Cloning, In Vitro Expression, and Tissue Distribution of a Human Prostaglandin Transporter cDNA (hPGT)

Run Lu, Naoaki Kanai, Yi Bao, and Victor L. Schuster

Departments of Medicine, Physiology, and Biophysics, Albert Einstein College of Medicine, Bronx, New York 10461

Abstract

We recently identified a cDNA in the rat that encodes a broadly expressed PG transporter (PGT). Because PGs play diverse and important roles in human health and disease, we cloned human PGT (hPGT) from an adult human kidney cDNA library. A consensus sequence (4.0 kb) derived from several clones, plus 3' polymerase chain reaction amplification, exhibited 74% nucleic acid identity and 82% amino acid identity compared to rat PGT. When transiently expressed in HeLa cells, a full-length clone catalyzed the transport of PGE₁, PGE₂, PGD₂, PGF_{2α} and, to a lesser degree TXB2. Northern blotting revealed mRNA transcripts of many different sizes in adult human heart, placenta, brain, lung, liver, skeletal muscle, pancreas, kidney, spleen, prostate, ovary, small intestine, and colon. hPGT mRNAs are also strongly expressed in human fetal brain, lung, liver, and kidney. The broad tissue distribution and substrate profile of hPGT suggest a role in the transport and/or metabolic clearance of PGs in diverse human tissues. (J. Clin. Invest. 1996. 98:1142-1149.) Key words: prostaglandins • carrier proteins • biological transport • molecular cloning

Introduction

PGs and TXs play ubiquitous and vital pathophysiological and therapeutic roles in human health and disease states, such as glaucoma: pregnancy, labor, delivery; and abortion: gastric protection and peptic ulcer formation; intestinal fluid secretion; liver protection and damage; airway resistance and asthma; blood pressure control; and modulation of inflammatory cells (1–15).

As charged organic anions at physiological pH (16, 17), PGs traverse biological membranes poorly (18). Accordingly, PG transport is carrier-mediated in many diverse tissues, including the lung (19-22), choroid plexus (23-28), liver (29), anterior chamber of the eye (23-25, 30), vagina and uterus (31-33), and placenta (34, 35). Indeed, our laboratory recently identified rat prostaglandin transporter (PGT), the first cloned carrier known to catalyze the transport of PGE and PGF_{2 α} (36).

Address correspondence to Victor L. Schuster, Renal Division, Ullmann 617, 1300 Morris Park Avenue, Bronx, NY 10461. Phone: 718-430-3158; FAX: 718-824-2765; E-mail: schuster@aecom.yu.edu

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1. Abbreviations used in this paper: hPGT, human PGT; oatp, organic anion transporting peptide; PGT, prostaglandin transporter.

mic compartment (37), it is unclear whether a carrier is necessary for their subsequent efflux into the extracytoplasmic space. Second, PGT might mediate epithelial PG transport. Vectorial PG transport occurs in many epithelia, including the

cells. Although PGs apparently do gain access to the cytoplas-

We have postulated three possible roles for PGT (36). First, PGT might mediate the efflux of newly-synthesized PGs from

liver, kidney, choroid plexus, anterior uvea, and uterus (22–24, 27, 33, 38, 39). The tissue distribution of PGT mRNA expression in the rat (36) suggested that its expression might be limited to epithelia, and therefore that epithelial transport of PGs may represent its primary function.

A third possible role of PGT is that of mediating PG clearance and degradation. After their release from cells, PGs and TXs bind to nearby surface receptors, where they signal a broad array of physiological functions. Because there is no enzymatic activity in the plasma capable of oxidizing PGE₁, PGE₂, or PGF₂ (38), these prostanoids could potentially bind to receptors at a substantial distance from their sight of release if they were not metabolized locally. Local clearance of PGE₁, PGE_2 , and $PGF_{2\alpha}$ occurs in a single passage through any of several vascular beds, such as the lung (38, 40, 41). In contrast, prostacyclin (PGI₂) is not cleared, and thus circulates as a hormone (42, 43). The differing metabolism of these prostanoids appears to be due to differences in the selective uptake of some PGs into cells.

