Effects of Parathyroid Hormone on Cytosolic Free Calcium Concentration in Individual Rabbit Connecting Tubules

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Abstract

PTH stimulates active Ca reabsorption in isolated perfused rabbit kidney connecting tubules (CNTs). The existence of PTH-sensitive adenylate cyclase and the reproduction of increased epithelial Ca transport by dibutyryl-cAMP suggest that cAMP is the mediator. Accordingly, we studied the effects of PTH and 8-bromoadenosine 3',5'-cAMP (8-Br-cAMP) on cytosolic free calcium concentration ([Ca$^{2+}$]) in individual rabbit CNTs. [Ca$^{2+}$] was estimated by continuous epifluorescence microscopy of single fura-2-loaded tubules during dual wavelength excitation. In nonperfused controls at 37°C, [Ca$^{2+}$] decreased with time. In contrast to vehicle controls, synthetic bovine (1–34) PTH (0.1 nM) increased [Ca$^{2+}$] within 4 min, produced a maximal effect in 7.2 min, and sustained its effect for at least 2 min after washout. 8-Br-cAMP (1 mM) mimicked the effect of PTH, but with an earlier onset of action. To test the hypothesis that lumen Ca is the predominant source of the rise in [Ca$^{2+}$], we studied singly perfused CNTs. In the absence of bath and lumen Ca, PTH elicited no rise in [Ca$^{2+}$], implying that intracelluar Ca stores are not the major source. In contrast, there was a rise when Ca was replenished in both media. In the continuous presence of bath Ca, lumen Ca was estimated to contribute 65% of the total rise in [Ca$^{2+}$] in response to PTH when it was first deleted and then replenished. However, when the sequence of lumen Ca manipulation was reversed, the contributions by lumen and bath Ca were found to be essentially equal. We conclude (a) at a physiologic concentration, PTH increases [Ca$^{2+}$] in rabbit CNTs, (b) 8-Br-cAMP mimics this action, implicating cAMP as a second messenger, and (c) the PTH-stimulated rise in [Ca$^{2+}$] depends importantly on both bath and tubular luminal fluid Ca.

Introduction

PTH increases Ca reabsorption by connecting tubules (CNTs) of the rabbit (1, 2). Since Ca$^{2+}$ is transported against a transepithelial electrochemical potential difference (1, 2), it has been inferred that this absorption is active and occurs via a cellular pathway (3). Definitive evidence for and analysis of a presumed transcellular route of Ca$^{2+}$ movement from lumen to bath are lacking, in part because of difficulties in measuring cytosolic free calcium concentration ([Ca$^{2+}$]). One model of a cellular pathway for epithelial Ca transport would predict a rise in [Ca$^{2+}$], in response to PTH (4). The availability of fura-2, an intracellular Ca$^{2+}$-sensitive fluorescent dye, has made it possible for us to investigate this thesis in freshly dissected individual rabbit CNTs that were suffused continuously with a physiologic solution in vitro. We found that [Ca$^{2+}$] increased 2–4 min after exposure to PTH. We reasoned that if the PTH-induced elevation in [Ca$^{2+}$], is related to its Ca transport effects, another agent known to mimic such actions also should increase [Ca$^{2+}$]. Given the PTH-sensitive adenylate cyclase in rabbit CNTs (5) and the stimulatory effect of dibutyryl-cAMP on Ca transport in this nephron segment (2), we examined the effects of 8-bromoadenosine 3',5'-cAMP (8-Br-cAMP). The results implicate cAMP as a messenger. Hypothetically, if the rise in [Ca$^{2+}$], is related to the known stimulation of Ca reabsorption by these agonists, lumen Ca ought to be the predominant source. To test this hypothesis, experiments were performed in singly perfused CNTs in which we manipulated lumen and bath Ca. Contrary to the above prediction, our results demonstrate that bath Ca is equally as important as lumen Ca in causing the PTH-induced rise in [Ca$^{2+}$].

