Protein tyrosine phosphatase 1B is located with glucagon vesicles, and its concentration is inversely correlated with the rate of glucagon secretion of INR1G9 cells

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Abstract

High concentrations of protein tyrosine phosphatase (PTP) were found with secretory vesicles of glucagon-producing INR1G9 cells by electron microscopic immunocytochemistry, using a polyclonal antiserum specific for the PTP1B/T-cell (TC)PTP subfamily of PTP. Since TCPTP protein and mRNA were below the detection limit in the cells but significant amounts of PTP1B and mRNA were recognised by a specific monoclonal antibody and a mRNA probe we conclude, that the PTP associated with the vesicles is PTP1B. Only reverse transcriptase (RT)-PCR with primers specific for PTP1B yielded a product of the expected nucleotide sequence. Thus, we conclude that the PTP associated with the vesicles is PTP1B. The presence of vanadate for 48 h attenuated PTP1B expression and caused reduction of steady-state levels of the phosphatase. These conditions also led to a continuing increase in the steady-state rate of glucagon release by the cells. This rate and tyrosine phosphatase levels showed an inverse relationship, suggesting a suppressive role of PTP1B on the regulated secretion of glucagon by INR1G9 cells.

Introduction

The secretory vesicles of glucagon-secreting cells of Langerhans islets of the pancreas contain high amounts of a protein exhibiting immunoreaction with an antiserum raised against the conserved N-terminal part of protein tyrosine phosphatase (PTP) of the PTP1B/T-cell (TC)PTP family (Wimmer et al. 1999). The immunoreaction was confined to secretory vesicles, thus suggesting a potential, but so far undefined role, of the PTP in the function of these vesicles.

Insulin-producing cells did not react with this antiserum. On the other hand, it has been reported that other proteins that are structurally related to transmembrane tyrosine phosphatase, namely, ICA512/IA-2 and phogrins/IA-2β (Kawasaki et al., 1996, Payton et al. 1995, Cui et al. 1998), are major autoantigens involved in the generation of insulin-dependent diabetes mellitus (reviewed by Pietropaolo et al. 1997, Zanone et al. 2003). These proteins are present not only in pancreatic B-cell secretory granules but also on neurosecretory granules (Dirks et al. 1998) and human lung cancer cell lines of neuroendocrine phenotype. It has been questioned whether these proteins exhibit PTP activity, but they apparently become phosphorylated (Wasmeier & Hutton 1999) and cleaved from membranes (Ort et al. 2001) upon stimulation of insulin secretion, and they may be involved in the mobilisation and exocytosis of secretory granules (Wasmeier & Hutton 2001, Wasmeier et al. 2002). Since involvement of PTP has been discussed not only for pancreatic islets (Gogg et al. 2001) but also for the secretion of gonadotrophin (Marantz et al. 1995) and for the exocytic process in exocrine pancreatic acinar cells (Feick et al. 1998, 1999), tyrosine dephosphorylation may be a rather general signal transducing mechanism in protein secretion. The presence of tyrosine phosphatases of distinct subfamilies in islet cells which secrete either glucagon or insulin would indicate different types of tyrosine dephosphorylation mechanisms involved in secretion of these hormones.

To avoid the cellular heterogeneity of pancreatic islets, we used INR1G9 cells derived from a hamster glucagonoma (Takaki et al. 1986) for the present study. This cell line is well characterised by stable basal secretion of glucagon and has been used in studies of proglucagon processing (Mineo et al. 1995, Dhanvantari & Brubaker 1998) and calcium oscillations (Drucker et al. 1988, Bode et al. 1994, 1999). The cells display ultrastructure similar to that of native A cells of islets, and they reacted with the antiserum against PTP1B/TCPTP in the same way as pancreatic A cells. Thus, the cell line seems to be an appropriate tool for studying the mechanisms of glucagon
secretion. Here we show the localisation of PTP1B with secretory vesicles of INR1G9 cells. We also demonstrate that treatment of the cells with low concentrations of vanadate for 48 h leads to reduced expression of PTP1B, and that attenuation of tyrosine phosphatase activity caused by vanadate treatment is correlated with strong stimulation of glucagon secretion.