Of these three possibilities, we currently favor the clearance role for PGT, because (a) the substrate specificity of the cloned transporter is very similar to that reported previously for PG clearance in the isolated perfused rat lung preparation (20–22, 36); and (b) we have been unable to obtain PGT-mediated PG efflux using the Xenopus oocyte expression system (Chan, B., and V.L. Schuster, unpublished observations).

Using a rat PGT probe on Northern blots of human kidney RNA, we found evidence for the presence of a likely human PGT homologue. Accordingly, we screened a human kidney cDNA library and expressed a full-length human cDNA clone in cultured cells. We now report that both rat PGT and human PGT (hPGT) transport PGD₂, as well as PGE₁, PGE₂, and PGF_{2a}. Although hPGT has cDNA and deduced amino acid sequences similar to those of the rat, the tissue distribution of the mRNA transcripts is substantially broader, the diversity of transcripts is greater, and the affinity for TXB₂ is greater. In addition, we find strong hPGT mRNA expression in the human fetus. The findings are important as a first step in understanding the transport of endogenous and synthetic PGs in humans.

Methods

Human kidney cDNA library screening. A DNA fragment, obtained from the rat PGT cDNA by cutting with the restriction endonuclease Hinc1 followed by gel purification, was labeled by the random primer method (44) with $[\alpha^{-32}P]dCTP$, and was used to probe a human kidney cDNA library in the phage vector $\lambda gt10$ ($\sim 10^8$ pfu/ml; Clontech,

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Palo Alto, CA), generated by mixed deoxythymidine and random hexamer oligonucleotide primers. After library plating and replicatransfer to Nytran Plus filters (Schleicher & Schuell, Keene, NH), filters were hybridized overnight at 40°C with the $^{32}\text{P-labeled}$ Hinc1 fragment in $5 \times \text{SSC}$, 2° blocking reagent, 0.1° *N*-laurylsarcosine, and 0.02° SDS (Genius System, Boehringer Mannheim Biochemicals, Indianapolis, IN). Filters were washed as follows: twice with $1 \times \text{SSC}$, 0.1° SDS at 40°C ; twice with $0.5 \times \text{SSC}$, 0.1° SDS at 40°C ; twice with $0.1 \times \text{SSC}$, 0.1° SDS at 40°C ; and twice with $1 \times \text{SSC}$, 0.1° SDS at 50°C . Filters were exposed overnight to X-Omat AR film at -70°C (Eastman Kodak Co., Rochester, NY). Positive plaques on duplicate filters were picked for secondary screening using the same hybridization and washing conditions. Of 800,000 colonies screened, seven positive clones were isolated and subcloned into the plasmid pSPORT-1 (Gibco BRL, Gaithersburg, MD).

DNA sequencing, restriction analysis, and computer analysis. All seven positive clones were subjected to restriction mapping. Using the dideoxy chain termination method (Sequenase version 2.0 DNA sequencing kit; U.S. Biochemical Corp., Cleveland, OH), all of both strands of clone 3 and parts of the other clones were sequenced by primer walking. In addition, parts of clones 5, 7, 9, and 10 were also sequenced. The data were aligned and analyzed by MacVector (Eastman Kodak Co., Rochester, N.Y.) and GeneWorks (Intelligenetics, Inc., Campbell, CA) software programs.

3' rapid amplification of cDNA ends (RACE) by PCR. First strand cDNA was reverse-transcribed from human kidney poly-A (+) RNA (0.5 µg; Clontech) using a Not1 primer-adapter (SuperScript Plasmid System, Gibco-BRL). The initial gene-specific 18-mer primer, generated from the region 3260-3277 of clone 3, and the Not1 primeradapter were used in a first round of PCR. The initial denaturation at 94° for 1.5 min was followed by 20 cycles of denaturation at 94° for 30 s, annealing at 50° for 30 s, and extension at 72° for 1.5 min. A final extension at 72°C was performed for 10 min. 1 µl of this product mixture was diluted into a second 50-µl reaction volume and nested PCR was performed with a second gene-specific 18-mer primer (region 3434-3451 of clone 3) and a 3' anchor primer (5'-TAGTTCTA-GATCGCGAGCGG-3') generated from the Not1 primer adapter. This second PCR was similar to the first except that the annealing temperature was 65°C and the number of cycles was 35. The PCR product, which migrated at \sim 600 bp, was subcloned into the TA cloning vector (Stratagene, La Jolla, CA) and sequenced as above from the 3' end.

