Continuous and Transient Vesicle Cycling at a Ribbon Synapse

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Optical methods were used to study the Ca\(^{2+}\) dependence of vesicle cycling in bipolar cells isolated from goldfish retinas. Uniformly raising the Ca\(^{2+}\) concentration to between 0.8 and 20 \(\mu\)M produced a continuous vesicle cycle of balanced exocytosis and endocytosis with a maximum rate equivalent to the turnover of the entire surface membrane of a terminal every 2 min (or \(\sim 900\) vesicles sec\(^{-1}\)). Increasing the Ca\(^{2+}\) concentration above 20 \(\mu\)M inhibited continuous vesicle cycling. In contrast, influx of Ca\(^{2+}\) through voltage-gated channels produced a transient burst of exocytosis that increased the surface area of a terminal by a maximum of 12\% (equivalent to the addition of 13,000 vesicles). Endocytosis was delayed until after Ca\(^{2+}\) influx stopped and the average Ca\(^{2+}\) concentration in the terminal declined. Hence, a single terminal has mechanisms for both continuous and transient vesicle cycling.

Key words: exocytosis; endocytosis; vesicle cycling; synaptic vesicle; bipolar cell; retina

Ca\(^{2+}\) influx has long been known to be an effective trigger for regulated exocytosis (Katz and Miledi, 1967). The coupling between Ca\(^{2+}\) and fusion has been studied extensively in neuroendocrine cells. Most studies have demonstrated that Ca\(^{2+}\) concentrations above a few micromolar are sufficient to trigger exocytosis (see, for example, Jankowski et al., 1992). Less is known about vesicle cycling in presynaptic nerve endings, which are usually small and consequently have been experimentally inaccessible. Only recently have the Ca\(^{2+}\) transiently released vesicles containing ribbon synapses. A change in membrane capacitance has been used to monitor vesicle fusion in retinal bipolar and photoreceptor cells. Surprisingly, two different views have emerged. Goldfish bipolar cells (Heidelberger et al., 1994; von Gersdorff and Matthews, 1994a) transiently released vesicles when the Ca\(^{2+}\) concentration exceeded 20 \(\mu\)M. In contrast, salamander rod photoreceptors (Rieke and Schwartz, 1996) sustained a continuous vesicle cycle when the Ca\(^{2+}\) concentration rose above 1 \(\mu\)M. A similar conclusion was reached after staining goldfish bipolar cells with a fluorescent dye and following the fate of tagged vesicles (Lagnado et al., 1996) (however, see the Appendix). Thus, the experiments paint pictures of synaptic vesicle cycling that differ in two important respects. First, the Ca\(^{2+}\) concentrations required for release differ >20-fold. Second, the entire process of release and vesicle resupply is either transient or continuous.

We have now reconciled these divergent views. Vesicle cycling was observed after bipolar cell membranes were stained with FM1–43 or FM4–64. Relatively low Ca\(^{2+}\) concentrations (0.8–20 \(\mu\)M) stimulated exocytosis and sustained a continuous vesicle cycle with maximal rates sufficient to replace the entire surface of a terminal every 2 min. Higher concentrations (>20 \(\mu\)M) inhibited continuous cycling. Large increases in the local concentration near voltage-gated channels produced a transient burst of exocytosis. A rapid wave of compensatory endocytosis was initiated after Ca\(^{2+}\) influx stopped and the average Ca\(^{2+}\) concentration declined.

MATERIALS AND METHODS

Cell isolation. Goldfish (Carassius auratus) were killed by severing the cervical spinal cord and pithing the brain, following guidelines of the University of Chicago Institutional Animal Care and Use Committee. The procedure for dissociating a retina and maintaining solitary cells was similar to that described by Bader et al. (1982). Solitary bipolar cells were identified by their characteristic morphology (Fig. 1). In our initial experiments, dissociated cells were plated into dishes whose bottom was a glass coverslip. A styrly dye stained cells and adsorbed to the glass coverslip. Unbinding of the dye from the coverslip was very slow. Quantitative measurement of vesicle cycling was difficult when images were contaminated by a large background signal. Fortunately, unbinding of the dye was much faster when cells were plated onto a coverslip that had been coated with a thin layer of silicone plastic (Sylgard 184; Dow Corning, Midland, MI). Cells adhered to the silicone plastic if it was first covered with 0.25 mg ml\(^{-1}\) concanavalin A in 1 m NaCl for 20 min, rinsed with distilled water, and air dried.

The extracellular saline contained (in mM): NaCl, 120; KCl, 2.5; CaCl\(_2\), 0.2; MgCl\(_2\), 1; Na-pyruvate, 2; Na-lactate, 2; glucose, 10; and HEPES, 10; the pH was adjusted to 7.4 with NaOH, and the saline was supplemented with vitamins (1:100) and amino acids (1:100) formulated for Eagle’s minimum essential medium (Life Technologies, Gaithersburg, MD). The osmolality was \(\sim 260\) mOsm kg\(^{-1}\). Cells were loaded with fura-2 by incubating in normal extracellular saline supplemented with 2 \(\mu\)M fura-2 AM plus 16 \(\mu\)M pluronit acid for 1 hr. For some experiments (see Fig. 4), cells were loaded with mag-Fura-5 and fluo-3 by incubating in 1 \(\mu\)M mag-Fura-5 AM, 1 \(\mu\)M fluo-3 AM, plus 16 \(\mu\)M pluronit acid for 1 hr. Experiments were performed at 20–25\°C.

