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Dephosphorylation of tyrosine phosphorylated synthetic peptides by rat liver phosphotyrosine protein phosphatase isoenzymes

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Five phosphotyrosine-containing peptides have been synthesized by FMOC solid-phase peptide synthesis. These peptides correspond to the 411–419 sequence of the \textit{Xenopus} src oncogene, to the 1191–1220 sequence of the human EGF receptor precursor, to the 1146–1158 sequence of the human insulin receptor, to the 856–865 sequence of the human \(\beta\)-PDGF receptor, and to the 5–16 sequence of the erythrocyte human band 3. The peptides were used as substrates for activity assay of two isoforms (AcP1 and AcP2) of a low molecular weight cytosolic PTPase. The assay, performed in microtiter EIA plates using Malachite green to determine the released phosphate, was rapid, reproducible, and sensitive. Both PTPase isoforms were able to hydrolyze all synthesized peptides, though with different affinity and rate. The main kinetic parameters were compared and discussed with respect to the role of the two enzymes in the cell.

1. INTRODUCTION

A low Mr phosphotyrosine protein phosphatase (E.C. 3.1.3.48) is present in the cytosol of a number of tissues [1]. (The enzyme was previously referred to as low Mr acid phosphatase (E.C. 3.1.3.2).) This enzyme is active toward Tyr-phosphorylated protein substrates [2–4] although it also catalyzes the hydrolysis of low Mr compounds such as \(p\)-nitrophenylphosphate (\(p\)NPP), \(l\)-phosphotyrosine, and acylphosphates [5–7].

The mechanism of the enzymatic reaction of the bovine heart enzyme has recently been identified, demonstrating the formation of a cysteinylphosphate covalent intermediate during catalysis [8]. The formation of a thiolphosphate intermediate appears as a specific event in the reaction mechanism of all PTPase family members [9–11]. Other phosphatases, such as acid and alkaline phosphatases form different covalent intermediates during the catalytic process [12]. The active site of this PTPase contains the sequence motif C-X-X-X-X-X-R in the 12–18 zone (both Cys and Arg are essential for the activity) and thus resembles the active site motif of the classical PTPase family members [13].

The enzyme from bovine liver has been sequenced in our laboratory [14]. It consists of a single polypeptide chain of 157 amino acid residues among which are present 8 cysteines, all in the reduced form [14]; two of these cysteine residues are involved in the catalytic site [13–15]. In other tissues, two isoenzymes differing only in the sequence of the 40–73 region are present [16]; they probably arise from alternative splicing.

The enzyme has been expressed in \textit{E. coli} using a synthetic polydeoxyribonucleotide with a sequence derived from that of the enzyme from bovine liver [17]. Recently, an experiment concerning the overexpression of the low Mr PTPase in \textit{v}-erb-B transformed fibroblasts [18] seemed to confirm the role of this enzyme as a possible regulatory element of cell growth [19].

By looking for biologically relevant substrates which could be dephosphorylated by this enzyme, it was observed that the autophosphorylated EGF receptor was efficiently dephosphorylated by this enzyme, as was as for other PTPases, is hi-
dered by the substantial difficulty of providing suitable substrates, since tyrosine phosphorylation occurs at very low levels within cells and the phosphoproteins are difficult to isolate and purify. Moreover, this makes the assay of the activity of these enzymes on substrates biologically more significant than synthetic phosphate esters such as \( p \)-nitrophenylphosphate, or \( \alpha \)-phosphotyrosine not easy.

**O-Phosphonyl peptide synthesis** can help to overcome these difficulties by providing phosphopeptides the sequence of which can be derived from that of the phosphorylation site of any phosphotyrosine protein kinase target protein the sequence of which is known. These peptides can therefore be considered as useful tools for routinely measuring the activity of PTPases and for preliminary investigations on their substrate specificity, if it exists.

Methods have recently been reported of the use of phosphotyrosyl peptides in PTPase activity tests [20,21]; these methods allow rapid, sensitive, reproducible, and accurate activity measurements that do not require the use of \( ^{32} \)P. In particular, a Malachite green colorimetric assay for phosphoserine protein phosphatase activity has recently been reported [21]. We extended this method to the phosphotyrosine protein phosphatase activity assay, and improved it in order to make the phosphatase assay easier and quicker by performing the reaction in microtiter EIA plate wells in the presence of varying phosphotyrosine-containing peptides which have previously been synthesized.

