Protein Tyrosine Phosphatase α (PTPα) and Contactin Form a Novel Neuronal Receptor Complex Linked to the Intracellular Tyrosine Kinase fyn

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Abstract. Glycosyl phosphatidylinositol (GPI)–linked receptors and receptor protein tyrosine phosphatases (RPTPs), both play key roles in nervous system development, although the molecular mechanisms are largely unknown. Despite lacking a transmembrane domain, GPI receptors can recruit intracellular src family tyrosine kinases to receptor complexes. Few ligands for the extracellular regions of RPTPs are known, relegating most to the status of orphan receptors. We demonstrate that PTPα, an RPTP that dephosphorylates and activates src family kinases, forms a novel membrane-spanning complex with the neuronal GPI-anchored receptor contactin. PTPα and contactin associate in a lateral (cis) complex mediated through the extracellular region of PTPα. This complex is stable to isolation from brain lysates or transfected cells through immunoprecipitation and to antibody-induced coclustering of PTPα and contactin within cells. This is the first demonstration of a receptor PTP in a cis configuration with another cell surface receptor, suggesting an additional mode for regulation of a PTP. The transmembrane and catalytic nature of PTPα indicate that it likely forms the transducing element of the complex, and we postulate that the role of contactin is to assemble a phosphorylation-competent system at the cell surface, conferring a dynamic signal transduction capability to the recognition element.

Key words: PTPα • tyrosine phosphatase • glycosyl phosphatidylinositol • neural signal transduction

The neural cell adhesion molecule contactin (F3/F11) contains Ig-like domains and FN-III repeats, and is attached to the cell surface through a glycosyl phosphatidylinositol (GPI) anchor (Ranscht and Dours, 1988; Brümmendorf et al., 1989; Gennarini et al., 1989; Zisch et al., 1992). This mode of membrane attachment precludes direct transfer of signals into the cell, but allows high mobility in the plane of the membrane, and as such ideally suits contactin for responding to macromolecular stimuli presented to the cell (Vaughan, 1996). The modulation of signal transduction would critically depend on cis association with transmembrane components. Lateral cis interactions occur with the Ig superfamily transmembrane receptors Ng-CAM and Nr-CAM (Brümmendorf et al., 1993; Morales et al., 1993; Sakurai et al., 1997) and with Caspr (Peles et al., 1997). Intriguingly, contactin can be isolated in a complex with the intracellular src family tyrosine kinase fyn (Zisch et al., 1995). The mechanism of this association is unknown, but likely requires an intermediate membrane-spanning component.

As many signal transduction pathways are initiated through activation of intrinsic or associated receptor tyrosine kinase activity, the association of fyn with contactin may be central to contactin signaling. Several protein tyrosine phosphatases (PTPs) can activate src family kinases in various systems, suggesting a PTP to be a candidate component of a contactin signaling complex.

The transmembrane nature of the receptor-like PTPs (RPTPs) indicates that they transduce extracellular signals, although few such signals/ligands are known. The identification of extracellular ligands for RPTPs could provide critical insights into the specific functions and regulation of these enzymes. PTPα is one such RPTP for
which a ligand is sought. The short glycosylated extracellular region of PTPα, unlike many other RPTPs, lacks known adhesion motifs. It is linked via the transmembrane segment to the intracellular region that contains two homologous catalytic domains, both of which have intrinsic phosphatase activity (Wang and Pallen, 1991). Several properties of PTPα indicate that it may, among other things, be involved in neuronal signaling. PTPα is particularly abundant in the brain and implicated in mediating neuronal differentiation, cell proliferation, and transformation, by its ability to activate the tyrosine kinase src (Zheng et al., 1992; den Hertog et al., 1993). PTPα also associates with, dephosphorylates, and activates brain fyn (Bhandari et al., 1998), a src family kinase that plays a role in axonal growth, myelination, spatial learning and memory (Lowell and Soriano, 1996). Mice lacking PTPα have reduced activities of brain src and fyn, demonstrating that PTPα is a positive physiological regulator of these kinases (Ponniah et al., 1999; Su et al., 1999).

