Identification of a Carbonic Anhydrase-Like Domain in the Extracellular Region of RPTPγ Defines a New Subfamily of Receptor Tyrosine Phosphatases

G. BARNEA,¹ O. SILVENNOINEN,¹ B. SHAANAN,² A. M. HONEGGER,¹ P. D. CANOLL,¹ P. D'EUSTACHIO,³ B. MORSE,⁴ J. B. LEVY,¹ S. LAFORGIA,⁵ K. HUEBNER,⁵ J. M. MUSACCHIO,¹ J. SAP,¹ AND J. SCHLESSINGER^{1*}

Departments of Pharmacology¹ and Biochemistry,³ New York University Medical Center, 550 First Avenue, New York, New York 10016; Department of Biological Chemistry, The Life Sciences Institute, The Hebrew University of Jerusalem, Givat Ram, Jerusalem 91904, Israel²; Rhone-Poulenc Rorer, Collegeville, Pennsylvania 19426⁴; and Jefferson Cancer Institute, Thomas Jefferson Medical College, Philadelphia, Pennsylvania 19107⁵

Received 2 October 1992/Accepted 14 December 1992

The tyrosine phosphatase RPTP γ is a candidate tumor suppressor gene since it is located on human chromosome 3,14.2-,21 in a region frequently deleted in certain types of renal and lung carcinomas. In order to evaluate its oncogenic potential and to explore its normal in vivo functions, we have isolated cDNAs and deduced the complete sequences of both human and murine RPTPy. The murine RPTPy gene has been localized to chromosome 14 to a region syntenic to the location of the human gene. Northern (RNA) blot analysis reveals the presence of two major transcripts of 5.5 and 8.5 kb in a variety of murine tissues. In situ hybridization analysis reveals that RPTPy mRNA is expressed in specific regions of the brain and that the localization of RPTPy changes during brain development. RPTPy is composed of a putative extracellular domain, a single transmembrane domain, and a cytoplasmic portion with two tandem catalytic tyrosine phosphatase domains. The extracellular domain contains a stretch of 266 amino acids with striking homology to the zinc-containing enzyme carbonic anhydrase (CAH), indicating that RPTPy and RPTPB (HPTPY) represent a subfamily of receptor tyrosine phosphatases. We have constructed a model for the CAH-like domain of RPTP_y based upon the crystal structure of CAH. It appears that 11 of the 19 residues that form the active site of CAH are conserved in RPTPy. Yet only one of the three His residues that ligate the zinc atom and are required for catalytic activity is conserved. On the basis of this model we propose that the CAH-like domain of RPTPy may have a function other than catalysis of hydration of metabolic CO₂.

It is now well established that the phosphorylation of tyrosine residues in cellular proteins plays an important role in the control of cell growth and differentiation (reviewed in references 5, 15, and 36). Tyrosine phosphorylation is a reversible process; protein tyrosine kinases are responsible for the incorporation of phosphate on tyrosine residues of cellular proteins, while protein tyrosine phosphatases are responsible for removal of the tyrosine-bound phosphate groups (reviewed in reference 10). Subversion of normal growth control pathways leading to oncogenesis has been shown to be caused by activation or overexpression of tyrosine kinases, which constitute a large group of dominant oncogenic proteins (reviewed in reference 15). Consistent with this view is the notion that the underexpression or inactivation of protein tyrosine phosphatases potentially results in oncogenesis. For this reason, tyrosine-specific phosphatase genes are candidate recessive oncogenes or tumor suppressor genes (21). Similarly to protein tyrosine kinases, protein tyrosine phosphatases can be divided into two classes: low-molecular-weight cytoplasmic enzymes and membrane-linked enzymes. The membrane-linked enzymes have all the hallmarks of cell surface receptors; they contain a putative extracellular domain, a single transmembrane region, and a cytoplasmic domain which usually contains

Recent studies indicate that the action of tyrosine phosphatases is not necessarily only suppressive. It was shown that members of the Src family of cytoplasmic tyrosine kinases such as $pp56^{lck}$ contain inhibitory tyrosine phosphorylation sites in their carboxy-terminal tails (reviewed in reference 14). The phosphorylation of this site by a specific tyrosine kinase, denoted CSK, leads to inhibition of tyrosine kinase activity (24). It was proposed that in T cells the dephosphorylation of such an inhibitory site in $pp56^{lck}$ by the tyrosine phosphatase CD45 leads to enhanced tyrosine phosphorylation (22). Hence, tyrosine phosphatases may function as either switch-on or switch-off signaling enzymes.

The RPTP γ gene is located in human chromosome 3_p14.2-_p21, in a region found to be deleted in certain types of renal and lung carcinomas. It was therefore proposed that RPTP γ may function as a tumor suppressor gene in these cancers (21). A partial cDNA clone of the catalytic domain of RPTP γ /HPTP γ has been previously described (17, 20). In this report we describe the cDNA cloning and the complete amino acid sequence of human RPTP γ and its murine homolog. We show that RPTP γ contains a single transmembrane domain and two tandem tyrosine phosphatase domains and that the extracellular domain of RPTP γ contains a region of 266 amino acids with striking sequence similarity to the enzyme carbonic anhydrase (CAH). We also show that

two tandem catalytic domains (reviewed in references 10 and 29).

^{*} Corresponding author.

TABLE 1. DNA fragment length variant associated with the murine RPTP γ ge	gth variant associated with the murine RPTPy gene ^a
--	--

Allele	Size (kb)	Strains
a	2.7	DBA/2J; C3H/HeJ; AKXD-2, -3, -6, -8, -11, -12, -15, -20, -22, -23, -26, -27; BXD-5, -8, -9, -14, -15, -16, -18, -21, -22, -24, -29, -31, -32; BXH-2, -3, -4, -9, -12, -14, -19
b	2.0	C57BL/6J; 020/A; AKR/J; C57L/J; SWR/J; SJL/J; BALB/cJ; STS/A; AKXD-1, -7, -9, -10, -13, -14, -16, -18, -21, -24, -28; BXD-1, -2, -6, -11, -12, -13, -19, -20, -23, -25, -27, -28, -30; BXH-6, -7, -8, -10, -11

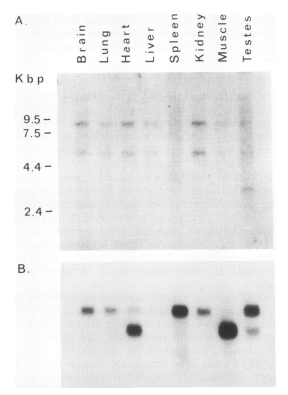
^a Liver or spleen genomic DNA (10 μg) was digested with TaqI enzyme and analyzed by Southern blotting with a probe from the first catalytic domain of murine RPTPγ.

