Protein tyrosine phosphatases expression during development of mouse superior colliculus

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Abstract   Protein tyrosine phosphatases (PTPs) are key regulators of different processes during development of the central nervous system. However, expression patterns and potential roles of PTPs in the developing superior colliculus remain poorly investigated. In this study, a degenerate primer-based reverse transcription-polymerase chain reaction (RT-PCR) approach was used to isolate seven different intracellular PTPs and nine different receptor-type PTPs (RPTPs) from embryonic E15 mouse superior colliculus. Subsequently, the expression patterns of 11 PTPs (TC-PTP, PTP1C, PTP1D, PTP-MEG2, PTP-PEST, RPTPj, RPTPl, RPTP6, RPTPRR, RPTPσ, RPTPκ and RPTPl) were further analyzed in detail in superior colliculus from embryonic E13 to postnatal P20 stages by quantitative real-time RT-PCR, Western blotting and immunohistochemistry. Each of the 11 PTPs exhibits distinct spatiotemporal regulation of mRNAs and proteins in the developing superior colliculus suggesting their versatile roles in genesis of neuronal and glial cells and retinocollicular topographic mapping. At E13, additional double-immunohistochemical analysis revealed the expression of PTPs in collicular nestin-positive neural progenitor cells and RC-2-immunoreactive radial glia cells, indicating the potential functional importance of PTPs in neurogenesis and gliogenesis.

Keywords   Superior colliculus · Protein tyrosine phosphatases · Real-time RT-PCR · Immunohistochemistry · Development

Introduction

The mouse retinocollicular system serves as an excellent model to explore the molecular and cellular mechanisms of axon growth and guidance as well as topographic map formation. Retinal ganglion cells (RGCs) differentiate within the retina at embryonic (E) day 11 (Cook 2003) and extend their axons toward the optic fissure. The first retinal axons travel out of the eye along the optic nerve and reach the optic chiasm at E14. After crossing, the retinal axons innervate their target, the superior colliculus, also known as the optic tectum in non-mammalian vertebrates, between E16 and postnatal (P) day 0 (Godement et al. 1984). Retinal inputs to the superior colliculus are organized in a precise topographic map, in which the temporal–nasal and dorsal–ventral axes of the retina correspond to the anterior–posterior and lateral–medial axes of the midbrain.
Extensive investigation of the developmental patterning of this topographic map has revealed gradients of transcription factors and cell surface receptors in the retina and tectum which guide its formation (Thanos and Mey 2001; O’Leary and McLaughlin 2005). Of interest here are the protein tyrosine phosphatases (PTPs), a large family of proteins that have been described to be implicated in axon growth and guidance (Garrity et al. 1999; Bixby 2000; Newsome et al. 2000; Stoker 2001) in different species. In RGC axons of Xenopus and chicken, RPTP-LAR, RPTPδ, RPTPμ and RPTPσ promote retinal neurite outgrowth (Burden-Gulley and Brady-Kalnay 1999; Ledig et al. 1999; Johnson et al. 2001), growth cone steering (Burden-Gulley et al. 2002) and targeting of retinal axons within the optic tectum (Rashid-Doubell et al. 2002). However, less is known regarding the potential role of PTPs in the development of the mammalian superior colliculus.

In order to identify PTPs that might contribute to signaling in the superior collicular development a degenerate primer-based reverse transcription polymerase chain reaction (PCR) approach was used to isolate cDNAs encoding PTPs from embryonic (E15) mouse superior colliculus. At this stage, neuronal and radial glial cells are generated (Gotz and Huttner 2005) as well as retinal ganglion axons first contact the superior colliculus. Using this approach seven different intracellular, non-transmembrane PTPs and nine different receptor-type PTPs (RPTPs) were identified. Subsequently, the expression pattern of 11 PTPs (TC-PTP, PTP1D, PTP1C, PTP- MEG2, PTP-PEST, RPTPj, RPTPe, RPTPRR, RPTPκ, RPTPγ and RPTPσ) was analyzed in more detail in embryonic (E13, E15, E18) and postnatal (P0, P4, P12, and P20) superior colliculus by real-time RT-PCR, Western Blotting and immunohistochemistry. With ongoing maturation, all 11 PTPs displayed a distinct spatiotemporal regulation of mRNAs and proteins in the pre- and postnatal superior colliculus correlating with different processes such as proliferation, differentiation, axonal innervation and arborisation.

Methods

Animals

Adult NMRI mice were obtained from Charles River Laboratories (Sulzfeld, Germany) and mated over night. Females were checked for the presence of a vaginal plug, which corresponds to the gestational day 0.5 (E0.5). For all analyses, embryonic (E13, E15, E18) and postnatal (P0, P4, P12, P16, P20) stages were determined according to the staging criteria of Theiler (Bard et al. 1998).

For RNA preparation, collicular tissue from each developmental stage was isolated, pooled, frozen in liquid nitrogen and stored at −70°C until RNA extraction. Total RNA was extracted according to the manufacturer’s instructions (RNeasy Mini or Midi kit, Qiagen, Hilden, Germany). Using a cDNA-synthesis kit (Fermentas GmbH, St. Leon-Rot, Germany) 1 µg of total RNA was used for reverse transcription.

PCR amplification of PTP sequences using degenerate primers

To produce PCR-generated DNA-fragments corresponding to PTP sequences in the conserved catalytic domain, cDNAs were reversely transcribed from E15 superior colliculus and were used as a template for the amplification with Taq polymerase (Eppendorf, Germany). Degenerate primers corresponding to amino acid sequences DFWQ(R/ K/E/G)MI(M/V)WD(E/Q/H) (upstream) and HCSAGI(V/M)G (downstream) were synthesized by Invitrogen (Carlsbad, CA, USA). Low stringency PCR-reaction conditions were as follows: 5 min 94°C, followed by 36 cycles of 1 min at 94°C, 1 min at 50°C and 1 min at 72°C. The reaction products were run on 1.5% agarose gels, ligated into pCRII-TOPO Plasmids (Invitrogen) and used to transform competent E. coli TOP10 cells (Invitrogen, Carlsbad, CA, USA). Clones that contained inserts were sequenced using automated DNA sequencing (Department of Molecular Neurobiochemistry, Ruhr-University-Bochum). Obtained PCR fragment sequences were compared to sequences covered in the NCBI databases.