Optical system and image analysis. Light from a 75 W Xe short arc lamp was used to excite dyes fluorescence. Excitation wavelength was selected with a filter changer (Lambda 10–2; Sutter Instruments, Novato, CA) containing narrow-band (15 nm) interference filters with transmission maxima at 331, 380, and 485 nm. The filtered light was reflected into a microscope with a 510 nm dichroic mirror and was imaged by an objective (Fluar 40x/1.3; Carl Zeiss) to a 150-\(\mu\)m diameter spot centered in the field of view. The maximum intensity within this spot was 1.4 W m\(^{-2}\) at 331 nm, 5.6 W m\(^{-2}\) at 380 nm, and 180 W m\(^{-2}\) at 485 nm. Fluorescent light collected by the objective passed through a second filter changer containing 520–580 nm bandpass filter or an RG630 Schott glass long-pass filter before being imaged onto a CCD camera (model AT200; Photometrics, Tucson, AZ). Each pixel imaged a 0.16 \(\mu\)m \(\times\) 0.16 \(\mu\)m region in the plane of focus. The operation of shutters and filter wheels was controlled by a computer. Because of overlap in both their excitation...
and emission spectra, fluo-3 and FM1–43 were not used together in the same experiment. Control experiments demonstrated that the combinations of wavelengths, light intensities, and dye concentrations actually used in the experiments allowed the interleaved measurement of Ca$$^{2+}$$ concentration and either FM1–43 or FM4–64 fluorescence without crosstalk between the signals. In some experiments we used both FM1–43 and FM4–64. In these experiments, we corrected for the crossover of light emitted by FM1–43 in the FM4–64 detection band (see Fig. 3).

At the beginning of each experiment, a Hoffman interference image of the entire cell was recorded (Fig. 1), and two regions of interest ~20 μm square were selected (Fig. 1, outlined by squares). One region included the synaptic terminal; the other region included a portion of the soma. Fluorescent images of the two regions produced by 100 msec exposures were collected for the required filter combinations and saved for later analysis.

Normally, several movies were made after an experiment was completed. For example, the measurement of Ca$$^{2+}$$ concentration and FM1–43 fluorescence required six movies. One movie (produced by 485 nm exciting light) imaged FM1–43 fluorescence in the terminal. Second (produced by 331 nm light) and third (produced by 380 nm light) movies imaged fura-2 fluorescence in the terminal and were used to calculate the average, cytoplasmic Ca$$^{2+}$$ concentration. A similar set of three movies imaged FM1–43 and fura-2 fluorescence in the soma. The ability to identify potential problems was greatly facilitated by recording a movie of fluorescent images instead of integrating all of the light in a microscope field with a photomultiplier tube. Preliminary experiments demonstrated that bright 331 and 380 nm lights adversely affected the morphology of stained cells. Consequently, light intensity and repetition rate for fura-2 and mag-fura-5 measurements were minimized. We rejected the rare experiments in which a cell significantly changed shape.

The total fluorescence in the terminal was measured by summing the intensities in pixels within a circular area slightly larger than the terminal (Fig. 1, circle designated t). The fluorescence in the soma was measured by summing the intensities of pixels within a circular subregion ~8 μm in diameter over a portion of the soma (Fig. 1, circle designated s). Background fluorescence of FM1–43 and FM4–64, measured in a region ~5 μm distant from a cell, was subtracted from each pixel. For the calcium indicator dyes, background fluorescence measured away from the cell was negligible.

We assumed that the fluorescence of dye molecules was unchanged during endocytosis. Results (see Figs. 5, 10) indicate that vesicles quickly diffused throughout a terminal. Because vesicles appeared to be uniformly distributed, we were able to use the fluorescence in an optical section to calculate the total amount of endocytosed membrane distributed throughout a terminal. A microscope objective collects light from an image depth of 0.8 μm, where μ is the numerical aperture of the objective. Hence the thickness of an optical section was 1.6 μm.

**Calcium measurement.**

The fluorescence of fura-2 or mag-fura-5 was excited by light of 331 and 380 nm. Emission was detected between 520 and 580 nm. Intracellular Ca$$^{2+}$$ concentration was calculated from the relation (Grynkiewicz et al., 1985): $[Ca^{2+}] = k \beta (F - F_{	ext{min}})/(F_{	ext{max}} - F)$, where $R$ is the ratio of the fluorescence intensity excited at 331 nm divided by the intensity excited at 380 nm, $F_{	ext{max}}$ is the ratio produced by a saturating Ca$$^{2+}$$ concentration, $F_{	ext{min}}$ is the ratio produced by a minimum concentration, $\beta$ is the ratio of emission intensity stimulated at 380 nm in the absence of Ca$$^{2+}$$ to that in the presence of an excess of Ca$$^{2+}$$, and $k$ is an apparent affinity constant. Affinity constants were determined by superfusing cells with either 10 μM ionomycin or 10 μM A23187 and a series of Ca$$^{2+}$$ concentrations set with Ca$$^{2+}$$ buffers (see below).

Each concentration was superfused until a steady state was achieved. The affinity constant was 237 nM for fura-2 and 10 μM for mag-fura-5. $R_{	ext{min}}$, $R_{	ext{max}}$, and $\beta$ were measured for each cell. $R_{	ext{min}}$ was estimated at the beginning of each experiment while a cell was superfused with a low Ca$$^{2+}$$ concentration; $R_{	ext{max}}$ and $\beta$ were measured by superfusing at the end of each experiment with saline containing 10 μM ionomycin, 4 mM Ca$$^{2+}$$, 40 mM K$$^+$$, and 1 μM BayK8646.

