Double Patch Clamp Reveals That Transient Fusion (Kiss-and-Run) Is a Major Mechanism of Secretion in Calf Adrenal Chromaffin Cells: High Calcium Shifts the Mechanism from Kiss-and-Run to Complete Fusion

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Transient fusion ("kiss-and-run") is accepted as a mode of transmitter release both in central neurons and neuroendocrine cells, but the prevalence of this mechanism compared with full fusion is still in doubt. Using a novel double patch-clamp method (whole cell/cell attached), permitting the recording of unitary capacitance events while stimulating under a variety of conditions including action potentials, we show that transient fusion is the predominant (>90%) mode of secretion in calf adrenal chromaffin cells. Raising intracellular Ca$^{2+}$ concentration ([Ca$^2+$]) from 10 to 200 μM increases the incidence of full fusion events at the expense of transient fusion. Blocking rapid endocytosis that normally terminates transient fusion events also promotes full fusion events. Thus, [Ca$^2+$] controls the transition between transient and full fusion, each of which is coupled to different modes of endocytosis.

Key words: exocytosis; endocytosis; kiss-and-run; fusion pore; membrane capacitance; chromaffin cell

Introduction

Since the development of the quantal hypothesis of transmitter secretion by Del Castillo and Katz (1954), it has been widely assumed that full vesicular fusion is the principal mechanism of exocytosis at central synapses. Recovery of vesicular components for recycling is postulated to occur through a clathrin-coated vesicle-based pathway (Heuser and Reese, 1973; De Camilli et al., 2000; Murthy and De Camilli, 2003; Dickman et al., 2005). Although this is still the prevailing dogma, an accelerating stream of evidence supporting alternative transient fusion mechanisms in neurons and neuroendocrine cells has emerged. Transient fusion (often known as "kiss-and-run") is envisaged to occur via a fusion pore between the vesicle and the surface membrane (Neher and Marty, 1982; Fernandez et al., 1984; Alvarez de Toledo et al., 1993). Transmitter or hormone can escape, often incompletely, through this aqueous channel between the vesicle interior and the extracellular space (Elhamdani et al., 2001; Aravanis et al., 2003; Taraska et al., 2003; Pawlu et al., 2004; Staal et al., 2004). The molecular nature of the fusion pore is still in doubt, although proteinaceous models involving SNARE (soluble N-ethylmaleimide-sensitive factor attachment protein receptor) proteins have been proposed recently (Han et al., 2004). Because both transient and full fusion evidently occur in most secretory cell types, a central question has been the relative preponderance of these alternate mechanisms, their relationship to stimulation conditions and indeed to each other, as well as the endocytotic mechanisms that terminate each process. In adrenal chromaffin (AC) cells that secrete the transmitter epinephrine from dense-core vesicles (DCVs), we have proposed that different endocytotic processes are evoked by distinct patterns of stimulation. Rapid endocytosis (RE), a non-clathrin-dependent event, occurs with mild stimulation (Artalejo et al., 1995, 1996, 2002), whereas stronger and sustained levels of stimulation trigger a clathrin-dependent slow endocytosis (SE) (Artalejo et al., 2002). We hypothesized that RE is the terminal step of transient fusion, whereas SE is activated after full fusion events in these cells (Palfrey and Artalejo, 1998; Elhamdani et al., 2001). Here, we use a double patch-clamp approach to assess at the unitary event level transient versus full fusion in AC cells and their relationship to the two modes of endocytosis. We show that transient fusion is the major mechanism of secretion and that RE is intimately associated with this process. Full fusion events increase at levels of stimulation that generate high intracellular Ca$^{2+}$ concentration ([Ca$^2+$]) or when RE is abrogated by ion substitution.