The shared and complementary features of PTPα and contactin suggest that they may form a signaling complex. Both proteins are expressed in some of the same neuronal cell types as migrating cerebellar granule cell neurons (Faivre-Sarrailh et al., 1992; Fang et al., 1996). At a molecular level, the physical association of PTPα and fyn raises the possibility that PTPα provides a transmembrane link between contactin and fyn. Furthermore, the ability of PTPα to activate fyn indicates that PTPα could thereby transduce a signal originating from the extracellular engagement of contactin. We demonstrate that PTPα and contactin associate and define certain requirements for interaction. These findings indicate that contactin is a novel extracellular partner of PTPα and that this complex may regulate aspects of neuronal development.

**Materials and Methods**

**Expression Plasmids**

Numbering of the human PTPα amino acid sequence is according to Krueger et al. (1990). The pX) 41 vectors expressing PTPα, VSVG-G-tagged PTPα, fyn, and CD-45 have been described (Zheng et al., 1992; Bhandari et al., 1998). To construct pXJ41-contactin-neo, two contactin CDNA fragments comprising nucleotides 38-1103 and 177-3961 were obtained from pSP72-contactin/11 (Zisch et al., 1995) and subcloned in two steps into pXJ41-neo to reconstitute the contactin coding sequence. To construct pXJ41-myrt-PTPα-neo, the PCR-amplified region of human src CDNA encoding the NH2-terminal myristylated sequence MGSDKSKPKA DQA was cloned into pGEM-T (Promega Corp.), and the reverse orientation was selected with the multiple cloning NotI site at the 5′ end of the myristylation signal. A NotI-XhoI fragment was excised and cloned into pXJ41-neo, creating pXJ41-myrt-neo. A PCR fragment of PTPα encoding the entire intracellular region (amino acids 148-774), and flanked with engineered Sall sites, was inserted in-frame into XhoI-cut pXJ41-myrt-neo. To construct pXJ41-PTPα-neoCD45-neo encoding the chimeric RPTPα, a region of the VSVG-G-PTPα CDNA encoding the signal peptide and amino acids 1-121 of PTPα was amplified by PCR using forward and reverse primers with engineered NotI and BglII sites, respectively. This NotI-BglII fragment was inserted into a pXJ41-neo intermediate vector. A CD45 CDNA fragment encoding 35 extracellular juxtamembrane amino acids, the transmembrane, and the entire intracellular regions was released from pXJ41-CD45-Hy with BglII and inserted into the intermediate vector.

**Cell Culture and Transient Transfections**

COS-1 cells were maintained and transiently transfected as described (Bhandari et al., 1998). The empty expression plasmid pXJ 41-neo was used to normalize the amount of DNA in each transfection. A 24 h of coculture, the cells were harvested and processed as described below. For tunicamycin treatment, 20 μg/ml tunicamycin (Boehringer Mannheim) in DME/10% FCS was added to the cells 6 h after transfection, and the cells were harvested after a further 42 h of culture.

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CO-S7 cells cultured on 1 ml polylysine or alcan blue-coated glass 12-mm-diam coverslips were processed for immunocytochemistry 24-48 h after transfection. Images were taken using a Zeiss Axiolab 100M equipped with a Hamamatsu C5810 3 color CCD cooled camera. Images were processed directly with a A dobe Photoshop.

**Antibody-mediated Copatching of Contactin and PTPα in Transfected Cells**

Copatching of contactin or VSVG-PTPα was performed on COS-7 cells 24–48 h after transfection as described (Zisch et al., 1995). Cells were incubated on ice with 10 μg anti-contactin (4D1) or anti-VSVG antibodies. Bound antibodies were clustered by incubation with a 1:100 dilution of FITC- or RITC-labeled goat anti-mouse antibodies (Tago), followed by fixation with 4% paraformaldehyde and permeabilization (when appropriate) with 0.2% Triton X-100. A affinity-purified rabbit antibodies to contactin or PTPα (see below) were used complementary to the mAb bs, together with goat anti-rabbit antibodies labeled with FITC or RITC (Tago).

