of antiserum was ade-
quate to detect the TPH band on Western blot.

Immunocytochemical localization of TPH
The new TPH antiserum localized TPH immunoreactive
cells in the DRN. Cells were below the aqueduct and ven-
tromedial area as well as in lateral portions of the DRN of
rat and mouse (Fig. 3A and C). Additionally, moderate
immunoreactivity was also observed in the medial raphe
nuclei. Immunostaining in adjacent tissue sections of rat

Fig. 1. Distribution of recombinant mouse TPH activity after
native polyacrylamide gel electrophoresis. Partially purified
enzyme preparation (2.78 mg protein) obtained from Sepharose
4B column chromatography was loaded on native slab gel (0.15
cm in thickness × 14 cm in width × 16 cm in length). Following
electrophoresis as described in Materials and Methods, the gel
was sliced into 0.2-cm sections (0.15 cm in thickness × 1 cm in
width × 0.2 cm in length). The sections were assayed as
described in Materials and Methods. The equivalent-sized gel
section was stained for protein with Coomassie blue. Note that a
single enzymatically active peak corresponding to a major
stained protein band was observed. The activity was expressed
in nM/15 min at 37 °C/section.

Fig. 2. Western blots of rat and mouse brain, rat pineal and
recombinant mouse TPH. Following SDS-polyacrylamide slab
gel electrophoresis and subsequent transferring to a membrane,
resolved proteins were analyzed as described in Materials and
Methods with polyclonal antiserum to recombinant mouse TPH.
Note that an identical TPH band (indicated by an arrow) was
detected with crude extract of recombinant mouse TPH (lane 1,
20 μg), rat dorsal raphe nuclei (lane 2, 200 μg), rat pineal gland
(lane 3, 200 μg), and mouse dorsal raphe nuclei (lane 4, 200 μg).
An additional immunoreactive band with smaller molecular
weight was also observed in the rat pineal gland.
Characterization of Tryptophan Hydroxylase

DRN by 5-HT antiserum revealed a similar distribution of 5-HT immunoreactivity (Fig. 3B). Intense immunolabeling by TPH antiserum was also observed in the rat (Fig. 3D) and mouse PG (data not shown). Detailed immunostaining patterns were illustrated in Fig. 3. As shown in the darkfield micrographs of TPH immunolabeling in Fig. 4, large caliber processes (Fig. 4A) can be seen in the DRN and only limited immunoreactive terminal fields (Fig. 4B) can be observed in the olfactory bulb.

Immunochemical crossreactivity

Antiserum directed against recombinant mouse TPH immunoprecipitated TPH protein from both rat and mouse DRN and recombinant mouse TPH preparations equally well, judging from the nearly identical slopes of the immunotitration curves (Fig. 5). Although a similar slope is observed in the immunotitration curve for TPH from rat PG as compared to other sources, the low TPH activity in homogenates from PG prevents a clear evaluation of the enzyme in this tissue. However, immunoprecipitation of TPH proteins from all four sources by the antiserum occurs in a concentration-dependent fashion. TPH protein in all four cases was almost completely immunoprecipitated by addition of 25 μL of the antiserum as shown in Fig. 5.

Effect of endogenous substances on TPH activity

In order to determine whether endogenous substances in the rat tissues inhibit TPH activity, assays of recombinant
mouse TPH were carried out in the presence of either DRN or PG extracts from rat. As indicated in Table 1, addition of PG extracts to recombinant mouse TPH protein led to no alteration in TPH activity, while DRN extracts produced a small increase of recombinant mouse TPH activity. The data do not indicate that the PG contains a TPH inhibitor.

Effect of end-products on TPH activity

It has been reported that rat TPH is inhibited in vivo by an end-product, 5-HT (7, 8). In the present study using 7.5 × 10^-4 M 6MPH4 as the cofactor and 2.5 × 10^-4 M tryptophan as the substrate, additions of various concentrations of end-products, either 5-HT, N-acetyl-5-hydroxytryptamine or melatonin were tested in our assay conditions for possible inhibition of enzyme activity in vitro. In contrast to the in vivo findings, TPH activity exhibited minimal inhibition in vitro with any of the above products, even when the concentration was as high as 5 × 10^-4 M. The data are not shown here but all the experimental values were within a few percentage of the control value.