The 485 nm light excited fura-3. Emission was detected between 520 and 580 nm. The Ca$$^{2+}$$ concentration was calculated from the relation (Grynkiewicz et al., 1985): $[Ca^{2+}] = k_1 F_{	ext{max}} - F_{	ext{min}})/F_{	ext{max}} - F$), where $F_{	ext{max}}$ is the fluorescence intensity, $F_{	ext{min}}$ is produced by a minimum Ca$$^{2+}$$ concentration, $F_{	ext{max}}$ is produced by a saturating Ca$$^{2+}$$ concentration, and $k_1$ is the apparent affinity constant. Minimum and maximum fluorescence were measured as described above for the ratiometric dyes. The affinity constant for fura-3, also measured as described above, was 0.9 μM (see also Rieke and Schwartz, 1996).

**Solutions and control of Ca$$^{2+}$$ concentration.** During an experiment, a cell was continuously superfused with solutions ejected from a four-barrelled pipette. Each barrel ended in a ~50 μm square opening. Each barrel was connected to a reservoir and pressure head through a solenoid valve. Solution changes were made by simultaneously opening and closing valves. Control experiments with fluorescent dyes demonstrated that solution changes were 95% complete in <100 msec. When an experiment required five different solutions, an additional single-barrelled pipette was introduced.

The superfused solutions were similar to the extracellular saline described above. Ca$$^{2+}$$ concentrations were set with Ca$$^{2+}$$ buffers; 1 mM bis(2-aminophenoxy)ethane-N,N,N',N'-tetra-acetic acid (BAPTA) was used for free Ca$$^{2+}$$ concentrations between 0.1 and 2 μM, and 1 mM

**Figure 1.** Hoffman modulation micrograph of a solitary bipolar cell. The squares indicate regions of interest that were saved during an experiment. The circles indicate areas in which the total fluorescence was measured (see text). Scale bar, 10 μm.
Kinetics of membrane staining depended on dye concentration. A, Image of a terminal stained with 20 μM FM1–43. B, The intensity of pixels along the solid line in A plotted as a function of radial distance. The arrow indicates the location of the surface membrane. C, The intensity in the surface membrane plotted as a function of time. The cell was superfused with two concentrations of FM1–43 as indicated by the horizontal bars. The peak intensity along a radial spoke (see B) was integrated around the circumference, except for the region from which the axon arose (as indicated by the dotted line in A). D, Intensity of membrane staining as a function of dye concentration. E, The t₁/₂ for the onset of staining as a function of dye concentration. F, Apparent exponential time constant for destaining as a function of dye concentration.

N-(2-acetamido)-2-iminodiacetic acid (ADA) was used for concentrations between 10 and 200 μM. Concentrations above 200 μM were unbuffered. Salines containing ionomycin and ADA lacked Mg²⁺. A high Ca²⁺ and K⁺ saline contained (in mM): NaCl, 82.5; KCl, 40; CaCl₂, 4; and MgCl₂, 1; pyruvate, lactate, glucose, and HEPES levels were as in the normal saline. Experiments were conducted on only one cell in a dish.

The concentration of BAPTA or ADA in stock solutions used to make Ca²⁺ buffers was determined by titrating with 1.0N CaCl₂ (BDH Laboratory Supplies) and measuring the free Ca²⁺ concentration with a Ca²⁺ electrode. After the concentration of each stock solution was known, salines with a specific free Ca²⁺ concentration were made according to recipes calculated with the computer program MAXC v1.70 (Bers et al., 1994).

**Kinetics of membrane staining.** The kinetics of membrane staining depended on the dye concentration. Our procedure for measuring the kinetics is explained in Figure 2. An image of a stained terminal is shown in Figure 2A. The intensity of pixels along the radial solid line is plotted in Figure 2B. The maximum intensity occurs at the surface membrane (Fig. 2B, arrow). To increase the accuracy with which the intensity of membrane fluorescence is measured, we rotated the inclination of the radial line and integrated the membrane signal around the terminal circumference excluding the segment containing the axon (as indicated by the dotted line in Fig. 2A). The intensity of membrane fluorescence was measured in a sequence of images and plotted as a function of time (Fig. 2C). The cell was exposed to two concentrations of FM1–43, first 5 and then 20 μM. When the dye was applied, the membrane stained with a nonexponential time course. When the dye was removed, the membrane destained with an approximately exponential time course.

The procedure illustrated in Figure 2, A–C, was used to obtain the data in Figure 2, D–F. In each experiment, a cell was exposed to two dye concentrations. To compare experiments, we normalized the steady state fluorescence to the intensity produced by 20 μM FM1–43. The intensity of membrane staining was proportional to dye concentration (Fig. 2D). The time to the half-maximum amplitude t₁/₂ decreased approximately in inverse proportion to the concentration (Fig. 2E). Destaining was approximately described by a single exponential with a time constant that also depended on concentration (Fig. 2F).

A linear relation between dye concentration and fluorescence intensity indicates that the number of dye molecules in the membrane is not limited by a small and fixed number of binding sites. A simple model would describe the movement of dye into the membrane as the partitioning between an aqueous and lipid phase accomplished in a single kinetic step. However, in this case, staining and destaining would have exponential time courses that are independent of concentration. This was not the case. The nonexponential time course for staining and a change in kinetics with dye concentration indicate that dye molecules equilibrate in a series of states.

The time course of membrane staining and destaining will obviously limit the time resolution for measurements of vesicle cycling. A concentration of 20 μM provided a reasonable resolution for membrane staining. However, to observe membrane destaining with a resolution finer than several tens of seconds, we needed an additional technique.

