Parathyroid Hormone–like Protein Is a Secretory Product of Atrial Myocytes

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Abstract

Parathyroid hormone-like protein (PLP) was originally identified from tumors associated with hypercalcemia. Recently, it has been found to be expressed in a stretch-responsive manner in several types of smooth muscle. We studied adult rat heart muscle for the presence of the PLP. Using immunohistology and the PCR, we demonstrated the presence of PLP and its mRNA in all heart chambers. Immunoelectron microscopy demonstrated PLP in secretory vesicles of atrial myocytes. Using immunohasay, we demonstrated that atria contained a higher concentration of PLP than ventricles. Furthermore, primary cultures of both chambers released PLP into conditioned medium, with atria secreting more than ventricles. Considered with studies of the role of PLP in other tissues, our observations suggest that the production and secretion of PLP by cardiac myocytes represents a calcium-related regulatory function for this stretch-responsive polypeptide in the cardiovascular system. PLP in the heart may be the calcium counterpart for the atrial natriuretic-sodium regulatory axis of the cardiovascular system. (J. Clin. Invest. 1993. 92:727–735.) Key words: secretion • neuroendocrine • secretory granules • smooth muscle • cardiac muscle

Introduction

The parathyroid hormone-like protein (PLP) was originally discovered as a product of tumors associated with hypercalcemia (1–6). The subsequent development and application of immunochemical and nucleic acid probes for PLP have demonstrated its presence in a wide variety of normal tissues and organs (1–14). The most extensive expression has been in fetal tissues, but the presence of PLP in normal adult tissues has been noted for several organs (8, 13, 14). Using immunohasay, immunohistology, and PCR, we have demonstrated the presence of PLP and its mRNA in cardiac myocytes. With immunoelectron microscopy, we have demonstrated that PLP is present in secretory vesicles of atrial cells.

Methods

Tissues. Fresh surgical specimens were collected from the heart of adult (100–150 g) rats (Harlan Sprague-Dawley Inc., Indianapolis, IN). Portions of the specimens were prepared for culture, PLP extraction, immunohasay, immunohistology, PCR, and immunoelectron microscopy, all with methods we have previously published (15–24).

Immunohasay. PLP was measured by two immunoassays that were conducted by modifications of previously described procedures (15–19). Tyrosinated human PLP 1-34 and PLP 107-138 were used to prepare tracer by chloramine T radioiodination and as respective assay standards, along with rabbit antisera to each of the respective peptides. Lack of cross-reaction in the assay for at least a 100-fold excess was demonstrated for PTH 1-34, the noncorresponding PLP peptides, calcitonin, calcitonin gene-related peptide, chromogranin A, and for rat atrial and brain natriuretic peptides (ANP and BNP). All samples were assayed in multiple dilutions that paralleled the corresponding PLP standard. Control media (nonconditioned) did not react in the assay. The assay incubations were conducted under standard, nonequilibrium conditions, and the intra- and interassay variations were 7 and 12%, respectively (15–19). All synthetic peptides were purchased from Peninsula Laboratories, Belmont, CA.

mRNA studies. Fresh surgical specimens were processed into total RNA by extraction with RNAzol™ (20–22). Preparations of cDNA from the total RNA of the heart muscle were evaluated by PCR with primers for the coding region of PLP as previously described (22). PCR amplifications of cDNA samples were performed with 100 ng RNA/cDNA and a cycle program that consists of an initial melting step at 95°C for 2.5 min, an annealing step of 55°C for 2 min, 30 cycles with a melting step at 95°C for 30 s, an annealing step at 55°C for 30 s, and a primer extension step at 72°C for 45 s. The program was complete after a 10-min final primer extension. Results of PCR amplifications were analyzed as previously described on 6% Tris-buffered EDTA acrylamide gels (22).

Immunohistology and electron microscopy. Immunohistology and immunoelectron microscopy were conducted on cardiac specimens as previously described with antibodies to human PLPI-34, 38-64, and 109-141 (15, 18, 19, 23, 24). Control studies for specificity were performed with irrelevant antibodies and with the test antibodies adsorbed with their antigen or control antigen (see figure legends for details). Whereas adsorption of antibodies with their corresponding peptide antigens blocked any positive immunohistological findings, adsorption with the previously indicated control peptides, including ANP and BNP, had no effect. Control, irrelevant antibodies (to salmon calcitonin) showed no reactivity.

Tissue incubation. Specimens of rat atria and ventricles were finely minced, dispersed, and incubated as primary cultures in minimal essential medium with 20% FBS (15, 16). After an overnight incubation, the media was changed and media samples were collected at the indicated times for PLP immunohasay, so that the PLP accumulated in the media for 18 and 44 h, respectively (15–19).