Transient expression in HeLa cells and transport assays. The full-length clone 3 cDNA, subcloned in pSPORT1 with coding strand downstream of the T7 promoter, was transfected into cultured mammalian cells as follows. HeLa cells grown on 35-mm dishes were infected for 30 min with recombinant vaccinia vTF7-3 (45), the medium was changed, and the cells were then transfected by adding premixed cDNA (10 μ g) and Lipofectin (20 μ g, Gibco-BRL), after which there was an additional medium change at 3 h. The plasmid pBluescript KS was used as a negative control.

After 20 h, cells were washed, ³[H]-PG was added in Waymouth's solution, and timed uptake at 27°C was determined. Uptake was stopped by one cold wash using Waymouth's with 5% BSA and three cold washes with Waymouth's only. Cells were scraped and counted by liquid scintillation. On Fig. 2, data are the mean±SEM of two dishes from each of two separate monolayer transfections.

For determining PG uptakes (see Fig. 2), the following 3 [H]-PGs and final concentrations were used (New England Nuclear, Boston, MA): PGE₂: 0.7 nM (176 cpm/fmol); PGE₁: 0.6 nM (62 cpm/fmol); PGD₂: 0.9 nM (126 cpm/fmol); PGF_{2 α}: 0.6 nM (185 cpm/fmol); and TXB₂: 1.0 nM (114 cpm/fmol). In addition, the PGI₂ analog 3 [H]-iloprost was used (Amersham Corp., Arlington Heights, IL) at 7.9 nM (14 cpm/fmol).

Inhibition of tracer PGE₂ uptake. Tracer PGE₂ uptakes at 10 min (0.2 nM ³[H]-PGE₂) with or without various concentrations of unlabeled prostanoids (100–500 nM; Cayman Chemical, Ann Arbor, MI) or inhibitors (10–100 μM, Sigma Chemical Co., St. Louis, MO)

were determined in duplicate on a given transfection for one or two separate transfections. Because the substrate concentration was at least 500 times less than the concentration of the unlabeled prostanoid, we defined an apparent affinity constant, $K_{1/2}$, from the equation:

$$K_{1/2} = [v_i/(v-v_i)][i]$$

where v = uptake without inhibitor, $v_i = uptake$ with inhibitor, and [i] = inhibitor concentration (46).

Northern blot hybridization. Based on preliminary data indicating that a rat probe lacking much of the 3' untranslated region gave lower backgrounds, the rat PGT cDNA was shortened by restriction digesting it with BamH1 and religating the remaining fragment, such that the resulting cDNA terminates at nucleotide 2055. An antisense digoxigenin-labeled RNA probe was generated from this clone using SP6 polymerase (Boehringer Mannheim Biochemicals) after linearizing with Sal1. A human PGT digoxigenin-labeled antisense probe was similarly generated from EcoR1-linearized clone 3 that had been previously shortened and religated at the 3' end using Acc1. As a house-keeping probe, human β -actin cDNA (Prime-It II, Stratagene) was labeled by the random primer method with $[\alpha^{-32}P]$ dCTP(3,000 Ci/mmol; New England Nuclear).

Rat whole kidney total RNA, prepared by the acid guanidinium method (47), and 1 μ g of human kidney poly-A+ RNA (Clontech, Palo Alto, CA) were separated by glyoxal denaturing agarose electrophoresis (44) and transferred to Hybond-N nylon membranes (Amersham Corp.). Separately, four multiple human tissue Northern blots were purchased from Clontech.