**Resonance energy transfer.** We have discovered an easy method to increase the time resolution of membrane destaining. FM4–64 rapidly quenched the fluorescence of FM1–43. An example is shown in Figure 3A. A bipolar cell was superfused first with FM4–64, next with FM1–43,
and finally with both dyes together. The emission intensities in the two detection channels are superimposed in Figure 3A (upper traces). Cross-over from FM1–43 into the FM4–64 detection channel (Fig. 3A, thin upper trace) was measured, and the corrected trace is plotted below. The essential observation came when the two dyes were applied together. Now, the fluorescence of FM1–43 was strongly quenched (Fig. 3A, asterisk). Dye intensity was reduced 94%. In addition, there was an apparent 14% increase in FM4–64 fluorescence. This behavior suggested the procedure in Figure 3B. First, a cell was superfused with a pulse of FM1–43. When the dye was removed, fluorescence in the FM1–43 channel (Fig. 3B, thick trace) declined with a time constant of 6.7 sec. Next, a pulse of FM1–43 was immediately followed by a pulse of FM4–64. Now, FM1–43 fluorescence was quickly quenched. The fluorescence in the surface membrane disappeared in <1.4 sec.

Sources and abbreviations. Ionomycin, FM1–43, FM4–64, fluo-3 AM, fura-2 AM, and mag-fura-5 AM were obtained from Molecular Probes (Eugene, OR). Papain was purchased from Worthington (Freehold, NJ). All other chemicals were obtained from Sigma (St. Louis, MO).

RESULTS

FM1–43 is a lipophilic dye that partitions into cell membranes. The dye has little fluorescence in aqueous solution and is much more fluorescent in a membrane environment. Positive charges are believed to prevent dye molecules from flipping between the outer and the inner membrane leaflet. Consequently, extracellular FM1–43 has been used to tag the outer leaflet of the surface membrane and then to visualize the bits that are pulled into the cytoplasm by endocytosis (Betz et al., 1992, 1996). However, as described in the Appendix, the ability of dye to cross the cell membrane can complicate its use as a membrane tag. We designed experiments to minimize this problem.

Ca2+ dependence of continuous vesicle cycling

To explore the relation between Ca2+ concentration and vesicle cycling, we adopted two strategies to produce a nearly uniform intracellular Ca2+ concentration. The first procedure relied on a Ca2+ ionophore, ionomycin, and the entry of Ca2+ from the extracellular saline. The second procedure relied on a proton ionophore, monensin, and the release of Ca2+ from intracellular, acidic organelles. Both procedures were able to raise the cytoplasmic Ca2+ concentration.

For the first procedure, cells were superfused with ionomycin (10 μM) and a Ca2+ buffer. Changing the free Ca2+ concentration in the extracellular saline was followed by a change in the intracellular concentration. During a typical 1–3 min application, the intracellular concentration reached 3–25% of the extracellular concentration. An example is illustrated in Figure 4. The cell was first superfused with ionomycin and 100 nm Ca2+. The addition of FM4–64 (Vida and Emr, 1995) stained the surface membrane and produced a fluorescence change (labeled s). Next, when the extracellular Ca2+ concentration was increased to 50 μM, the intracellular concentration followed with a rise to ∼1.7 μM. At the same time, the fluorescence of FM4–64 increased labeled v). We believe that the increase in fluorescence is produced as exocytosis adds vesicle membrane to the surface membrane. The rate of continuous exocytosis Cx was calculated as 

\[
C_x = \frac{\rho v_x}{\Delta t}
\]

where ρ is a geometrical constant (see Materials and Methods; Fig. 3). At this moment the Ca2+ concentration was ∼1 μM. The profile of the terminal is nearly round and smooth. Image b was taken 68 sec later, when the intracellular Ca2+ concentration had increased to 4 μM. Small patches of fluorescence are located immediately beneath the surface membrane, the surface contour itself is slightly irregular with sites of local expansion or dilation, and the interior of the terminal has a uniform weak fluorescence.

Finally, image c was taken 15 sec after the fluorescence in the surface membrane was quenched so that only the fluorescence of FM1–43 in endocytosed vesicles was detected. The fluorescence of FM1–43 was almost uniformly distributed, indicating that endocytosed vesicles quickly diffused throughout the cytoplasm.

The rate of steady endocytosis was estimated by first raising the intracellular Ca2+ concentration and then measuring the amount of FM1–43 accumulated during a 12 sec test pulse (Fig. 6). The rate of continuous endocytosis Cn was calculated as

\[
C_n = \frac{\rho v_n}{\Delta t}
\]
where \( v_n \) is defined in the figure. For the two measurements in Figure 6, the Ca\(^{2+} \) concentration was 1.3 \( \mu \)M, and \( C_n \) was 0.33 and 0.22 MEq min\(^{-1} \).

Figures 4 and 6 provide two methods for measuring rates of continuous vesicle cycling. A summary of 102 experiments is presented in Figure 7. Crosses were measured by the procedure in Figure 4; closed symbols were measured by the procedure in Figure 6. Vesicle cycling in the terminal was not observed at Ca\(^{2+} \) concentrations below 0.8 \( \mu \)M. A high rate of cycling was sustained at Ca\(^{2+} \) concentrations between 0.8 and 20 \( \mu \)M. Increasing the Ca\(^{2+} \) concentration above 20 \( \mu \)M decreased the rate of cycling. The maximum rate of vesicle cycling in Figure 7 was \( 0.5 \) MEq min\(^{-1} \).