Materials and Methods

Cell culture

Bovine calf (average age, 10–12 weeks) AC cells were prepared by collagenase digestion of adrenal medulla obtained from a local slaughterhouse. Cells were purified and cultured using previously described methods (Artalejo et al., 1995). AC cells, plated at a density of 3 × 10$^5$ cells on collagen-coated 35 mm diameter dishes, were used in all studies within 1 week of plating.
Electrophysiology

Cell-attached capacitance measurements. Sine-wave stimuli were generated from a lock-in amplifier (SR830 DSP; Stanford Research Systems, Stanford, CA); the signal-to-noise ratio was optimized using a sine wave of 50 mV amplitude and 20 kHz frequency (Debus and Lindau, 2000). The current evoked at the patch was filtered at 10 kHz and driven back from an EPC-10/2 (HEKA Elektronik, Lambrecht, Germany) to the lock-in amplifier input. The in-phase (real) and 90° out-of-phase (imaginary) outputs of the lock-in amplifier, corresponding to the conductance and capacitance of the patch, respectively, were digitized using a Digidata BNC-2090 (National Instruments, Austin, TX) and stored at a 1 kHz sampling rate in a personal computer. The output filter of the lock-in amplifier input was set to a time constant of 1 ms, 12 dB. Data were acquired and analyzed using Igor (WaveMetrics, Lake Oswego, OR), and macros were developed in Dr. M. Lindau’s laboratory (Cornell University, Ithaca, NY).

In cell-attached capacitance measurements, two parameters were monitored carefully. First, the formation of an Ω shape of the membrane inside the cell-attached pipette was avoided by applying minimum suction to obtain the seal between a freshly polished pipette and a clean cell membrane. Also, true changes of capacitance and conductance were assured by setting the correct phase and monitoring the effect of ion channel activity on the conductance trace. Two manipulations that affect the phase in a characteristic manner were used in setting the phase: a 100 mV jump of the fast capacitance (Cf) and a slight suction force on the patch applied by mouth. Both should increase the capacitance without affecting the conductance of the patch, and they proved to be ideal tools for phase adjustment. To preclude ion channel activity effects on the conductance trace, patch current was recorded in parallel with real and imaginary traces.

For transient fusion events, vesicle capacitance (Cv) and fusion-pore conductance (Gp) were calculated from the imaginary (Im) and real (Re) traces corresponding to the admittance of the patch. Cv = [(Re2 + Im2)/Im]/ω and Gp = (Re2 + Im2)/Re. For full fusion events, Cv = Im/ω, where ω is the angular frequency (ω = 2πf) of the sine-wave frequency. Occasionally, the fusion pore expanded slowly for a short period of time before full fusion. In this case, the conductance of the fusion pore was calculated from the Im trace only; Gp = Im/(Im/Im − 1) − 1/2 [see Dernick et al. (2003) for a full description].

Pipettes were constructed of thin-wall borosilicate patch pipette glass. The resistance of the pipette was ≤1 MΩ, and resistance of the seal was ≥20 MΩ. The bath solution consisted of (in mM) 140 NaCl, 10 glucose, 10 HEPES, 1 MgCl2, 5 KCl, and 5 CaCl2, pH adjusted to 7.3 with NaOH. The cell-attached pipette solution is the same as the bath solution. When 10 mM Sr2+ or Ba2+ was substituted for 5 mM Ca2+, NaCl was reduced to 130 mM, 1.1-Dimethyl-4-phenylpiperazinium isodide (DMPP) (final concentration was added daily to the pipette solution from a fresh 10 mM stock solution). For cell-attached capacitance experiments with DMPP, the tip of the cell-attached pipette was dipped for ~3 s in a DMPP-free pipette solution. Thereafter, the pipette was back filled with the same solution containing DMPP. The seal is made relatively fast, which reduces significantly the chance of the cell being exposed to DMPP from the pipette before the seal is secured.

Whole-cell capacitance measurements. For the double patch-clamp procedure, we used an EPC-10/2 amplifier to simultaneously perform cell-attached and whole-cell recordings. Whole-cell membrane capacitance was evoked by a 50 mV (root mean square) sine wave at 1500 Hz using the manufacturer’s Pulse software (HEKA Electronik). RE was evoked with a brief stimulation protocol (10 × 50 ms) pulse at 2 Hz, and SE was evoked with a sustained stimulation protocol (29 × 75 ms) pulse at 0.25 Hz. Each stimulation pulse corresponded to a depolarization of the cell from a holding potential of ~80 to +10 mV, which is preceded by a 50 ms prepulse to +120 mV to recruit the facilitation Ca2+ channels (Artalejo et al., 2002). The patch pipette contained (in mm) 100 K-glutamate, 12 NaCl, 30 HEPES, 5 MgCl2, 2 ATP, 0.35 GTP, pH 7.2 adjusted with KOH. The external solution consisted of (in mM) 140 NaCl, 10 glucose, 10 HEPES, 1 MgCl2, 4 KCl, and 5 CaCl2, pH 7.3 adjusted with NaOH. All experiments were performed at room temperature (24°C). Data are mean ± SEM. Statistical significance was evaluated using the Peritz test.

