Structural Requirements for Parathyroid Hormone Action in Mature Bone
Effects on Release of Cyclic Adenosine Monophosphate and Bone Gamma-Carboxyglutamic Acid-containing Protein from Perfused Rat Hindquarters

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Abstract

To determine the structural requirements for parathyroid hormone (PTH) activity in mature bone, we perfused the surgically isolated hindquarters of adult male rats with either native bovine PTH-(1-84) [bPTH-(1-84)] or the synthetic amino-terminal fragment, bovine PTH-(1-34) [bPTH-(1-34)]. Changes in the release of cyclic AMP (cAMP) and bone Gla protein (BGP) were monitored as evidence of bone-specific response to PTH; tissue specificity of the cAMP response was confirmed through in vitro examination of nonskeletal tissue response to PTH. Biologically active, monoiodinated 125I-bPTH-(1-84) was administered to determine if mature murine bone cleaves native hormone. We found that perfused rat bone continuously releases BGP, and that both bPTH-(1-84) and bPTH-(1-34) acutely suppress this release. In addition, both hormones stimulate cAMP release from perfused rat hindquarters. When examined on a molar basis, the magnitude of the cAMP release was dose-dependent and similar for both hormones, with doses yielding half-maximal cAMP responses. The response for bPTH-(1-34) was 0.5 nmol and for bPTH-(1-84) was 0.7 nmol. Moreover, biologically active 125I-bPTH-(1-84) was not metabolized in our hindquarter perfusion system. These findings indicate that PTH-(1-84) does not require extraskeletal or skeletal cleavage to an amino-terminal fragment in order to stimulate cAMP generation in, or suppress BGP release from, mature rat bone.

Introduction

Parsons and Robinson observed that perfusion of feline tibias with bovine parathyroid hormone (bPTH) stimulated calcium release from the bone only if the hormone had first been circulated through the whole animal (1). Their work led to the first suggestion that parathyroid hormone (PTH) requires extraskeletal processing to attain full potency in bone.

Later work showed intact 84-amino acid PTH [PTH-(1-84)] to be metabolized in liver and kidney to biologically inactive mid- and carboxyl (COOH)-terminal region fragments that enter the circulation (2). Under some circumstances, the hepatic processing may yield circulating biologically active amino-terminal (NH2)-fragments (3–5). Both NH2-fragments of PTH and the intact hormone are active in kidney, but information about structural requirements for PTH action in mammalian bone is contradictory. For example, Martin et al. (6) reported perfused canine tibias to selectively take up the amino-terminal fragment of bPTH [bPTH-(1-34)], but not bPTH-(1-84). Furthermore, only bPTH-(1-34) markedly stimulated release of cyclic AMP (cAMP) from the bones. Findings in other systems differ sharply from Martin's: bPTH-(1-84) is fully potent in activating adenyl cyclase in fetal rabbit calvarial membranes (7), isolated bone cells from rat calvarias (8), rat bone calvarias (9), chick bone (10), and isolated chick bone cells (11). Bone may (9) or may not (6) metabolize PTH-(1-84), but there are no data to verify skeletal production of bioactive NH2-fragments of PTH. Age and species differences in the models used may partly explain the above discrepancies. In any event, the concept that PTH-(1-84) requires extraskeletal cleavage to an NH2-terminal fragment for complete activity in bone is intriguing and possibly of great general importance.

Therefore, we have used the perfused rat hindquarters to test independently the hypothesis that bPTH-(1-34), but not bPTH-(1-84), can stimulate skeletal release of cAMP. The model excluded the possibility of PTH metabolism by visceral organs, but maintained the bone intact in situ. We also tested the effects of PTH on skeletal release of bone Gla protein (BGP, or osteocalcin), as a separate assessment of PTH action on bone. Finally, the studies clearly demonstrated the utility of the perfused rat hindquarters as a system for studies of mature mammalian skeletal physiology.

Methods

Perfusion technique. The surgically isolated hindlimb technique used in these studies was modified from Bliziotis et al. (12) and Ruderman et al. (13). The system is shown schematically in Fig. 1. Male Sprague-Dawley rats (300–350 g, Holtzman Co., Madison, WI) were anesthetized with sodium pentobarbital (5 mg/100 g body weight) i.p. After making a midline ventral incision, we ligated major branches of the aorta and inferior vena cava (the renal, spermatic, and ilolumbar vessels, and inferior mesenteric artery). The epigastric and hypogastric branches of the iliac vessels were ligated to restrict flow to the viscera and bone wall; ligatures were also placed around the distal colon, bladder neck, and vessels of the accessory sex organs. The colon was cut between two ligatures and the intestines removed. We next inserted a 22-gauge plastic catheter (Critikon, Tampa, FL) into the aorta anterior to the iliac bifurcation and immediately started the perfusion.