RPTP γ is expressed in specific regions of the rat brain and that its expression is developmentally regulated.

MATERIALS AND METHODS

Isolation of cDNA clones. (i) Human RPTP γ . The initial human RPTP γ gene clone was isolated from a λ gt11 cDNA library of 1-day-old human brain stem (obtained from the American Type Culture Collection; ATCC 37432) by screening approximately 300,000 plaques with a nick-translated leukocyte common antigen probe that spanned both conserved phosphatase domains, under relaxed hybridization conditions. The rest of the cDNA clones were isolated by screening the human brain stem library with randomly primed probes from the previously isolated clones of the RPTP γ gene under high-stringency hybridization conditions.

(ii) Mouse RPTP γ . The initial mouse RPTP γ gene clone was isolated from a λ gt11 mouse brain cDNA library (pur-



chased from Clontech, Palo Alto, Calif.) by screening approximately 10^6 plaques with a randomly primed probe from human RPTP γ that spanned all of the first catalytic domain, under high-stringency hybridization conditions. The rest of the cDNA clones were isolated by screening the mouse brain library with probes from the previously isolated clones of the mouse RPTP γ gene under high-stringency hybridization conditions.

Nucleotide sequence determination. DNA preparations of $\lambda gt11$ RPTP γ clones were digested with *Eco*RI and subcloned into Bluescript SK+ plasmid (purchased from Strat-

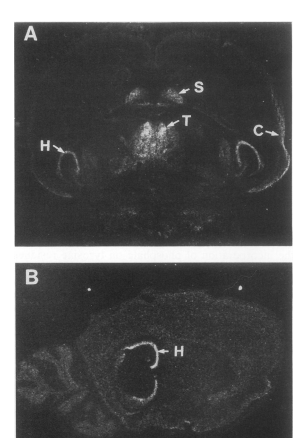


FIG. 1. Expression of RPTP_{γ} in different murine tissues. (A) Northern blot analysis of poly(A)⁺ RNA from different mouse tissues probed with an RPTP_{γ} probe encompassing the first catalytic domain, the juxtamembrane domain, the transmembrane domain, and the beginning of the extracellular domain. (B) The same blot probed with a β -actin probe.

FIG. 2. In situ hybridization analysis of RPTP γ in the newborn and adult rat brain. (A) A horizontal section through the newborn rat brain shows the highest level of expression in the hippocampal formation (H), the cortex (C), the septal nuclei (S), and the midline thalamic nuclei (T). (B) A sagittal section through the adult brain shows the highest level of expression in the hippocampal formation (H).

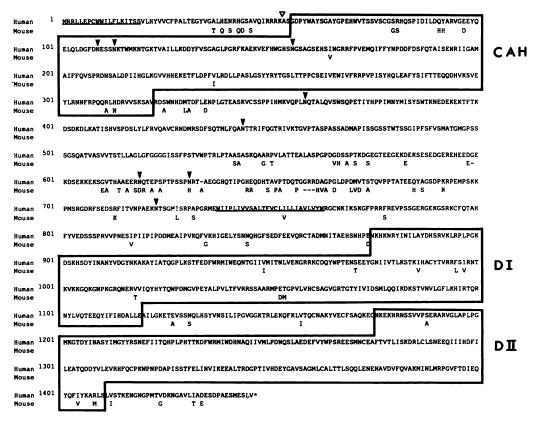


FIG. 3. Amino acid sequences of human and murine RPTP γ . The amino acid sequence of human RPTP γ is presented. Amino acids that are different in the murine gene are indicated. Dashes indicate amino acids that are not found in one of the sequences. The hydrophobic signal peptide (39) and the transmembrane domain are underlined. The potential N-glycosylation sites are indicated by arrowheads. The putative proteolytic cleavage site is indicated by an open triangle. The CAH-like domain and the phosphatase domains DI and DII are boxed.

agene, La Jolla, Calif.). Nucleotide sequences were determined by the dideoxynucleotide chain termination method (Sequenase; United States Biochemical, Cleveland, Ohio) with specific synthetic oligonucleotides as primers. All the clones were sequenced on both strands.

Sequence alignments. All DNA and protein data base searches were done with the Genetic Computer Group sequence analysis software package (8). The SWISSPROT and GenBank/EMBL data bases were searched with FASTA and TFASTA, respectively (25). Proteins were aligned with the Genetics Computer Group programs LINEUP, PILEUP, PRETTY, and BESTFIT.

Modeling of the CAH domain and energy minimization. After alignment of the CAH domain of RPTP_{γ} with the sequences of the soluble CAHs, the corresponding substitutions, deletions, and insertions were performed on an Evans & Sutherland and Silicon Graphic interactive displays, using the mutate options provided in program O (16). Inserted peptides and peptides flanking deleted segments were given an initial conformation which best fitted similar peptides taken from proteins with known three-dimensional structures and which are stored in the program as a data bank of conformational information. The initial side chain conformations of inserted and substituted residues were selected according to the rotamer library of Ponder and Richards (26). This rebuilt model was subjected to several cycles of energy minimization by using the program X-PLOR (4) and the energy parameter sets param19.pro and toph19.pro of the program CHARM (3). C α atoms were constrained to remain close to their original positions in the CAH structure. The root mean square deviation between the main-chain atoms in the model thus derived and the original CAH structure is 0.7 Å (0.07 nm). All the phi/psi angles of the resulting model fall within the allowed regions of the Ramachandran plot.