Ionomycin allowed Ca\(^{2+} \) to enter the cytoplasm from the extracellular saline. An alternative procedure was to raise intracellular Ca\(^{2+} \) without a change in extracellular Ca\(^{2+} \) concentration. For this purpose we used monensin, an ionophore that disrupts the pH-dependent accumulation of Ca\(^{2+} \) in intracellular organelles (Fasolato et al., 1991). The subsequent release of Ca\(^{2+} \) raises cytoplasmic Ca\(^{2+} \) concentration. The procedure is illustrated in Figure 8. The cell was continuously superfused with an extracellular saline containing 100 nm Ca\(^{2+} \) (and no ionomycin). When 2 \( \mu \)M monensin was added, the Ca\(^{2+} \) concentration in the terminal cytoplasm increased transiently to nearly 6 \( \mu \)M. This increase in Ca\(^{2+} \) concentration was not produced by an influx from the extracellular saline (where the Ca\(^{2+} \) concentration was only 100 nM) but was instead produced by the release of Ca\(^{2+} \) from intracellular organelles. The rise in Ca\(^{2+} \) concentration was accompanied by an increase in FM4–64 fluorescence. A similar result was observed in eight experiments.

Both the experiments with ionomycin and monensin demonstrate that a rise in intraterminal Ca\(^{2+} \) concentration to between...
0.8 and 20 μM produced a continuous cycle of vesicle exocytosis and endocytosis. The maximum rate was sufficient to turn over the surface membrane of a terminal every 2 min.

**Transient vesicle cycling initiated by Ca^{2+} entry through voltage-gated channels**

Normally, fusion is triggered when Ca^{2+} enters through voltage-gated channels. Ca^{2+} first accumulates at the cytoplasmic end of each open channel and then diffuses into the cytoplasm. The result is many small domains of high Ca^{2+} concentration that quickly dissolve. The following experiments investigated vesicle cycling triggered by depolarization and Ca^{2+} influx. In each experiment, cells were initially superfused with a saline containing 100 nM Ca^{2+} to minimize premature Ca^{2+} entry during a control period. Exocytosis was triggered by superfusion with a saline containing 4 mM Ca^{2+} and 40 mM K^{+}.

Exocytosis induced by depolarization and Ca^{2+} influx is illustrated in Figure 9. First, FM4–64 stained the surface membrane (labeled s). Next, a 4 sec exposure to the high Ca^{2+} and K^{+} saline depolarized, increased the average Ca^{2+} concentration in the terminal, and produced a step increase in FM4–64 fluorescence (labeled v). We assume that the abrupt increase in the fluorescence of the terminal was proportional to the membrane added to the surface by exocytosis. The amount of exocytosis $T_x$ was calculated as $T_x = \rho_v / s$. In Figure 9, $T_x$ was 0.13 and 0.08 MEq. In contrast, depolarization did not increase fluorescence in the soma (Fig. 9, thin trace). In 25 experiments $T_x$ varied from being barely detectable to ~0.13 MEq.

The previous measurements were confirmed by observing endocytosis. Cells were first superfused with a saline containing 100 nM Ca^{2+} to minimize Ca^{2+} entry and to prevent Ca^{2+}-triggered vesicle cycling. An 18 sec exposure to FM1–43 stained the surface membrane of the terminal (Fig. 10, thick upper trace). As expected, a 6 sec exposure to high Ca^{2+} and K^{+} saline produced a Ca^{2+} influx and raised the average cytoplasmic Ca^{2+} concentration (thick lower trace). After FM1–43 was removed from the extracellular saline, the surface membrane destained, and fluorescence remained trapped in the cytoplasm of the terminal (labeled $v_n$). The amount of endocytosis $T_n$ was measured as $T_n = \rho_v / s$. In this case, $T_n$ was 0.08 MEq. In contrast, little or no dye was trapped in the soma (Fig. 10, thin upper trace). In addition, no endocytosis occurred when FM4–64 was applied as a control at the end of the experiment (Fig. 10, middle traces). In 22 experiments $T_n$ was 0.049 ± 0.030 MEq (equal to 5500 vesicles). The largest values were ~0.12 MEq (equal to 13,000 vesicles).

Images of the terminal are shown in two insets in Figure 10. Image a was taken during the time the membrane was stained with FM1–43. Image b was taken after the dye was removed and the membrane destained. There is no evidence of localization of newly endocytosed vesicles beneath the surface membrane. Instead, stained vesicles appear to be uniformly distributed throughout the interior of the terminal.

A second application of high Ca^{2+} and K^{+} saline without dye did not release a significant fraction of the labeled vesicles. We were never able to label vesicles, destain the surface membrane, and then release the accumulated dye in a second bout of vesicle cycling. Of course, this is not surprising if newly endocytosed vesicles diffuse rapidly and mix with the entire intraterminal

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*Figure 8.* The release of Ca^{2+} from intracellular organelles raises the cytoplasmic Ca^{2+} concentration and stimulates exocytosis. The cell was continuously superfused with an extracellular saline buffered to contain 100 nM free Ca^{2+}. The lower timing trace indicates the application of 20 μM FM4–64. The upper timing trace indicates the addition of 2 μM monensin.

*Figure 9.* Exocytosis was induced by depolarization and Ca^{2+} influx. A bipolar cell was superfused with an extracellular saline containing 100 nM Ca^{2+}. The lower timing trace indicates the application of 20 μM FM4–64. The upper timing trace indicates 4 sec applications of a high Ca^{2+} and K^{+} saline (4 mM Ca^{2+} and 40 mM K^{+}). $v$ is the intensity increase produced by vesicle exocytosis. Transient exocytosis $T_x$ was measured as $T_x = \rho_v / s$. For the two applications of high Ca^{2+} and K^{+} saline, $T_x$ was 0.13 and 0.08 MEq.
membrane. The gain in average intraterminal Ca\textsuperscript{2+} was eight times the gain in a. Scale bar, 10 \( \mu \text{m} \).