The aorta and vena cava were ligated above the arterial catheter and a bolus of heparin (100 U, Liquaemin sodium, Organon Diagnostics, West Orange, NJ) in 2 ml saline was delivered via the perfusate to ensure

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1. Abbreviations used in this paper: BGP, bone Gla protein, or osteocalcin; bPTH, bovine parathyroid hormone, 1,25(OH)2D3, 1,25 dihydroxyvitamin D3; HFBA, heparitobutyrinic acid; HPLC, high-performance liquid chromatography; IBMX, isobutylmethylxanthine; KHB, Krebs-Henseleit bicarbonate buffer; NaOAc, sodium acetate; ODS-silica, octadecylsilil silica; PTH, parathyroid hormone.


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thorough flushing. The posterior vena cava was then quickly cannulated (18 gauge, Critikon) and the venous return collected. After removal of the kidneys, the perfused lower portion of the rat was surgically isolated from the forebody just above the renal arteries. The spinal stump was ligated with surgical steel and the preparation was transferred to a warmed, humidified plexiglass chamber. Finally, we inserted a temperature probe (Yellow Springs Instrument Co., Yellow Springs, OH) into the rectal stump to monitor body temperature.

We continuously perfused the hindlimbs with oxygenated Krebs-Henseleit bicarbonate buffer (KHB) containing fresh washed bovine red blood cells (hematocrit 30-35%), 3% bovine serum albumin (BSA) (Cohn Fraction V, Miles Laboratories, Kankakee, IL), and 10 mM glucose, as described by Schultz et al. (14). The nonrecirculating perfusion system, shown in Fig. 1, used a peristaltic pump to deliver perfusate to a Silastic tubing lung (15) (0.058 in. i.d. and 0.077 in. o.d., Dow Corning, Midland, MI) at an average flow rate of 3 ml/min. The tubing lung allowed oxygenation and achievement of physiologic pH (7.4) without foaming, PH, PO2, and PCO2 were regulated by adjustment of gas mixture (95% O2/5% CO2 vs. 100% O2). The oxygenated perfusate was warmed to 37°C by a condenser perfused with hot water (Lauda Div., Brinkmann Instruments Co., Westbury, NY). Perfusion then passed a mixing chamber with a hormone delivery portal, an arterial sampling portal, and a bubble trap proximal to the arterial cannula. Not shown in Fig. 1 is a line connected to a pressure transducer for monitoring changes in arterial line pressure. We saw no evidence of PTH-induced vasodilation in arterial pressure tracings or effluent flow rates.

The hindlimb preparation was flushed for 20 min (washout), after which four 5-min basal collections of effluent were made before hormone delivery. At the time of hormone or vehicle delivery (2-ml vol over 30 s), 10 1-min collections were started, followed by four or more 5-min collections of venous effluent. At several points during each perfusion, arterial and effluent medium pH, PCO2, and PO2 were measured (model 313 blood gas analyzer, Instrumentation Laboratory, Inc., Lexington, MA) as indicators of tissue viability. Degree of tissue perfusion was assessed at the end by introducing a vital dye into the perfusate and observing its distribution.

We collected all venous effluent samples on ice and removed the red blood cells by centrifugation. 0.5 ml of the cell-free perfusate was mixed with an equal volume of 0.05 mM sodium acetate (NaOAc) buffer (pH 5.1) containing 0.5 mM 3-isobutylmethylxanthine (IBMX) (Sigma Chemical Co., St. Louis, MO) and heated for 10 min at 90-95°C. The denatured protein was removed by centrifugation at 3,000 g for 10 min, and the supernatant frozen until assayed.

Assay for cAMP and BGP. Cell-free perfusate samples were assayed in triplicate for cAMP using the method of Steiner et al. (16-17), as modified by Wray et al. (18). We used a goat antiserum to cAMP (G-829, 1:18,000 final dilution) recently developed in our laboratory against 2'-0-monomosuccinyl cAMP. The assay was highly sensitive and specific for cAMP; no significant displacement of tracer (125I-cAMP) resulted when we added adenosine, ATP, ADP, AMP, guanosine, GDP, GMP, 23'GMP, uric acid, theophylline, inosine, 5'TP, and 5IDP. Any displacement of tracer by these analogs occurred at concentrations > 10^-6 M while 10^-12 M 35cAMP effectively displaced the tracer. Without sample acetylation, the assay limit of detection was 0.25 pmol/tube. All samples from a given rat were centrifuged for phase separation at the same time to minimize intraassay variation; interassay coefficient of variation was 13%.