Materials and Methods

Materials
INR1G9 cells, a generous gift from Dr U. Wulbrand (Department of Medicine, Philipps-University, Marburg, Germany), were cultured in RPMI 1640 medium supplemented with 10% fetal calf serum (FCS), 1% penicillin/streptomycin at 37°C in the presence of 5% CO₂. For electron microscopic examination, INR1G9 cells were fixed in 2% paraformaldehyde/0.25% glutaraldehyde and embedded in Unicryl (Polyscience, Eppelheim, Germany).

The antisera against TCPTP/PTP1B used for immunocytochemical demonstration as well as for immunoblots were raised in rabbits against a 35 kDa N-terminal fragment of human TCPTP (Cool et al. 1989) comprising the catalytic domain (amino acids 1–319) expressed in E. coli (Schmid et al. 1996, Wimmer et al. 1998). Monoclonal antibodies selective for TCPTP (Cat. no. PH03L) and PTP1B (Cat. no. PH02) were purchased from Oncogen Research. The guinea pig antisera against glucagon was purchased from Biotrend (Cologne, Germany). Specificity of antibodies against tyrosine phosphatases was checked with His6-fusion proteins of PTP1B and TCPTP expressed with pQE30 vectors (Qiagen) in the E. coli strain AD202 and purified by Ni-agarose chromatography.

Immunocytochemistry

Immunocytochemical staining was performed according to a standard protocol either with cells fixed on the culture slide or with ultrathin sections of the Unicryl-embedded cells. Double immunostaining was done with an antibody to TCPTP/PTP1B (1:250) in combination with an antibody to glucagon (1:2000) or to synaptophysin (1:250) as indicated. Anti-PTP was detected by gold-conjugated secondary goat antibody to rabbit IgG (particle size 10 nm; British Biocell, Cardiff, UK), and anti-glucagon was visualised by gold-conjugated secondary goat antibody to guinea pig IgG (particle size 20 nm; British Biocell). For double staining with synaptophysin, anti-PTP was marked by 20 nm gold particles, while anti-synaptophysin was detected by gold-conjugated antibody to mouse IgG (particle size 10 nm). Contrast enhancement was achieved by treatment with 5% uranyl acetate.

Monoclonal antibodies to TCPTP and to PTP1B were used in 1:125 dilution for light microscopic immunocytochemistry. Both reactions were visualised by fluorescein isothiocyanate (FITC) conjugated to antibodies raised against mouse IgG (1:80).

Biochemical analysis

PTP activities were determined after homogenisation of the cells in buffer I (50 mM triethanolamine-HCl, pH 7.5, 15 mM 2-mercaptoethanol, aprotinin 2000 units/ml, and pepstatin 10 µM) in a glass homogeniser. The supernatant was separated from the pellet by centrifugation (23,000 g, 10 min), and the pellet was re-extracted, first with buffer I and then twice with buffer I containing 1% Triton X-100.

Enzyme assay

PTP assays were performed in 50 mM triethanolamine-HCl (pH 7.5) containing 15 mM 2-mercaptoethanol, 0.1% Triton X-100, bovine serum albumin (1 mg/ml) and 4 µM reduced, maleylated and carboxymethylated lysozyme (Tonks et al. 1988) labelled with 32P on tyrosine (Batzer et al. 1990) at 22°C in a volume of 40 µl.

After 15 min of incubation, the reactions were terminated by addition of an equal volume of 20% trichloroacetic acid. The samples were spun in an Eppendorf microcentrifuge, and 40 µl of the supernatant were added to 1 ml H₂O for Cerenkov counting. The assay procedure was developed for PTP of the PTP1B/TCPTP subfamily (Tonks et al. 1988) but shows low activities for the enzymes of the SH-PTP subfamily (A Voges, PhD thesis, University of Konstanz, 2003).

Immunoblotting

Samples were separated on a 10% SDS–PAGE gel. Transfer on an Immobilon P membrane (Millipore, Bedford, MA, USA) was performed according to Towbin et al. (1979). The membrane was blocked with 10% fat-free milk powder in PBS and subsequently incubated for 1 h with antibodies diluted in the 0.5% fat-free milk powder in PBS. Incubation with monoclonal anti-PTP1B or anti-TCPTP was done overnight at 4°C. Alkaline phosphatase-conjugated anti-mouse immunoglobulins was applied for 1 h at room temperature, and the reaction was visualised with the SigmaFast BCIP/NBT substrate. A mixture of prestained molecular weight markers (BioRad Low Range and Invitrogen SeeBlue) was used for calibration.