Results

Previously, we used whole-cell capacitance measurements in calf AC cells to assess global exocytosis and endocytosis events as well as amperometry to detect the kinetics of unitary fusion events (Artalejo et al., 1995, 1996, 2002; Elhamdani et al., 1998, 1999, 2001). Here, we have improved this procedure to double patch-clamp cells to precisely control the intracellular milieu with a whole-cell pipette while simultaneously recording unitary capacitance events in the cell-attached configuration. In the latter procedure, noise is sufficiently low that fusion of single DCVs can be resolved readily. To determine the efficacy of cell-attached recordings in the present setup, we first stimulated cells with the nicotinic agonist DMPP (10 μM) using a single cell-attached electrode. As shown in Figure 1, individual fusion events were clearly resolved. It was immediately apparent that the majority of the events represented transient fusion. In fact, from a total of 29 cells in which at least one secretory event was recorded, 169 transient fusion events, but only 18 full fusion events, were obtained. That each transient event is derived from a single vesicle fusion is predicated on the observation that the coefficient of correlation between the up and subsequent down step is close to 1 (data not shown). Full fusion was classified as a jump in the capacitance trace that did not return to baseline during the course of the recording, accompanied by no change or a brief transient in the resistance trace. Three parameters are used to describe the characteristics of the transient events: vesicle capacitance (Cv), fusion pore conductance (Gp), and the event duration. Cv and Gp were calculated from the imaginary (Im) and real (Re) traces (Fig. 1B) using equations described in Materials and Methods, and the distributions of these parameters for transient fusion events are shown in Figure 1C. The average Cv was 2.1 ± 0.44 fF (range, 1–3.5 fF), close to the estimated average for a single DCV in AC cells (Neher and Marty, 1982). Assuming a membrane capacitance value of 9 fF/μm2, the mean vesicle radius was 130 ± 1.3 nm, also in line with previous microscopic determinations of this parameter in AC cells stimulated with nicotine (Ornberg et al., 1995). These results support the idea that we are examining fusion of single vesicles with the surface membrane in the cell-attached recordings. The mean Cv was 250 ± 5.4 pS (range, 70–500 pS), which is slightly lower than the values reported for DCVs in posterior pituitary cells (~300 pS [Klyachko and Jackson, 2002]) and adult bovine AC cells (~330 or ~400 pS [Alès et al., 1999; Dernick et al., 2003]). A histogram distribution of unitary event duration exhibits a mean value of 223 ± 15 ms (range, 26–962 ms). In this study, we did not separate between burst of events from the same or different vesicles. Henkel et al. (2000)
defined a burst from the same vesicle as at least three events occurring in <3 s. However, as shown in Figure 3Bb, four successive events from at least two different granules (two transients and two full fusion events) did occur in <1 s.

Full fusion events had a similar size ($C_v = 1.98 \pm 0.17$ fF; $n = 18$) to transient fusion events (data not shown), but before merging completely with the plasma membrane, some vesicles (8 of 18) formed a significantly larger fusion pore ($G_p = 358 \pm 50$ pS) that stabilizes for a few milliseconds before full expansion.