Figure 10. Endocytosis was induced by depolarization and Ca\textsuperscript{2+} influx. A bipolar cell was superfused with an extracellular saline containing 100 nm Ca\textsuperscript{2+}. Timing traces indicate an 18 sec application of 20 \( \mu \text{M} \) FM1–43, an 18 sec application of 20 \( \mu \text{M} \) FM4–64, and 6 sec applications of a high Ca\textsuperscript{2+} and K\textsuperscript{+} saline. \( s \) is the fluorescence intensity of the surface membrane. \( v_n \) is the fluorescence of endocytosed vesicles that remain trapped in the cytoplasm. Transient endocytosis was measured as in von Gersdorff et al., 1996. Thus, only 1–2\% would be labeled, and a decrease in fluorescence by a second bout of endocytosis would be difficult to measure.

Vesicle cycling induced by depolarization depended on Ca\textsuperscript{2+} entry through voltage-gated Ca\textsuperscript{2+} channels and stopped when Ca\textsuperscript{2+} channels were blocked. An example is shown in Figure 11. The cell was exposed to high Ca\textsuperscript{2+} and K\textsuperscript{+} saline three times. The first exposure produced a bout of endocytosis with \( T_n = 0.08 \text{ MEq} \). After superfusion with the high Ca\textsuperscript{2+} and K\textsuperscript{+} saline, \( T_n \) was 0.08 MEq. \textit{Insets}. Images of the terminal taken at the times marked a and b are shown. The gain in b is eight times the gain in a. Scale bar, 10 \( \mu \text{m} \).

Figure 11. Pharmacologically blocking voltage-gated Ca\textsuperscript{2+} channels blocked Ca\textsuperscript{2+} influx and endocytosis. A bipolar cell was superfused first with an extracellular saline containing 100 nm Ca\textsuperscript{2+} and then three times with FM1–43 and high Ca\textsuperscript{2+} and K\textsuperscript{+} saline. Each application was a sequence of 20 \( \mu \text{M} \) FM1–43 in control saline for 4 sec, FM1–43 in high Ca\textsuperscript{2+} and K\textsuperscript{+} saline for 4 sec, and finally FM1–43 in control saline for 12 sec. Nisoldipine (50 \( \mu \text{M} \)) was added to the extracellular salines during the second and third applications. \( T_n \) (measured as illustrated in Fig. 10) for the three sequential applications was 0.08, 0.006, and 0.003 MEq.

The same result was observed in 11 experiments. Depolarization stimulated vesicle cycling only when Ca\textsuperscript{2+} was able to influx through voltage-gated channels.

We were next interested to know whether Ca\textsuperscript{2+} influx through voltage-gated channels produced a relatively steady rate of vesicle cycling or a burst of cycling that developed quickly and soon subsided. Long and short episodes of Ca\textsuperscript{2+} entry were promoted by superfusing with a saline containing high Ca\textsuperscript{2+} and K\textsuperscript{+}. An example is shown in Figure 12. After a 3 sec exposure to high Ca\textsuperscript{2+} and K\textsuperscript{+} saline, \( T_n(3) \) was 0.11 MEq. Lengthening the depolarization to 15 sec produced a very similar value; \( T_n(15) \) was 0.12 MEq. Finally, the 15 sec exposure was repeated with 1 \( \mu \text{M} \) BayK8466 added to the high Ca\textsuperscript{2+} and K\textsuperscript{+} saline to prolong the open time of Ca\textsuperscript{2+} channels (see Heidelberger and Matthews, 1992). As expected, the average cytoplasmic Ca\textsuperscript{2+} concentration rose to a higher level, but \( T_n \) was still 0.11 MEq. In 15 experiments the ratio \( T_n(3)/T_n(15) \) was 1.06 ± 0.49. Each pulse of high Ca\textsuperscript{2+} and K\textsuperscript{+} produced an approximately equal increment in dye accumulation. An experiment described above (see Fig. 10) demonstrates that labeled vesicles quickly mixed with a 100\times larger vesicle pool. Consequently, only \~{}1\% of the vesicles labeled during one bout of cycling were released in a subsequent bout of cycling, and each increment represents labeled vesicles added in a new round of exocytosis with little loss of previously accumulated vesicles. Thus, equal steps of accumulation indicate that the majority of exocytosis initiated by superfusion with a high Ca\textsuperscript{2+} and K\textsuperscript{+} saline is completed in a single, transient burst.
Capacitance measurements by von Gersdorff and Matthews (1994b) indicate that endocytosis does not occur while the Ca²⁺ concentration is elevated. The following experiment (Fig. 13) provides additional insight. A cell was exposed to two pulses of high Ca²⁺ and K⁺ saline. The first application included a 3 sec pulse of high Ca²⁺ and K⁺; \( T_n = 0.11 \text{ MEq} \). The second application included a 15 sec pulse of high Ca²⁺ and K⁺; \( T_n = 0.12 \text{ MEq} \). The third application was similar but added 1 \( \mu \text{M} \) BayK8466 to the high Ca²⁺ and K⁺ saline to prolong the open time of Ca²⁺ channels; \( T_n = 0.11 \text{ MEq} \). Increasing the duration or amount of Ca²⁺ entry did not alter transient exocytosis. The control saline contained 100 \( \text{nM} \) Ca²⁺. Each pulse of FM1–43 extended 12 sec beyond the end of the high Ca²⁺ and K⁺ application.

**DISCUSSION**

Electrical and optical experiments now provide different perspectives of synaptic vesicle cycling. Both methods have been used to study ribbon synapses in the large synaptic endings of goldfish bipolar cells. Capacitance measurements are best suited to detect rapid events. Heidelberger et al. (1994) have measured the change in electrical capacitance produced when a jump in Ca²⁺ concentration promotes the rapid fusion of a limited number of vesicles. In contrast, fluorescence from styryl dyes cannot resolve rapid events but is best suited to detect slow, continuous, or maintained changes. As we explain below, the two methods provide complementary views of synaptic vesicle dynamics.