**Western Blots and Immunoprecipitation**

Six embryonic chick brains (15-d-old) were mechanically suspended in 30 ml of ice-cold PBS buffer, filtered through a nylon mesh, homogenized by 10 strokes in a glass Dounce homogenizer, and washed three times in PBS. Chick brain cells or transfected COS cells were lysed in 10 mM Tris-Cl, pH 8, 150 mM NaCl, 1 mM EDTA, 1% Brij-96, 20 μg/ml aprotrin, and 2 mM PMSE, and the lysates were clarified by centrifugation. A nti-bodies toward VSVG (Sigma Chemical Co.), contactin (4D1) (Zisch et al., 1995), CD45 or fyn (Transduction Laboratories), PTPα (antisemur 2205, raised against PTPα-D1), and NCA M (Chemicon International, Inc.) were used for immunoprecipitation and/or immunoblotting.

**Phosphatase Assays**

The phosphatase activity of 5 μl immunoprecipitate toward 2 μM phosphotyrosyl-RR-src peptide was measured at 30°C for 15 min as described (Lim et al., 1997).

**Results**

**PTPα and Contactin Are Associated in Chick Brain**

Contactin and PTPα immunoprecipitates from embryonic chick brain lysates were probed for the presence of contactin and PTPα. PTPα was present in anti-contactin immunoprecipitates (Fig. 1A), and contactin was present in PTPα immunoprecipitates, but was not precipitated by preimmune serum (Fig. 1B). Thus, PTPα and contactin exist in a complex in brain lysates. To check the specificity of the association of PTPα and contactin, we examined the interaction of PTPα with NCA M, another fyn-associated (Beggs et al., 1997) cell adhesion molecule highly expressed in the brain. The 120- and 140-kD NCA M isoforms were detected in anti-NCA M immunoprecipitates from mouse brains, but PTPα was not present (Fig. 1C). Likewise, NCA M could not be detected in anti-PTPα immunoprecipitates prepared from these lysates (Fig. 1D) using the same anti-PTPα antisemur as employed with the chick brain lysates (raised to the species conserved intracellular D1 region of PTPα).
Association of Ectopically Expressed PTPα and Contactin

To investigate the molecular basis of the association between PTPα and contactin, we used a transient expression system where the interaction of different forms could be manipulated. PTPα and contactin coimmunoprecipitated with one another from COS cells coexpressing PTPα (tagged in its extracellular region with an epitope of VSVG to facilitate immunoprecipitation [Bhandari et al., 1998]) and contactin (Fig. 2 A). In other experiments, the immunocomplexes were assayed for PTP activity (Fig. 2 B). Anti-VSVG immunoprecipitates from cells expressing PTPα alone or with contactin contained comparable levels of phosphatase activity, whereas virtually no activity was detectable in anti-VSVG immunoprecipitates from cells expressing contactin alone. Anti-contactin immunoprecipitates from coexpressing cells contained about a fivefold higher phosphatase activity than those from cells expressing contactin or PTPα alone. These results indicate that the contactin-PTPα complexes are functionally active.

The levels of PTPα protein and phosphatase activity were much lower in anti-contactin immunoprecipitates than in anti-VSVG immunoprecipitates from cells coexpressing contactin alone. Anti-contactin immunoprecipitates from coexpressing cells contained about a fivefold higher activity than those from cells expressing contactin or PTPα alone. These results indicate that the contactin-PTPα complexes are functionally active.

Similar experiments were carried out with contactin and CD45, a receptor-like PTP with structural similarity to PTPα. No coimmunoprecipitation of contactin and CD45 from coexpressing cells was detected (Fig. 2 C), indicating that the interaction of PTPα and contactin is specific and not merely due to heterologous expression.

Colocalization and Coclustering of Contactin and PTPα

In situ localization of contactin and PTPα in cotransfected COS cells revealed a similar distribution for both proteins within the plane of the plasma membrane (Fig. 3, A and B). In contrast, control transfections of contactin together with the RPTP CD45, gave a completely different pattern of distribution to that of contactin (data not shown), indicating that they do not associate in the same cellular complexes.