Results

Our studies demonstrate the presence of PLP and its mRNA in adult rat heart muscles. Fig. 1 demonstrates the presence of PLP in heart muscle by immunohasay. PLP-containing cells are identified by antibodies that span the linear sequence of the molecule. The staining is diffuse in the atria but patchy in the ventricles, and we could not discern any differences between the staining and nonstaining cells in the latter. Fig. 2 shows the presence of PLP mRNA in heart as demonstrated by...
PCR. Although atrial and ventricular PLP mRNA were present in indistinguishable amounts in these qualitative studies, quantitative PCR studies will be necessary to demonstrate any differences between the chambers. Fig. 3 summarizes the immunoassays of PLP in extracts of heart muscle. PLP concentration was significantly higher in the atria than in the ventricles. The predicted PCR product was 325 bp and spanned the linear sequence of rat PLP from -32 to +71 residues.

**Discussion**

Our studies demonstrate by immunohistochemical and nucleic acid probes the presence of PLP in adult rat heart (Figs. 1–3) and in atrial secretory vesicles (Fig. 5). PLP was originally discovered as a product of cancers associated with hypercalcemia (1–6). Subsequent studies demonstrated the widespread production of PLP in fetal tissues (8, 13, 14). These two early observations led to the consideration of PLP as an oncofetal protein (1). However, more extensive studies with improved methodologies continued to demonstrate PLP expression in some normal adult tissues, with the placenta as a notable example (8–14, 22, 25). Many early studies of PLP production either did not include the heart or failed to demonstrate the polypeptide in this organ, although some studies did demonstrate low expression of PLP in heart (1–6, 12). By contrast, significant PLP production has been demonstrated by antibody and nucleic acid probes in several types of smooth muscle, including vascular, breast myoepithelial, urinary bladder, uterus, and avian ovoduct (11, 26–31). A regulatory role for PLP at these muscle sites has been suggested by the observation that the production of PLP and/or its mRNA was increased by muscle stretch (26, 28). Correspondingly, receptor and end-organ studies demonstrated that PLP 1-34 was able to specifically bind to and relax smooth muscle or to inhibit its contraction (32–35).

Our findings of robust PLP expression in heart muscle are the first to demonstrate its presence in secretory vesicles (36) (Fig. 5). Rather than the weak or variable expression of PLP previously observed in the heart, we found consistent and significant expression of the polypeptide in this organ. Our observations were made with antibody probes that span the linear sequence of PLP1-141. These observations suggest a local regulatory action for PLP in the heart. Direct evidence for a regulatory effect of PLP on cardiac function comes from the observations of Nickols et al. (37–39), who have shown that PLP administration increases heart rate, contractility, and coronary

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*Figure 2.* Demonstration by PCR of the presence of PLP-specific mRNA in heart muscle. PCR was conducted with cDNA prepared from total RNA. Lane 1, left ventricle. Lane 2, right ventricle. Lane 3, left atrium. Lane 4, right atrium. The migration positions of the molecular size standards are at the left. PCR reactions were performed with 100 ng of RNA/cDNA, 10 μl cDNA, 0.5 μM primers, 10X Taq buffer (Promega Corp., Madison, WI). 0.5 μCi [32P]dCTP (NEN, Boston, MA) and 2.5 U Taq DNA polymerase (Promega Corp.) were added to each reaction. Amplification was performed in a thermal cycler (model 9600; Perkin-Elmer Cetus Instruments, Norwalk, CT). Dried gels were exposed to Kodak XAR films for 90 min. The 5' primer was CTGGTT-CACGATGGAGCGTCT, the 3' primer was CTTGAGTTAGGGTAT-CTGCCCTGCCC. The predicted PCR product was 325 bp and spanned the linear sequence of rat PLP from -32 to +71 residues.

*Figure 3.* Concentration of PLP by immunoassay in chambers of rat heart (mean±SEM). Measurements were made in extracts of chambers from nine animals by the two immunoassay systems described in the text, one based on PLP1-34 (left panel, 20 chambers), the other on PLP109-141 (right panel, 36 chambers). The concentrations in the atria were significantly (P < 0.001) higher than in the atria. The concentrations in the ventricles were not significantly different from skeletal muscle (100±10 pmol/mg protein). There were no significant differences between the left and right heart chambers.

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Figure 4. Release of immunoreactive PLP into conditioned medium by primary cultures of rat atria and ventricles as measured by immunoreactivities based on PLP1-34 (circles) and PLP109-141 (squares). Four chambers were finely minced and dispersed into 2-4 35-mm tissue culture cells with minimal essential medium supplemented with 20% FBS. After 3 days of culture, the media were collected at the indicated time intervals for PLP radioimmunoassay. Atria released significantly more PLP than ventricles at each time point.