The hormone-stimulated release of cAMP from perfused rat hindlimb was calculated as follows: cAMP = cAMPi - [(1 - hct/100)(cAMPf/100 g body wt) where cAMPi is cAMP release (pmol/min per 100 g hindlimb wt), cAMPf is venous cAMP (picomoles per milliliter), hct is the perfusate hematocrit, F is the perfusate flow rate (milliliters per minute), and body wt is the weight of the hindlimbs (grams).

Samples of undiluted cell-free perfusate were frozen for radioimmunoassay of BGP content, as previously described by Price and Nishimoto (19).

Hormone preparations. In order to assure biological potency of all parathyroid hormone preparations used in our bone perfusion experiments, we assessed their biological activity with the guanylnucleotide-amplified canine renal plasma membrane adenylate cyclase assay of Nisenson et al. (20). The system included the GTP analog, 5-guanylimidodiphosphate [Gpp(NH)p] (100 µM), to augment adenylate cyclase sensitivity to PTH, and a phosphodiesterase inhibitor (1.0 mM IBMX). Synthetic bPTH-(1-34) was used as standard. This method produced consistent results in assays repeated over a year. We used bPTH-(1-84) generously provided by Dr. H. T. Keutmann, Massachusetts General Hospital, Boston, MA and synthetic bPTH-(1-34) (Beckman Instruments, Inc., Spincio Div., Palo Alto, CA). The biological activities of these hormones in the canine membrane assay (20) were 535 and 3,470 U/mg, respectively; however, these values differ markedly from the manufacturer's potency estimates of 3,540 U/mg for bPTH-(1-84) (Dr. Keutmann, personal communication) and 6,800 U/mg for synthetic bPTH-(1-34). Inconsistency in the estimated biological activity of these hormones is probably attributable to differences in the bioassay procedures and conditions. To avoid confusion, and because no rational choice could be made among the bioassay values available, we chose to express the PTH dosages on a molar basis.

A range of molar doses of PTH was used to examine differences in potency of intact hormone and the synthetic fragment. At a given time, we simultaneously infused two hormone preparations in parallel perfusion systems. All hormone doses were combined in 2 ml of KHB buffer containing 3% BSA (vehicle); control perfusates were carried out with vehicle alone.

Hormone cleavage. Biologically active radioiodinated bPTH-(1-84) was prepared and purified as described by Rosenberg et al. (21) and Nissenson et al. (22). Our objective was to determine whether or not rat bone cleaves native PTH, potentially generating bioactive fragments. The bPTH-(1-84) is labeled with 125I on tyrosine-43 so that metabolism of this molecule to an NH2-terminal fragment would also yield a labeled COOH-terminal fragment. Before iodination, the biological potency of the bPTH-(1-84) was estimated to be 2,650 U/mg using the rat renal membrane adenylate cyclase assay system (21). The specific activity of the preparation used in our studies was 0.012 mU/10^6 dpm. Biological activity of the 125I-bPTH-(1-84) preparation was established before use by PTH receptor binding to rat osteosarcoma cells and chicken renal membranes (23); it was previously shown that the electrolytic iodination procedure does not decrease the biological potency of PTH (21).

The lyophilized labeled hormone was reconstituted with 0.01 M acetic acid and aliquoted into doses containing ~10^6 cpm. Each dose was brought up to 2 ml total volume with vehicle for injection into the hindlimb preparation. After delivery of the labeled hormone (over 30 s), we made ten 3-min collections of venous effluent. Approximately 60% of the radioactivity of the injected dose was recovered in the first three venous collection tubes. The cell-free perfusate from these tubes was
weighed, and incubated with bPTH-(1-84) (Sigma Chemical Co.). The supernatant was assayed for cAMP as described above, and the results expressed as picomoles cAMP released per gram muscle. We have extended the perfusion beyond 120 min with good results. The arterial pH and P02 showed little variation over the course of a perfusion. Maintenance of arterial-venous differences in P02 indicated tissue viability, and were confirmed at the end of each perfusion.

We also examined the ability of PTH to stimulate cAMP release from rat hindlimb vessels. Sections of the external iliac arteries and veins from 350-g male rats were excised and freed of adherent connective tissue while immersed in 4°C KHB buffer. The vessels were blotted, weighed, and incubated with bPTH-(1-84) (2.8 nmol/ml), prostaglandin E2 (10 ng/ml), or PTH vehicle (0.01 mM acetic acid) for 5 min. The incubation media were processed and analyzed as described for muscle, and the results reported as picomoles cAMP released per gram vascular tissue. In addition, we examined the ability of PTH to stimulate cAMP release from isolated murine adipocytes (31) and bone marrow cells (32).