Blots probed with either the rat PGT or the clone 3 hPGT riboprobes were hybridized overnight at 65°C in 5 \times SSC, 50% formamide, 2% blocking reagent, 0.1% N-laurylsarcosine, 0.02% SDS, and 0.01 M EDTA (Genius System; Boehringer Mannheim Biochemicals). This was followed by sequential washes at 65°C: twice in 1 \times SSC, 0.1% SDS, 0.01 M EDTA; twice in 0.5 \times SSC, 0.1% SDS, 0.01 M EDTA; and twice in 0.1 \times SSC, 0.1% SDS, 0.01 M EDTA. For the β -actin probe of the same membranes, hybridization was in 5 \times SSC, 2% blocking reagent, 0.1% N-laurylsarcosine, 0.02% SDS, and 0.01 M EDTA overnight at 65°C. These blots were washed at 65°C, twice in 1 \times SSC, 0.1% SDS, 0.01 M EDTA; twice in 0.5 \times SSC, 0.1% SDS, 0.01 M EDTA; and twice in 0.1 \times SSC, 0.1% SDS, 0.01 M EDTA.

Detection was performed using a horseradish peroxidase–coupled antidigoxigenin antibody (Fab fragment) (Boehringer Mannheim Biochemicals). Signals were visualized by chemiluminescence autoradiography (ECL, Amersham Corp.).

Results

To search for human homologues of rat PGT, we first examined adult human kidney RNA by Northern blotting it with a rat PGT antisense RNA probe. Fig. 1 shows that the rat probe revealed two bands of ~ 4.0 and 4.7 kb in total rat kidney RNA, as previously reported (36). When applied to human adult kidney poly A+ RNA at the same high stringency, the rat probe hybridized strongly to bands of 2.3 and 4.2 kb, and weakly to bands of 6.0 and 7.0 kb (Fig. 1). These results suggested that there was a human PGT homologue, with perhaps several PGT transcripts expressed in human kidney.

To understand the human homologue further, we screened an adult human kidney cDNA library with a ³²P-labeled rat PGT DNA probe. Seven hybridizing clones were isolated. After preliminary DNA sequencing and restriction mapping, six of the seven clones appeared to be related to each other and to rat PGT, and so were subjected to a more detailed analysis.

The longest clone isolated (No. 3) was 3.5 kb in length. Preliminary sequencing of this clone at the 5' end revealed simi-

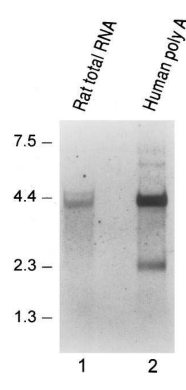


Figure 1. Northern blot hybridization of rat kidney total RNA (35 μ g) and human kidney poly A+ RNA (1 μ g) using a rat PGT antisense RNA probe. Hybridization and washing conditions as per Methods.

larity to rat PGT, including an ATG codon in good context for translation initiation (48). Before proceeding with sequencing, we tested the function of clone 3 using the vaccinia-T7 system (36, 45), on the assumption that mediation of PG transport would indicate that the clone did not contain splice artifacts causing frame shifts or premature stop codons.

Fig. 2 shows that clone 3 catalyzed the rapid and time-dependent uptake of several PGs. The 2-min rank-order of uptakes is $PGE_2 > PGE_1 > PGF_{2\alpha} > PGD_2 > TXB_2$. By 10 and 20 min, there is essentially no difference between the transport

Table I. Inhibition of hPGT-mediated Tracer PGE₂ Transport

Inhibitor*	10 μΜ	100 μΜ
Furosemide	91.5±3.3	63.3±1.7
Probenecid	104 ± 5.2	92.8 ± 2.0
Indomethacin	114 ± 3.8	82.5 ± 1.0
Unlabeled Prostanoid‡	100 nM	500 nM
PGE ₁	50.8±3.4	
PGE_2	45.2 ± 1.4	_
PGD_2	47.9 ± 0.9	_
PGF_{2a}	45.0 ± 1.3	_
TXB_2	_	26.7 ± 0.1

Uptakes were determined \pm various inhibitors or unlabeled prostanoids. Results are shown as the percentage of the control 10-min tracer PGE₂ uptake. *Mean \pm SEM from four separate determinations. \pm Mean \pm SEM from two separate determinations.

rates of PGE_2 , PGE_1 , $PGF_{2\alpha}$, or PGD_2 . In contrast, the prostacyclin analogue iloprost was not transported above baseline (data not shown). With the exception of the information on the transport of PGD_2 , which is new to the present study, this rank order is very similar to that of rat PGT, as we recently reported (36). Because of the new finding here of PGD_2 transport by hPGT, we reexpressed rat PGT in HeLa cells, and found that rat PGT also transports tracer PGD_2 at a rate equivalent to that of PGE_2 (data not shown). Taken together, the data of Fig. 2 indicate that clone 3 very likely represents a human homologue of rat PGT.