RT-PCR

RNA was prepared from 200 µg INR1G9 cells frozen in liquid nitrogen using the Roti-Quick Kit (Roth, Karlsruhe, Germany). An amount of 3-4 µg mRNA was reverse-transcribed with 40 units M-MuLV reverse
transcriptase (MBI Fermentas, Riga, Latvia) in the presence of 1 mM dNTP and 20 pmol specific reverse primer in a total volume of 20 µl.

Reactions were performed for 60 min at 37 °C followed by inactivation of the enzyme at 72 °C for 10 min. PCR was performed with either Pfu-polymerase (Promega) or Taq-polymerase (Peqlab, Erlangen, Germany), using 1 µl reverse transcriptase reaction product, 12 pmol specific forward and reverse primers, and 0·2 mM dNTP in a total volume of 50 µl through 30 cycles of 30 s at 94 °C, 30 s at 57 °C and 2 min at 72 °C for Pfu-polymerase or 1 min for Taq-polymerase in a Biometra thermocycler (Göttingen, Germany). The forward and reverse primers were chosen for conserved regions of PTP-1B and TC-PTP, using Windows 32 PrimerSelect – Lasergene DNASTAR software. The following primers were used: for PTP-1B: forward, 5′-CCCACTGATCCTGCACTGACGAG-3′, and reverse, 5′-TGTTGCATCAGGCCACCTG-3′; for TCPTP: forward, 5′-ACCATCGAGCGGGAGTTC-3′, and reverse, 5′-ATCTTGGCCTTTTTTCTTTTTCGTTTCAGTTT-3′.

The PCR products were purified from agarose gel using an extraction kit (Machery & Nagel, Düren, Germany) and ligated into pCR-Script plasmid. Sequencing of DNA inserts was done by SeqLab (Göttingen, Germany) with T3 and T7 primers.

Additional PCR reactions were done, using the Taq polymerase protocol to check for the presence of transcripts SHPTP1 and SHPTP2 with the following primers: for SHPTP1, 5′-CGGAATTCCTCGAGCGGGAGTTC-3′ and 5′-GGAATTCCACCGAGACCTCAG TGGGCTGGAT-3′, and for SHPTP2, 5′-CGGAATTCCACCGAGACCTCAG TGGGCTGGAT-3′ and 5′-ACCATCGAGCGGGAGTTC-3′.

This protocol was also used for quantitation, using the following primers for glyceraldehydephosphate dehydrogenase: 5′-GCCAGGGTTTCTCAGGATAATGAGC-3′ and 5′-TCAGAGTTGGAGGGCCACATTC-3′.

Northern blotting

An amount of 10 µg RNA, prepared as described above, was separated on 1% agarose gels in the presence of formaldehyde and formamide according to Lehrauch et al. (1977), blotted to positively charged nylon membrane (Boehringer, Mannheim, Germany), and fixed by 5-min irradiation with UV-light (285 nm). Hybridisation with digoxigenin-labelled probes was done in Dig-Easy-Hyb buffer (Boehringer) at 62 °C and 65 °C for 12 h. The DIG Wash and Block buffer set (Boehringer) was used for the subsequent washing steps. The blots were repetitively used after stripping with formamide (50%) and SDS (1%) in 50 mM Tris-HCl (pH 8·1). The antisense probes recognised coding porcine mRNA base segments 1–1311 for TCPTP, 106–1260 for PTP1B, 318–1385 for PTP1C and 1–1749 for PTP1D porcine proteins (R. Simpfendorfer & H.W. Hofer, unpublished). They were synthesised with RNA polymerase T3 or RNA polymerase T7 respectively and labelled with digoxigenin with the DIG RNA Labelling Mix (Boehringer). Purity and labelling of the probes were checked by electrophoresis and immunoblotting. Chemiluminescent detection of blots employed antidigoxigenin–AP Fab fragments (Boehringer) and CSPD (Roche). The lanes on films were scanned, and the intensity of bands was evaluated with the ScanPack II program (Biometra, Göttingen, Germany). The amount of 18 S rRNA was used as an internal standard.

Determination of glucagon secretion

Cells were grown to confluence as described above. For the determination of the glucagon secretion rate, the medium was replaced by Hanks’ balanced salt solution, and the cells were incubated for 30 min. The supernatant was collected, and the concentration of glucagon was determined by a standard RIA method (Biermann, Bad Nauheim, Germany).