Statistical analyses. Data are presented as mean±SEM, unless stated otherwise. Student's t test was used to determine significance of differences between results of control and experimental tissue incubations. In experiments involving serial effluent sampling after hormone stimulation, the data were reduced to one response per rat by summation of all responses from the time of hormone delivery to 30 min. Statistical significance of differences was assessed by analysis of variance. When significant differences were indicated, Student-Newman-Keul's test was used to determine which differences were significant.

Results

Viability and system specificity. Fig. 2 illustrates the ability of the model to maintain tissue uptake and exchange of oxygen for 80 min; however, we have extended the perfusion beyond 120 min with good results. The arterial pH and PO2 showed little variation over the course of a perfusion. Maintenance of arterial-venous differences in PO2 indicated tissue viability, and were confirmed at the end of each perfusion.

We found that incubation of nonskeletal tissue with PTH had no effect on cAMP release or tissue content. Fig. 3 shows typical results from incubation of rat epitrochlearis muscles, wherein we found no stimulation by PTH of cAMP release to the medium or changes in muscle cAMP content. In all other tissues examined (external iliac vessels, bone marrow, and adipose tissue) PTH also failed to stimulate release of cAMP (data not shown).

The responses to increasing doses of bPTH-(1-34) and bPTH-(1-84) are shown in Fig. 4. For both hormones, peak cAMP release occurred within 10 min of hormone delivery over a wide range of doses. cAMP concentrations in effluent were essentially back to pretreatment levels at 20–30 min. For both dose-response studies, we estimated the areas under individual response curves by summing the values of all time points. The reduced data, plotted against hormone dose, allowed a comparison of the
bioactivity of the two hormones (Fig. 5). Increasing the dose of PTH gave a biphasic cAMP response, with declining total cAMP release above 6.8 nmol/preparation for bPTH-(1-84) and 11.6 nmol/preparation for bPTH-(1-34). The concentrations of bPTH-(1-34) and bPTH-(1-84) required for half maximal release of cAMP were estimated to be 0.5 and 0.7 nmol, respectively.

The rat hindlimbs released immunoreactive BGP, with the release slowly declining over the 120-min perfusion (Fig. 6). Treatment with bPTH-(1-84), 6.85 nmol, produced a significantly greater decrease of BGP release than in vehicle perfused hindquarters. Synthetic bPTH-(1-34), 2.78 nmol, also suppressed BGP release but to a lesser degree. The different magnitude of effects by the two peptides probably resulted from the nearly 2.5-fold difference in dose.

Metabolism of biologically active 125I-bPTH-(1-84). The fate of biologically active 125I-bPTH-(1-84) in our hindquarter perfusion system is illustrated in Fig. 7. The profile shown represents the average of values from two rats perfused with similar doses of 125I-bPTH-(1-84). We carried out a total of four separate perfusions with this labeled hormone, all of which yielded the same results. Chromatography of pooled venous effluent containing the greatest radioactivity revealed one major peak which coeluted with intact bPTH-(1-84). The HPLC profiles of labeled bPTH-(1-84) were identical before and after perfusions through the rat hindquarters. Approximately 60% of the radioactive label infused was contained in the venous effluent collected within the first 12 min. Subsequent collections were also pooled and chromatographed, but they contained no new peaks of activity.

Fig. 7 also illustrates the ability of our method (ODS-silica extraction followed by reverse-phase HPLC) to distinguish 125I-bPTH-(1-84) from cleavage products generated enzymatically in vitro. From these results, we are confident that the ODS-silica extraction procedure did not cause loss of fragments possibly generated in the hindquarter perfusion.

Discussion

The present studies verify that PTH evokes rapid, dose-related increases of cAMP release from perfused rat hindquarters (12), and suggests that PTH acutely suppresses release of BGP. We believe that both responses reflect the actions of PTH on bone. There is no serious question about specificity of the BGP response, since BGP occurs only in skeletal tissue (33). On the other hand, adenylate cyclase occurs in many tissues; muscle (27) and fat cells (31), and fat cell ghosts (34) reportedly respond to PTH with increased cAMP generation. However, Bliziotes et al. (12) found no effect of bPTH-(1-34) on rat muscle cAMP content or release, and repeated trials in our laboratory also showed no effect. In addition, we found no effect of PTH on cAMP release from murine fat cells (31, 34), marrow cells (32), and blood vessels. Despite the heterogeneity of perfused tissue in this system, we conclude that the BGP and cAMP responses were specific for bone.