Quantitative real-time RT-PCR

Real-time-PCR using Syber Green I (Eurogentec) was performed on an Opticon-Cycler (MJ Research). Primer sequences of both housekeeping genes β-actin and cyclophilin and of the identified five intracellular PTPs and six RPTPs were designed (Horvat-Brocker et al. 2008). Their sequences, expected amplicon sizes and accession numbers are listed in Supplemental Table 1.

Primer concentration was optimized to a final concentration of 0.6 µM and combined with 20 ng RNA per well. Total three reactions per sample RNA (triplets) were set with a final volume of 20 µl per single reaction. Real-time PCR was performed as described previously (Ray et al. 2005; Horvat-Brocker et al. 2008). Briefly, each RT-PCR was performed from a pool of tissues originating from 15 different animals per each developmental stage. The average Ct values of three independent experiments (triplicates) were used to calculate the ratios for intracellular PTPs as described before (Pfaffl et al. 2002). In order to obtain amplification
Cloning and identification of intracellular and receptor-type protein tyrosine phosphatases (PTPs) from embryonic E15 mouse superior colliculus using a degenerate primer-based RT-PCR approach

| PTPs      | No. of PTP clones = 310 (%) |
|-----------|-----------------------------|
| RPTPα     | 158 (51)                    |
| RPTP-LAR  | 62 (20)                     |
| TC-PTP    | 23 (7.4)                    |
| PTP1B     | 17 (5.5)                    |
| RPTPJ     | 12 (4)                      |
| RPTPRR    | 9 (3)                       |
| RPTPδ     | 5 (1.6)                     |
| RPTPε     | 3 (0.9)                     |
| PTP-MEG2  | 3 (0.9)                     |
| PTP36     | 2 (0.6)                     |
| PTP1C     | 2 (0.6)                     |
| PTP1D     | 2 (0.6)                     |
| RPTPγ     | 1 (0.3)                     |
| RPTPγ     | 1 (0.3)                     |
| PTP-PEST  | 1 (0.3)                     |

The reamplified intracellular PTPs and receptor-type protein tyrosine phosphatases (RPTPs) cloned by RT-PCR were ranked by the relative abundance of the PCR products. The seven different intracellular PTPs (TC-PTP, PTP1B, PTP-MEG2, PTP36, PTP1C, PTP1D and PTP-PEST) recovered in the E15 mouse superior colliculus screen represented 50 (16%) from a total of 310 clones. The nine different PTPs (RPTPRR, RPTPJ, RPTPJ, RPTPR, RPTP, RPTP, RPTP and RPTPβ) were represented by 260 clones (84%).

efficiencies of different primer sets, we generated standard curves by a twofold dilution series with template amounts ranging from 8 to 0.125 ng DNA per well. The efficiency of the PCR reaction was calculated for each primer pair according to the equation $E = 10^{(-1/slope)}$ (Pfaffl et al. 2002).

The relative expression ($R$) of PTPs was calculated based on real-time PCR efficiency ($E$) and the threshold cycle ($Ct$) deviation of PTP at each developmental stage versus a control (embryonic stage 13, E13) according to the following equation: $R = \left( \frac{E_{target}^{ACTarget} (\text{MEAN control} - \text{MEAN sample})}{E_{reference}^{ACTarget} (\text{MEAN control} - \text{MEAN sample})} \right)$.

For Western blotting, mouse collicular tissues from each developmental stage (E13, E15, E18, P0, P4, P12, P16 and P20) were isolated, pooled and immediately homogenized in ice-cold lysis buffer (50 mM HEPES, pH 7.5, 150 mM NaCl, 1.5 mM MgCl2, 5 mM EGTA, 10% v/v glycerin, 1% v/v Triton X-100, 0.1 mM Na3VO4, 10 μg/ml aprotinin, 10 μg/ml leupeptine, 1 mM phenylmethylsulfonyl fluoride). Lysates were clarified by centrifugation at 12,000×g for 20 min at 4°C. The protein concentration was determined using a BCA Protein Assay Kit (Pierce, Rockford, USA). 20 μg of each protein lysate was separated on 4–10% SDS-polyacrylamide gels (Roth, Karlsruhe, Germany). Immunoblotting and immunodetection were performed as described previously (Horvat-Brocker et al. 2008) using the following primary PTP-antibodies: rabbit anti-TC-PTP (1:2,000), mouse anti-PTP1C (1:2,000), rabbit anti-PTP1D (1:2,000), rabbit anti-PTP-MEG2 (1:1,000), rabbit anti-PTP-PEST (1:2,000), rabbit anti-RPTPJ (1:1,000), rabbit anti-RPTPPR (1:1,000), rabbit anti-RPTPP (1:1,000), rabbit anti-RPTPγ (1:1,000) and rabbit anti-RPTPα (1:1,000). Immunoreactivities were detected using goat anti-rabbit-HRP-coupled or goat anti-mouse-HRP-coupled antibodies (all from Dianova, Hamburg, Germany).

Antibodies

The following primary antibodies were used for Western blot analyses and immunohistochemical stainings: rabbit anti-TC-PTP (Lammers et al. 1993), mouse anti-PTP1C (Santa Cruz Biotechnology, Santa Cruz, United States), rabbit anti-PTP1D (Stein-Gerlach et al. 1995; Tomic et al. 1995), rabbit anti-PTP-MEG2 (Sugen, Redwoodcity, CA, USA), rabbit anti-PTP-PEST (Eurogentec, Köln, Germany), rabbit anti-RPTPJ (Jallal et al. 1997) rabbit anti-RPTPPR (Fuchs et al. 1996; Anders et al. 2006), rabbit anti-RPTPPγ (Horvat-Brocker et al. 2008), rabbit anti-RPTPP (Moller et al. 1995), rabbit anti-RPTPPR (Horvat-Brocker et al. 2008), rabbit anti-RPTPR (Aicher et al. 1997), mouse anti-β-actin (Sigma, St. Louis, MO, USA), mouse anti-nestin (Chemicon, Temecula, CA, USA) and mouse anti-RC2 (Developmental Studies Hybridoma Bank, University of Iowa, Iowa City, IA, USA). The secondary antibodies used for Western blot analyses and immunohistochemical stainings were as follows: goat anti-mouse-CY3-coupled, goat anti-mouse-CY2-coupled, goat anti-rabbit-CY3-coupled, goat anti-rabbit-HRP-coupled and goat anti-mouse-HRP-coupled antibodies (all from Dianova, Hamburg, Germany).