Tissue expression. Poly(A)⁺ RNA was prepared from adult mouse tissues by oligo(dT) selection as described elsewhere

Fbn III-7	LSPPTNLHLBANPDTGVLTVSWERSTTPDITGYRITTTPTNGQQGTALEEVVHADQSSCTFDNLSPGLEYNVSVYTVKDDKESVPISDTIIP
Contac	PGPPGGIRIBEIRDTAV.ALTWSRGTDNHSPISKYTIQSKTFLSEEWKDAKTEPSDIEGNMESARVIDLIPWMEYEFRIIATNTLGTGEPSMPSQRI
HLAR	SGPPRKVEVEPLNSTAV.HVYWKLPVPSKQHGQIRGYQVTYVRLENGEPRGLPIIQDVMLAEAQWRPEESEDYETTISGLTPETTYSVTVAAYTTKGDGARSKPKIVT
нртрб	SGPPRKVEVBAVNSTSV.KVSWRSPVPNKQBGQIRGYQVHYVRMENGEPKGQPMLKDVMLADAQWEFDDTTEHDMIISGLQPETSYSLTVTAYTKGDGARSKPKLVS
RPTPY	SSPPIHMKVQPLNQTAL.QVSWSQP.ETIYHPPIMNYMISYSWTKNEDEKEKTFTKDSDKDLKATISHVSPDSLYLFRVQAVCRNDMRSDFSOTMLF.
Consensus	PPBT-VV-WHIY

FIG. 4. Fibronectin type III repeat in RPTP γ . The sequence of the fibronectin type III repeat of human RPTP γ is aligned with those of typical fibronectin type III repeats of the human tyrosine phosphatases LAR (32) and HPTP δ (20), domain III-7 of human fibronectin (Fbn III-7) (18), and a fibronectin domain of chicken contactin (contac) (28). Amino acid residues that are shared by four repeats or more are printed in boldface type and indicated at the bottom as the consensus sequence.

A .	
PIFase Y human 50 PTPase Y mouse CAH : human CAH : human CAH : human CAH VI sheep CAH-r.p. mouse CAH-r.p. vac.v	. G D P Y W A Y S G A Y G P B . B W V T S S V S C G G R H Q B P Y D I L D Q Y A R V G E E Y Q E L Q
PTPase y human ¹⁰⁴ PTPase y mouse CAH I human CAH II human CAH II human CAH VI sheep CAH-r.p. mouse CAH-r.p. vacc.v	L D G F D N E S S N K T M M K W T G K T V A I L L K D D Y F V S G A G L P G R F K A E K V L D G F D N E S S N K T M M K W T G K T V A I L L K D D Y F V S G A G L P G R F K A E K V V . S T N P A T A K E I I W V G H S F H V N F E D N D N R S V L K G G P F S D S Y R L F G F V . S T D Q A T S L R I L W N G H A F N V E F D D S O D R A V L K G G P L D G T Y R L I O F V . S T D G G S A L R I L W N G H R F N V E F D D S O D R A V L K G G P L D G T Y R L I O F V . S T D G G S A L R I L W N G H R T C R V V F D D T Y D R S M L R G G P L D G T Y R L R O F L T G Y G L M H G E . F P V T W N G H T V Q I S L P S T M S M T T S D G T O Y L A K O M L . S P N Y V V C R D C E V T W D G H Y I O V I L K S K S V L S G G P L O G O E F E V Z V I . H Y N E S K P T T I O W T G K V V R I N F K G G Y I S G G F I P N E Y V L S S L
Consensus	
PTPase y human ¹⁴⁹ PTPase y mouse CAH : human CAH II human CAH II human CAH VI sheep CAH-r.p. mouse CAH-r.p. vacc.v	V EVEND. HENGSAGEESINGRRPVENOIPFYDPDDPDBPDTAISENRII ¹⁹⁷ EFENG. HENGSAGEEEVNGRRPVENOIPFYDPDDPDBPOTAISENRII HFENG. STNEHGEEEVDGVKYSAELAUVAEWDT.KISJELAEAASKADGL EFENG. SLDGGEEETVDGKKKYAAELELVEWDT.KIGDFGAAVOOPDGL ELENG. SLDGGEEETVDGVKYAAELELVEWDF.KINTFKEALKORDGI BFENGGABSEISGEEEVDGNRYVIEIEUVEWDF.KYNNFYEBAOREDGI RFENG. RENORGSEEVDGNRYVIEIEUVEWDF.KYNNFYEBAOREPDGL RFENG KEDDYGENELIDVYKYSGEINLVEWDKKEYSBYEEAAKKHODGL
Consensus	<u> </u>
PTPase Y human ¹⁹⁹ PTPase Y mouse CAH I human CAH II human CAH III human CAH VI sheep CAH-r.p. mouse CAH-r.p. vacc.v	G A M A X Y P Q V S P . R D W S A L D P I I H G L K G V V H H E R B T P L D . P Y U R D L D A S G A M A Y P Q V S P . R D W S A L D P I I H G L K G V V H H E R B T P L D . P Y U R D L D A S A V I G V L M K V G E A W P K L Q K V L D A L Q A Y K T K G E R A P P T . N P D P S T L D S S A V I G Y L M K V G S A K P G L Q K V U D A L Q A Y K T K G E R A P P T . N P D P S T L D S S A V I G Y L M K V G S A K P G L Q K V V D V L D S X K T K G E S A D P T . N P D P R G L D E S A V I G Y L K Y G H E W G E F Q I F L D A L D K Y K T K G E S A D P T . N P D P R G L D E S A V I G Y L K Y G H E W G E F Q I F L D A L D K Y K T K G E S A D P T . K P D P S T L D S C A V I A A L V E V K D Y T E W A Y S K F I S H L E D X K T K G E S K T I P . C P N P N T L D B C A I I I A L P V Q I G K E H V G L K A V T E I L Q D X Q Y K G E S K T I P . C P N P N T L D P I I I S T P L Q V S D H . K W V Y F Q K I V N Q L D S T R S A N T S A P F D S V P Y L D N L P S T
C08968888	aifva1gkff
PTPase y human ²⁴⁶ PTPase y mouse CAH I human CAH III human CAH III human CAH VI sheep CAH-r.p. mouse CAH-r.p. vacc.v	LG. SYYRY RYTG SLTTPPCSEITWEWIVFR RPVPISYHQLE APYSIFTTEQQD 295 LG. SYYRYTG SLTTPPCSEITWEWIVFR RPVPISYHQLE APYSIFTTEQQD 205 L DFM TYPG SLTHPPCSEIFUTICKESISV8SEQULAQPESLLS NVEGD L DYM TYPG SLTHPPLLESUTWIICKESISV8SEQULAQPESLLS NVEGD R DYM TYQG SFTTPPCEECUTWILLLKEPMIVSSOOMAKLESILSSAENF R DYM TYQG SFTTPPCEECUTWILLLKEPMIVSSOOMAKLESILSSAENF LR R. YYYSXLG SLTTPPCEECUTWILLF AFFENTVSSOOMAKLESILSSAENF LL R DYM TYCG SCTTPPCEECUTWILF AFFENTVSSOOMAKLESILSSAENF LL R DYM TYCG SCTTPPCEECUTWILF AFFENTVSSOOMAKLESILSSAENF LC R DYM TYCG SCTTPPCEECUTWILF AFFENTVSSOOMAKLESILSSAENF LC R DYM TYCG SCTTPPCEECUTWILF AFFENT LC R DYM TYCG SCTTPPCESEGUTWIIFFTPIN N HSDQLSKEPRTLSSSNHD
Consensus	1 d y Y - G B 1 T t p p c - o - V - W i p s Q f r - 1
PTPase Y human ²⁹⁵ PTPase Y mouse CAH I human CAH II human CAH III human CAH VI sheep CAH-r.p. mouse CAH-r.p. vacc.v Consects	H V K S V E Y L R N W F R P Q Q R L H D R V S K S A V ³²² H V K S V E Y L R N W F R P Q Q A L N D R V S K S A V ³²² N A V P M Q H W N R P T Q P L K G R T V R A S F .