APs trigger predominantly transient fusion events

Having established the ability of the cell-attached pipette to monitor single fusion events in calf AC cells, we then turned to a double patch-clamp approach in which the second whole-cell electrode is used to stimulate the cell with APs and control $[\text{Ca}]_i$, in a precise manner via intracellular buffer infusion (Fig. 2A). In this configuration, we found that APs at 1 or 7 Hz trigger single exocytotic events almost entirely by a transient fusion mechanism (Fig. 2B). APs at 1 Hz triggered 69 transient fusion events gathered from 15 cells, with no full fusion events that were apparent at this frequency. APs at 7 Hz triggered more full fusion events ($n = 14$), but transient fusion still predominated ($n = 150$ in 19 cells). Cumulative data from transient fusion events are shown in Figure 2C as histograms. The mean values of $C_v$ are statistically indistinguishable in each case, but APs at 7 Hz slightly increased the mean value of $G_p$ from 159 ± 4.3 to 179 ± 5.1 pS ($p < 0.02$) and reduced the mean unitary event duration significantly from 195 ± 28 to 82 ± 6 ms ($p < 0.001$). Previously, amperometry data in AC cells showed that the duration of the entire amperometric spike was shortened, and the total charge (amount of catecholamine released per spike) was significantly elevated when AP frequency was raised from 1 to 7 Hz (Elhamdani et al., 2001) or 0.5–15 Hz (Fulop et al., 2005). Furthermore, Fulop et al. (2005) showed that bigger fluorescent molecules (7 and 12 nm diameter) could be uptaken into the cell via a bigger fusion pore opening at 15 Hz but not 0.5 Hz. Together with the present results, this suggested the possibility that higher $[\text{Ca}]_i$, consequent after an increase in stimulation frequency, increases the fusion pore conductance and speeds up its closure, as well as promotes full fusion events. To test these hypotheses, we stimulated cells in the presence of different levels of free $[\text{Ca}]_i$, clamped with buffers.

Figure 1. DMPP triggers burst and isolated single secretory events. A. Real and imaginary traces, evoked by 10 μM DMPP, correspond, respectively, to the conductance and capacitance of the patch under a cell-attached pipette, as shown in the inset. B. Histogram of the conductance (Re) and capacitance (Im) of transient fusion events are calculated from real and imaginary traces (see Materials and Methods). C. Histogram of the $G_p$, $C_v$, and event duration ($n = 169$; 18 cells) with their corresponding mean values. When DMPP was omitted from the pipette solution, from 141 cells tested, only 6 cells had a total of three transient fusion events and seven full fusion events (data not shown), indicating that cells were primarily quiescent before stimulation. DMPP triggered the first event with a mean delay time of 167 ± 28 s (see Materials and Methods). Increasing DMPP to 100 μM did not decrease the delay time or increase the number of events. Re, Real; Im, imaginary.

Figure 2. APs trigger predominantly transient fusion events. A. Diagram showing the double patch-clamp approach: the left patch pipette in the cell-attached configuration (CA) to record capacitance and conductance of single secretory events and the right pipette in the whole-cell configuration (WC) to stimulate the cell with APs. N, Nucleus. B. Conductance (Re) and capacitance (Im) traces show transient fusion events evoked by APs shown underneath at 1 Hz (69 events, 15 cells) and 7 Hz (150 events, 19 cells). Each cell was stimulated with different rounds of 150 or 500 APs, separated by a 4 min resting period, at 1 or 7 Hz, respectively.

Different levels of free $[\text{Ca}]_i$ dictates the fate of fusing vesicles

Different levels of free $[\text{Ca}]_i$ were delivered to the cell interior through the whole-cell pipette, and exocytosis was monitored by concurrently recording both whole-cell and cell-attached capac-
endocytosis. Indeed, during these experiments, it was evident that increasing [Ca]i led to significant changes in the kinetics of RE as detected in whole-cell capacitance recording. Our standard protocol that triggers exocytosis coupled to RE involves short trains of stimuli (10 × 50 ms) and in calf AC cells can be reproducibly evoked at 1 min after establishing whole-cell configuration and every 5 min thereafter (Artailejo et al., 1995, 1996). Indeed, RE is readily apparent and quite reproducible at 10 μM [Ca]i but is already compromised by the second stimulation at 102 μM [Ca]i and absent altogether at 210 μM [Ca]i (Fig. 3, compare insets in a1, b1, and c1). Analysis of transient fusion events show that increasing the [Ca]i from 10 μM (n = 17) to 102 μM (n = 20) does not significantly affect the Cc3 (2.17 ± 0.19 vs 2.08 ± 0.09 fF) or Gv (250 ± 17 vs 246 ± 18 pS). However, the mean duration was significantly reduced from 227 ± 60 ms (range, 26–780 ms) to 62 ± 13 ms (range, 21–230 ms). Given that transient events were virtually absent at 210 μM [Ca]i (Fig. 4A) and this pattern paralleled the disappearance of RE as a function of [Ca]i, it immediately raised the question of whether RE is mechanistically associated with transient fusion and that when RE is abrogated full fusion becomes the norm. To answer this question, we used two stimulation protocols that selectively and reproducibly trigger RE or SE.

