SECRETAGOGUE INDUCED CALCIUM MOBILIZATION
IN SINGLE PANCREATIC ACINAR CELLS

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Received October 12, 1988

Summary: Microspectrofluorometry of fura-2 was utilized to monitor \([\text{Ca}^{2+}]_i\) in single acinar cells stimulated with a cholinergic agonist and cholecystokinin. A similar amplitude of agonist induced Ca mobilization between single cell and populational approaches was observed. New findings in single cells not observable in populations of cells include: 1) the maintenance of a sustained elevation in \([\text{Ca}^{2+}]_i\) above basal levels throughout agonist application, 2) the reloading of the agonist-sensitive Ca pool only following removal of the agonist and 3) the presence of oscillations of \([\text{Ca}^{2+}]_i\) in response to agonist application which is enhanced at lower agonist concentrations.

Acetylcholine and cholecystokinin stimulate pancreatic acinar cell enzyme secretion through generation of inositol 1,4,5-trisphosphate and 1,2-diacylglycerol which induce mobilization of calcium from intracellular stores and activation of protein kinase C respectively (1). In addition, sustained secretagogue stimulation apparently causes an activation of Ca influx across the plasma membrane. This latter mechanism is believed to remain activated following secretagogue removal thereby allowing entry of extracellular Ca and refilling of the intracellular agonist-sensitive Ca pool (2). Characterization of these changes in \([\text{Ca}^{2+}]_i\) have recently been studied using Ca-sensitive fluorescent indicators such as quin2 and fura-2 (3-8). To date, however, the use of fluorescent probes has been performed on populations of isolated acini under static conditions. This arrangement results in a number of experimental shortcomings. These include: 1) changes in \([\text{Ca}^{2+}]_i\) reflect an integrated response over a large number of cells which may not be representative of temporal changes in \([\text{Ca}^{2+}]_i\) in single acinar cells. 2) removal of agonist activation relies on application of an antagonist and 3) an increasing background dye signal may result from dye leakage and mechanical damage of a percentage of the stirred acini.

The purpose of the present study was to utilize dual wavelength microspectrofluorometry of fura-2 to examine the secretagogue induced changes in \([\text{Ca}^{2+}]_i\) in superfused single acinar cells. The results indicate that individual acinar cells possess a releasable Ca\(^{2+}\) pool responsive to multiple agonists. Heretofore unreported oscillations of intracellular Ca\(^{2+}\) were also observed.
MATERIALS AND METHODS

Acini were prepared from pancreas of male Sprague-Dawley rats (200-250 g) according to the collagenase digestion procedure detailed by Williams et al. (9). Isolated acini were incubated with 2 μM fura-2/AM at 37 °C in a physiological salt solution (PSS) containing (in mM): NaCl, 145; KCl, 5; CaSO_4, 1.8; MgSO_4, 1.0; glucose, 10; HEPES, 10; bovine serum albumin, 0.5 mg/ml; soybean trypsin inhibitor, 0.1 mg/ml; which was adjusted to pH 7.4 with NaOH. During the 15 min incubation period the acini received continuous oxygenation and shaking (30 cycles/min). Acini loaded with fura-2 were collected by centrifugation at 50 x g and resuspended in saline. PSS utilized during the fluorescence measurements was identical to that given above except that BSA and trypsin inhibitor were omitted. In addition nominally Ca-free PSS was prepared by omitting the CaSO_4 and adding 1 mM EGTA.

Measurement of [Ca^{2+}]_i. Fluorescence measurements of fura-2 loaded acini were performed on both populations of acinar cells as well as at the level of single acinar cells. Populational studies were performed by attaching acini to cover slips with Cell-Tak (Biopolymers; Farmington, CT) which were then inserted into a cuvette along the diagonal axis and placed into the spectrofluorometer chamber regulated to 37 °C. A superfusion system with a flow rate of 10 ml/min provided rapid exchange of the cuvette volume (2 ml). To measure fluorescence of single acinar cells, acini were attached to glass cover slips mounted in a chamber (0.1 ml volume) regulated at 37 °C which allowed continuous superfusion at 1 ml/min with switching of solutions. The chamber was mounted on the stage of a Nikon Diaphot inverted microscope equipped with a 40X oil immersion epifluorescence lens (NA = 1.3). Single acinar cells were optically masked using a pinhole diaphragm stopped down to an optical diameter of 10 μm.