Β.

PERCENT IDENTITY: CAH-r.p. vacc.virus RPTP Y RPTP Y CAH I CAH II CAH III CAH VI CAH-r.p mouse human human human human sheep mouse 34.5 100.0 37.0 35.3 RPTP Y human 98.1 35.6 38.9 39.0 RPTP y mouse 100.0 35.6 39.2 40.2 37.0 35.3 34.9 40.1 CAH I 100.0 64.2 63.3 39.1 47.1 human 41.7 CAH II human 100.0 63.0 42.7 47.3 CAH III human 100.0 44.4 45.6 42.6 CAH VI sheep 100.0 40.0 36.2 CAH-r.p. mouse 100.0 33.6 CAH-r.p. vacc.v 100.0

(38), fractionated (5 μ g per lane) on a formaldehyde-containing gel, transfered to Nytran (Schleicher & Schuell) by standard procedures, and probed with mouse clones MB-8 and MB-122 that encompass all of the first phosphatase domain, the juxtamembrane domain, the transmembrane domain, and the beginning of the extracellular portion.

In situ hybridization. Fresh-frozen rat tissue was cut on a cvrostat into 20-um-thick sections and thaw mounted onto gelatin-coated slides. The sections were fixed in 4% paraformaldehyde in 0.1 M sodium phosphate (pH 7.4) for 30 min and rinsed three times for 5 min each time in 0.1 M sodium phosphate and twice for 10 min each time in $2 \times SSC$ (1× SSC is 0.15 M NaCl plus 0.015 M sodium citrate). Two different oligonucleotide probes were used in the hybridization analysis: a 51-base oligonucleotide complementary to a portion of the cytoplasmic domain and a 52-base oligonucleotide complementary to a portion of the extracellular domain. The oligonucleotides were labeled with [35S]dATP (NEN DuPont) by using terminal deoxynucleotidyltransferase (Boehringer Mannheim) and purified by using Sephadex G-25 quick-spin columns (Boehringer Mannheim). The specific activity of the labeled probes was between 5×10^8 and 1×10^9 cpm/µg. Prehybridization and hybridization were carried out in a buffer containing 50% deionized formamid, $4 \times$ SSC, $1 \times$ Denhardt's solution, 500 µg of denatured salmon sperm DNA per ml, 250 µg of yeast tRNA per ml, and 10% dextran sulfate. The tissue was incubated for 12 h at 45°C in hybridization solution containing the labeled probe (10^6 cpm per section) and 10 mM dithiothreitol. Controls for specificity were performed on adjacent sections by adding a 30-fold concentration of the unlabeled oligonucleotide or by hybridization with the sense probe. After hybridization the sections were washed in two changes of $2 \times$ SSC at room temperature for 1 h, $1 \times$ SSC at 55°C for 30 min, 0.5× SSC at 55°C for 30 min, and 0.5× SSC at room temperature for 15 min and dehydrated in 60, 80, and 100% ethanol. After air drying, the sections were exposed to X-ray film for 5 to 10 days.

Chromosomal localization of murine RPTPy. To define the genetic locus encoding mouse RPTP γ , we used a probe that contains the first catalytic domain of mouse RPTPy to search for a restriction fragment length variant among inbred strains of mice. Southern blotting analysis of TaqI-digested DNA revealed two DNA fragments, of 2.9 and 1.8 kb, shared by all strains examined plus a fragment of either 2.7 kb (C3H/ HeJ and DBA/2J) or 2.0 kb (all other strains examined). The inheritance of this DNA variant in the AKXD, BXD, and BXH recombinant inbred strains of mice defined a genetic locus linked to Odc-9 (8 recombinants among 61 strains) and Plau (11 recombinants among 49 strains) near the centromeric end of the linkage map of chromosome 14 (Table 1). We propose *Ptpg* (phosphotyrosine phosphatase gamma) as the symbol for the locus, consistent with the symbol Ptpa previously assigned for mouse RPTP α (30).

Nucleotide sequence accession numbers. The sequence data for human and murine RPTP γ have been deposited in the GenBank data base under accession numbers L09247 and L09562, respectively.

RESULTS AND DISCUSSION

Resolving the issue of whether RPTP γ may function as a tumor suppressor gene (21) will require a detailed screening of tumors for genomic rearrangements and point mutations and reintroduction of wild-type RPTP γ into tumor cells. Since so far the genomic analysis of RPTP γ has been performed only with the partial sequence that has been described previously (17), we have cloned the full-length human cDNA. In addition, we have cloned the murine homolog of the RPTPy gene to facilitate an analysis of its tissue expression, as well as its normal in vivo function. The murine RPTPy gene was localized to chromosome 14 (Table 1) to a region syntenic to the chromosomal localization of the human RPTP γ gene (21). Northern (RNA) blot analysis shows that RPTP γ is widely expressed in different murine tissues (Fig. 1). Two major RPTPy transcripts of 5.5 and 8.5 kb were detected in the murine brain, lung, kidney, heart, skeletal muscle, liver, spleen, and testes. An additional shorter transcript of approximately 3.0 kb was detected in testes.