Methods

Female albino rabbits, 0.9–1.2 kg, were fed a nutritionally complete synthetic diet containing 0.9% Ca, 0.4% P, and 60 IU vitamin D$_3$/100 g of feed (6) until each gained at least 200 g. They were decapitated and the left kidney was excised. Blood was collected from vessels of the neck for analyses.

Individual tubules were dissected and studied in vitro in simple electrolyte solutions. Unless stated otherwise, these contained (in millimolar): CaCl$_2$, 2; NaCl, 148.5; NaH$_2$PO$_4$, 2.0; KCl, 5; MgSO$_4$, 1.2; D-glucose, 5.6; and L-alanine, 6.0. CaCl$_2$ was replaced isosmotically with NaCl in Ca-free solutions. The EGTA solution for the minimal value of R (ratio of net fluorescence emitted at 500 nm for excitation wavelengths of 340 and 380 nm, respectively) measured in the absence of Ca (R$\text{min}$) and the 2.5 mM Ca solution for the maximal value of R measured in the presence of saturating Ca (R$\text{max}$) were identical to the standard solutions except that CaCl$_2$ was replaced by 2 mM EGTA in the former and by 2.5 mM CaCl$_2$ in the latter. Each solution was titrated to a pH of 7.40 with 1 M NaOH and bubbled for 15 min with 100% O$_2$, Na, K, and Ca and osmolality were measured in all solutions (7).

A thin slice of the kidney was immersed in chilled (4°C) dissection solution containing 1 mg/ml of defatted BSA. The medulla was am-

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1. Abbreviations used in this paper: 8-Br-cAMP, 8-bromoadenosine 3',5'-cAMP; [Ca$^{2+}$], cytosolic free calcium concentration; CNT(s), connecting tubule(s); R, ratio of net fluorescence emitted at 500 nm for excitation wavelengths of 340 and 380 nm, respectively; R$\text{min}$, minimal value of R measured in the absence of Ca; R$\text{max}$, maximal value of R measured in the presence of saturating Ca.

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putated and the cortex was microdissected (at 20–22°C) under a stereo microscope. CNTs were teased from the cortical labyrinth. They were identified (Fig. 1) by their location adjacent to interlobular arteries, their attachment to distal convoluted tubules or arcades, and their characteristic size and microscopic appearance.

$[^{\text{Ca}^{2+}}]$, was measured with fura-2 as described by Tsien et al. (8). Fura-2-AM (Molecular Probes, Inc., Eugene, OR) was dissolved in DMSO and stored at −20°C in a desiccator. Isolated CNTs were incubated in a 5-μM fura-2-AM solution and loaded at 20–22°C for 4.5–7.5 h. A fluorometer system (Spex Industries, Inc., Edison, NJ) was interfaced with a modified Diaphot inverted microscope (Nikon, Inc., Garden City, NY). Exciting radiation was split between two monochromators, each with entrance and exit slits of 1.8 nm. One beam at $\lambda = 340$ nm and another at $\lambda = 380$ nm were alternately directed by a reflective chopper into the nosepiece of the microscope. The incoming beams were reflected into a UV-F 40×, glycerin immersion, objective lens (Nikon, Inc.; numerical aperture = 1.30) by a 400-nm dichroic mirror. Epifluorescence from the tubule was collected by the same objective lens and passed through the dichroic mirror and a 420-nm barrier filter. It was projected by a ×1 lens through a rectangular aperture and an interference filter (500-nm peak, 20–30-nm bandwidth) onto a photon-counting detector (Hamamatsu Photonics K.K., Shizuoka, Japan).

A black aluminum suffusion chamber (White Instrument Co., Sunitland, MD) was mounted on the stage of the microscope and water-jacketed to 37°C. The tubule was suffused continuously at 2 ml/min with a syringe pump. Solutions were prewarmed and temperatures were controlled between 36.7 and 37.1°C. The effluent bathing medium was aspirated continuously. The bottom of the chamber was a 0.15-mm-thick quartz coverslip. The suffusion apparatus was surrounded by a black enclosure, and during fluorescence measurements the room was illuminated solely with red light.