We examined whether an enforced redistribution of either contactin or PTPα, induced by incubating the live cells with antibodies to the extracellular domains of these molecules, would lead to coclustering of the respective partner. Clustering of PTPα (Fig. 3 E) caused contactin to redistribute to closely match the PTPα pattern (Fig. 3 F). The close similarity of these two patterns supports the efficacy of clustering via the free-standing VSVG tag on the NH₂-terminal of PTPα, leading to little or no interference with the interactions between PTPα and contactin. Clusters of contactin and PTPα could also be induced by 4D1 mAb specific for contactin (Fig. 3, C and D). Clusters of various sizes were induced in individual cells, with the PTPα localization largely matching the contactin pattern.

The Extracellular Region of PTPα Is Required for Association with Contactin

To identify the region of PTPα involved in the interaction with contactin, we generated a membrane-associated intracellular form of PTPα by replacing its extracellular and transmembrane regions with the myristylation signal of src (myr-PTPα). Expressed myr-PTPα was associated with the membrane fraction (data not shown), however, contactin only associated with wild-type PTPα and not with myr-PTPα (Fig. 4 A). Thus, contactin does not interact with the PTPα lacking the extracellular and transmembrane regions.

Since CD45 does not associate with contactin, we created a PTPα/CD45 hybrid molecule where most of the extracellular region of CD45 was replaced with that of PTPα. PTPα or the PTPα/CD45 hybrid was coexpressed with contactin. Anti-VSVG-immunoprecipitated PTPα/CD45 hybrid and PTPα were complexed with contactin (Fig. 4 B), demonstrating that contactin associates with the extracellular region of PTPα. Furthermore, the transmembrane region of PTPα is not specifically involved in the associa-
N-linked Glycosylation Is Not Required for the Association of PTPα and Contactin

The mature PTPα protein contains both N- and O-linked oligosaccharides (Daum et al., 1994). Chick contactin has nine potential sites for N-linked glycosylation (Ranscht and Dours, 1988; Brümmendorf et al., 1989). When COS cells coexpressing PTPα and contactin were cultured with tunicamycin, an inhibitor of N-linked glycosylation, faster migrating, less diffuse forms of PTPα and contactin were detected on SDS-PAGE (Fig. 4 C), which is consistent with a loss of N-linked oligosaccharides. Nevertheless, contactin was present in anti-VSVG immunoprecipitates from cells treated with or without tunicamycin (Fig. 4 C, lanes 7 and 8), indicating that the association of contactin and PTPα occurs independently of N-linked glycosylation of either protein.

PTPα and Contactin Associate In Cis but Not In Trans

Two experiments were carried out to address the question of whether PTPα and contactin associate in a cis or trans conformation. First, anticontactin precipitates were prepared from lysates of COS cells expressing either PTPα or contactin, or from cells coexpressing both PTPα and contactin (Fig. 5 A, lanes 1–3), as well as from another sample made by mixing lysates from the cells expressing either contactin or PTPα (Fig. 5 A, lane 4). A nontarget immunoprecipitates prepared from coexpressing cells contained PTPα, but those from mixed cell lysates did not (Fig. 5 A, bottom, lanes 7 and 8). The lack of detectable association of PTPα and contactin in the mixed lysates suggests that interaction cannot take place in a trans conformation. Still, this may require a particular presentation of these cell surface molecules in growing cells that cannot form in solubilized cell lysates. Therefore, contactin-expressing cells were trypsinized 24 h after transfection and replated in dishes containing PTPα-expressing cells (these were not trypsinized for replating because this resulted in a large decrease in PTPα expression). After 24 h of coculture, the cells were lysed and immunoprecipitates were prepared. As a positive control for contactin–PTPα association, PTPα- and contactin-cotransfected cells were cultured for 48 h, harvested, and processed the same way. PTPα and contactin coimmunoprecipitated from cotransfected cells (Fig. 5 B, top, lanes 3 and 5), but not from cocultured cells (Fig. 5 B, top, lanes 4 and 6). Thus, even when cells expressing contactin are cultured together with other cells expressing PTPα, no association in trans of these two receptor proteins occurs.