Table I presents results using three known PG transport inhibitors. Although furosemide was a moderately potent inhibitor of hPGT, probenecid and indomethacin were much less effective. Table I also shows the inhibition of tracer PGE₂ by an excess of unlabeled prostanoids, assessed using clone 3. The apparent affinity constants ($K_{1/2}$, in nM) were: PGE₂ = 100, PGD₂ = 83, PGF_{2 α} = 92, PGE₁ = 82, and TXB₂ = 182. (It

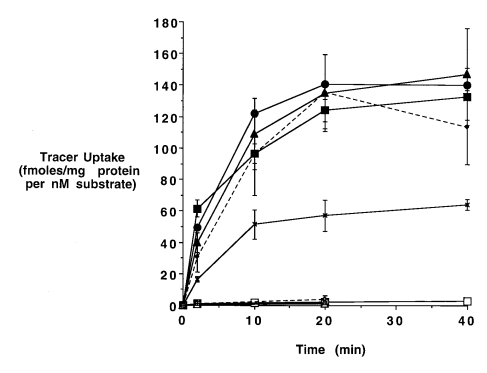


Figure 2. Time-dependent uptake of tracer prostaglandins into HeLa cells expressing clone 3. *PGT*, vector contained clone 3 cDNA; *pBS*, vector was pBluescript control without any PGT sequence.

- -■-, PGE2, PGT; -□-, PGE2, pBS;

- \longrightarrow TxB2, PGT; \longrightarrow , TxB2, pBS.

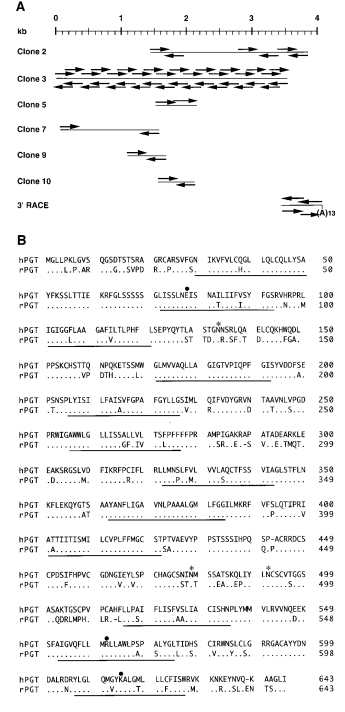


Figure 3. (A) Sequencing map of hPGT clones. Arrows indicate direction and length of dideoxy DNA sequencing runs. (B) Deduced amino acid sequence of hPGT compared to that of rat PGT. Dots in the hPGT sequence represent identity to rat PGT. (Underlines) putative membrane-spanning domains. (Black circles) charged residues within putative membrane spans that are conserved between hPGT, rat PGT, and the related transporter oatp (50). *Potential glycosylation sites.

should be noted that, because tracer uptake in the presence of unlabeled PGs was determined at 10 min and does not represent an initial value, $K_{1/2}$ approximates, but does not equal, the Michaelis–Menten constant K_m).

Fig. 3 A shows the sequencing strategy and alignments of

the six cDNA clones examined from the human kidney cDNA library. Clones No. 2 and 3 overlap and cover a total distance of 3.8 kb (Fig. 3 A). The 3' end of clone 3 was used to generate two gene-specific primers for a 3' RACE strategy to clone the remaining 3' untranslated region. As described in Methods, and as illustrated in Fig. 3 A, a PCR product of 612 bp was amplified, cloned, and sequenced. This yielded 400 bp that overlap with the 3' end of clone 2, as well as 212 new nucleotides that terminate in a poly-A tail. The total length of the cDNA sequence assembled from clone 2, clone 3, and the 3' RACE sequence comprised 4,046 bp, which is in reasonable agreement with the size of the major transcript in most tissues (~ 4.2 kb, Figs. 4–6 below).