For Western blotting, mouse collicular tissues from each developmental stage (E13, E15, E18, P0, P4, P12, P16 and P20) were isolated, pooled and immediately homogenized in ice-cold lysis buffer (50 mM HEPES, pH 7.5, 150 mM NaCl, 1.5 mM MgCl2, 5 mM EGTA, 10% v/v glycerin, 1% v/v Triton X-100, 0.1 mM Na3VO4, 10 μg/ml aprotinin, 10 μg/ml leupeptine, 1 mM phenylmethylsulfonyl fluoride). Lysates were clarified by centrifugation at 12,000×g for 20 min at 4°C. The protein concentration was determined using a BCA Protein Assay Kit (Pierce, Rockford, USA). 20 μg of each protein lysate was separated on 4–10% SDS-polyacrylamide gels (Roth, Karlsruhe, Germany). Immunoblotting and immunodetection were performed as described previously (Horvat-Brocker et al. 2008) using the following primary PTP-antibodies: rabbit anti-TC-PTP (1:2,000), mouse anti-PTP1C (1:2,000), rabbit anti-PTP1D (1:2,000), rabbit anti-PTP-MEG2 (1:1,000), rabbit anti-PTP-PEST (1:2,000), rabbit anti-RPTPJ (1:1,000), rabbit anti-RPTPPR (1:1,000), rabbit anti-RPTPP (1:1,000), rabbit anti-RPTPγ (1:1,000) and rabbit anti-RPTPα (1:1,000). Immunoreactivities were detected using goat anti-rabbit-HRP-coupled or goat anti-mouse-HRP-coupled secondary antibodies (1:12000). Protein bands were detected using enhanced chemiluminescence ECL (Roth). After stripping with Restore Western blot Stripping Buffer (Pierce) all blots were reprobed using mouse anti-β-actin antibody (Sigma, 1:5,000), which indicates nearby equal amounts of proteins per each analysed developmental stage. Protein levels were quantified by using image J analysis software.
Identification of PTPs expressed in the mouse superior colliculus at embryonic stage 15 (E15) by degenerate primer RT-PCR

In order to identify PTPs that might contribute to the signaling in the embryonic (E15) mouse superior colliculus, a degenerate primer-based RT-PCR approach was used. At E15, retinal ganglion axons start to innervate the superior colliculus (Edwards et al. 1986; Herrera et al. 2003) and intensive genesis processes (differentiation, proliferation, migration) of neuronal and radial glial cells occur (Altman and Bayer 1981; Edwards et al. 1986; Galileo et al. 1990). The primers used in this study were synthesized based on the catalytic domains of PTPs (DFWSQ(R/K/E/G)M(M/V)WD(E/Q/H) (upstream) and HCSAGIQ (downstream)). Single PCR products were isolated and ligated into a pCRII-TOPO Vector (Invitrogen). Three hundred and ten clones were sequenced, and fragments containing PTP-like domains were identified. These 310 cDNA-fragments encoded 16 different PTPs in the mouse superior colliculus at E15. Novel PTP sequences were not found. From a total of 310 clones 50 clones (16%) coded for seven different intracellular PTPs, whereas 260 clones (84%) coded for nine different RPTPs. The relative abundance of different intracellular PTPs and RPTPs among the RT-PCR-generated cDNAs is shown in Table 1. The four most abundant cDNAs were RPTPσ (158 clones), RPTP-LAR (62 clones), TC-PTP (23 clones) and PTP1B (17 clones). The fifth most abundant cDNAs (12 clones) encoded RPTPj. Nine cDNAs encoded RPTPRR and RPTPε, while five cDNAs encoded RPTPδ, RPTPζ and PTP-MEG2 were represented by three clones each. For the intracellular protein tyrosine phosphatases PTP36, PTP1C and PTP1D each two clones were identified. RPTPκ, RPTPγ and PTP-PEST were represented by only one single clone each.

Quantitative real-time RT-PCR and Western blotting revealed differential regulation of PTP mRNAs and proteins during development of mouse superior colliculus

Since antibodies for all PTPs, which were identified with degenerate primers in the E15 mouse superior colliculus, were not available in this study, we focused on the 11 PTPs: five intracellular PTPs (TC-PTP, PTP1C, PTP1D, PTP-MEG2 and PTP-PEST) and six receptor-type PTPs (RPTPj, RPTPPR and RPTPε). To study the role of these PTPs in the developing mouse superior colliculus in detail, the regulation of the corresponding mRNAs and proteins was first investigated by quantitative real-time RT-PCR and Western blotting at different developmental time points (E13, E15, E18, P0, P4, P8, P12, P16 and P20). The embryonic and postnatal stages from E13 to P12 cover different developmental processes including genesis of neuronal and glial cells, formation of synapses and retinocollicular topographic mapping. Sequences of primer pairs used in real-time RT-PCR experiments are shown in Table 1 (Supplemental data).

The results of the quantitative real-time RT-PCRs revealed equal amounts of mRNAs (C<sub>T</sub> values = mean threshold cycle) for the reference genes β-actin and cyclophilin during collicular development (Supplemental data, Table 2). At E13, the C<sub>T</sub> values for the investigated PTPs varied between 22 and 29, indicating different expression levels of the PTPs at this early developmental stage (Supplemental data, Table 2). With ongoing development of the superior colliculus, the PTPs showed differential mRNA patterns when compared to the expression level at E13 (set to 0) (Fig. 1a, b, Supplemental data, Table 2).

In comparison to E13, the expression of TC-PTP mRNA decreased slightly during collicular development and reached a significant (P < 0.05) 3(2<sup>1.5</sup>)-fold downregulation
at P20 (Fig. 1a). In Western Blot analysis, TC-PTP antibody recognized three protein bands at 45-, 48- and ~50 kDa (Fig. 2a). The 48 and 45 kDa bands correspond to the previously described two TC-PTP human isoforms (Tiganis et al. 1998, 1999; Lam et al. 2001). The 50 kDa band corresponds probably to the intracellular protein tyrosine phosphatase PTP1B, which shows more than 70% homology to TC-PTP (Ibarra-Sanchez et al. 2000; Bourdeau et al. 2005). During development of the superior colliculus only the 48 kDa TC-PTP isoform revealed almost the same regulation pattern as TC-PTP mRNA, whereas the 45 kDa isoform was uniformly expressed. Strong signals of the 50 kDa protein, corresponding probably to PTP1B, were detected at E13 and E15 before dropping between E18 and P16. At P20 a faint 50 kDa band was detected. However, it has to be mentioned that primer pairs for TC-PTP amplification in real-time RT-PCR experiments did not amplify PTP1B.