Martin et al. (6) found only a weak cAMP response to bPTH-(1-84) in perfused canine tibia, while bPTH-(1-34) markedly stimulated cAMP release. A major finding of the present study was that bPTH-(1-84) and bPTH-(1-34) were potent stimulators of cAMP release.
of cAMP release from mature rat bone. For both hormones, the increase of cAMP release occurred within 3 min of hormone delivery and attained maximal values by 7–8 min. The magnitude of response was dose-dependent for both hormones. These findings strongly suggest that intact PTH does not require extraskelatal cleavage to an amino-terminal fragment to stimulate adenylate cyclase in mature rat bone.

Other work supports our findings. For example, cultured rat bone cells (8) and fetal rat calvariae (9, 35) are fully responsive to bPTH-(1-84). Rat calvariae may be able to cleave PTH (9), but there is no evidence for skeletal production of bioactive PTH fragments. Furthermore, fetal rabbit calvarial membranes responded to bPTH-(1-84) without apparent ability to metabolize the hormone (7). Our data clearly demonstrated that perfused murine bone did not cleave intact bPTH-(1-84) during a single pass. In addition, chromatography of effluent collected later than 12 min after hormone delivery revealed no new peaks, indicating that any delay in the release of the hormone was not due to skeletal cleavage. These findings further suggest that native PTH requires no metabolic activation for full potency in adult rat bone.

Effects of the two PTH peptides on cAMP release were indistinguishable over the dose range 0.1–2 nmol. However, increasing doses of bPTH-(1-34) beyond this range continued to stimulate cAMP release, finally reaching a plateau at 11.6 nmol, while the responses to doses of bPTH-(1-84) beyond this range plateaued or declined. We have no ready explanation for this phenomenon, but it strongly suggests that potency comparisons of PTH and its analogs must encompass full dose-response curves, to obtain complete information. These findings also demonstrate that potency comparisons of PTH preparations made in kidney may not accurately reflect their relative potencies in bone. The synthetic bPTH-(1-34) was 2.8-fold more potent than the native bPTH-(1-84) when bioassayed in the canine renal plasma membrane adenylate cyclase system, while these hormone preparations were essentially equipotent in our perfused bone system. These differences in potency estimates strongly suggest that careful consideration be given to the choice of organ system and species used to test the efficacy of PTH analogs.

Another important finding in these studies was that BGP was released continuously from perfused rat bone; moreover, this release acutely responded to PTH. Administration of bPTH-(1-84) and bPTH-(1-34) suppressed BGP release to below that of vehicle-perfused rats. The difference in the degree of suppression observed between the two hormonal treatments suggests that the magnitude of BGP suppression by PTH may be dose-dependent in perfused whole bone, as it was in bone cells (36).

Beresford et al. (36) reported that cultured osteoblast-like human bone cells stimulated with 1,25 dihydroxyvitamin D3 [1,25(OH)2D3] released BGP. When these cells were incubated with 1,25(OH)2D3 and PTH, there was a dose-dependent decrease in BGP release over a 48-h period. The reduction in BGP release was very slow and highly dependent upon stimulation with 1,25(OH)2D3 for initial release of BGP. By contrast, in the present study, perfused bone responded within minutes to a single dose of PTH by markedly reduced BGP release. PTH action may suppress BGP release by altering the rate of BGP synthesis or degradation, presumably by osteoblasts (37). The rapidity of
this response supports the proposed function of BGP as an informational protein responsible for regulation of cellular activities involved in bone turnover (38). BGP release from vehicle-perfused preparations also declined substantially with time, for unknown reasons; because of this, we believe our findings with respect to PTH inhibition of BGP release should be interpreted with caution.

In summary, the isolated perfused rat hindquarter system is a very useful model for the study of acute PTH actions on mature mammalian bone; we have only begun to explore its potential. The preparations also released cAMP in response to salmon calcitonin (unpublished results), but specificity of the response to bone has not been established. Both synthetic bPTH(1-34) and native bPTH(1-84) produced dose-dependent increases of cAMP release over a wide range of hormone doses. Mature murine bone clearly did not require extraskelatal cleavage of intact PTH to an NH2-terminal fragment for biological activity, and it could not cleave the intact hormone under the conditions of our perfusion system. BGP was also released from perfused rat hindquarters, and the administration of PTH produced a significant decline in BGP release. These findings suggest that PTH may acutely suppress the release of BGP from osteoblasts.

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