The two RPTPases most related to RPTPy are RPTPB/ HPTP((19, 22a) and a Drosophila phosphatase, DPTP99A (13, 34, 40). Since these phosphatases are specifically expressed in the central nervous system, we have analyzed the expression pattern of RPTPy in the brains of newborn and adult rats by in situ hybridization. In newborn rats the highest level of expression was detected in the hippocampal formation, in the septal and midline thalamic nuclei, and in the cortex (Fig. 2A). However, in the adult rat brain RPTP γ is highly expressed in the hippocampal formation (Fig. 2B) but not in the septal and midline thalamic nuclei or in the cortex (data not shown). Hybridization with probes derived from either the cytoplasmic or the extracellular domain gave similar results. The addition of a 30-fold concentration of unlabeled oligonucleotides completely blocked the labeling in all areas. Furthermore, no signal was observed in adjacent sections hybridized with the sense probe. These results demonstrate that the two probes hybridize to mRNA in a sequencespecific manner. The transient expression in the septal and midline thalamic nuclei and in the cortex of the newborn rat brain indicates that the expression of RPTP γ is developmentally regulated and may play a role in the development of these regions. It is noteworthy that the patterns of expression of RPTPy and the closely related RPTP β are totally different. In the embryo, RPTP β is expressed in the ventricular and subventricular zones of the brain and spinal cord. In the adult brain, RPTP_β is expressed in the Purkinje cell layer of the cerebellum, the dentate gyrus, and the subependymal layer of the anterior horn of the lateral ventricle (22a).

The complete amino acid sequences of the human and murine RPTP γ deduced from cDNA clones are presented in Fig. 3. Translation of the cDNA sequences reveals the

FIG. 5. Alignment of the CAH-like domain in RPTP γ with different forms of CAH. (A) The sequences of the CAH-like domains of human and murine RPTP γ are aligned with representative sequences of the CAH family: human CAH-1, CAH-2, and CAH-3 (sequences can be found in SWISSPROT accession numbers P00915, P00918, and P07451, respectively), sheep CAH-6 and vaccinia virus (vacc.v) CAH-like protein (CAH-r.p.) (SWISSPROT accession numbers P0860 and P04195, respectively), and mouse CAH-related protein (CAH-r.p.) (GenBank accession number X6197). Residues conserved in at least five of the eight sequences are boxed. The positions in human RPTP γ of the first and last amino acids in each line are indicated. The three His residues involved in Zn binding in CAH are indicated with arrowheads. (B) A matrix of the percent identity between the CAH-like domains in human and murine RPTP γ and the CAH sequences shown above derived from the alignments shown in panel A.

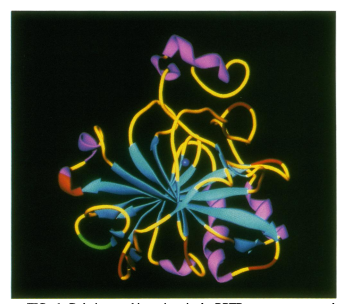


FIG. 6. Deletions and insertions in the RPTP γ sequence mapped onto the CAH structure. A ribbon diagram (6) of human CAH-2 with regions corresponding to deletions and insertions in the sequence of RPTP γ shown in green and red, respectively. Helices are shown in magenta, β strands are in blue, turns are in yellow, and the zinc atom is in dark blue. Note that all the deletions and insertions fall in exposed regions of CAH.

existence of an open reading frame of 1,445 amino acids for human RPTP γ and 1,442 amino acids for the murine homolog. The initiation Met in both human RPTP γ and the murine homolog are preceded by in-frame stop codons. Both proteins contain a signal sequence (underlined in Fig. 3) and a putative extracellular domain of 736 amino acids for human RPTP γ and 733 amino acids for the murine gene. The extracellular domain of human RPTP γ contains eight putative N-linked glycosylation sites, six of which are conserved in the murine homolog (marked with arrowheads in Fig. 3). The amino-terminal region of the extracellular domain of RPTP_γ (residues 56 to 322) has a striking sequence similarity to the enzyme CAH. This CAH-like domain is preceded by 4 basic amino acids (ArgArgArgLys) that resemble the cleavage site in the extracellular domain of the insulin receptor (35) separating the α and β subunits of the insulin receptor. This sequence motif may function as a cleavage site for proteolytic enzymes (1). It is of note that a similar cleavage site has been identified in the extracellular domain of another receptor-type phosphotyrosine phosphatase, denoted LAR (31, 41).

The CAH-like domain is followed by one fibronectin type III repeat, a motif found in many cell surface proteins. Alignment of the fibronectin type III sequence of RPTP γ with typical fibronectin type III repeats of other proteins is presented in Fig. 4. The remaining 293 amino acids of the extracellular domain are devoid of any Cys residues and can be subdivided into three regions: a Ser/Thr-rich region in which 32% of the amino acids are Ser and Thr (residues 442 to 560), followed by a region composed of 90% charged and polar amino acids (residues 561 to 662) and a region with no similarity to any known sequence motif. Hence, the region downstream from the fibronectin repeat may function as a spacer separating the CAH-like domain and the fibronectin type III repeat from the transmembrane region.

The extracellular domain of RPTP γ is followed by a typical transmembrane domain of 23 amino acid residues. As in most known receptor tyrosine phosphatases, the intracellular domain of RPTP γ contains two tandem phosphatase domains (10). It is noteworthy that the second phosphatase domain of RPTP γ has an Asp residue at position 1351 instead of a conserved Cys residue thought to be essential for catalytic activity (11, 27). Interestingly, an Asp residue is also found in a similar position in RPTP β (17) and in the *Drosophila* phosphatase 99A (13, 34, 40). Moreover, the second catalytic domain of RPTP γ contains an insert of 15 amino acids (residues 1299 to 1313) identical to the insert found in corresponding position in RPTP β (17) and so far

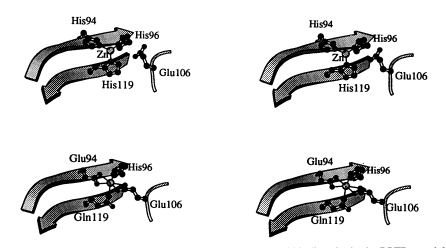


FIG. 7. Comparison between the zinc-binding site of CAH and the putative metal-binding site in the RPTP_Y model. (Top) Stereo view of the zinc-binding site in human CAH-2 viewed roughly from the direction of the water molecule which is the fourth zinc ligand in this enzyme (OHH263 [9]). Note the tetrahedral coordination of the zinc. Bonds from the zinc to ligand atoms are shown in thin lines. (Bottom) Stereo view of the putative metal-binding site in the RPTP_Y domain (same view as top drawing), after applying the substitutions His-94 to Glu and His-119 to Gln and energy minimization (see text). Glu-106, which is part of the active site in human CAH-2, has been rotated to become one of the potential ligands. Note the planar constellation of atoms around the putative metal. Bidentate coordination is shown for Glu-94 and Glu-106. All the numbers of residues are according to CAH-2.

appears to be unique to these two phosphatases. This insert contains a Tyr residue in position 1308 which is preceded by two Asp residues and followed by a Val residue, suggesting that it may function as a phosphorylation site for tyrosine kinases.