The electrical measurements reveal that a rapid, saturating increase in the intracellular Ca²⁺ concentration triggers the fusion of ~3000 vesicles with a time constant of 0.35 msec (Heidelberger et al., 1994), too short for an individual fusion site to be used more than once. Consequently, all 3000 vesicles must be pre docked and ready for release. The rate constant for the last step in vesicle fusion accelerates from 1 sec⁻¹ when the Ca²⁺ concentration is ~10 \( \mu \text{M} \) to 1000 sec⁻¹ when the concentration reaches 200 \( \mu \text{M} \) [see Heidelberger et al. (1994), their Fig. 3a]. The optical measurements reported here demonstrate that a Ca²⁺

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**Figure 12.** Vesicle cycling triggered by Ca²⁺ influx was transient. The cell was superfused three times with a 30 sec pulse of 20 \( \mu \text{M} \) FM1–43. Transient endocytosis \( T_n \) was measured as shown in Figure 10. The first application included a 3 sec pulse of high Ca²⁺ and K⁺; \( T_n = 0.11 \text{ MEq} \). The second application included a 15 sec pulse of high Ca²⁺ and K⁺; \( T_n = 0.12 \text{ MEq} \). The third application was similar but added 1 \( \mu \text{M} \) BayK8466 to the high Ca²⁺ and K⁺ saline to prolong the open time of Ca²⁺ channels; \( T_n = 0.11 \text{ MEq} \). Increasing the duration or amount of Ca²⁺ entry did not alter transient exocytosis. The control saline contained 100 \( \text{nM} \) Ca²⁺. Each pulse of FM1–43 extended 12 sec beyond the end of the high Ca²⁺ and K⁺ application.

**Figure 13.** Endocytosis was delayed until after Ca²⁺ influx stopped. The timing of endocytosis was measured with 18 sec applications of 20 \( \mu \text{M} \) FM1–43 combined with 6 sec pulses of high Ca²⁺ and K⁺ saline. The first applications of FM1–43 and high Ca²⁺ and K⁺ saline ended together and were immediately followed by a pulse of 20 \( \mu \text{M} \) FM4–64 in a saline containing 100 \( \text{nM} \) Ca²⁺. The FM4–64 quenched the fluorescence of FM1–43 in the surface membrane (see Fig. 3) but not in vesicles trapped in the cytoplasm. \( T_n \) measured (as shown in Fig. 10) with FM1–43 was 0.004 MEq and indicated that little endocytosis occurred during superfusion with the high Ca²⁺ and K⁺ saline. \( T_n \) measured with FM4–64 was 0.05 MEq and indicated that significant endocytosis occurred after return to the control saline. The second application of FM1–43 measured the total amount of endocytosis; \( T_n = 0.07 \text{ MEq} \).
concentration between 0.8 and 20 μM can maintain a continuous cycle of balanced exocytosis and endocytosis with a maximal rate of 900 vesicles sec\(^{-1}\) (Fig. 7) or ~0.3 vesicle sec\(^{-1}\) at each release site. The continuous cycle slows when the Ca\(^{2+}\) concentration rises above 20 μM. Both the electrophysiological and optical measurements demonstrate that Ca\(^{2+}\) influx through voltage-gated channels produces a burst of exocytosis but little or no endocytosis until the average Ca\(^{2+}\) concentration falls. Moreover, the maximum number of vesicles released appears to be greater than the number of fusion sites (i.e., 10,000 vesicles vs 3,000 fusion sites), implying that several vesicles fuse at each site before cycling stops.

Our results differ from those of Lagnado et al. (1996). We have observed that (1) depolarization by high K\(^+\) produces a transient burst of vesicle cycling, (2) newly endocytosed vesicles rapidly diffuse and mix with the total intraterminal vesicle population, and (3) a high rate of continuous cycling is sustained at concentrations between 0.8 and 20 μM intracellular Ca\(^{2+}\) and stops at higher concentrations. Lagnado et al. (1996) claim that (1) depolarization by high K\(^+\) produces continuous vesicle cycling, (2) newly endocytosed vesicles accumulate beneath the surface membrane, and (3) continuous cycling is sustained at all Ca\(^{2+}\) concentrations above 0.3 μM Ca\(^{2+}\). We cannot explain these differences.

We expect the function of fusion sites to depend critically on the location of Ca\(^{2+}\) channels and illustrate our view with two examples. One extreme behavior would occur in a "synapse" with only one Ca\(^{2+}\) channel and an immediately adjacent fusion site. In this case, the Ca\(^{2+}\) concentration in a microdomain at the cytoplasmic end of the channel will tightly control vesicle cycling. In contrast, another extreme behavior would occur in a "synapse" with a Ca\(^{2+}\) channel far from a fusion site. In this case, release would be controlled by the average Ca\(^{2+}\) concentration in the cytoplasm. Many channel openings would raise the average cytoplasmic Ca\(^{2+}\) concentration and sustain a continuous cycle of repeated exocytosis and endocytosis. Of course, these idealized synapses illustrate extreme behaviors. But the examples indicate how dynamic behavior may depend on the number and relative spatial distribution of channels and fusion sites. Many central synapses have only one, or a few, fusion sites (see Redman, 1990; Korn and Faber, 1991; Ryan et al., 1997) and consequently may approach either of the two extreme behaviors described above. However, cells with ribbon synapses have many fusion sites (Raviola and Gilula, 1975; von Gersdorff et al., 1996) and, combining the behavior of both models, should mediate both transient and continuous release.