Nonperfused tubules. Fura-2-loaded CNTs were transferred to the suffusion chamber, which contained ~0.5 ml of 2-mM Ca solution at room temperature. After the tubule was mounted and immobilized on the coverslip with glass micropipettes it was flushed with 25 ml of the suffusate to remove residual dye. When the temperature reached 37°C, final positioning and focusing were performed. Fluorescence at 500 nm of an ~200 × 40 μm region of the tubule was measured continuously and integrated every 0.5 s. From each kidney, three to six CNTs unexposed to fura-2-AM were studied identically to measure autofluorescence.

In time controls fluorescence was measured for 80–90 min at 37°C during continuous suffusion with 2-mM Ca. Thereafter, a calibration was performed in situ by changing the suffusate sequentially to 2 mM EGTA for 5 min, 2 mM EGTA + 5-μM ionomycin until $R_{max}$ was reached, and 2.5-mM Ca until $R_{max}$ was obtained. With PTH, 8-Br-cAMP, or acetic acid, fluorescence was first measured for 14±1 min after reaching 37°C while the tubule was suffused with 2-mM Ca solution to provide baseline data. Thereafter, the tubule was exposed to the agonist or reagent in the same suffusate for 10 min, followed by a 5-min recovery period. In three rabbits we studied an acetic acid control and a PTH-treated tubule from the same kidney.

Perfused tubules. Fura-2-loaded CNTs were studied similarly to nonperfused tubules with two differences. First, the lumen was cannulated and perfused at ~75 nl/min by gravity flow (9). Perfusate composition was changed without interrupting perfusion (10). The general protocol is shown in Fig. 7. Baseline $[^{\text{Ca}^{2+}}]$, was measured for 10 min at 37°C, during which both perfusate and suffusate contained 2-mM Ca. Thereafter, Ca was deleted from the lumen. After 10 min, 0.1 mM PTH was added to the suffusate. After another 10 min, luminal Ca was restored in the continued presence of PTH. Finally, PTH was withdrawn, followed by a 30-min recovery period with 2-mM Ca in both perfusate and suffusate. Second, during the calibration period the lumen also was perfused with 2 mM EGTA followed by 2 mM Ca for measurements of $R_{min}$ and $R_{max}$, respectively. Ionomycin was absent from all of these perfusates. Controls were studied identically but without PTH. In experiments to assess the role of intracellular Ca stores, both suffusate and perfusate Ca were deleted for 20 min after the baseline period. In all figures of perfused CNTs the times of perfusate exchanges, 4.5±0.25 min, were omitted for clarity. Net fluorescence was calculated by subtracting the autofluorescence of nonloaded tubules dissected from the same kidney.

Synthetic bovine PTH 1–34 tetracontapeptide (Peninsula Laboratories, Inc., Belmont, CA) was dissolved in 10 mM acetic acid and lyophilized. Aliquots were stored in a desiccator at −20°C. 8-Br-cAMP was obtained from Sigma Chemical Co., St. Louis, MO; EGTA, puriss P.A., from Fluka AG, Buchs, Switzerland; and ionomycin from Calbiochem-Behring Corp., San Diego, CA.

$[^{\text{Ca}^{2+}}]$, was calculated as (11): $[^{\text{Ca}^{2+}}] = K_d \cdot (R_{max}/R_{min} \cdot R_{max} - R(F_o/E_o))$. $K_d$, the dissociation constant of the fura-2:Ca complex at 37°C, was assumed to be 220 nM (11). $F_o/E_o$, is the ratio of net fluorescence at $\lambda = 380$ nm excitation for $R_{max}$ to that for $R_{min}$.

The tracings in Figs. 2–6 and 8 represent 10-s averages. The absolute values of $[^{\text{Ca}^{2+}}]$, are given in Table 1 represent averaged fluorescence in the indicated time intervals.

Statistical comparisons were made by analysis of variance of repeated measures (12) or by paired or unpaired t test (13) as appropriate. $P$ values < 0.05 were considered to be significant.