We used the above method to check the behavior of five phosphopeptides (derived from the sequence near the phosphorylation sites of the src gene product, the EGF receptor, the insulin receptor, the \( \beta \)-PDGF receptor, and the human erythrocyte band 3) as substrates of the low \( M_r \) phosphotyrosyl protein phosphatase isoforms from rat liver (AcP1 and AcP2).

2. MATERIALS AND METHODS

FMOC-Tyr(PO_3Me_2)OH was from Bachem, all other amino acid derivatives, DMF, piperidine, diisopropylcarbodiimide, and TFA were from MilliGen, as well as FMOC-Val-PepSynKA, FMOC-Gln-PepSynKA, and FMOC-Leu-PepSynKA resins. The C18 RP HiPore column was from Bio-Rad. The DEAE-HPLC column was from MilliGen. Membranes for automatic peptide sequencing were from MilliGen. All other reagents and solvents were of the highest available purity. The peptide sequences were taken from the PC-Gene data bank.

2.1. Phosphopeptide synthesis

Tyr(P)-containing peptides were prepared using a MilliGen PepSynthesizer by FMOC-solid-phase synthesis through dimethylphosphate protection using a Kieselguhr-polymethylacrylamide resin functionalized with \( \beta \)-alanine as internal reference amino acid, the acid labile 4-hydroxymethylphosphonoxastic acid linkage agent, and containing the FMOC-protected COOH-terminal residue. Subsequent peptide bond forming reactions utilized a fourfold excess of the OPfp-ester of the appropriate FMOC-amino acid (Dhbt-ester in the case of Ser and Thr) (4 equiv.) in the presence of 1 equiv. of HOBr. FMOC-Tyr(PO_3Me_2)-

OH was coupled in the presence of DIPCDI/HOBt. Side chain protection was carried out with \( t \)-butyl ester and ether for Asp, Glu, Ser, Thr, and Tyr, with the BOC group for Lys, and with the Mtr group for Arg. FMOC group deprotection was carried out with 20% piperidine in DMF. All other synthesis steps were carried out in DMF.

At completion of the synthesis, the resin-bound peptide was cleaved from the resin and deprotected with 1 M TMSBr-thioanisole/TFA (m-cresol, 10 mequiv. Tyr) at 4°C for times varying from 8 to 15 h, depending upon each peptide. The crude peptide was either low pressure evaporated, precipitated and washed with cold diethyl ether or directly treated with it. The peptide was then purified by semipreparative C18 RP-HPLC which, in the case of srcIP, was preceded by DEAE-HPLC.

2.2. Peptide purity assessment

The peptide purity was determined by amino acid analysis, FAB-mass spectrometry, and amino acid sequencing. Amino acid analysis was carried out as previously described [22]. Values for serine and threonine were corrected for loss during hydrolysis. The amino acid sequence was determined by Edman degradation by using a MilliGen Protein Sequencer mod. 6600 on peptides covalently immobilized on polyvinylidene difluoride membranes derivatized with 1,4-phenylene disothiocyanate (Sequen DITC) or alyamine groups (Sequen AA). Owing to its particular amino acid sequence, B3P was sequenced manually by Edman degradation. FAB mass spectra were determined using a VG Analytical 70-70 EQ instrument as previously described [23]. The peptide concentration was determined on the basis of the amino acid content. Phosphate analysis in the synthesized peptides was performed on a peptide sample of known concentration by the method of Fiske and Subbarow [24].

2.3. PTPase purification

The two isoforms of the low molecular weight PTPase, indicated by AcP1 and AcP2, were purified from rat liver as previously described [16].

3. RESULTS AND DISCUSSION

The synthesis and deprotection method used allowed us to obtain pure phosphotyrosine-containing peptides, even though extensive purification of the crude peptides was necessary, probably owing to the presence of impurities arising mainly from the deprotection step, which uses trimethylbromosilane as a hard acid source. For this reason we found that, in order to obtain crude peptides with minor impurities, the time of deprotection was critical and different for the varying synthesized phosphopeptides. The purified peptides were checked for purity by amino acid analysis, FAB mass spectrometry, amino acid sequencing, and inorganic phosphate determination by both combustion and enzymic hydrolysis. The correct amino acid composition was obtained...
Fig. 1. Standard curve for phosphate determination by the Malachite green assay. Each point was the average of four determinations.

for each peptide (the hydrolysis product, Tyr, was observed for PTyr). The amino acid sequence data for each phosphopeptide were consistent with the expected sequence (Table I). The inorganic phosphate determination in both ways showed a phosphate/peptide molar ratio of about 1.