The consensus hPGT cDNA sequence, derived from all five clones, was 74% identical to that of the available comparable rat PGT cDNA sequence. There was also strong homology to several expressed sequence tags in the human database (Genbank accession numbers H63820, H63772, T85296, R25350, T85507, and R26541).

Fig. 3 *B* compares the deduced amino acid sequences of hPGT and rat PGT starting at comparable ATG translation initiation codons (position 88 in the rat and 92 in the human). The predicted rat and human proteins are highly similar (82% identity). hPGT is predicted to have 12 membrane-spanning domains based on the Kyte–Doolittle hydropathy algorithm using a window of 13 residues (49). Glycosylation of hPGT could occur at any or all of three extracytoplasmic asparagine consensus sites (Fig. 3 *B*).

In addition to its similarity to rat PGT, the hPGT deduced protein sequence is 32% identical to human organic anion transporting polypeptide (oatp). Like PGT, oatp has been shown to be an organic anion transporter, primarily of bile salts and conjugated steroids (50, 51). It is therefore of interest that three charged residues in hPGT (E77, R561, and K614, see Fig. 3 *B*), which lie within putative membrane spans, are highly conserved between hPGT, rat PGT, and oatp (50), suggesting that these amino acids may play an important role in anion transport.

We examined the tissue distribution of hPGT expression in adult human tissues by Northern blot analysis of poly A+RNA (Fig. 4, A and B). A diversity of hybridizing transcripts was observed. The most strongly hybridizing RNA bands were 1.8–2.0 kb in skeletal muscle, prostate, testis, ovary, small intestine, and colon; 2.5–2.9 kb in heart and skeletal muscle; 4.0 kb in ovary; 4.4–5.1 kb (often a doublet) in heart, whole brain, placenta, lung, liver, skeletal muscle, kidney, spleen, thymus, prostate, testis, ovary, small intestine, colon, and peripheral blood leukocytes; 8.8 kb in testis and colon; and 10.1 kb in testis. (The RNA derived from pancreas appeared to undergo some degradation, so statements about the exact PGT message in that organ remain limited).

Further examination of specific regions of the human brain by Northern blotting revealed major transcripts of 2.1, 3.5, 4.8, and 7.5 kb in most regions, with an additional 10-kb band seen in the caudate nucleus (Fig. 5). Although it is possible that the 2.1-kb bands represents the riboprobe probe binding to 18S RNA, this seems unlikely because this blot utilized poly-A+RNA, not total RNA, and because we do not see carry-over of ribosomal RNA in the other blots. Moreover, the presence of strong bands of similar size in fetal lung, and in adult prostate, testis, ovary, small intestine, colon, and skeletal muscle argue that the 2.1-kb bands in brain derive from mRNA.

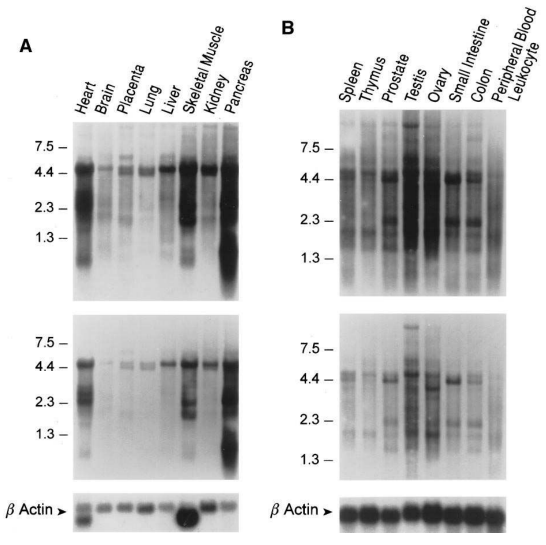


Figure 4. (A and B) Northern blot hybridization of human poly A+ RNA using a human PGT clone 3 antisense RNA probe. Hybridization and washing conditions as per Methods. Top panels represent exposure overnight, middle panels represent exposure for 15 min.