DISCUSSION

TPH is highly labile during purification and storage, as a result of its very low tissue concentration (14-16, 20, 30). Because of its instability and low abundance, previous purification methods that involved numerous steps could not successfully produce large amounts of enzymatically active TPH protein from brain tissues (14, 16). Recently, we expressed abundant amounts of enzymatically active recombinant mouse TPH in E. coli, using a bacterial expression vector, pKSTPH, containing the full coding region of mouse TPH. Subsequently, a specific antiserum to the subunits of the recombinant mouse TPH was produced by injecting the TPH band cut from SDS-polyacrylamide slab gels (20). This antiserum detected a single band of predicted molecular weight for TPH from rat DRN, and PG by Western blotting. However, this antiserum immunostained not only TPH-containing neurons but also some nonspecific staining was observed, perhaps because the antiserum was raised to TPH protein containing SDS. In the present study we described a simple procedure to produce large quantities of recombinant mouse TPH purified to an apparent homogeneity by a two-step purification method, Sepharose 4B column chromatography and non-denaturing gel electrophoresis. Subsequently, we produced a specific antibody to recombinant mouse TPH using this purified enzyme protein. To our knowledge, this is the first report of the production and characterization of antibody to TPH using recombinant full-length mouse TPH, which immunohistochemically as well as immunocytochemically recognizes native mouse and rat TPH from the brain as well as the PG. The specificity of this TPH antiserum was determined by three independent methods, Western blot analysis, immunostaining and immunoprecipitation.

In our Western blot analysis, this specific antiserum recognized an identical protein band equivalent to molecular weight of 54,000 Dalton in crude extracts of rat and mouse DRN, rat PG as well as recombinant mouse TPH preparations. This molecular weight corresponds to the one reported previously by others (14-16, 20) for native TPH from rat
brain, mouse mastocytoma, and recombinant mouse TPH. However, an additional lower molecular weight band was also detected only in the crude extract of rat PG and the intensity of the two bands appeared almost equal. The presence of this additional enzyme protein band in the PG extract was not detected using any of the previously reported TPH antisera. Detection of TPH on Western blot from rat and mouse DRN and rat PG extracts required loading of more protein and less dilution of the antiserum, compared with in the case of recombinant mouse TPH preparation. The differences can be attributed to the very low level of TPH protein in the crude tissue extracts as compared with the recombinant TPH preparation.

Earlier reports described the production of antiserum to purified rat TPH but this antiserum also contained anti-catalase antibodies (10, 16) in addition to anti-TPH antibodies. Thus, prior to use this antiserum had to be purified to remove anti-catalase antibodies by column chromatography. In addition, their affinity-purified TPH antibody (10) reportedly recognized tyrosine hydroxylase protein in the PG extracts, but detected only one TPH band from extracts of rat raphe on Western blot. This antiserum was produced against native rat TPH protein which may fold differently from recombinant full-length mouse TPH. Thus, the specific epitope recognizing the lower molecular weight band may be hidden. Recently, the production of anti-peptide antibodies to rat TPH was reported. These antibodies detected one principal (TPH protein band judging from its size) and three additional higher molecular weight bands from extracts of rat raphe areas on immunoblot (17) unlike our TPH antiserum used in this study, i.e., TPH antiserum directed against the subunits of recombinant mouse TPH (20) and the affinity-purified rat TPH antibodies (16). Therefore, it seems that the peptide antiserum contains additional non-defined antibodies that interact with antigens other than TPH protein. They have not tested whether these anti-peptide antibodies to TPH crossreact with the pineal TPH. Unlike the new polyclonal antiserum to recombinant full-length mouse TPH, our previously reported polyclonal antiserum to the subunits of the recombinant mouse TPH (20) also detected only a single predicted molecular weight (54 kDa) band for TPH from rat DRN as well as PG by Western blotting. Therefore, the antiserum to recombinant full-length mouse TPH likely contains an antibody to a different TPH epitope, which perhaps resulted from the differences in protein folding. Thus, the following possibilities must be taken into consideration with respect to this tissue specific presence of the lower molecular weight band on Western blot. Firstly, it seems unlikely that this antiserum contains an antibody against a minor contaminant protein since the crude DRN extract does not show an equivalent lower molecular weight band. Also, the relative abundance of the two bands in the crude PG extract is equal and this antiserum is also immunohistochemically specific as described below.