Figure 5. The demonstration by electron microscopic immunocytochemistry of PLP in secretory vesicles of the rat atria. Monoclonal antibody 9H7 was used in an immunogold format. This antibody was selected because of its consistent performance in non-electron microscopy (light level) immunohistology studies. Random areas of rat cardiac muscle were surgically removed and fixed in 3.0% paraformaldehyde, 1.0% glutaraldehyde, in 0.1 M sodium cacodylate buffer, pH 7.4. Tissues were then incubated in 100 mM glycine in sodium cacodylate buffer followed by dehydration in a graded series of ethanold solutions. Embedding and polymerization were conducted in LR white resin (medium grade; Ted Pella Inc., Redding, California). Thin sections (60-90A*) were placed on nickel grids coated with a 0.5% parlodian film. Grids were incubated with 20 μg/ml ligand affinity purified monoclonal antibody 9H7 to PLP 109-141 in Tris-buffered saline, 0.8% BSA, 0.1% gelatin, and 0.01% azide. Control grids were incubated with antibody after absorption for 24 h with 3 μg/ml of its antigen, PLP 109-141, or in buffer without the primary antibody. After incubation and washing, the samples were transferred to a 1:10 dilution of protein A conjugated to 15 nm colloidal gold (E-Y Laboratories, San Mateo, CA). Samples were then fixed in 2% glutaraldehyde and poststained with 2% aqueous uranyl acetate followed by bismuth subsalinate. Examination was with a Zeiss EM10B electron microscope.

The dark granules within the secretory vesicles correspond to the electron-dense gold particles. No specific granules were seen in any other locations, although occasional background granules could be observed. Adsorption of the antibody with its peptide blocked immunoreactivity, and there was no immunoreactivity with an irrelevant antibody and without the primary antibody. PLP could not be demonstrated in secretory vesicles in the ventricle. Magnification: 5A-17,325; 5B-110,250; 5C-320,000.

These observations regarding PLP in heart muscle are reminiscent of those relating to ANP and BNP, two other stretch-responsive peptides present in cardiac myocytes (44, 45). The distribution of PLP in various heart chambers (Figs. 1 and 3) roughly corresponds to the distribution of ANP and BNP in the heart. It is notable that in atrial secretory vesicles ANP is colocalized with chromogranin A, a protein that is also colocalized with PLP at several neuroendocrine sites (15, 15, 46). Even though there are no structural similarities between PLP and these two natriuretic molecules at either the amino acid or nucleotide level, their distribution pattern made it imperative for us to rule out any suprinos immunocytochemical cross-reactivity between PLP and ANP or BNP for immunocytochemical immunohistology studies, (Figs. 1, 3, and 4) and any nucleotide identity for PCR studies (Fig. 2).

The actions of PLP on the heart should be reconciled with its effects on other cell systems and their systemic effect on calcium and mineral homeostasis (47). A common denominator for all of these effects is calcium flux, transcellular and intracellular (1-6). The change in ambient calcium produced by PLP may initiate a tissue-specific cascade that leads to the respective and distinctive final action of PLP in each cell type, perhaps via the sarcoplasmic reticulum and actin–myosin complex in heart muscle (48). This view may be too simplistic, however, because it considers only the well studied effect of the amino terminus of PLP and does not accommodate the growing body of evidence that PLP may be processed into peptides with unique biological actions (1-6). Tissue-specific mRNA splicing may further account for the diversity of the actions of PLP (20-22).

The presence of PLP in secretory vesicles of cardiac myocytes suggests an important regulatory role for the polypeptide in cardiovascular function (48). Further studies will be necessary to define conclusively its role in the normal heart. We have also demonstrated in preliminary studies the presence of PLP and for its mRNA in human, pig, and rabbit heart (unpublished observations of the authors). Additionally, PLP may participate in the stretch, hypertrophy, and corresponding oncocyto gene expression that is exhibited by the failing heart (49, 50). It should be recalled that PLP 1-34 acts like PTH to decrease the renal clearance of calcium (1, 41). Thus, PLP could represent a second stretch-responsive regulatory axis in cardiac function that, in contrast to ANP and BNP, subserves calcium rather than sodium translocation. These two regulatory systems may also function in the utero-placental complex to regu-
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late blood flow (50). The presence in heart of a PLP-calcium axis that corresponds to an ANP/BNP-sodium axis may explain some aspects of the role of both of these two ions in hypertension and congestive heart failure (51–53).

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References