Secretion experiments with vanadate or okadaic acid

The cells were grown to confluence in culture flasks under routine conditions. Then the culture medium was replaced by fresh medium plus vanadate or okadaic acid at the indicated concentrations. After 4 or 24 h for okadaic acid and 48 h of incubation with vanadate, the secretion of glucagon during a 30-min period was determined as described in the previous paragraph.

Results

Co-localisation of glucagon and tyrosine phosphatase in secretory vesicles

Immunofluorescence studies confirmed the existence of a tyrosine phosphatase of the TCPTP/PTP1B type co-localised with glucagon in glucagon-producing INR1G9 cells (data not shown). Immunocytochemical analysis of the distribution of PTP and glucagon in the cells at electron microscopic level is shown in Fig. 1. The small (10 nm) gold particles identify the tyrosine phosphatase of the TCPTP/PTP1B type and the bigger (20 nm) particles represent the localisation of glucagon.

Both proteins were obviously present on or in the immediate vicinity of vesicles. The size of the vesicles in INR1G9 cells was between 140 and 240 nm, and they corresponded in size, shape and intracellular distribution to secretory vesicles of glucagon-secreting alpha-cells in the Langerhans islets of the Syrian golden hamster (Takaki et al. 1986) from which the cells were derived.
To verify the localisation of PTP on secretory granules, the cells were checked for the co-existence of synaptophysin, an integral membrane protein of secretory granules, and PTP. Light microscopic examination of double immunofluorescence (Fig. 2a and b), and electron microscopic studies (Fig. 2c) revealed co-localisation of both antigens on secretory granules.

For more information about intracellular localisation of PTP, the cells were probed with an antibody to golgin 97, a protein localised at the cytoplasmic side of the Golgi complex, in combination with the polyclonal antibody to PTP. As shown by Fig. 3, PTP and golgin 97 were co-localised around the nucleus.

Analysis for tyrosine phosphatases in INR1G9 cells

The polyclonal antibodies used for immunocytochemical analysis were directed against the N-terminal and catalytic domain conserved in PTP1B and TCPTP, and they detected both members of this PTP subfamily. Therefore, commercially available monoclonal antibodies specific for either PTP1B or TCPTP were used for differentiation between the two phosphatases (Fig. 4a and b). Specific recognition of the phosphatases by their respective monoclonal antibodies was verified by in situ hybridisation (Fig. 4c and d) as well as immunoblotting of the bacterially expressed TCPTP and PTP1B after denaturing electrophoresis (Fig. 5a). However, these antibodies, although specific, were not sufficiently sensitive for electron microscopic immunocytochemistry but were useful only for immunofluorescence studies. INR1G9 cells exhibited a distinct reaction with the monoclonal antibody directed against PTP1B (Fig. 4a), whereas the reaction with the monoclonal antibody against TCPTP was at the level of background fluorescence (Fig. 4b). In situ hybridisation assays were in agreement with these observations since they confirmed the presence of mRNA encoding PTP1B, whereas the reaction failed for TCPTP (Fig. 4c and d).

Immunoblotting of proteins in extracts from INR1G9 cells on nitrocellulose membranes with a monoclonal antibody against PTP1B showed a single band of an estimated molecular mass of about 50 kDa (Fig. 5a). Probing of the blots for members of the SH2-domain containing tyrosine phosphatase subfamily (PTP1C and PTP1D or SHPTP1 and 2 respectively) showed negative results.

We also qualitatively checked for the mRNA of PTP1B, TCPTP and the SH2 tyrosine phosphatase subfamily members SHPTP1 and 2 by PCR after reverse

Figure 1 Immunocytochemical localisation of PTP and glucagon in ultrathin sections of INR1G9 cells. (a) Overview showing parts of three cells (original magnification ×6000). N: nucleus; bar = 1 µm. (b) Closeup of the cell in the upper left corner of panel a, showing part of the nucleus and the area of secretory vesicles (original magnification ×12930). Bar = 0.5 µm. (c) Area of the secretory vesicles at higher magnification (×35970). PTP is marked by gold particles 10 nm in size, and glucagon is marked by gold particles 20 nm in size. Bar = 0.1 µm.
transcription of the mRNA. Primers derived from the base sequences of highly conserved regions of the corresponding mouse enzymes in the case of PTP1B and TCPTP were used for amplification with proofreading Pfu-polymerase (Fig. 5b). A PCR product of the expected size (836 bp) was obtained with primers for PTP1B, while no amplification product of the expected size (1126 bp; Fig. 5b, arrow) was found for TCPTP with Pfu polymerase.