When compared with E13, the quantification of PTP1C-mRNA using real-time RT-PCR revealed a slight downregulation at E15 (P < 0.05) and E18 (Fig. 1a). At P0 the PTP1C mRNA expression returned to the level found at E13. From P4 to P20, a slight upregulation of the gene was observed (statistically not significant). A similar expression pattern was found at protein level, as revealed in the immunoblot with specific anti-PTP1C antibody, which recognized a single protein band at 67 kDa (Fig. 2b). This band represents the full PTP1C protein as calculated from the amino acid sequence.

The quantification of PTP1D mRNA using real-time RT-PCR revealed a 2(2^1)-4(2^2)-fold decrease in the expression at the late embryonic stage E18 (P < 0.001) and the early postnatal stages P0 and P4 (P < 0.01) when compared to E13 (Fig. 1a). From P12 to P20, PTP1D mRNA expression returned to the level found at E13. Western Blot analysis revealed a similar expression pattern (Fig. 2c). Using a specific PTP1D-antibody, a single 68 kDa band corresponding to the predicted molecular weight of PTP1D was detected. A large amount of 68 kDa protein was found at E13 before dropping between E15 and P8. The signal intensity of 68 kDa protein increased in the later postnatal stages (from P12 to P20).

The MEG2 mRNA and protein revealed constant expression levels from E13 to P20 (Fig. 1a). In Western Blot analysis, a specific PTP-MEG2 antibody recognized a single protein band at 67 kDa (Fig. 2d), corresponding to the predicted size.

In contrast to other intracellular PTPs, the PTP-PEST mRNA expression was characterized through a continuous 2(2^1)-5(2^2.5)-fold decrease in superior colliculus development (P < 0.05 at E15, P4 and P12; P < 0.001 at E18, P0 and P20, Fig. 1a). Note that the regulation of the PTP-PEST protein, which was detected at ~120 kDa using Western blot analyses (Fig. 2e), correlates with the mRNA expression as detected by quantitative real-time RT-PCR. The predicted molecular weight of the murine PTP-PEST is 86 kDa. As revealed by other groups (Davidson et al. 1997; Sirosi et al. 2006; Halle et al. 2007) and as shown by our analyses, PTP-PEST migrates in the range of 120 kDa.

When compared with E13, the mRNA regulation of the receptor protein tyrosine phosphatase RPTPj was characterized through a 2(2^1)-4(2^2)-fold increase in the expression from late embryonic stage E18 until P20 (at E18 and P0, P < 0.05; at P4, P12 and P20, P < 0.01) (Fig. 1b). Nearby the same expression pattern was found at the protein level, as revealed in the immunoblot with specific anti-RPTPj antibody, which recognized a 159 kDa band (Fig. 3a). Although the calculated molecular weight of RPTPj is 134 kDa (Kuramochi et al. 1996), our data indicate that during collicular development RPTPj might be glycosylated.

The quantification of RPTPκ mRNA using real-time RT-PCR revealed a 2(2^1)- to 4(2^2)-fold increase of the expression level from E15 to P20 (P < 0.05) in comparison with E13 (Fig. 1b). Western blot analysis using specific anti-RPTPκ antibody (Fig. 3b) resulted in the detection of four bands of 164, 115, 105 and 91 kDa. The 164 kDa band represents the full RPTPκ protein (Jiang et al. 1993), while the 115-, 105- and 91 kDa bands correspond to the proteolytic cleavage products of RPTPκ (Anders et al. 2006). Weak signals of the full RPTPκ protein (164 kDa) were detected at E13 and E15 before being gradually upregulated between E18 and P20. At E13, the 115-, 105- and 91 kDa proteins were weakly expressed. A strong upregulation of the 115 kDa protein was detected between E15 and P8 prior to its gradual downregulation between P12 and P20. The 105 kDa protein band was weakly but nearby constantly present during collicular development. In contrast, the 91 kDa protein was prominently detectable at E15 and E18 and remained constantly present during postnatal collicular development until P20. Taken together, only the complete RPTPκ protein (164 kDa) and the 91 kDa protein revealed the same regulation pattern as RPTPκ mRNA.

When compared with E13, the RPTPγ mRNA revealed a significant increase 2(2^1)-fold to 3(2^1.5) of the expression level from E15 to P4 (P < 0.05). At P20 the expression returned to the level found at E13 (Fig. 1b). In Western blot analysis, the anti-RPTPγ antibody recognized five bands of 161, 120, 114, 110 and 80 kDa (Fig. 3c). The 161 kDa band corresponds to the predicted size deduced from the amino acid sequences (Barnea et al. 1993; Shintani and Marunaka 1996). The 120, 114, 110 and 80 kDa bands might represent the cleaved products of RPTPγ (van Niekerk and Poels 1999). The full RPTPγ protein (161 kDa) showed the same regulation pattern as RPTPγ mRNA. The large amount of 120 kDa protein was first detected in later postnatal stages.
The regulation pattern of RPTPε was detected at the protein level (Fig. 3d). The RPTPε antibody (Moller et al. 1995) recognized a 160 kDa band that became gradually upregulated with the maturation of the superior colliculus. The observed 160 kDa band corresponds to the highly glycosylated form of RPTPε, which has been reported in the mouse brain (Elson and Ledor 1995).

Since several transmembrane and cytoplasmatic isoforms of the RPTPRR gene are known (Augustine et al. 2000a; Chirivi et al. 2004), primer pairs which span the sequence between the extracellular and intracellular part of the transmembrane RPTPBr7 isoform were used in the quantitative real-time-PCR (Horvat-Brocker et al. 2008). When compared with E13, a significant 2(2^1)-fold upregulation of RPTPBR7 mRNA was found at P0 and P4 (P < 0.05), while in other embryonic and postnatal stages the expression level was almost the same as at E13 (Fig. 1b). The anti-RPTPBR7 antibody, which was raised against the intracellular domain (Horvat-Brocker et al. 2008), recognized in Western blot analyses (Figs. 2b–e, 3a–f) were applied to sagittal sections of the mouse superior colliculus. The specificity of the immunohistochemical signals was determined in control experiments using species-specific secondary CY3-labeled antibodies. In these control experiments, no signals were detected (data not shown). With the exception of RPTPBR7, that displayed a decreasing anterior–posterior protein gradient (insert, Fig. 5a), the examination of the superior colliculus in both anteroposterior and dorsoventral orientation revealed no significant protein expression gradients of the investigated PTPs (Figs. 4a–d, 5a–f).