Kiss-and-run is linked to RE, whereas full fusion is linked to SE

Previously, we showed that brief (10 × 50 ms) and sustained (29 × 75 ms) physiological stimulations (square voltage depolarization) trigger two modes of endocytosis dubbed RE and SE, respectively (Artailejo et al., 2002). We also found that replacement of extracellular Ca2+ with Ba2+ or Sr2+ did not affect the extent of secretion or SE significantly but inhibited RE completely (Artailejo et al., 1995, 1996, 2002). Both of these paradigms were used to investigate the connection between RE and transient fusion events. Control stimulation with 10 × 50 ms pulses again showed reproducible RE in the whole-cell mode (Fig. 5Aa1) and only transient fusion events in the cell-attached patch (Fig. 5Aa2). From a total of 25 transient fusion events gathered from 12 cells, the Cc1 was 1.9 ± 0.1 fF, the Gv was 135 ± 9 pS, and the event duration was 70 ± 8 ms. Interestingly, these transient events have a similar duration as in the two previous conditions, raising [Ca]i to 102 μM or increasing the frequency of AP stimulation to 7 Hz. All three conditions speed up the closure of the fusion pore to a similar magnitude (∼70 ms). By analogy to the release of catecholamines from DCVs, the mean duration of the amperometric spikes (rising phase plus falling phase) evoked at 7 Hz APs was 63 ms (Table 1) (Elhamdani et al., 2001).

In contrast, when Sr2+ replaced bath Ca2+, the 10 × 50 ms pulse train resulted in only exocytosis with RE being completely blocked (Fig. 5Bb1) (cf. Artailejo et al., 1995, 1996). At the same time in the cell-attached patch, only full fusion events were recorded (Fig. 5Bb2). The blockade of RE does not affect fusing vesicle size (Cv = 1.85 ± 0.13; n = 23 events from nine cells).
However, the conductance of the fusion pore stabilizes at 415 ± 40 pS for a few milliseconds before full expansion. Furthermore, when more sustained stimulation was used (29 × 75 ms pulses) in the presence of Ca^{2+} as charge carrier, the robust C_m increase was followed by a slow decrease in C_m (SE) that takes ~11 min to return to the baseline value (Fig. 5C). Again, the blockade of RE does not affect the vesicle size (C_v = 1.8 ± 0.1 fF; n = 34 gathered from 14 cells), and in 14 of 34 events, which occurred between stimulations, the G_p stabilizes at 486 ± 47 pS for a few milliseconds before full fusion. All full fusion events recorded in this condition were evoked between the 20th and 29th stimulation pulses. These results strongly suggest that, at moderate physiological stimulations, RE is intimately coupled to the kiss-and-run type of secretion, whereas full fusion events are followed by SE at more sustained stimulations.

**Discussion**

Use of the double patch-clamp approach described in this study has allowed us to determine the kinetics of different modes of secretion under precisely controlled conditions in calf AC cells. Although the success rate of the procedure is quite low (5–12% of cells patched), necessitating the testing of large numbers of cells (>1300 cells were patched, 128 cells were used in the present study), the advantages conferred by this configuration are substantial. The major conclusions we draw are as follows: (1) transient fusion (kiss-and-run) is the major mechanism of vesicle exocytosis under moderate stimulation conditions; (2) sustained stimulation favors the incidence of full fusion events at the expense of transient events; (3) stimulation frequency controls the open time and the conductance of the fusion pore; (4) high [Ca^{2+}] favors full fusion, not kiss-and-run; and (5) RE is associated with kiss-and-run events when RE is abrogated, full fusion takes place and vesicles are recovered by SE), a process that we previously showed is similar to clathrin-coated vesicle endocytosis.