Results

Nonperfused CNTs

These studies demonstrate the feasibility of measuring $[^{\text{Ca}^{2+}}]$, with fura-2 in individual microdissected rabbit CNTs. At a suffusion rate of 2 ml/min, it is unlikely that any extracellular fura-2 that might have resulted from leakage out of the cells would contribute significantly to the measured fluorescence.

TIME CONTROLS

The autofluorescence was similar whether it was studied before or after fura-2-loaded tubules on a given day. In a few unloaded tubules evaluated extensively autofluorescence was stable for 80–100 min and was essentially unaffected by either agonists or reagents.

Morphologically the tubules appeared intact throughout the entire experiment, irrespective of exposure to PTH or 8-Br-cAMP. The functional integrity of the cells was evidenced by the return of $[^{\text{Ca}^{2+}}]$, with time at 37°C (Fig. 2), to values reported in other kidney cells (14–23).

Although body weights, food intake, blood pH, and serum ionized [Ca] values (not shown) of the rabbits from which these CNTs had been dissected were comparable, individual tubules varied considerably in their basal $[^{\text{Ca}^{2+}}]$. After several hours of exposure to the loading medium at 20–22°C in vitro, estimated $[^{\text{Ca}^{2+}}]$, was initially higher than the apparent

Figure 1. Locations from which connecting tubules were dissected: (1) subcapsular CNT not associated with arcades; (2) initial portion of CNT within mid- and deep-cortical arcades; and (3) joining segments between the branch points within arcades. Solid, white, and stippled structures represent thick ascending limbs of Henle’s loop, distal convoluted tubules, and cortical collecting ducts, respectively. C, cortex; OS, outer stripe of the outer medulla; G, glomerulus; md, macula densa.

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steady-state value of ~ 200 nM attained after equilibration at 37°C in time-control tubules (Fig. 2). The progressive decline was a direct function of time after having reached 37°C, with the [Ca^{2+}]_i (nM) = 347 - 2.11 (min), r = -0.94, P < 0.0001. Our results suggest that absolute values for resting [Ca^{2+}] are relevant and interpretable only under defined experimental conditions, notably temperature.

EFFECTS OF PTH (FIG. 3)
In six tubules treated with 10 μM acetic acid, the vehicle for PTH, mean [Ca^{2+}]_i tended to fall from a basal value of 267±18 to 251±14 nM after 10 min of suffusion. In contrast, 0.1 nM PTH increased [Ca^{2+}]_i from a mean value of 278±35 to 311±36 nM (Δ = 33 nM, P < 0.01) within 4 min, and by 7 min a maximal increment of 68 nM (P < 0.005) was attained (Fig. 3). The effects of PTH were sustained at least 2 min beyond the removal of hormone from the suffusate, with an average increase in [Ca^{2+}]_i of 40 nM during the 4-min postexposure period, similar to the increment of 36 nM during the entire 10 min of PTH administration.

EFFECTS OF 8-Br-cAMP (FIG. 4)
To examine the potential role of adenylate cyclase, we added 8-Br-cAMP to the suffusate (Fig. 4). Suffusates without any cAMP, administered to six other tubules, were associated with a tendency for [Ca^{2+}]_i to fall during this 10-min period. 8-Br-cAMP mimicked the effects of PTH, increasing [Ca^{2+}]_i within the first 2 min of exposure (Δ = 31 nM, P < 0.001), and by 6 min a maximal increment of 84 nM (P < 0.005) was attained (Fig. 5). Similar to PTH, the effects of 8-Br-cAMP were sustained beyond the 10-min exposure: the mean increment in [Ca^{2+}]_i during the 4-min postexposure (Δ = 61 nM) was comparable to the average increment during the entire 10 min of infusion (Δ = 64 nM).