The PCR product representative for PTP1B was ligated into the pPCRSCRIPT plasmid and sequenced. It encoded 280 amino acids that were 98.5% identical with a corresponding sequence of PTP1B from rat. All four amino-acid exchanges within this region were conservative and confirmed the identity of the PCR product with cDNA encoding hamster PTP1B. While, in contrast to PTP1B, the PCR reaction for TCPTP was negative when Pfu-polymerase was used, the sequence of an amplification product obtained with non-proofreading Taq polymerase was not related to that of a tyrosine phosphatase. Primers for SHPTP1 and SHPTP2 did not even generate detectable amplification products with Taq polymerase (data not shown).

Tyrosine phosphatase and glucagon secretion

Cultured INR1G9 cells secreted glucagon into the culture medium. The average secretion rate under standard conditions (see Materials and Methods) was about 6 ng per 30 min per 10⁶ cells. To check for possible involvement of P-type serine/threonine phosphatases, the secretion rate of the cells was also studied in the presence of either 2 nM or 10 nM okadaic acid, a well-known inhibitor of PP1 and PP2a (Cohen 1989, Cohen et al. 1990). No significant change of the secretion rate was observed after 4 or 24 h of exposure.

To check for a potential influence of PTP activity on glucagon release, we reduced the activity of the enzyme by the use of PTP inhibitors. Addition of vanadate, a well-known inhibitor of tyrosine phosphatases, to INR1G9 cultures elevated glucagon secretion in a concentration-dependent manner and also decreased the tyrosine phosphatase activity extracted from the cells. Reduced PTP activity in the assays, however, was not due to kinetic inhibition of the enzyme by residual intracellular vanadate because this had undergone more than 1000-fold dilution, but was representative of a decrease in the amount of active phosphatase.

Figure 6a shows the rate of glucagon secretion plotted against extracellular vanadate concentrations to which the
cells had been subjected for 48 h. Subsequently, the medium was replaced for glucagon determinations, and the cells were incubated for another 30 min, as described in Materials and Methods. There was an obvious relationship between the vanadate concentration in the culture medium and the rate of glucagon release. Glucagon secretion increased to 615 ng/30 min per 10^6 cells in the presence of 33 µM vanadate in the medium compared to 6 ng/30 min per 10^6 cells in untreated controls. Figure 6b depicts the tyrosine phosphatase activities determined in extracts of the cells from the same series of experiments. Apparently, there was an inverse relationship between cellular phosphatase activity and vanadate concentration in the culture medium. As shown by the inset in Fig. 6b, the rate of glucagon secretion of INR1G9 cells was well correlated with the reciprocal of cellular tyrosine phosphatase activity (correlation coefficient r=0.95).

As demonstrated by the experiment depicted in Fig. 7, the effect of 10 µM vanadate on tyrosine phosphatase activity in INR1G9 cells was fully reversed to the value of untreated cells by removing vanadate from the incubation medium. Tyrosine phosphatase of cells remained constant during the first 48 h of incubation in the absence of vanadate and gradually decreased by 28% during another 48 h of incubation. In the presence of vanadate, however, the tyrosine phosphatase activity decreased by 66% during the first 48-h period but returned to the values of control cells during a second 48-h period of incubation after removal of vanadate.

Rescue of PTP activity from vanadate inhibition by dithiothreitol (Huyer et al. 1997) also reversed the effect of

**Figure 4** (a) Localisation of PTP1B in INR1G9 cells demonstrated with a specific monoclonal antibody (original magnification × 400). (b) Localisation of TCPTP as probed with a monoclonal antibody (× 400). (c) *In situ* hybridisation of PTP1B mRNA in INR1G9 cells. (d) *In situ* hybridisation of TCPTP in mRNA INR1G9 cells.
vanadate on glucagon secretion. Incubation of cells in the presence of 10 µM vanadate led to an increase in glucagon secretion to 622 ng/30 min per 10⁶ cells and to a decrease in PTP activity to 1.8 nmol/min per mg. Subsequent addition of 1 µM DTT reduced the secretion rate to 18.5 ng/30 min per 10⁶ cells and increased the activity to 6.5 nmol/min per mg. Moreover, inhibitors of PTP activity other than vanadate also led to increased glucagon secretion by INR1G9 cells. A 4-h incubation of cells with 50 µM phenylarsine oxide (Zhang et al. 1992) reduced PTP activity to 26%, 8% and 0% respectively of that in the control cells, while glucagon release increased 1.8-, 4.9- and 350-fold. As shown in Fig. 8, there was an immediate onset of the stimulating effect of vanadate on glucagon secretion by INR1G9 cells, with an approximately fivefold increase in the secretion rate within the first hour. Maximum secretion was observed about 40–50 h after the addition of vanadate. The presence of vanadate in cultures of INR1G9 cells reduced transcription of PTP1B mRNA.