At E13, immunolabeling using antibodies for the intracellular PTPs PTP1C, PTP1D, PTP-MEG2, PTP-PEST, RPTP1, RPTPκ, RPTPγ, RPTPe, RPTPrr and RPTPr during development of the mouse superior colliculus, immunohistochemical analyses were performed at the early embryonic stage E13 and the late postnatal stage P20 (Figs. 4a–d, 5b–f).

To determine the spatial and temporal localization of PTP1C, PTP1D, PTP-MEG2, PTP-PEST, RPTP1, RPTPκ, RPTPγ, RPTPe, RPTPrr and RPTPr during development of the mouse superior colliculus, immunohistochemical analyses were performed at the early embryonic stage E13 and the late postnatal stage P20 (Figs. 4a–d, 5a–f). Due to the cross-reactivity of the anti-TC-PTP antibody, which also recognized PTP1B protein (Fig. 2a), no immunohistochemical analysis was performed for this PTP. For the other PTPs, appropriate antibodies already used in Western blot analyses (Figs. 2b–e, 3a–f) were applied to sagittal sections of the mouse superior colliculus. The specificity of the immunohistochemical signals was determined in control experiments using species-specific secondary CY3-labeled antibodies. In these control experiments, no signals were detected (data not shown). With the exception of RPTPBr7, that displayed a decreasing anterior–posterior protein gradient (insert, Fig. 5a), the examination of the superior colliculus in both anteroposterior and dorsoventral orientation revealed no significant protein expression gradients of the investigated PTPs (Figs. 4a–d, 5a–f).
Fig. 2 Temporal regulation of intracellular PTP proteins as revealed by Western blot analysis in the developing mouse superior colliculus. Protein lysates of superior colliculus tissues from E13, E15, E18, P0, P4, P8, P12, P16 and P20 were separated by SDS-PAGE, and immunoblotted with TC-PTP (a), PTP1C (b), PTP1D (c), PTP-MEG2 (d) and PTP-PEST (e) antibodies, respectively. Molecular weights of intracellular PTP proteins are indicated on the right. All blots were reprobed for β-actin to demonstrate the equal protein loading. Changes in the relative protein expression (normalized protein level) of each PTP protein are shown in the graphical representations. In the case of PTP1C, PTP1D, PTP-MEG2 and PTP-PEST each PTP antibody recognized only one band, which corresponded to the predicted molecular weights of these PTPs. However, the TC-PTP antibody recognized three protein bands at 45, 48 and ~50 kDa (Fig. 2a). The 48 kDa and 45 kDa bands correspond to the previously described two TC-PTP human isoforms, whereas the 50 kDa band corresponds most probably to the intracellular protein tyrosine phosphatase PTP1B. Note that proteins of the investigated intracellular PTPs are differentially regulated during superior collicular development.
addition, all intracellular PTPs displayed stronger immunoreactivities in the ventricular zone (VZ) (Fig. 4a–d). At P20, PTP1C-, PTP1D- and PTP-MEG2-immunoreactive cells were found dispersed throughout the stratum griseum superficiale (SuG) and the stratum opticum (Op) of the superior colliculus (Fig. 4a, b and c). In contrast, no PTP-PEST immunoreactivity was detected at P20 (Fig. 4d). Here, rabbit non-immune serum labels only the pial surface non-specifically.

In comparison to the intracellular PTPs, the receptor-type PTPs RPTPJ and RPTPκ revealed unique immunostainings at E13. The RPTPJ-antibody strongly labeled cells in the ventricular zone (VZ) and subventricular zone (SVZ) of the embryonic superior colliculus (Fig. 5a), whereas the RPTPκ epitope was located on radial processes that span the whole tissue (Fig. 5b). The most prominent RPTPγ- (Fig. 5c), RPTPε- (Fig. 5d) and RPTPσ-immunoreactivities (Fig. 5f) were found in the intermediate zone (IZ) of the E13 superior colliculus. In comparison to the other RPTPs, RPTPRR immunoreactivity was found uniformly distributed throughout the E13 superior colliculus (Fig. 5e). At P20, single RPTPγ- (Fig. 5a), RPTPγ- (Fig. 5c) and RPTPσ-immunoreactive cells (Fig. 5f) seemed to be restricted to the stratum griseum superficiale (SuG) and stratum opticum (Op), whereas immunofluorescence signals of RPTPκ (Fig. 5b), RPTPε (Fig. 5d) and RPTPRR (Fig. 5e) revealed a more dispersed staining pattern within all visual layers of the P20 superior colliculus.

Cell-type specific expression of PTPs in collicular neural progenitors and radial glia cells at E13

Based on our observations of a prominent PTP expression in the early embryonic superior colliculus at E13, a period of intense neurogenesis and gliogenesis, we hypothesized a possible involvement of PTP proteins in the regulation of early collicular neural progenitors and radial glia cells. To determine whether PTPs are expressed by these specific cell types, we performed double-immunohistochemistry on E13 superior colliculus slides using antibodies against PTPs and two different markers of early neuronal precursor cells, nestin, and a subtype of radial glia cells, RC-2 (Figs. 6–9). Within the superior colliculus, both nestin- and RC-2 antibodies strongly labeled radially oriented processes, which span the whole tissue. The most prominent immunoreactivities for nestin as well as for RC-2 positive cells/processes were found in close association with the subventricular and ventricular zones. In comparison to the observed nestin-immunoreactivity, decreased label intensity from the subventricular/ventricular zone (SVZ/VZ) to the intermediate zone (IZ) was found for RC-2.

Double-immunofluorescence analysis revealed that the immunoreactivity for the intracellular PTPs PTP1D, PTP-MEG2 and PTP-PEST colocalized partially with the precursor cell marker nestin (Fig. 6) and the radial glia cell marker RC-2 (Fig. 8). The most prominent signals were observed in the VZ and SVZ. PTP1C immunoreactivity, which was found most prominently in the outer edge of the intermediate zone and in single cells of the ventricular zone, showed reduced co-localization with RC-2 positive cells in the ventricular zone and did not co-localize with minor labeled cells in the intermediate zone. Considering the species of the nestin and PTP1C antibodies (both mouse, IgG) it was not possible to determine the co-localization of the PTP1C protein with nestin-positive precursor cells. We speculate that the overlap should be minor, if any. This interpretation is in accordance with the PTP1C-immunoreactivity, which was mostly found in more mature cells of the IZ.