All studies utilized a Spex fluorolog spectrofluorometer system to generate 340 and 380 nm excitation wavelengths and for collection and analysis of emitted photon counts. Emitted light was monitored at an emission wavelength of 505 nm for populational studies or following passage through a barrier filter (480 nm) for single cell experiments. In both cases photon counts were averaged over intervals of 0.5 sec at each wavelength. For populational studies intensity ratios (340/380 excitation) were calculated after subtracting autofluorescence of acini without fura-2 at each wavelength. Autofluorescence at the level of single acinar cells was, however, found to constitute less than 6% of the emitted signal and was not subtracted. For all studies, the fluorescence ratio (340/380) was converted to [Ca^{2+}]_i using the equation of Grynkiewicz et al. (10). For single cell studies, Kd, Fmin, Fmax, and Fo/Fs values of 224 nM, 0.8, 18.62 and 4.61 respectively, were used, as determined using cytoplasmic resembling solutions at fixed [Ca^{2+}]_i containing 25 M fura-2. For populational studies the formula 340/380 = -3.72(pCa) + 29.05 was used to calculate [Ca^{2+}]_i, as determined from the linear portion of the relationship, using solutions of fixed [Ca^{2+}]_i containing 1.2 μM fura-2.

RESULTS

Comparison of populational and single cell measurements.

A comparison of fluorescent measurements from the cover slip/cuvette approach, which averages fluorescence over a large number of cells, with those obtained from single acinar cells and of their responsiveness to the cholinergic agonist carbachol (CCh) is shown in Fig. 1. Mean basal and CCh stimulated [Ca^{2+}]_i values for the populational measurements were 114 ± 12 nM (n = 7, + SE, 6 preparations) and 420 ± 78 nM respectively. Basal and stimulated values determined in single cells were 136 ± 15 nM (n = 28, 6 preparations) and 754 ± 77 respectively. CCh stimulated increases in [Ca^{2+}]_i were reproducibly higher using the single cell approach. In addition both noise and
Figure 1. Comparison of CCh evoked changes in \([\text{Ca}^{2+}]_i\) between populational and single cell approaches. A) CCh induced change in \([\text{Ca}^{2+}]_i\) using a populational approach where the fluorescence signal is averaged from a large number of acini. B) Change in \([\text{Ca}^{2+}]_i\) in a single acinar cell in response to a similar pulse of CCh. In both instances 10 \(\mu\text{M}\) CCh was superfused over the cell preparation for a period of 4 min before return to PSS as indicated below the traces. Subsequent addition of 100 \(\mu\text{M}\) atropine elicits no further decrease in \([\text{Ca}^{2+}]_i\) substantiating complete washout of CCh.

Application of 10 \(\mu\text{M}\) CCh for 4 min resulted in a transient rise in \([\text{Ca}^{2+}]_i\) which returned to near basal values within 2 - 3 min although remaining elevated at plateau values averaging 61 \(\pm\) 7 nM (\(n = 28\)) above basal values for single cells. Removal of CCh rapidly returned \([\text{Ca}^{2+}]_i\) to prestimulatory values. Subsequent application of atropine after removal of the CCh showed no further change in \([\text{Ca}^{2+}]_i\) providing evidence for complete washout of the CCh stimulus.

Recovery of agonist-sensitive Ca pool following CCh stimulation.