In human fetal tissues (Fig. 6), the RNA transcripts seen by Northern blot hybridization were 2.1 kb in brain, lung, liver, and kidney; 2.5 kb in lung; 4.8–5.1 kb in brain, lung, liver, and kidney; 6.1 kb in liver; and 10 kb in lung and liver.

Discussion

We have cloned, sequenced, and heterologously-expressed a human homologue (hPGT) of PGT, a newly-recognized prostaglandin transporter that we previously identified in the rat (36). The hPGT nucleic acid and deduced amino acid sequences are similar to that of the rat transporter. In vitro expression of a full-length hPGT cDNA causes the transport of PGE₁, PGE₂, PGD₂, PGF_{2 α}, and to a lesser extent TXB₂. The mRNA expression for hPGT in native human tissues is very broad. Indeed, hPGT mRNA was seen in essentially every tissue or cell type examined, and there is a rich diversity of mRNA transcripts ranging from 1.8 to over 10 kb in length.

Although there is a substantial literature on carrier-mediated PG transport in other species (19–36), there is surprisingly little information about this phenomenon in humans,

perhaps because of difficulty in obtaining tissue for such studies. PGE_2 has been shown to be transported by the isolated perfused human placental cotyledon (52) and by the isolated human amnion (35). Despite this paucity of direct information, the literature in animals, taken with the present functional profile and broad tissue expression pattern of hPGT, make it very likely that extensive carrier-mediated PG transport occurs in diverse human tissues.

As discussed in the Introduction, it appears that PGT most likely plays a role in the uptake of newly released PGs before intracellular oxidation. In this regard, the affinities of hPGT for its substrates are of interest. Despite the observation that hPGT and rat PGT both transport TXB₂ at a rate about half that of PGE₂ or PGF_{2 α} (36) (Fig. 2), the $K_{1/2}$ of hPGT for this substrate is half that of the rat affinity constant (182 vs 423 nM, respectively, Table I and [36]), suggesting that PGT may play a more prominent role in thromboxane transport in the human than in the rat. Further studies will be required to address this possibility directly.

The apparent affinities $(K_{1/2})$ of hPGT for PGE₂ and PGF_{2 α} are in the range of 80–100 nM. PG concentrations as presented

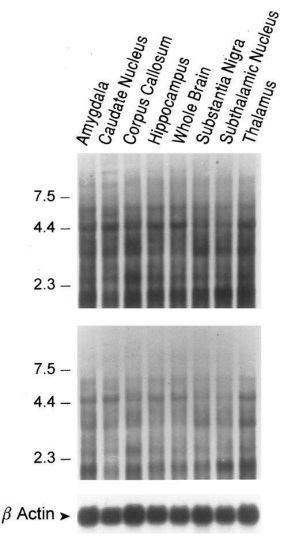


Figure 5. Northern blot hybridization of human brain poly A+ RNA using a human PGT clone 3 antisense RNA probe. Hybridization and washing conditions are as per Fig. 4. Top panels represent exposure for 1 h, middle panels represent exposure for 5 min.

to the transporter in tissues are probably at least this high. In human lung, for example, the PGE $_2$ concentration is about 25 ng/gram tissue, or ~ 70 nM; in human semen this value is ~ 25 µg/ml, or ~ 70 µM (53). On the other hand, plasma PGE and PGF concentrations in humans are ~ 0.25 ng/ml, or 0.7 nM (54). Assuming that hPGT is the mechanism for reuptake of PGs after their release, this plasma concentration would represent the residual from such a process.

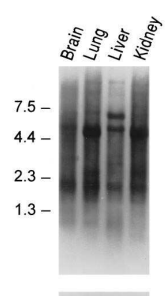
Many transporters exist in families whose individual members recognize structurally related, but distinct, substrates. It is therefore possible that transporters for other prostanoids exist. Indeed, PGT appears to be part of a gene family, the other currently recognized member of which is oatp (36). Work is currently underway to identify and characterize such PGT-related cDNAs.

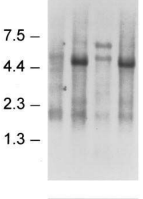
The presence of many different mRNA transcripts on high stringency Northern blots suggests that several functionally distinct mRNAs may arise from a single hPGT gene. Such diverse hPGT transcripts could arise by alternative splicing and/ or alternative promoters. At present, we have no definitive in-

formation with which to distinguish between these possibilities. However, we have recently isolated human genomic PGT clones (Lu, R., and V.L. Schuster, unpublished observations), a step that should aid in addressing the origin of the diverse transcripts.