In addition, no co-localization with nestin-positive progenitor cells (Fig. 7) and RC-2 positive radial glia cells (Fig. 9) was observed for the receptor-type PTP RPTPJ. In contrast, the RPTPκ epitope was strongly co-expressed by nestin- and by RC-2-positive cells, as revealed by a nearby completely overlapping expression pattern (Figs. 7, 9). RPTPγ, RPTPε, RPTPRR and RPTPσ were partially expressed in nestin- and RC-2- double-positive cells/processes (Figs. 7, 9).

Discussion

Our study of the PTP family highlights the identification and differential expression pattern of eleven PTPs during critical periods of mouse superior colliculus histogenesis. Quantitative real-time PCR and Western blotting of TC-PTP, PTP1C, PTP1D, PTP-MEG2, PTP-PEST, RPTPJ, RPTPκ, RPTPγ, RPTPε, RPTPRR and RPTPσ revealed distinct temporal changes of gene and protein expression during collicular development. Moreover, double-immunohistochemical analysis demonstrated for the first time that the intracellular PTPs PTP1D, PTP-MEG2 and PTP-PEST and the receptor-type PTPs RPTPκ, RPTPγ, RPTPε, RPTPRR and RPTPσ are co-expressed in a diverse pattern by collicular neural progenitors and radial glia cells at E13. In the light of these particular regulatory patterns and drawing on reports relating to other systems, we propose that the tyrosine phosphatases play an important regulatory role for the neural stem/progenitor cells in the superior colliculus.

Using a degenerate primer-based RT-PCR approach we identified TC-PTP and PTP1B as the most abundant intracellular PTPs. As reported by previous studies, TC-PTP shares 71% amino-acid identity with PTP1B.
Proteolytic cleavage or alternative splicing can result in differential regulation at the level of posttranslational modification (Bourdeau et al. 2005; Ibarra-Sanchez et al. 2000). Based on the cross-reactivity of the TC-PTP antibody used in Western blotting, PTP1B was identified as a 50 kDa protein band. PTP1B has been detected in RGCs where it may function as a positive modulator of nerve fiber outgrowth (Pathre et al. 2001). Therefore, it is conceivable that PTP1B could play a crucial role for the establishment of topographic retino-collicular projections. Consistent with this view, PTP1B protein exhibited the most prominent expression at early stages (E13 and E15), when the first RGC fibers begin to grow out and innervate the superior colliculus. Different from the case of PTP1B, the expression of TC-PTP is described here for the first time for the murine CNS, where two alternatively spliced isoforms were visualized by Western blot. The constitutively expressed 48 kDa TC-PTP isoform was gradually downregulated during development, whereas the 45 kDa isoform remained constant. Previously, two alternatively spliced variants have been described in the human: a 48 kDa form, which is localized to the endoplasmatic reticulum by a hydrophobic C-terminus, and a 45 kDa form which lacks the hydrophobic C-terminus and is located in the nucleus (Tiganis et al. 1998, 1999; Lam et al. 2001). Earlier investigations reported a regulatory involvement of TC-PTP in EGF- and PDGF-signal transduction pathways (Tiganis et al. 1997; Persson et al. 2004; Mattila et al. 2005), which are implicated in cell proliferation and cell adhesion processes. Furthermore, TC-PTP has been described as a negative regulator of colony-stimulating factor 1 signaling and macrophage differentiation (Simoncic et al. 2006). Consistent with these reports, our data suggest different potential roles of TC-PTP during collicular development, especially in early developmental stages.

PTP1C mRNAs and proteins were weakly expressed during the embryonic stages and at the day of birth, while a strong expression was detected from P4 onward. In addition, immunohistochemistry revealed a prominent expression of PTP1C in the intermediate zone (IZ) of the superior colliculus, which consists of more mature cells at E13. Furthermore, PTP1C was detected in cells scattered throughout the SuG and Op of the superior colliculus at P20. As shown by Jena et al. (1997), PTP1C is associated with synaptic vesicles and interacts directly with the vesicular protein synaptophysin, which suggests an involvement of PTP1C in synaptic transmission. Indeed, the expression profile of PTP1C in the superior colliculus correlates well with synaptogenesis, which starts approximately at P1 (Lo and Mize 1999). Furthermore, synaptophysin is up-regulated during neuronal development and maturation (Becher et al. 1999a, b). In addition, previous studies reported that PTP1C is involved in proliferation and differentiation of astrocytes, microglia (Horvat et al. 2001; Wishcamper et al. 2001) and oligodendrocytes (Massa et al. 2004). These cell types are generated at late embryonic and early postnatal stages.

The peak amount of PTP1D mRNA and protein was detected at E13 and at late postnatal stages. The strong expression of PTP1D at E13, the sharp drop at late embryonic and early postnatal stages and the subsequent increased expression at late developmental stages are in agreement with the assumption that PTP1D could adopt different functions during the early and late development of the superior colliculus. PTP1D may affect cell growth, cell adhesion and cell motility either as a positive or a negative regulator of distinct signal transduction pathways (Li et al. 1994; Matozaki et al. 1994; Noguchi et al. 1994; Miao et al. 2000; Poliakov et al. 2004; Zhang et al. 2004). In addition, PTP1D directly interacts with EphA2 and influences directional cell migration. In this context, the impact of the Eph family in patterning and establishing the topographic mapping within the visual system is well described (Flanagan and Vanderhaeghen 1998; Feldheim et al. 2000, 2004). Furthermore, we could show that PTP1D is expressed in RC-2 positive radial glia cells during early collicular development, which could hint at a potential functional role of this PTP in neuronal migration processes.

PTP-MEG2 mRNA and protein were continuously expressed from E13 to P20, indicating its important role during development as well as in the maintenance of a
mature superior colliculus. A strong PTP-MEG2 immunoreactivity was detected throughout all layers of E13 superior colliculus, whereas at P20 the protein was restricted to single cells of the SuG and Op. Interestingly, PTP-MEG2 deficient mice display several severe neurodevelopmental deficits such as neural tube defects associated with craniofacial abnormalities, exencephaly, encephalocoeles and meningomyelocoeles. In addition, the brains of neonatal PTP-MEG2 null-mice are grossly smaller in comparison to wild type animals (Wang et al. 2005). Extrapolating from the severe developmental defects that were observed in the null-mice, PTP-MEG2 might be involved in a diversity of signal transduction pathways that regulate neurulation processes. Consistent