The rapid reversibility of CCh application afforded by the superfusion system provides the opportunity to examine the responsiveness of single acinar cells to repetitive CCh stimuli following variable periods of recovery. Representative examples taken from single acinar cells receiving repetitive CCh pulses are shown in Fig. 2A-C. Recovery of the Ca response is clearly dependent on the length of the intervening interval between CCh tests with 65% recovery occurring after approximately 3 min (Fig. 2B) and greater than 90% recovery requiring 10 to 15 min (Fig. 2C and 2D). Reloading of the agonist-sensitive Ca pool following agonist removal occurs without an elevation in the cytoplasmic \([\text{Ca}^{2+}]_i\) suggesting either highly synchronized extracellular Ca entry and sequestration or of direct reloading from extracellular Ca into the agonist-sensitive pool. Particularly interesting in Fig. 2B (see also Fig. 4D) is the presence of oscillations in \([\text{Ca}^{2+}]_i\) during secretagogue application. Oscillations of this type at 10 \(\mu\text{M}\) CCh were observed in approximately 40% (\(n = 34\)) of cells examined at the level of the single cell but were never observed during secretagogue application in the populational studies.

Effect of extracellular Ca on CCh responses.

The effect of external Ca on the CCh induced increase in \([\text{Ca}^{2+}]_i\) and on responsiveness to a second CCh application is shown in Fig. 3. Application of CCh in Ca-free medium resulted in an increase in \([\text{Ca}^{2+}]_i\) whose amplitude showed close similarity to
that of acini in Ca-containing medium. Mean basal and stimulated \([Ca^{2+}]_i\) under these conditions were 129 ± 10 nM and 619 ± 69 nM (n = 5, 3 preparations) respectively. A second CCh challenge 10 min later, under maintained Ca-free conditions, resulted in a response on average only 12% (n = 3) that of the initial response. Similar results were

Figure 2. Time course of recovery of the agonist-sensitive Ca pool following CCh stimulation. Application of an initial CCh stimulation (4 min) is followed by a period in PSS of 0.5 min (A), 3 min (B) or 10 min (C) before application of a second stimulation with CCh. D) Pooled results from 3 preparations (different symbols) showing the time course of recovery of the agonist sensitive Ca pool. Percent recovery is calculated from the CCh induced change from basal \([Ca^{2+}]_i\) of the second CCh pulse to that of the first. Note oscillations of \([Ca^{2+}]_i\) in B which are augmented during the second CCh stimulus.

Figure 3. Effect of extracellular Ca on CCh evoked changes in \([Ca^{2+}]_i\), and recovery of the agonist sensitive Ca pool. A) Removal of extracellular Ca 0.5 min prior and during application of 10 \(\mu\)M CCh is ineffective in altering the rise in \([Ca^{2+}]_i\). Sustaining Ca-free PSS for 10 min following removal of the initial CCh pulse reduces reloading of the agonist-sensitive Ca pool as assessed by the amplitude of a second CCh pulse. B) Removal of extracellular Ca following an initial CCh pulse reduces reloading of the agonist-sensitive Ca pool to an extent similar to that shown in A.
Comparison and interaction between CCK and CCh induced changes in [Ca\(^{2+}\)]\(_i\). A) CCK (1 nM) evokes an increase in [Ca\(^{2+}\)]\(_i\) similar in amplitude to that exhibited by CCh. Note, however, that addition of CCh following CCK does not elicit a change in [Ca\(^{2+}\)]\(_i\). B) Application of CCK subsequent but coincident with a CCh induced increase in [Ca\(^{2+}\)]\(_i\) evokes no further change in [Ca\(^{2+}\)]\(_i\). C) Insertion of an interval in PSS following CCh stimulation allows partial recovery of the agonist sensitive Ca pool which responds to CCK stimulation. D) Application of CCK at a reduced concentration (0.1 nM) results in enhancement of oscillations of [Ca\(^{2+}\)]\(_i\) which are sustained throughout agonist application.

Interaction and comparison of CCh and CCK responses.