Figure 5 (a) Immunoblots of extracts of INR1G9 cells probed with monoclonal antibodies against PTP1B (right) and TCPTP (left) (see Materials and Methods for details). (b) RT-PCR of mRNA prepared from INR1G9 cells using Pfu-polymerase and primers for the conserved regions of PTP1B and TCPTP (see Materials and Methods for details). Markers (MBI Fermentas): M: λDNA EcoRI/HindIII. The positions indicated at the right side correspond to the following sizes (from top): 21 226, 4973, 2027, 1584, 947 and 831 bp. The arrow indicates the expected position of a PCR product encoding TCPTP (1126 bp, lane empty). The PCR product encoding PTP1B visible at the expected position (836 bp) corresponded to its predicted size. Reactions for mRNA of PTP1C and PTP1D were negative.

Figure 6 (a) Rate of glucagon secretion by INR1G9 cells treated with various concentrations of vanadate for 48 h. Experimental procedures as described in Materials and Methods. (b) PTP activity in cellular extracts obtained from the cells of the same series of experiments shown in panel a. Inset: correlation between the rate of glucagon secretion shown in panel a and the reciprocals of the tyrosine phosphatase activities depicted in panel b. The regression line indicates an almost inverse relationship between secretion rate and tyrosine phosphatase activity (r=0.95).
The result of quantitative evaluation of Northern blots is depicted in Fig. 9. A transcript of 3·1 kb size was identified on the blots, and the signal persisted through high-stringency washings. The intensity of the band from cells incubated with 10 µM vanadate for 48 h was only 38% of that of untreated cells when referred to the same amount of ribosomal 18S RNA. Removal of vanadate and growth in normal medium for another 48 h led to a recovery of mRNA to 94% of the mRNA level of cells grown for 48 h in the absence of vanadate, corresponding to an increase in mRNA concentration by almost 250%, although the intensity of the band was still somewhat less than that from cells grown in the absence of vanadate for 96 h. The concentrations of mRNA transcripts of PTP1B in the presence of vanadate, therefore, changed in a similar manner as PTP activity. The signal of TCPTP on the Northern blots was too weak to allow determination of its quantity. PTP1C (synonym SHPTP1) and PTP1D (synonym: SHPTP2) showed no detectable hybridisation.

Discussion

Glucagon-secreting INR1G9 cells show intensive immunoreaction with a polyclonal antiserum raised against the conserved moiety of the PTP1B/TCPTP subfamily of PTP in a manner similar to that previously observed in A cells of Langerhans islets (Wimmer et al. 1999). Most of the immunoreactive tyrosine phosphatase was located on secretory granules and co-localised with synaptophysin, an integral membrane glycoprotein of secretory vesicles in endocrine pancreas (Kalina et al. 1991, Redecker et al. 1991). The polyclonal antibodies used for these immunocytochemical studies recognised both members of the PTP1B/TCPTP subfamily of PTP. Yet, solely monoclonal antibodies against PTP1B showed a positive reaction on immunoblots of SDS–PAGE of cell extracts. The preponderance of PTP1B over TCPTP in INR1G9 cells was also supported by experiments that proved the presence of specific mRNA (RT-PCR, in situ hybridisation, Northern blots), whereas none of these assays were positive for TCPTP.

This was in agreement with the signals obtained on Western blots using the same monoclonal antibodies. The PTP inhibitor vanadate led to a concentration-dependent 100-fold increase of glucagon release by INR1G9 cells. Vanadate is known as a kinetic inhibitor of PTP1B binding to the catalytic centre of the enzyme (Barford et al. 1994), but it may also exert other actions in cells. Distinct stimulation of glucagon secretion has already been observed in the presence of 0·3 µM vanadate in vitro. Although there was a rapid onset of the stimulation of glucagon secretion following the addition of vanadate, the maximum effect was visible only after about 2 days. Vanadate also reduced PTP activity in cell extracts when assessed by an assay which preferentially detected PTP of the PTP1B/TCPTP subfamily. The reduction of tyrosine phosphatase activity in INR1G9 cells was reversed by the removal of vanadate and by addition of dithiothreitol. The vanadate effect was possibly due to reduced expression of PTP1B protein, since the vanadate concentrations present in the activity assays were very low and unable to account for the observed reduction of phosphatase activity. This view is corroborated by the reduced levels of PTP1B mRNA after treatment with vanadate and its recovery after vanadate removal.