Secondly, TPH mRNA levels are greater in PG than DRN, while enzyme activity is higher in the DRN. This difference is possibly due to the presence of an endogenous activator or inhibitor in the DRN and the PG. In order to verify this possibility, TPH activity assays of partially purified recombinant mouse TPH preparation were carried out in the presence of a crude extract either from the rat DRN or from the rat PG. The results indicated the absence of either an activator or an inhibitor. Therefore, another explanation must be found for the differences in activity observed in these tissues. The in vivo inhibition of TPH activity by an end product, 5-HT has been reported (7, 8), but addition of up to $5 \times 10^{-4}$ M of end-products, either serotonin, N-acetyl-5-hydroxytryptamine, or melatonin did not affect in vitro TPH activity in our assay conditions. These findings were consistent with the observations by Youdim et al. (9) with 5-HT in vitro. Therefore, the difference in the TPH activity of the DRN and the PG may not originate from end-product inhibition. If the lower molecular weight band from the PG represents another form of TPH, it is likely to be inactive, since only one enzymatically active peak is observed by chromatofocusing column chromatography of the crude PG extract (11, 20). As mentioned above, we are tempted to speculate that the lower molecular weight band, if it represents a less or inactive form of TPH from the PG, may be responsible for the difference in the TPH activity. However, the nature of the additional band and its role in the lowered PG TPH activity remains to be resolved.

The TPH antiserum used in this study intensively immunostained serotonergic neurons located in dorsomedial, ventromedial as well as lateral portions of the rat and mouse DRN and the rat PG. These immunocytochemical distribution patterns of TPH-immunopositive cells in the DRN are in good agreement with the localization of serotonin-immunopositive cells in the same areas as illustrated in Fig. 3A, B, and C. The intensity of serotonin-immunoreactivity in the DRN appears stronger than that for TPH-immunostaining. This difference may be due to the differences in the concentration of 5-HT and TPH protein present in the cells. These results are in good agreement with previous observations made by others, using affinity-purified TPH antibodies (6, 10), anti-peptide TPH antibodies (17), serotonin-antibodies (4, 5, 11) or formaldehyde-induced fluorescence histochemistry (31). Our TPH antiserum also immunostained rat and mouse PG, consistent with previous studies (10), while antibodies to peripheral TPH did not recognize brain TPH (12). The TPH antiserum used in this study also immunostained large caliber processes in the DRN and terminals in the olfactory bulb. The relative ineffectiveness of this TPH antiserum for immunostaining serotonergic processes and terminals may be attributed to a lower concentration of TPH protein compared with the concentration of 5-HT in processes and terminals.
The TPH antiserum described in the present study appears to equally immunoprecipitate TPH protein from all four sources used in a concentration-dependent manner. The immunocrossreactivity between species and tissues was expected from the almost identical amino acid sequences deduced from nucleotide sequences (11, 32-34).

In summary, we have purified large quantities of recombinant mouse TPH by a simple two-step purification procedure and subsequently produced a specific antiserum using this purified TPH. This antiserum recognizes one predicted TPH protein band (54,000 Dalton) on immunoblot of crude extracts from rat and mouse dorsal raphe, and rat PG and also detect an additional lower molecular weight band only from the pineal extract. This lower band may be involved in the pineal low TPH activity despite its higher mRNA levels since there is no evidence for an endogenous inhibitor in PG or for end-product inhibition. Further studies are needed to characterize the nature of the lower molecular weight band observed in the PG extract in view of its low TPH activity. This TPH antiserum specifically immunostained serotonergic neurons of rat and some serotonergic processes in the DRN and terminals in the olfactory bulb as well as immunolabeling rat raphe and some serotonergic processes in the DRN and terminals in the olfactory bulb as well as immunolabeling rat and mouse PG. It also immunoprecipitated TPH protein from the DRN and PG in a concentration-dependent fashion. These data indicate that TPH protein from DRN and PG appears immunologically identical as expected from the identical coding sequence of TPH cDNA from DRN and PG.

ACKNOWLEDGMENTS

This work was supported by NIH Grant MH 44043. We would like to express our gratitude to Ms. Nan Min for her technical assistance in tryptophan hydroxylase immunostaining and Mr. Charles Carver for his outstanding artwork in preparing the figures and the photographs.

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