Perfused CNTs

SOURCES OF PTH-STIMULATED INCREMENTS IN [Ca^{2+}]
Role of intracellular Ca stores (Fig. 6). Deletion of Ca from both the bath and lumen reduced [Ca^{2+}]_i from mean baseline values of 146±18 nM (averaged over the first 5 min) and 139±18 nM (the second 5 min) to 70±17 nM and 58±15 nM (P < 0.05), respectively (Fig. 6). In the absence of media Ca, PTH produced no rise in [Ca^{2+}]_i, which tended to fall during the first (51±14 nM) and second (53±16 nM) 5-min periods of PTH exposure. In contrast, with 2 mM Ca in both the suffusate and perfusate, PTH promptly raised [Ca^{2+}]_i to 219±57 nM and 231±53 nM, respectively, in two subsequent 5-min periods. For the second 5-min period the increment was 177±38 nM when compared with the corresponding period in the preceding Ca-deletion phase (P < 0.05). Despite withdrawal of PTH the increase in [Ca^{2+}]_i persisted for another 5 min (237±55 nM), followed by a progressive decline, respectively, to 200±54, 154±44, 143±40, 124±34, and 121±31 nM in the ensuing consecutive 5-min periods (Fig. 6). Thus, similar to nonperfused tubules, PTH also increased [Ca^{2+}]_i in perfused CNTs. Importantly, this effect was absent when Ca was deleted from the media.

In two CNTs we suffused 5 μM ionomycin after similar durations of media Ca deletion to assess whether intracellular Ca stores were intact under these conditions. The Ca-sensitive fluorescence ratio [340:380] rose with ionomycin (from 0.58 to 0.63 in one tubule and from 0.58 to 0.67 in another) despite

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continued media Ca omission, arguing indirectly against depletion of intracellular Ca stores as the reason for a lack of response to PTH.

Roles of lumen and bath Ca (Figs. 7 and 8, Table I). By manipulating lumen Ca we evaluated the contributions of lumen and bath Ca (Fig. 7). During the two consecutive 5-min baseline periods, mean [Ca2+]i's were comparable between the control and PTH groups (Fig. 8). Selective deletion of lumen Ca reduced mean [Ca2+]i by 44 and 50 nM, respectively. Despite the continued absence of lumen Ca, PTH significantly increased [Ca2+]i by 24±6 nM in the second 5-min period, whereas [Ca2+]i tended to decline in the controls. Their divergent changes were highly significant. When 2 mM Ca was reintroduced in the perfusate, PTH produced a prompt additional increment in [Ca2+]i. This was sustained not only throughout the 10 min of exposure, but also for 5 min after PTH withdrawal. In control tubules lumen Ca repletion caused a numerical rise in [Ca2+]i (0.05 < P < 0.10). This was followed by a steady and continuous decline in the ensuing 30 min, similar to the nonperfused time-control tubules (Fig. 2).

Estimates of the contribution by bath Ca. The tendency for [Ca2+]i in control tubules to fall during lumen Ca deletion beyond the first 10 min (Table I) suggests that the increment in PTH-treated tubules, observed under identical conditions,
probably underestimated the contribution by bath Ca. Specifically, the calculated increment of 24 nM induced by PTH in the absence of lumen Ca assumed a steady value of 183 nM, as in the pre-PTH phase (Table I). This increment should be augmented by adjusting for the concomitant, though small, fall associated with sustained luminal Ca deletion, demonstrable in control but not in PTH-treated CNTs.

By extrapolating a linear regression (using 30 consecutive 10-s average [Ca$^{2+}$], values) from the last 5-min period of the second phase into the third phase (i.e., dotted line between minutes 20 and 30, as illustrated in Fig. 7), we derived a predicted value of 174±41 nM if no hormone had been given to PTH-treated tubules. After this adjustment the average increment in [Ca$^{2+}$], stimulated by PTH in the absence of lumen Ca, and hence attributable to bath Ca ($\delta_b$ in Fig. 7), was 33±8 nM (derived from the difference between the observed result of 207±47 [Table I] and the predicted value of 174±41 nM). This is greater than the similarly derived difference (−7±4 nM; $P < 0.005$) in control tubules manipulated identically but not exposed to PTH.

Estimates of the total increments in [Ca$^{2+}$], due to PTH in the presence of bath and lumen Ca. To derive these estimates a baseline value of [Ca$^{2+}$], temporally corresponding to no hormone treatment in the same tubules (200±43 nM) was obtained by interpolation between values measured during the last 5 min of the baseline and recovery phases (Fig. 7, dashed line). The total increment due to PTH ($\delta_{t}$ in Fig. 7) was 94±22 nM, derived from the difference between the observed result of 293 nM (Table I) and this predicted value of 200 nM. This is greater than the similarly derived difference (8±8 nM; $P < 0.01$) in control tubules manipulated identically but not treated with PTH.