Human RPTPy and its murine homolog are 90% identical at the nucleotide level and 95% identical at the amino acid level, and most of the differences are conservative substitutions (Fig. 3). This strikingly high conservation is found in both the cytoplasmic and extracellular domains. This degree of conservation between human and mouse RPTP γ is higher than in some of the other phosphatases, such as the extracellular portion of RPTP α (17, 20, 23, 30). The most conserved parts are the fibronectin repeat (100% identity), the CAH-like domain (97% identity), and the Ser/Thr-rich domain (97% identity). The rest of the extracellular domain is the least conserved (70% identity). It is noteworthy that the cluster of charged and polar amino acids that follows the Ser/Thr-rich domain is found in both human RPTPy and the murine homolog in spite of the lower degree of sequence conservation. The high degree of identity between the human and murine sequences in the CAH-like domain, the fibronectin type III repeat, and the Ser/Thr-rich domain may suggest that these regions are required for a conserved biological functions.

As already mentioned, the amino-terminal region of the extracellular domain of RPTPy contains a region of 266 amino acids with a striking sequence similarity to the enzyme CAH. CAH catalyzes the hydration of metabolic CO_2 or the dehydration of HCO_3^- in the following reaction: CO_2^- + $H_2O \Leftrightarrow H^+ + HCO_3^-$. CAHs are ubiquitously expressed enzymes with extremely efficient turnover rate of 10^6 s^{-1} for CO₂ hydration. It has also been shown that CAHs are able to hydrolyze certain esters and to hydrate specific aldehydes. All CAHs are zinc metalloenzymes in which the zinc atom is required for the catalytic activity. Seven types of CAH have been identified so far (reviewed in reference 33). CAH-1, CAH-2, and CAH-3 are cytoplasmic enzymes, CAH-4 is an extracellular glycoprotein, CAH-5 is a mitochondrial enzyme, CAH-6 is a secreted enzyme, and CAH-7 is a membrane-bound enzyme. In addition, vaccinia virus also contains a transmembrane protein with a CAH-like domain in its extracellular portion.

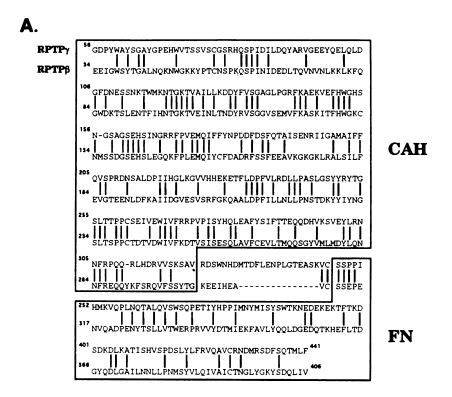
The CAH-like domains in human and murine RPTPy were aligned with the amino acid sequences of the different forms of CAH (Fig. 5A). Comparison of the sequences revealed clusters of identical amino acids that usually match regions of high conservation of sequences between the different forms of CAH. Very few insertions and deletions were required in order to align the CAH domain in RPTP γ to that of the different forms of CAH. It is noteworthy that 11 of the 19 residues that form the active site of CAH (9) are also found in RPTP γ . Of the three His residues that ligate the zinc atom in CAH (indicated with arrowheads in Fig. 5A), only one, in position 151, is conserved in RPTP γ , whereas the other two are replaced by Glu and Gln residues (positions 149 and 175, respectively). The CAH domain of RPTP γ shares 35 to 40% sequence identity with all known CAHs (Fig. 5B)

On the basis of the striking sequence similarity of this region with CAHs, we have constructed a model for the structure of this region in RPTP γ by substitution of amino acids from the RPTP γ sequence into equivalent positions in the known three-dimensional crystal structure of CAH (9) (entry lca2 in the Brookhaven Protein Data Bank). This was followed by energy minimization (4) in which the C α atoms

were restrained to their positions in the original CAH structure. In the absence of direct structural evidence it is difficult to assess the correctness of the resulting model, but three features point strongly to its relevance to the actual fold of the RPTPy domain. First, as all the deletions and insertions relative to the CAH sequence map to surface regions and all but one fall in stretches of the polypeptide chain lacking any well-defined secondary structure (Fig. 6), they could all be accommodated without distorting the original model. Second, substitutions of internal residues in the RPTP γ domain follow a pattern that preserves the tight packing of the two hydrophobic cores in CAH (9), as is often encountered in families of homologous proteins (2). Although replacement of some aromatic residues by aliphatic ones in the RPTPy domain (Phe-66 in CAH to Val-124 in RPTPy, Phe-70 to Leu-128, Phe-93 to Val-148, and Phe-176 to Leu-233) reduces somewhat the aromatic character of the hydrophobic core proximal to the zinc-binding site (9), the tight packing is maintained through the substitutions Ile-59 in CAH to Trp-116 in RPTPy, Val-160 to Ile-217, and Ser-56 to Asn. Third, inspection of the sequence of the CAH domain of mouse RPTP γ reveals that substitutions relative to the human gene also follow the pattern of conservation of a tightly packed hydrophobic core (e.g., Ile-165 in human RPTPy to Val in the murine protein and Val-237 in the human protein to Ile in the murine homolog).

Particularly intriguing is the fate of the zinc-binding site and the active site of CAH in the RPTPy domain. Sequence alignment shows that two of the conserved histidines ligating the zinc in all known CAHs have been replaced in RPTPy (His-94 in CAH to Glu-149 in RPTPy and His-119 to Gln-175). Inspection of zinc-binding sites in proteins whose three-dimensional structure is known (37) reveals that none of them contains Gln. Furthermore, loss of zinc-binding capability as a result of His-to-Gln mutation was reported in the growth hormone family (7). Moreover, inspection of the energy-minimized model of the CAH domain of RPTPy suggests that the residues Glu-149, His-151, Glu-162, and Gln-175 can form a planar constellation of atoms (Fig. 7) that is often found in the binding sites of octahedrally coordinated transition metals such as manganese (e.g., see reference 12).