Figure 7 Rescue of tyrosine phosphatase activity in vanadate-treated INR1G9 cells. Tyrosine phosphatase activity measured in cells treated with 10 µM vanadate for 48 decreased from 6·24 nmol min⁻¹/mg protein to 2·10 nmol min⁻¹/mg protein whereas the activity in control cells remained constant with ±5%. When the vanadate-containing medium was replaced by vanadate-free medium, the tyrosine phosphatase activity determined after a second 48 h period was the same (4·48 nmol min⁻¹/mg cells) as assayed in control cells cultured for 96 h.

Figure 8 Dependence on incubation time of the effect of 10 µM vanadate on the glucagon secretion of INR1G9 cells. Experimental procedures as described in Materials and Methods.
mechanism of the influence of vanadate on the transcription of PTP1B mRNA is unknown at present; however, a similar effect has been previously observed in lymphocytes (R. Simpfendorfer & H. Hofer, unpublished).

It should be emphasised that stimulation of glucagon secretion was not a transient or short-term effect and was unlikely to be due to vanadate-induced overall damage of cellular function, since the cells were cultivated in the presence of vanadate for 2 days before the rate of glucagon secretion was measured. Therefore, the cells under study must have been fully in balance with the effects of the inhibitor after 48 h of exposure. The enhancement of glucagon release in response to vanadate under the steady-state conditions of our experiments, therefore, suggests a direct influence of vanadate on secretion rather than a secretory burst potentially caused indirectly by the interference of vanadate with other cellular functions. The inverse relationship between the effect of increasing vanadate concentrations on glucagon secretion and the effect on tyrosine phosphatase activity in the cells suggests a link between the depletion of phosphatase and elevated secretion. One reason to prefer the use of the potentially less specific vanadate to the unstable, but more specific and putatively irreversible, tyrosine phosphatase inhibitor pervanadate (Fantus et al. 1989) was to avoid the possibility of acute but transient toxicity and to establish steady-state conditions under which the inhibitor was expected to create a stable response of the cells – that is, a persistent increase in the level of tyrosine phosphorylation due to reduced phosphatase activity. Since a large amount of PTP1B was found in contact with secretory vesicles, we hypothesise that its function is to prevent inappropriate secretion of glucagon by attenuating a secretory trigger mechanism or by inhibiting a key step of exocytosis.

In contrast to tyrosine-specific phosphatases, the involvement of serine/threonine phosphatases in the regulation of glucagon secretion was not supported by our experiments. Okadaic acid, a potent inhibitor of type 1 and type 2A protein phosphatases, which led to increased or reduced secretion rates in other studies (Arufe et al. 1999, Matovcik et al. 1999, Sjoholm et al. 2002), had no influence on glucagon secretion by INR1G9 cells.

The proposed effect of PTP1B on secretory granules of glucagon-secreting A cells appears to be different from that proposed for the receptor-like PTP IAR, or phogrin, on insulin-producing B cells. These phosphatases were found to be activated by secretagogues (Cui et al. 1998). Loading of a recombinant PTP to permeabilised pancreatic acini also stimulated Ca\(^{2+}\)-dependent amylase secretion (Jena et al. 1991). An opposite function of tyrosine phosphatase in amylase secretion, however, was indicated by the report that the increase in cytosolic paxillin-dephosphorylating tyrosine phosphatase activity caused by disruption of the actin cytoskeleton led to an inhibition of amylase secretion in a pancreatic acinar cell line (Feick et al. 1999). In agreement with this observation, treatment of the
cells with pervanadate stimulated amylase release and phosphorylation of paxillin (Feick et al. 1998). Probably tyrosine phosphorylation is involved in more than one branch of the secretory pathway, and these pathways may even be subjected to variable regulatory mechanisms depending on the cell type. Nevertheless, the observations reported here suggest a suppressive effect of PTP1B on the release of glucagon and an important role of the enzyme in the regulation of glucagon secretion.

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