Fig. 4 Spatiotemporal expression pattern of PTP1C (a), PTP1D (b), PTP-MEG2 (c) and PTP-PEST (d) in the embryonic (E13) and postnatal (P20) mouse superior colliculus. Sagittal sections through the embryonic (E13) and late postnatal (P20) superior colliculus were labeled with appropriate intracellular PTP antibodies. Dorsal and posterior orientation is schematically presented. a–d At E13, immunofluorescent staining using antibodies against PTP1C, PTP1D, PTP-MEG2 and PTP-PEST reveals a more uniform distribution throughout the whole superior colliculus. (b–d) The strongest signals for PTP1C, PTP1D and PTP-PEST were obtained in the outer dorsal edge of the intermediate zone (IZ). a–d In addition, a prominent immunoreactivity for all intracellular PTPs was found in the ventricular zone (VZ) of the embryonic superior colliculus. a–e At P20, PTP1C-, PTP1D- and PTP-MEG2-immunoreactivities were found in single cells scattered throughout the stratum griseum superficiale (SuG) and the stratum opticum (Op) of the superior colliculus. d In contrast, no immunoreactivity was found for PTP-PEST at P20. Here, rabbit non-immune serum labels only the pial surface non-specifically. IZ intermediate zone, VZ/SVZ ventricular zone/subventricular zone, ZO stratum zonale, SuG stratum griseum superficiale, Op stratum opticum. Scale bar 50 μm

Fig. 5 Spatiotemporal expression pattern of RPTPJ (a), RPTPκ (b), RPTPγ (c), RPTPε (d), RPTPRR (e) and RPTPr (f) in the embryonic (E13) and mature (P20) mouse superior colliculus. Sagittal sections through the embryonic (E13) and late postnatal superior colliculus (P20) were labeled with appropriate RPTP-antibodies. Dorsal and posterior orientation is schematically presented. a At E13, a polyclonal antibody against RPTPJ detects single immunoreactive cells in the subventricular zone (SVZ) of the embryonic superior colliculus. In addition we found RPTPJ expressing cells in a decreasing anterior–posterior gradient (insert, Fig. 5a). Insert: lower magnification of a RPTPγ stained sagittal section of the E13 superior colliculus (scale bar = 200 μm). In the P20 superior colliculus, single cells, scattered throughout the stratum griseum superficiale (SuG) and stratum opticum (SO), were labeled. b The RPTPκ epitope is located on radial processes in the embryonic superior colliculus, whereas in the P20 superior colliculus a broad RPTPκ-immunoreactivity was found within all visual layers. (c, d and f) The strongest RPTPγ-, RPTPε- and RPTPπ-immunoreactivities were found in the dorsal part of the E13 superior colliculus (IZ intermediate zone). At P20, RPTPγ and RPTPπ immunoreactive cells were restricted to the stratum griseum superficiale (SuG) and stratum opticum (SO), whereas RPTPκ immunoreactivity displays a more dispersed distribution pattern. e Immunolabeling using a polyclonal RPTPRR antibody reveals a weak and uniform staining pattern in the embryonic as well as in the P20 superior colliculus. IZ intermediate zone, VZ/SVZ ventricular zone/subventricular zone, ZO stratum zonale, SuG stratum griseum superficiale, Op stratum opticum. Scale bar 50 μm
with this view, PTP-MEG2 controls the development and growth of erythrocyte precursors cells (Xu et al. 2003). A potential neuro-developmental role of PTP-MEG2 is supported by its expression in nestin-positive radial glia cells and neuronal collicular progenitors. Therefore, we believe that this PTP might act as a molecule which impacts on the differentiation pathway of these specialized cell types.

In contrast to other intracellular PTPs, PTP-PEST mRNA and protein levels were dramatically downregulated from P0 onward. Immunohistochemical analysis revealed a prominent expression of this PTP in early collicular progenitor- and radial glia cells, while it was absent at P20. These findings suggest that PTP-PEST might be involved in the regulation of early developmental processes, such as cell proliferation, differentiation and migration. Indeed, similar to PTP1D, PTP-PEST acts as a negative regulator of cell migration processes (Davidson et al. 1997; Garton et al. 1997; Shen et al. 1998; Garton and Tonks 1999; Davidson and Veillette 2001).

Fig. 6 Co-localization of PTP1D, PTP-MEG2 and PTP-PEST proteins in nestin-positive neural progenitor cells in the E13 mouse superior colliculus. Sagittal sections through the embryonic E13 superior colliculus were double-labeled with antibodies against PTPs and the neural progenitor marker nestin. Arrowheads indicate examples of double-labeled cells. PTP1D, PTP-MEG2 and PTP-PEST epitopes partially co-localize with nestin-positive neural progenitor cells. Because the nestin- and PTP1C antibodies originated from the same species (both mouse, IgG), it was not possible to analyse the co-localization of the PTP1C protein with nestin-positive precursor cells. Note that the antibodies directed at intracellular PTPs label mainly nuclei/perikarya of collicular cells; both markers reveal an incomplete overlapping expression pattern (merge). IZ intermediate zone, VZ/SVZ ventricular zone/subventricular zone. Scale bar 50 μm

Fig. 7 Co-localization of RPTPJ, RPTPx, RPTPy, RPTPz, RPTPRR and RPTPr proteins in nestin-positive neural progenitor cells in the E13 mouse superior colliculus. Sagittal sections through the embryonic E13 superior colliculus were double-labeled with antibodies against RPTPs and the neural progenitor marker nestin. Arrowheads indicate examples of double-labeled cells. Note that RPTPy, RPTPz, RPTPRR and RPTPr epitopes partially co-localize with nestin-positive progenitors. In contrast, RPTPJ was not co-localized with nestin-positive cells. Since nestin produces strictly radial staining whereas RPTPJ, RPTPy, RPTPx, RPTPRR and RPTPr antibodies label mainly nuclei/perikarya of collicular cells, both markers reveal a faint overlapping expression pattern (merge). As shown for RC-2 immunostainings, RPTPz immunoreactivity directly overlaps with nestin-intermediate filament staining (merge). In comparison to the RC-2/RPTPz overlap, the correspondence of nestin and RPTPz appears less extensive. IZ intermediate zone, VZ/SVZ ventricular zone/subventricular zone. Inserts: Higher magnification of immunopositive cells (merge figures represent co-immunoreactive cells). Scale bar 50 μm
The receptor PTPs RPTPRR and RPTP$\sigma$, RPTPJ RPTPK, RPTP$\gamma$ and RPTPe displayed a constant up-regulation of mRNAs and proteins during collicular development, suggesting an important role both in developmental processes and in homeostasis of the mature superior colliculus.