Discussion

We demonstrate that a receptor protein tyrosine phosphatase forms a membrane-spanning complex with a neuronal GPI-anchored receptor. PTPα and contactin associate with one another in a lateral (cis) forming complex mediated through the extracellular region of PTPα. This complex is sufficiently stable to permit its isolation from brain lysates or transfected cells through immunoprecipitation of either component, and to permit coclustering of PTPα with contactin upon antibody-induced clustering of contactin within cells and vice versa. Our findings that contactin–PTPα complexes form in cotransfected cells, but not upon coculture or in mixed lysates of PTPα-expressing cells and contactin-expressing cells, provides compelling evidence that PTPα and contactin can only associate within the same cell; thus, forming a receptor complex rather than a ligand–receptor pair.
This is the first identification of an extracellular partner and potential regulator of PTPα. Contactin lacks any intracellular region and is tethered to the external face of the plasma membrane through a GPI linkage. Contactin could, thus, link an extracellular contactin-mediated signal to an intracellular response. Fyn is complexed with contactin and is transiently activated upon aggregation of contactin (Zisch et al., 1995). Likewise, Fyn is associated with PTPα and is dephosphorylated and activated by PTPα (Bhandari et al., 1998; Harder et al., 1998; Ponniah et al., 1999; Su et al., 1999). The association of PTPα and contactin suggests that PTPα might act as an intermediary molecule in a tripartite complex of contactin, PTPα, and Fyn, in accord with its proposed role as a transducer. Zisch et al. (1995) reported that antibody-mediated cross-linking of contactin results in the activation of associated Fyn, raising the possibility that cross-linking of contactin–PTPα and contactin–Fyn complexes brings PTPα into proximity with Fyn, allowing the subsequent dephosphorylation and activation of Fyn. In view of the constitutive activity of PTPα (and other RPTPs), such a contactin-regulated access of PTPα to its substrate provides an attractive mechanism for mediating RPTP activity and the action of GPI-anchored receptors as modulators in larger signaling complexes.

Contactin interacts with multiple ligands and the identification of PTPα as a novel contactin-associated protein with its proposed role as a transducer.
extends the number and the nature of possible contactin-containing receptor complexes. Contactin is found in regions of active neuronal migration or outgrowth in the developing brain and in areas of synaptic development and activity (Faivre-Sarrailh et al., 1992; D’Alessandri et al., 1995). The role of this mobile GPI-anchored receptor may well be to deliver the components of a phosphorylation-competent machinery to receptor complexes mediating neuronal motility or synaptic activity. Evidence for such a role has been obtained for the closely related GPI receptor axonin-1/TAG-1, where neu- rite fasciculation mediated by complexes of axonin-1 and NgCAM is accompanied by a rapid downregulation of fyn phosphorylation (Kunz et al., 1996).

An intriguing possibility arising from this study is that contactin acts as an adapter to bring together two RPTPs, namely RPTPζ/β and PTPα. The NH2-terminal carbonic anhydrase-like region of the transmembrane and secreted extracellular forms of glial cell RPTPζ/β associates in trans with contactin, and in doing so promotes neurite growth (Peles et al., 1995). If PTPα functions as a signaling component of a RPTPζ/β-bound contactin–PTPα complex, this would represent a novel mode of RPTP interaction and regulation.

The proposed signaling through a contactin–PTPα complex represents a new paradigm of receptor-mediated tyrosine kinase activation. Receptors with intrinsic tyrosine kinase activity or directly associated with active nonrecep-

tor tyrosine kinases have been well documented. Contactin, lacking the intracellular region required for either of these mechanisms, may utilize an associated RPTP, PTPα, to effect intracellular activation of tyrosine kinases. This is reminiscent of the recent finding that the GPI-anchored cell surface receptors GDNFR-α and NTNR-α form functional coreceptor complexes with the transmembrane tyrosine kinase Ret (Buj-Bello et al., 1997; Klein et al., 1997) and underlines the concept that GPI-anchored receptor signaling is achieved by modulation of protein tyrosine phosphorylation. Our study unites previous progress in two areas of research: the interaction of extracellular ligands with the neural cell adhesion molecule contactin, and the intracellular signaling events mediated by PTPα. The components of a novel signal transduction pathway, thus, have been identified and can now be tested for function and physiological relevance to aspects of neuronal development.

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