The positive FAB mass spectra of the target peptides contained a distinct molecular ion at m/z values one unit higher with respect to the calculated molecular weight of each peptide, corresponding to the M + H+ species.

The PTPase activity of the two purified enzymes (AcP1 and AcP2) was determined by a Malachite green test. 400 ml of 0.1 M acetate buffer, pH 5.5, containing 1 mM EDTA and varying concentrations of each phosphopeptide were incubated at 37°C for 20 min in the presence of 5 U/ml of either AcP1 or AcP2. The reaction was stopped by adding 100 ml of the staining solution previously described [20,24] and left standing for 10 min at room temperature. The mixture was then centrifuged, 400 ml of the supernatant transferred in wells of microtiter EIA plates and the absorbance at 600 nm measured. The amount of inorganic phosphate released was determined by comparison with a standard curve prepared together with the experiments using an inorganic phosphate standard solution.

Fig. 1 shows the standard curve of the phosphate determination. As can be seen, the limit of sensitivity of this test is about 0.5 nmol of phosphate. The test was highly reproducible, with minor differences among absorbance values relative to different experiments.

Using this assay procedure, we have been able to determine the main kinetic parameters of AcP1 and AcP2 for varying phosphorylated substrates. Table II reports the kinetic data of the two enzyme isoforms. It can be noted that the two enzymes show pH optimum values in the 4.5–5.5 range on phosphotyrosine and PNPP, as well as on all phosphorylated peptides, except B3Pi, on which they elicit a pH optimum in the 4.0–5.0 range, probably due to the sharp acidic nature of this peptide. This further confirms a positive regulation of the enzyme activity by the H+ ion. As for affinity for the varying substrates tested and catalytic rate of their hydrolysis by the two enzymes, it can be pointed out that the apparent $K_m$ values of the two enzymes for the phosphorylated peptides, as well as for phosphotyrosine and PNPP, are very similar and in the 0.2–0.6 mM range, except for the apparent $K_m$ value of AcP2 for t-phosphotyrosine, which is one order of magnitude higher. Exceptions are represented by the low affinity values of both enzymes for PDGFRP2, as well as of AcP1 for B3Pi. The low affinity of AcP1 and AcP2 for PDGFRP2 could be due to the fact that, differently from the other peptides, the phosphotyrosine residue is located near the N-terminus in this peptide; this could render the positioning of the phosphotyrosine residue into the active site more difficult. The absence of a free α-amino group near the phosphotyrosine residue of the other peptides could be the cause of the lowering of the apparent $K_m$ values to levels similar to those of PNPP (Table I). AcP1 shows a low affinity also for B3Pi; in this case, the high apparent $K_m$ value could tentatively be ascribed to the acidic
nature of B3P, which is strongly negative at pH values near the pH optimum, lacking any basic residue.

The different affinities of AcP1 and AcP2 for both free phosphotyrosine and B3P, probably reside in peculiar differences in the sequence 40–73, the only segment which is different in the two enzymes. On the other hand, the $K_m$ values of each enzyme for the different tested substrates suggest that there is a variability in phosphopeptide substrate affinity, src1P showing the highest affinity and hence, probably, the best fitting into the substrate binding site.

PDGFRP1 and B3P are the most efficiently hydrolyzed peptides, and in this case it can be pointed out that AcP1 shows a higher hydrolysis rate with respect to AcP2. The PDGFRP1 and B3P, hydrolysis rate by AcP1 is very high, almost two orders of magnitude higher than that of the other peptides and the same order of magnitude as that of PNPP and free phosphotyrosine; the hydrolysis rate of the two peptides by AcP2 is quite lower, though higher than that of any other tested peptide. This observation is noteworthy, indicating that PDGFRP1 and B3P are good peptide substrates for AcP1 and, to a lesser extent, for AcP2, and suggests that the catalytic efficiency and affinity of the two enzymes on the purified, tyrosine-phosphorylated protein substrates for every specific PTPase. We think that the use of synthetic peptides containing phosphotyrosine and the availability of a suitable method to measure their dephosphorylation rate (like the one described in this paper) are useful for screening among synthetic substrates in the search for the physiological phosphorylated protein substrates for every specific PTPase.

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