A strong expression of the RPTPJ-epitope was restricted to single cells within the subventricular and ventricular zones (SZV/VZ) of the E13 superior colliculus. These areas contain predominantly immature proliferative and early postmitotic cells. Interestingly, in C. elegans RPTPJ regulates signaling of the EGF-receptor and enables Notch-dependent determination and differentiation of certain cell types (Berset et al. 2005); in the human and mouse, RPTPJ exhibits tumor-suppressor activity (Keane et al. 1996; Trapasso et al. 2000; Ruivenkamp et al. 2002). Thus, we assume that the expression of RPTPJ in the embryonic superior colliculus might be necessary to promote neurogenic/gliogenic determination and differentiation of specific collicular progenitors. In fact, we found no co-localization of RPTPJ with early immature neural progenitors or radial glia cells, which supports the notion of a potential role in cell fate determination and specification. In addition, RPTPJ was expressed in a decreasing anterior-posterior gradient-like pattern, which suggests that the determination of specific cell types starts in the most anterior part during superior colliculus development. This gradient like pattern was restricted to the early E13 superior colliculus and was not observed in other CNS structures.

In contrast to other PTPs, RPTPK displayed a completely overlapping expression with nestin-positive progenitor cells and RC-2-positive radial glial cell processes in the E13 superior colliculus. Recently, we showed that RPTPK is expressed on radial processes in the embryonic retina (Horvat-Brocker et al. 2008) and co-localizes with nestin-positive retinal progenitor cells. Because radial glia processes serve as cellular scaffold during CNS development and function as guide for neuronal migration (Newman and Reichenbach 1996; Bauch et al. 1998), the expression on radial processes might implicate RPTPK in the process of neuronal migration within the developing superior colliculus. In support of this view, it has been shown that RPTPK stimulates the migration of cancer cells (Kim et al. 2006) and acts as cell adhesion molecule with homophilic adhesion properties (Jiang et al. 1993; Sap et al. 1994). Western blot analysis for RPTPK demonstrated a complex temporal regulation of the full RPTPK protein (164 kDa) and the proteolytic cleavage products (115-, 105- and 91 kDa). Thus, we surmise that the integral RPTPK protein and its isoforms are functionally relevant during mouse collicular development.

As shown for RPTPe and RPTP$\sigma$, RPTPK exhibits a predominant expression in the intermediate zone (IZ) of the embryonic mouse superior colliculus. At E13, this layer consists of more mature post-mitotic cells, which are generated in an inside-out sequence (Altman and Bayer 1981; Edwards et al. 1986). Indeed, as revealed by previous studies in the rat, RPTPK mRNA is most prominently expressed by post-mitotic neurons in the superficial layers of the postnatal cortex (Sahin et al. 1995). However, our data additionally indicate that the RPTPK protein is expressed by a subset of neural progenitor and radial glia cells within the ventricular zone. With regard to the described prominent expression of RPTPK in post-mitotic cells, real-time PCR and Western Blot analysis revealed an up-regulation of the RPTPK mRNA and RPTPK proteolytic cleavage products (120-, 114-, 110- and
80 kDa) during collicular development. Nevertheless, considering our observation that RPTPγ is subject to marked proteolytic cleavage, we suppose a diversity of functions of RPTPγ during collicular development. For example, RPTPγ is involved in the differentiation of oligodendrocytes (Runjan and Hudson 1996; Fraser et al. 2006) and inhibits NGF-induced neurite outgrowth of PC12D cells (Shintani et al. 2001).

Our analyses of RPTPe revealed a minor mRNA- and protein-expression within the embryonic superior colliculus, which was followed by a gradual up-regulation of mRNA and protein during collicular maturation. In some systems, RPTPe can downregulate mitogenic signaling by inhibiting MAP-kinase activity (Wabakken et al. 2002; Toledano-Katchalski et al. 2003) and JAK-STAT-signaling in M1 leukemia cells (Tanuma et al. 2000, 2001, 2003). Furthermore, RPTPe negatively regulates proliferation of endothelial cells (Thompson et al. 2001). Based on these studies, we assume that the faint expression of RPTPe is required to permit proliferation of early collicular cells.

RPTPe was widely expressed with ongoing maturation, while at early embryonic stages it was mainly confined to the superficial layer of the superior colliculus. In fact, studies of the physiological roles of RPTPe in the brain discuss the biological significance of this PTP in neuronal differentiation processes (Mukouyama et al. 1997). This assumption is in accordance with the observed minor co-expression of RPTPe in immature, partially proliferative, nestin- and RC-2-positive precursor/radial glia cells within the ventricular zone.

Previous studies pointed out that several transmembrane and cytoplasmatic isoforms of the RPTPRR gene are known (Augustine et al. 2000a; Chirivi et al. 2004). In the developing superior colliculus, our Western blot results indicate the differential expression of three alternative splicing variants (RPTPBr7 (74 kDa), RPTPPBSδ (51 kDa) and RPTPPBSδ (47 kDa)) of the RPTPRR gene. In addition, we identified a 155 kDa protein band, which might represent a glycosylated form of one of the described receptor-like isoforms. With the exception of RPTPPBSδ ( + ), all RPTPRR proteins were found prominently expressed in early embryonic collicular stages. A similar developmental regulation of the RPTPRR gene and the corresponding alternatively spliced variants has been reported in the retina (Horvat-Brocker et al. 2008), in Purkinje cells (van den Maagdenberg et al. 1999) and during chondrogenesis (Augustine et al. 2000a). Considering the generation and developmentally distinct regulation of multiple isoforms from a single gene through alternative splicing, it seems plausible that the identified RPTPRR proteins regulate various biological and physiological processes at different developmental time points.
In contrast to the other RPTPs, RPTPσ mRNA was downregulated in the late postnatal stages (at P12 and at P20), correlating well with the completion of the retino-collicular map. On protein level, we observed a nearby constant expression during development. In RGC axons of Xenopus and chicken, RPTPγ promotes retinal neurite outgrowth (Burden-Gulley and Brady-Kalnay 1999; Ledig et al. 1999; Johnson et al. 2001), growth cone guidance (Burden-Gulley et al. 2002) and targeting of retinal axons within the optic tectum (Rashid-Doubell et al. 2002). We assume an analogous functional role of RPTPσ in the ventricular and subventricular zones of the developing CNS (Walton et al. 1993; Meathrel et al. 2002), where RPTPσ plays an important role in neural stem cell differentiation. Accordingly, the expression of RPTPs in early collicular progenitors and radial glia cells is suggestive of a functional relevance of this PTP in stem cell development.

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