Although transient fusion has recently become more accepted as an alternative mode of secretion to the traditional full fusion model, the lack of direct measurement of fusion pore kinetics under physiological conditions has hindered an evaluation of the prevalence of this mechanism. For example, patch-amperometry studies in rat chromaffin cells claimed that only nonphysiological maneuvers, such as raising external [Ca] to very high levels (90 mM), promoted the incidence of kiss-and-run events, suggesting that it may be a minor player under normal physiological circumstances (Alés et al., 1999). A major difference between the present study and the study of Alés et al. (1999) is that the latter results were based solely on spontaneous release, whereas our data reflect release characteristics of cells stimulated in a physiological manner. Thus, although differences between the two sets of observations may be partly attributable to species considerations, our ability to fully control stimulation and [Ca] parameters lends credence to our conclusions. Indeed, calf AC cells manifest insignificant spontaneous release, as monitored at the level of a single secretory event in the cell-attached mode (present data) or as found in amperometric recordings (Elhamdani et al., 1998, 1999, 2001). A similar controversy between the relative importance of transient and full fusion mechanisms has emerged in the synaptic literature. On the one hand, cell-attached capacitance experiments in nonstimulated posterior pituitary nerve terminals suggested that transient fusion represented only 5% of the total number of fusion events under spontaneous release conditions. However, 55% of the patches tested in that study did not show any further activity after depolarization with high KCl, suggesting that initial [Ca] levels evoking release were already high (Klyachko and Jackson, 2002). Similarly, vesicle recycling in synaptic pHluorin transgenic mice studies claimed that fast recycling in hippocampal neurons is rare based only on one sustained stimulation protocol, APs at 20 Hz (Li et al., 2005). In contrast, recent dye-release experiments performed under mild stimulation conditions (low probability of release) in hippocampal neurons showed that transient fusion may comprise 70–85% of all events at this excitatory synapse (Richards et al., 2005) (see also Aravanis et al., 2003; Gandhi and Stevens, 2003). Similarly in AC cells, a combination of amperometry and imaging of the uptake of different fluorescent molecules of different sizes, evoked by APs at 0.5 and 15 Hz, was used to probe the size of the vesicle fusion pore (Fulop et al., 2005). Fulop et al. (2005) showed the existence of mainly the kiss-and-run type of secretion at 0.5 Hz and full fusion at 15 Hz.

We suggest that many studies showing a low incidence of the transient fusion mode of secretion rely mostly on spontaneous release or sustained stimulation (e.g., bath application of high KCl) and do not accurately represent normal stimulation conditions.

Another important conclusion that we demonstrate here is a direct connection between RE and kiss-and-run events. Although
we hypothesized that this was likely true based on circumstantial considerations in our previous work (Elhamdani et al., 2001; for review, see Artalejo et al., 1998), the direct correlation between loss of RE and transient fusion events in cells stimulated with Sr2+ as charge carrier or sustained stimulations indicates that this form of endocytosis is normally part of the kiss-and-run mechanism.

The present data prompts us to elaborate a model of transient and full fusion of DCVs evoked by physiological stimulation in calf AC cells (supplemental Fig. 1, available at www.jneurosci.org as supplemental material). Previous studies have shown the existence of a pool of DCVs in AC cells that are already docked and primed and fuse with the plasma membrane after an elevation in [Ca2+]. However, it is becoming clear that the mode and pattern of Ca2+ elevation also plays a role in determining the type of fusion that takes place. In our model, different [Ca2+]i governed by different patterns of stimulation give rise to at least three modes of fusion and vesicle recycling. (1) At low stimulation frequency (low [Ca2+]), vesicles fuse with plasma membrane for a longer duration (~200 ms) through a narrow fusion pore. This is in line with our previous amperometry data showing that the amperometric spikes evoked at low frequency or low external [Ca2+] are smaller in amplitude but wider than those evoked at higher frequencies (Elhamdani et al., 2001). Interestingly, these spikes were not affected by antagonism of dynamin function, suggesting that the closure of the fusion pore in “mode 1” may simply be a reversal of the fusion reaction, possibly similar to that found in synaptic terminals (Ceccarelli et al., 1973; Stevens and Williams, 2000). AC cells (Graham et al., 2002), and PC12 cells (Holroyd et al., 2002; for review, see Südhof, 2004). (2) At moderate stimulation (medium [Ca2+]), DCVs fuse transiently but for a shorter duration (~70 ms) and higher pore conductance. Because RE involves Ca2+ and dynamin-1, it is most likely that this level of [Ca2+] corresponds to a specific threshold, which could trigger the formation of a dynamin-1 ring around the neck of the retracting vesicle, facilitating the closure of the fusion pore and initiating fission. This hypothesis is also supported by the fact that when RE is abrogated with sustained stimulations or high free [Ca2+], only full fusion events were seen at the expense of kiss-and-run events. (3) At sustained stimulation (high [Ca2+]), dynamin-1-dependent RE is blocked. Therefore, the absence of dynamin-1 rings around the neck of the vesicle might allow the fusion pore, which is already at high conductance in presence of high Ca2+, to expand until full collapse of the vesicle with the plasma membrane. These vesicles will be retrieved by the clathrin- and dynamin-2-dependent SE and recycled through the endosomal compartment.