Estimates of the contribution by lumen Ca. Because our results (Fig. 6) suggested a negligible role for intracellular Ca stores in mediating the PTH-induced rise in [Ca$^{2+}$], the total increment in [Ca$^{2+}$], was resolved into two components: that due to bath Ca and that due to lumen Ca. Subtraction of the bath Ca component from the total provides an estimate of the lumen Ca component, 61±16 nM, which is greater than that found in controls, 14±6 nM ($P < 0.04$). Analysis of data derived by this protocol suggests a greater contribution by lumen Ca than by bath Ca (61 vs. 33 nM). However, in two additional perfused CNTs in which the PTH-induced increment in [Ca$^{2+}$] was measured first in the presence of perfusate Ca and then in its absence, we found the contributions of lumen and bath Ca to be very similar (mean of 31 vs. 35 nM, respectively).

**Discussion**

We found that at concentrations similar to values seen in Ca or vitamin D-deprived rabbits (6, 24) PTH causes a sustained but reversible increase in [Ca$^{2+}$], in rabbit CNT cells. 8-Br-cAMP mimics PTH, but with an earlier onset of action, consistent with a messenger role for cAMP. Our studies demonstrate the feasibility of measuring [Ca$^{2+}$], with fura-2 in freshly dissected rabbit CNTs. During equilibration at 37°C, [Ca$^{2+}$], fell to levels reported in other kidney cells (14–23, 25). The slightly higher resting [Ca$^{2+}$] in CNTs might be explained by differences in species, methods of harvesting cells, nephron sites, and protocols, among others.

The effects of PTH on [Ca$^{2+}$] have been examined in other kidney-derived tissues (14, 19–21, 26). Except for two studies (21, 22), PTH was reported to increase [Ca$^{2+}$]. Qualitatively the increase in [Ca$^{2+}$], produced by PTH in the CNT is similar to these observations, since 0.1 nM PTH also elicited an effect in cultured proximal tubule cells (14, 20).

There are, however, several differences. First, the onset of PTH action in CNTs (requiring 4 min of exposure) was slower than the response (within seconds) in proximal tubules (14, 19, 20, 26). Second, studies in proximal tubules (19, 20, 26) using a superfusion apparatus and epifluorescence microscope (19) suggest a transient 1- to 3-min (26) spike in [Ca$^{2+}$], despite continued hormone exposure. In contrast, the PTH effect in CNTs was sustained during prolonged exposure and persisted 3 to 4 min after removal (Figs. 5, 6, and 8). Third, contrary to proximal tubule cells (14, 26), 8-Br-cAMP mimicked PTH in the CNT, similar to the increased 45Ca uptake by cultured monkey kidney cells when exposed to either PTH or dibutyryl-cAMP (27, 28). In the proximal tubule cells neither dibutyryl-cAMP, 3-isobutyl-1-methyloxanthine, nor forskolin reproduced the effects of PTH (14). These divergent responses to cAMP analogues may reflect differences in species, cell preparation, experimental protocol, or origin and function of the nephron segment. In addition, it is possible that PTH raises [Ca$^{2+}$], by both cAMP-dependent and -independent mechanisms, similar to PTH-sensitive osteosarcoma cells (29). The former produced a delayed but sustained rise in [Ca$^{2+}$], related directly to intracellular cAMP concentration. The latter caused a rapid but short-lived (1-min) rise in [Ca$^{2+}$]. In proximal tubules an immediate stimulatory effect of PTH on the hydrolysis of phosphatidylinositol 4,5-bisphosphate produces a rapid rise in inositol 1,4,5-trisphosphate and diacylglycerol, which are postulated to release intracellular Ca and elicit a 60-s Ca transient (20). These effects were not mimicked by cAMP. Despite the 0.5-s resolution of our measurement system we did not observe a spike response to PTH in the CNT. The delayed and sustained increase in [Ca$^{2+}$] in the CNT was, therefore, more comparable to the cAMP-dependent rise in [Ca$^{2+}$], in osteosarcoma cells (29) and to the increased 45Ca uptake in cultured kidney cells (28). Finally, it is possible that effects of PTH to alter solute transport and to signal cellular metabolism are mediated by different mechanisms. For example, glucagonosis (30) and membrane topography (31) may be al-