Application of 1 nM CCK (Fig. 4A) resulted in an increase in [Ca\(^{2+}\)]\(_i\) (683 ± 150 nM; n = 5, 3 preparations) similar in amplitude and shape to that of a maximal dose of CCh. Note, however, that the [Ca\(^{2+}\)]\(_i\) does not fully return to baseline on switching perfusion to normal saline indicating ineffective washout of CCK. A subsequent test with CCh resulted in no change in [Ca\(^{2+}\)]\(_i\) as would be expected for agonists operating to release Ca from a single intracellular pool that has previously been depleted by the CCK application. Similarly initial application of CCh followed, without an intervening interval in normal saline, with CCK resulted in no further change in [Ca\(^{2+}\)]\(_i\) (Fig. 4B). Insertion of an interval in normal saline subsequent to CCh application but prior to CCK stimulation resulted in an increase in [Ca\(^{2+}\)]\(_i\) by the second stimulus (Fig. 4C). The resulting increase by CCK likely occurs from partial refilling of the agonist sensitive Ca pool prior to CCK
stimulation. As shown in Fig. 4D exposure of acinar cells to 100 pM CCK resulted in a clear enhancement of oscillatory behavior of \([Ca^{2+}]_i\) in these cells which was sustained throughout the 10 min period of agonist application. Similar enhanced oscillations were also seen in response to a submaximal concentration of CCh (1 μM).

**DISCUSSION**

The present study compares measurements using the Ca-sensitive fluorescent probe fura-2, of \([Ca^{2+}]_i\) in populations of cells and by microspectrofluorometry of single acinar cells. Differences in basal and CCh stimulated \([Ca^{2+}]_i\) of acinar cells between the two approaches are not different from those of a number of previous reports on populations of acinar cells (3-8). Our results on single cells also substantiate the results of a number of others suggesting that CCh and CCK act to release Ca from the same intracellular agonist sensitive Ca pool as addition of a second agonist following a first elicits no additional rise in \([Ca^{2+}]_i\) (Fig. 4). These results, however, conclusively show for the first time that different agonists mobilize Ca within the same cell.

There are also several differences between \([Ca^{2+}]_i\) measurements obtained from single acinar cells and those obtained in prior studies from a fluorescence signal averaged over a large number of acini. One of these differences is the clear maintenance of an increased \([Ca^{2+}]_i\) above basal levels following the initial rapid rise in \([Ca^{2+}]_i\) induced by CCh or CCK. Previous reports utilizing fura-2 in stirred cells in a cuvette were interpreted as showing a return to basal \([Ca^{2+}]_i\) with sustained secretagogue application (2,6,8,11). Utilizing the single cell approach we have observed a sustained increase above basal \([Ca^{2+}]_i\) in every cell in which agonist induced Ca mobilization was observed. The increase above basal is maintained for agonist applications up to 10 min (data not shown). The elucidation of these finer changes in \([Ca^{2+}]_i\) near basal values was the result of a significantly higher signal to noise characteristic occurring as a result of reduced light scattering and autofluorescence in the superfused immobile acinar cells. A second difference we have observed relates to reloading of the agonist sensitive Ca pool. The lack of a significant difference between second responses to CCh, either with Ca present only during the initial CCh challenge, or absent throughout, suggests that reloading of the agonist sensitive Ca pool occurs only following removal of the agonist. This result is in contrast to that of Muallem et al. (11) who have reported that partial reloading of the agonist sensitive pool occurs during CCh application if extracellular Ca is present during agonist application. [This conclusion, however, was based on application of ionomycin, the effects of which are not clearly understood in pancreatic acinar cells.] A third unique finding is that of oscillations in \([Ca^{2+}]_i\) in response to agonist application. Clearly such oscillatory behavior is largely masked in populational studies as a result of the asynchrony of oscillations over the large number of cells which the fluorescent signal is measured. Oscillatory changes in \([Ca^{2+}]_i\) have recently been observed in single cell recording of
several cell types (12-15). We have further observed that this oscillatory behavior becomes more prominent at lower doses of Ca mobilizing agonists as shown by Jacob et al. (16) for endothelial cells.

ACKNOWLEDGMENTS: Supported by NIH grants DK-32994 (JAW) and fellowship DK-07692 (ELS).

REFERENCES