We also have no information at present as to whether transcripts that are either substantially larger or smaller than the \sim 4.0 kb cDNA encode functional transporters. It is of interest, however, that in neither the rat nor the human cDNA is the first methionine codon preceded by a stop codon (36). This raises the interesting possibility that the larger mRNA transcripts (e.g., the strong \sim 10-kb band in testis) might encode proteins that are extended at the amino-terminus relative to the present hPGT consensus cDNA. Confirmation of this hypothesis, as well as functional evaluation of any additional encoded proteins, must await isolation and characterization of additional cDNAs.

The presence of strongly hybridizing hPGT mRNA transcripts in human heart and skeletal muscle stands in contrast to our findings in the rat, where there was no detectable expression, at least by Northern blot analysis, in these two tissues (36). At present, we have no clear explanation for this difference other than species. The unequivocal expression of PGT mRNA in human heart and skeletal muscle argues against our





 β Actin >

Figure 6. Northern blot hybridization of human fetal poly A+RNA using a human PGT clone 3 antisense RNA probe. Hybridization and washing conditions, and film exposures are as per Fig. 4.

previous conclusion that PGT is restricted to tissues containing epithelia (36). Because human hearts synthesize PGE₂, PGF_{2 α}, and 6-keto-PGF_{1 α}, especially in response to cold pressor testing (55–58), it is possible that PGT plays an as yet unidentified role in cardiac prostanoid metabolism.

The novel observation in the present paper of PGD₂ transport by both rat and human PGT is of considerable physiological interest. PGD₂ has been implicated in several processes in the central nervous system (CNS), including the control of body temperature (59) and of olfactory function, hormone release, and pain perception (60). In addition, both PGE₂ and PGD₂ have been postulated to mediate central sleep-wake cycles (59, 61, 62). PGE₂ receptors are very widely expressed in diverse structures of the rat brain (63). The equally broad expression of hPGT mRNA in the human brain (Fig. 6), and the ability of cloned hPGT to transport PGE2 and PGD2 approximately equally well (Fig. 2), suggest that the transporter may play a role in the release, epithelial transport, and/or degradation of these prostanoids in the CNS. The present findings provide probes that should be useful in further studies on the CNS function of these eicosanoids.

There are other possible clinical implications of the present work. First, circulating PG levels are elevated in several clinical circumstances, including bone marrow transplantation (64) and the hyperprostaglandin syndrome. The latter is a Bartter syndrome–like complex tubular disorder found in premature infants, characterized by increased systemic PGE $_2$ activity, fever, diarrhea, and osteopenia with hypercalciuria (65, 66). One hypothesis to explain these conditions would be a failure to clear circulating PGE $_2$ secondary to abnormal regulation or functional impairment of hPGT.

Second, various synthetic PGE₁ or PGE₂ analogs have been used to treat glaucoma (10, 11, 67) and impotence (68), to terminate pregnancy (12), and to provide gastric protection (13–15). The clinical use of these agents is predicated on certain assumptions about their pharmacokinetics, yet with few exceptions (69) detailed information about the clearance and metabolism of these agents in humans is lacking. Only recently, for example, was exogenous PGE₁ shown to be rapidly converted to circulating 13,14-dihydro-PGE₁ and 15-keto-PGE₁ in humans (70, 71). The role of hPGT, if any, in the clearance from the circulation of medicinal PGs can now be approached experimentally by expressing the transporter in vitro and evaluating its interactions with these agents. Moreover, in theory, at least, it should be possible to use the cDNA to design synthetic PGs that are not transported by hPGT, and thus might be cleared more slowly than those that are transported.

In summary, we have cloned, sequenced and heterologously expressed a novel human PG transporter cDNA (hPGT), and have characterized the expression of hPGT mRNA in a large number of human tissues. The substrate specificity and broad tissue distribution suggest a role in the clearance and metabolism of several important endogenous prostanoids.

Acknowledgments

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