A CAH-like domain was also found in the amino-terminal region of RPTP β /HPTP ζ (19, 22a). It appears that the salient features of the CAH-like domain in RPTP γ , such as the tightly packed hydrophobic core and the replacement of two of the three conserved His residues, are also observed in the CAH domain of RPTPB. This similarity may reflect functional parallels between these two domains. The biological role of the CAH domains of RPTP γ and RPTP β is not known. In view of the fact that only one of the three His residues that ligate zinc and are crucial for CAH activity is conserved, it is conceivable that the CAH domains of RPTP γ and RPTP β may lack the ability to bind zinc. It is possible, however, that the site occupied by zinc in CAH will be capable of binding other transition metals and thus have a function other than hydration of metabolic CO_2 . Interestingly, the homology between RPTP γ and RPTP β extends into the downstream fibronectin type III repeat which is flanked in both proteins by two conserved Cys residues. An alignment of the sequences of these two domains in RPTPy and RPTP β (Fig. 8A) shows that they share approximately 37% sequence identity. The fibronectin type III repeat in RPTP γ is followed by a stretch of 293 amino acids that is characterized by a lack of Cys residues and thus may serve as a spacer that separates the CAH-like domain





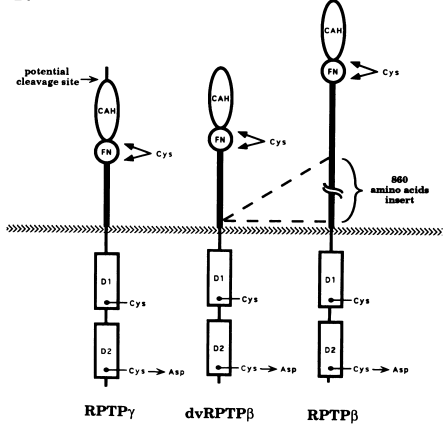


FIG. 8. RPTP γ and RPTP β define a new subfamily of receptor tyrosine phosphatases. (A) Alignment of the CAH-like domains and the fibronectin type III repeats of RPTP γ and RPTP β (22a). The sequence in this region of RPTP β is identical to that reported by Krueger and Saito (19) for RPTP ζ . The aligned CAH domains and fibronectin type III repeats (FN) are boxed. Identical amino acids are indicated by a connecting line. (B) Schematic diagram summarizing the conserved features that define the subfamily of RPTP γ and RPTP β . The extracellular regions of RPTP γ and the two forms of RPTP β (RPTP β and the deletion variant [dvRPTP β]) contain CAH-like domains (CAH), fibronectin type III repeats (FN), and spacers of variable length that are characterized by a very low content of cysteine residues (indicated by thick lines). The conserved cysteine residues that flank the fibronectin type III repeat are marked. The cytoplasmic regions of RPTP γ and RPTP β contain two typical phosphatase domains (D1 and D2). A conserved Cys residue in the first phosphatase domain and an Asp residue that replaces the Cys residue in the second phosphatase domain are indicated. The potential cleavage site in RPTP γ and the 860-amino-acid insert in RPTP β which is not present in the deletion variant are denoted.

and the fibronectin repeat from the transmembrane domain. Two forms of RPTP β have been identified (22a); both contain intact CAH-like domains and fibronectin type III repeats followed by a Cys-free region. The two forms of RPTP β differ in the length of the Cys-free spacer, which contains 1,048 amino acids in the long form and 384 amino acids in the deletion variant. As mentioned earlier, the high similarity between RPTP γ and RPTP β extends to the cytoplasmic domains of RPTP γ and RPTP β .

On the basis of these similarities we suggest that these two phosphatases define a new subfamily of receptor tyrosine phosphatases (Fig. 8B). The elucidation of the biological function of RPTP γ and RPTP β and the role of the CAH-like domain in these proteins may require the identification of their putative natural ligands and the binding region of these orphan receptors.

ACKNOWLEDGMENTS

This work was supported by grants from SUGEN, Inc. (to J. Schlessinger), and from HFSPO (to J. Schlessinger); J. Sap is a recipient of a postdoctoral fellowship from HFSPO. Computing was supported by NSF under grant DIR-8908095.

We thank Kiki Nelson for synthesis of peptides and oligonucleotides.

REFERENCES

- Barr, P. J. 1991. Mammalian subtilisins: the long-sought dibasic processing endoproteases. Cell 66:1-3.
- Bordo, D., and P. Argos. 1990. Evolution of protein cores constraints in point mutations as observed in globin tertiary structures. J. Mol. Biol. 211:975–988.
- Brooks, B. R., R. E. Bruccoleri, B. D. Olfason, D. J. States, S. Swaminathan, and M. Karplus. 1983. CHARM: a program for macromolecular energy minimization and dynamics calculations. J. Comput. Chem. 4:187–217.
- 4. Brunger, A. T. 1992. X-PLOR (version 3.0) manual. Yale University, New Haven, Conn.
- Cantley, L. C., K. R. Auger, C. Carpenter, B. Duckworth, A. Graziani, R. Kapeller, and S. Soltoff. 1991. Oncogenes and signal transduction. Cell 64:281-302.
- Carson, M. 1987. Ribbon models of macromolecules. J. Mol. Graphics 5:103-106.
- 7. Cunningham, B. C., S. Bass, G. Fuh, and J. A. Wells. 1990. Zinc mediation of the binding of human growth hormone to the human prolactin receptor. Science 250:1709–1712.
- 8. Devereux, J., P. Haeberli, and O. Smithies. 1989. A comprehensive set of sequence analysis programs in the VAX. Nucleic Acids Res. 12:387–396.
- 9. Ericksson, A. E., T. A. Jones, and A. Liljas. 1988. Refined structure of human carbonic anhydrase II at 2.0 A resolution. Proteins 4:274–282.
- Fischer, E. H., H. Charbonneau, and N. K. Tonks. 1991. Protein tyrosine phosphatases: a diverse family of intracellular and transmembrane enzymes. Science 253:401–406.
- Guan, K., and J. E. Dixon. 1991. Evidence for protein-tyrosinephosphatase catalysis proceeding via a cysteine-phosphate intermediate. J. Biol. Chem. 266:17026–17030.
- 12. Hardman, K. D., R. C. Agarwal, and M. J. Freiser. 1982.

Manganese and calcium binding sites of concanavalin A. J. Mol. Biol. 157:69–86.