What is the mechanism by which [Ca2+] regulates the switch between RE and SE, or between transient and full fusion? Although this is still unknown, a prime candidate might be the Ca2+-sensitive phosphatase calcineurin. We suggested previously (Artalejo et al., 2002) that calcineurin-dependent dephosphorylation of dynamin-1 might reduce its ability to mediate RE (key factors here may be the dependence of dynamin-1 GTPase activity on phosphorylation state and/or the ability of dynamin-1 to reassociate with its anchoring protein amphiphysin when dephosphorylated). Thus, high Ca2+ might promote rapid dephosphorylation of dynamin-1 and compromise RE, thus biasing the system toward other means of endocytosis. Indeed, our previous studies using calcineurin antagonists showed that blocking this enzyme in calf AC cells caused an acceleration of RE as reflected in whole-cell membrane capacitance recordings (Artalejo et al., 1996). Similarly, blockade of calcineurin function accelerates vesicle recycling in nerve terminals of Drosophila larvae (Kuromi et al., 1997, Kuromi and Kidokoro, 1999).

We believe the present results might be relevant to synaptic transmission. Until recently, the bulk of evidence relating to the plasticity of neurotransmission concerned the postsynaptic and synaptic cleft compartments. Now that attention has shifted to the presynaptic arena (Krupa and Liu, 2004), it is apparent that regulation of the fusion pore may be a key element in the regulation of quantal size (Elhamdani et al., 2001; Aravanis et al., 2003; Pawlu et al., 2004; Staal et al., 2004) and synaptic strength (Choi et al., 2000; Renger et al., 2001; Zharkova et al., 2002). In light of our present data, we suggest that the opening duration and the conductance of the fusion pore are governed by the pattern of stimulation and thereby dictate the rate and amount of transmitter release. As shown in supplemental Figure 1 (available at www.jneurosci.org as supplemental material), mode 1 is mediated by a smaller pore and longer open time. Therefore, a small amount of transmitter will be released for a longer duration. At low activity, this mode of release is important to maintain a low basal level of release with few vesicles, especially if the release ready pool contains a small number of vesicles. In synaptic terminals, this mode of release has been suggested to play an important role in maintaining the synapse silent by desensitization of the receptors at the postsynaptic level (Choi et al., 2000). During mode 2, secretion is very efficient, and the same vesicle is retrieved intact and could be reused after refilling with transmitter from the cytoplasm. Indeed fluorescent “timer” proteins show that the youngest vesicles in the release ready pool are released first by nicotinic stimulation of AC cells (Duncan et al., 2003). In synaptic terminals, a brief release of glutamate at a high rate could turn a glutamatergic silent synapse into an active one (Choi et al., 2000). Under sustained stimulation (mode 3), vesicles will dump their entire content into the bloodstream or the synaptic cleft and the fusion pore expands irreversibly promoting full vesicle fusion with the surface membrane. As shown in many classical studies on vesicle recycling under tetanic conditions, vesicle retrieval is via a coated vesicle route with an obligate endosomal intermediate required to regenerate the synaptic vesicle population (Heuser and Reese, 1973). We have shown that this pathway coexists with other mechanistically distinct routes for vesicle retrieval that may play key roles in regulating transmitter release under physiological conditions.

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