**Figure 8.** Response of [Ca$^{2+}$], to PTH in the absence and presence of luminal Ca in singly perfused CNTs. Suffusate [Ca$^+$] = 2 mM throughout. The tracings represent the means for six PTH-treated and five time-control CNTs.

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tered by PTH using [Ca²⁺], as a messenger, whereas inhibition of Na:K exchange, Na-dependent phosphate uptake, and increased absorptive Ca flux may be mediated by cAMP and thus reproducible by cAMP analogues (2, 21, 32–37), forskolin (32), and 3-isobutyl-1-methylxanthine (32). In isolated perfused proximal convoluted tubules, increasing [Ca²⁺], (23) by raising bath [Ca] did not alter lumen-to-bath phosphate flux (38), whereas the same maneuver stimulated gluconeogenesis (38).

The distal nephron is the primary site regulating Ca excretion (37, 39–42). The CNT is the most distal rabbit nephron segment demonstrating PTH-stimulated Ca reabsorption (2), which presumably occurs transepithelially. Hypothetically, if the stimulatory action of PTH on Ca transport is related to the increase in [Ca²⁺]], that we observed, lumen Ca ought to be the predominant contributor. Though our estimates indicate that bath Ca contributes substantially, lumen Ca does account for 50–65% of the PTH-induced increment in [Ca²⁺]], consistent with the above hypothesis.

A finding inconsistent with this hypothesis, however, was the increase in [Ca²⁺]], caused by PTH in the absence of lumen Ca (Fig. 8), implying that basolateral membrane Ca entry is also stimulated. Although this component was smaller than that due to lumen Ca when estimated by perfusing the tubule with zero Ca first, it was comparable when estimated by reversing the sequence of manipulation. The physiologic role of basolateral Ca entry is presently unknown, because it is in a direction opposite to PTH-stimulated Ca reabsorption. We speculate that it may “prime” basolateral plasma membrane Ca²⁺ extrusion by the high-affinity Ca-Mg-ATPase (and/or Na:Ca exchange) or may serve as a signal for these or other processes.

Our estimates of the roles of bath and lumen Ca in the PTH-induced rise in [Ca²⁺]], must be considered tentative for two reasons. First, our approach to correct for the effect of lumen Ca depletion and for the transient overshoot in [Ca²⁺]], during acute Ca repletion is only a first approximation. Second, the deionized water used to prepare the nominally 0-mM Ca perfusate was estimated to contain 0.9 μM Ca, so that the increment in [Ca²⁺]], assigned to bath Ca may be an overestimate.

Similar to other renal epithelia (14, 16, 19, 23), we observed that media Ca depletion rapidly and markedly reduced [Ca²⁺]], in the CNT, with a mean fall of ~ 80 nM (Fig. 6). Eliminating perfusate Ca while keeping sulfusate [Ca] at 2 mM attenuated this fall in [Ca²⁺]] (44–50 nM; Table I and Fig. 8), implying that Ca influxes across both apical and basolateral plasma membranes are important in regulating [Ca²⁺]], in the CNT. Without media Ca, PTH had no effect on [Ca²⁺]], (Fig. 6), mitigating against release of Ca from intracellular stores.

In conclusion, this study demonstrates that physiological concentrations of PTH cause a significant increase in [Ca²⁺]], in cells of the rabbit CNT. This action is mimicked by 8-Bromo-cAMP, implicating cAMP as an intracellular messenger. The PTH-stimulated rise in [Ca²⁺]], depends on extracellular fluid Ca, with bath and tubular luminal fluids both playing important roles.

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