- Hariharan, I. K., P. T. Chuang, and G. M. Rubin. 1991. Cloning and characterization of a receptor-class phosphotyrosine phosphatase gene expressed on central nervous system axons in *Drosophila melanogaster*. Proc. Natl. Acad. Sci. USA 88: 11266–11270.
- 14. Hunter, T. 1987. A tail of two src's mutatis mutandis. Cell 49:1-4.
- Hunter, T. 1991. Cooperation between oncogenes. Cell 64:249– 270.
- Jones, T. A., J. Y. Zou, S. W. Cowan, and M. Kjeldgaard. 1991. Improved methods for binding protein models in electron density maps and the location of errors in these models. Acta Crystallogr. Sect. A47:110-119.
- 17. Kaplan, R., B. Morse, K. Huebner, C. Croce, R. Howk, M. Ravera, G. Ricca, M. Jaye, and J. Schlessinger. 1990. Cloning of three human tyrosine phosphatases reveals a multigene family of receptor-linked protein-tyrosine-phosphatases expressed in brain. Proc. Natl. Acad. Sci. USA 87:7000-7004.
- Kornblihtt, A. R., H. K. Umezawa, K. Vibe-Pedersen, and F. E. Baralle. 1985. Primary structure of human fibronectin: differential splicing may generate at least 10 polypeptides from a single gene. EMBO J. 4:1755-1759.
- Krueger, N. X., and H. Saito. 1992. A human transmembrane protein-tyrosine phosphatase, RPTPζ, is expressed in brain and has an N-terminal receptor domain homologous to carbonic anhydrase. Proc. Natl. Acad. Sci. USA 89:7417–7421.
- Krueger, N. X., M. Streuli, and H. Saito. 1990. Structural diversity and evolution of human receptor-like protein tyrosine phosphatases. EMBO J. 9:3241–3252.
- LaForgia, S., B. Morse, J. Levy, G. Barnea, L. A. Cannizzaro, F. Li, P. C. Nowell, L. Boghosian-Sell, J. Glick, A. Weston, C. C. Harris, H. Drabkin, D. Patterson, C. M. Croce, J. Schlessinger, and K. Huebner. 1991. Receptor protein-tyrosine phosphatase γ is a candidate tumor suppressor gene at human chromosome region 3p21. Proc. Natl. Acad. Sci. USA 88:5036-5040.
- Ledbetter, J. A., N. K. Tonks, E. H. Fischer, and E. A. Clark. 1988. CD45 regulates signal transduction and lymphocyte activation by specific association with receptor molecules on T or B cells. Proc. Natl. Acad. Sci. USA 85:8628–8632.
- 22a.Levy, J. B., et al. Submitted for publication.
- Matthews, R. J., E. D. Cahir, and M. L. Thomas. 1990. Identification of an additional member of the protein-tyrosinephosphatase family: evidence for alternative splicing in the tyrosine-phosphatase family: evidence for alternative splicing in the tyrosine phosphatase domain. Proc. Natl. Acad. Sci. USA 87:4444-4448.
- Nada, S., M. Okada, A. MacAuley, J. A. Cooper, and H. Nakagawa. 1991. Cloning of a complementary DNA for a protein-tyrosine kinase that specifically phosphorylates a negative regulatory site of p60^{c-src}. Nature (London) 351:69–72.
- Pearson, W. R., and D. J. Lipman. 1988. Improved tools for biological sequence comparison. Proc. Natl. Acad. Sci. USA 85:2444-2448.
- Ponder, J. W., and F. M. Richards. 1987. Tertiary templates for proteins: use of packing criteria in the enumeration of allowed sequences for different structural classes. J. Mol. Biol. 193:775– 791.

- Pot, D. A., and J. E. Dixon. 1992. Active site labeling of a receptor-like protein tyrosine phosphatase. J. Biol. Chem. 267: 140-143.
- Ranscht, B., and M. T. Dours. 1988. Sequence of contactin, a 130-kD glycoprotein concentrated in areas of interneuronal contact, defines a new member of the immunoglobulin supergene family in the nervous system. J. Cell Biol. 107:1561–1573.
- Saito, H., and M. Streuli. 1991. Molecular characterization of protein tyrosine phosphatases. Cell Growth Differ. 2:59–65.
- Sap, J., P. D'Eustachio, D. Givol, and J. Schlessinger. 1990. Cloning and expression of a widely expressed receptor tyrosine phosphatase. Proc. Natl. Acad. Sci. USA 87:6112–6116.
- Streuli, M., N. X. Kreuger, P. Dariniello, M. Tang, J. M. Munro, W. A. Blattler, D. A. Adler, C. M. Disteche, and H. Saito. 1992. Expression of the receptor-linked protein tyrosine phosphatase LAR: proteolytic cleavage and shedding of the CAM-like extracellular region. EMBO J. 11:897–907.
- 32. Streuli, M., N. X. Kreuger, L. R. Hall, S. F. Schlossman, and H. Saito. 1988. A new member of the immunoglobulin superfamily that has a cytoplasmic region homologous to the leukocyte common antigen. J. Exp. Med. 168:1523–1530.
- Tashian, R. E. 1989. The carbonic anhydrases: widening perspectives on their evolution expression and function. Bioessays 10:186–192.
- 34. Tian, S. S., P. Tsoulfas, and K. Zinn. 1991. Three receptorlinked protein-tyrosine phosphatases are selectively expressed

on central nervous system axons in the drosophila embryo. Cell 67:675–685.

- 35. Ullrich, A., J. R. Bell, E. Y. Chen, R. Herrera, L. M. Petruzzelli, T. J. Dull, A. Gray, L. Coussens, Y. C. Liao, M. Tsubokawa, A. Mason, P. H. Seeburg, C. Grunfeld, O. M. Rosen, and J. Ramachandran. 1985. Human insulin receptor and its relationship to the tyrosine kinase family of oncogenes. Nature (London) 313:756-761.
- 36. Ullrich, A., and J. Schlessinger. 1990. Signal transduction by receptors with tyrosine kinase activity. Cell 61:203-212.
- Vallee, B. L., and D. S. Auld. 1990. Zinc coordination, function, and structure of zinc enzymes and other proteins. Biochemistry 29:5647-5659.
- Vennstrom, B., and J. M. Bishop. 1982. Isolation and characterization of chicken DNA homologous to the two putative oncogenes of avian erythroblastosis virus. Cell 28:135-143.
- 39. von Heijne, G. 1986. A new method for predicting signal sequence cleavage sites. Nucleic Acids Res. 14:4683-4690.
- 40. Yang, X., K. T. Seow, S. M. Bahri, S. H. Oon, and W. Chia. 1991. Two drosophila receptor-like tyrosine phosphatase genes are expressed in a subset of developing axons and pioneer neurons in the embryonic CNS. Cell 67:661–673.
- 41. Yu, Q., T. Lenardo, and R. A. Weinberg. 1992. The N-terminal and C-terminal domains of a receptor tyrosine phosphatase are associated by non-covalent linkage. Oncogene 7:1051-1057.