Bipolar cells, like rod photoreceptors (Rieke and Schwartz, 1996), sustain high rates of vesicle cycling at relatively low (0.8–20 μM) Ca\(^{2+}\) concentrations. This is the result of having many release sites that operate in parallel. Although the rate of turnover at an individual release site is estimated as only 0.3 vesicle sec\(^{-1}\) the net effect of simultaneously maintaining release at a large number of sites is an enormous membrane flux, equal in a bipolar cell to the replacement of the entire surface area of a terminal every 2 min. Neurons with ribbon synapses—photoreceptors, hair cells, and retinal bipolar cells—appear to be specialized for the massive, parallel release of many vesicles.

An endocytosed vesicle passes through an uncertain number of intermediate steps before being ready for a new round of fusion. If newly endocytosed vesicles had to be immediately reused, then continuous release would be limited by the time required to produce a mature, fusion-competent vesicle. Significantly, endo-

cytosed vesicles are not immobilized near release sites but quickly diffuse throughout the terminal (Figs. 5, 10). A total of 5–10 × 10\(^5\) vesicles (von Gersdorff et al., 1996) is equal to ~500–1000 vesicles per release site. Hence, vesicles may continuously move to docking sites from an abundant reserve. A large number of release sites and a surfeit of vesicles are required to maintain a high rate of continuous release.

**APPENDIX: FM DYES CROSS THE SURFACE MEMBRANE**

When a bipolar cell was superfused with FM1–43 for several minutes, dye sometimes accumulated in the cytoplasm even without stimulation or a change in experimental condition. For example, in Figure 14A, a cell was permeabilized with a Ca\(^{2+}\) ionophore (10 μM ionomycin) and continuously superfused with 100 nM Ca\(^{2+}\). The intracellular Ca\(^{2+}\) concentration remained constant at ~100 nM throughout the experiment. Nonetheless, FM1–43 initially stained the surface membrane and then, after a significant delay, entered the cytoplasm.

The spontaneous entry of dye did not depend on vesicle cycling. Instead, the following experiment demonstrates that it was pro-
duced by dye molecules crossing the surface membrane. Cells were first fixed with 2% formaldehyde for 1 hr and afterward superfused with a saline containing both FM1–43 and FM4–64. The result is shown in Figure 14B. There are two essential observations. First, FM1–43 accumulated in the cytoplasm of a fixed cell. Thus, spontaneous entry did not require vesicle cycling. This conclusion is reinforced by the second observation; FM1–43 accumulated in the cytoplasm with a different time course than did FM4–64. Of course, the two dyes would be accumulated with the same time course if the two dyes tagged the surface membrane and were internalized together in the membrane of endocytosed vesicles. Instead, the accumulation of FM4–64 was much slower. The results are easily explained if FM1–43 molecules cross the lipid bilayer and the other dye, FM4–64, crosses more slowly.

Images recorded during spontaneous dye entry (Fig. 14) differed in two ways from images recorded during vesicle cycling (Fig. 5). First, fluorescent foci did not develop beneath the surface membrane, and the surface contour did not become irregular. Second, dye preferentially accumulated in the center of a terminal and formed a fluorescent spot occupying perhaps half the internal cross-sectional area. A similar central spot developed when FM1–43 was applied to cells that had been permeabilized by a brief exposure to 20 μM digitonin (data not shown). Thus, the central spot was a region that avidly stained when free dye was available in the cytoplasm.

Several observations indicate that styril dyes form a series of states when they partition into membranes. The complex kinetics of membrane staining and destaining observed in Figure 2 indicates that the dye has more than one state in a membrane. In addition, the glitches seen in Figure 3 (at the arrowhead) are produced because spectral components of FM1–43 fluorescence leave the membrane at different rates. The correlation of spectral and kinetic components indicates distinct states. Finally, the ability of FM4–64 to quench FM1–43 indicates that dye molecules can be closely packed (see Stryer, 1978). Hence, dyes may aggregate. Thus, we were not surprised to find that FM1–43 facilitated its own entry. For example, Figure 15 shows the result from a cell superfused with two concentrations of FM1–43. When the cell was first superfused for several minutes with 5 μM FM1–43, there was a steady, slow rate of cytoplasmic accumulation (Fig. 15, lower trace). Next, when the concentration was increased to 40 μM, the rate of entry (normalized by the dye concentration) increased fivefold (viz., from 0.074 to 0.38 M Eq min⁻¹).

We believe that dye molecules equilibrate between a series of states. Some of these transitions may be fast and others quite slow. Moreover, some states may depend on or be nucleated by specific membrane molecules. Eventually, dye molecules might cross the surface membrane either by flipping from the outer to the inner leaflet or by interacting with the lipids that surround channels, opening channels, and then passing through the open pore. The experiments in Figure 16. A and B, provide a hint for the mechanism of dye entry. Cells were loaded with fura-2 AM and superfused alternately with salines that contained either 2 mM Mg²⁺ or 2 mM Cd²⁺. Fura-2 was used to detect the entry of Cd²⁺. Normally, the permeability of Cd²⁺ is low. Thus, in each of seven control experiments, there was only a small change in the ratio of fura-2 fluorescence elicited by 331 and 380 nm light (Fig. 16A). In contrast, a different picture was observed in eight experiments when a cell was also superfused with FM1–43 (Fig. 16B). Now, fura-2 fluorescence reported a significant increase in intracellular divalent ion concentration each time the cell was exposed to Cd²⁺. Because the extracellular saline lacked Ca²⁺, the change in fura-2 fluorescence is attributed to the entry of Cd²⁺. Thus, continuous exposure to FM1–43 increased the membrane permeability of Cd²⁺. In addition, each exposure to Cd²⁺ decreased the simultaneous entry of FM1–43 and, after the removal of Cd²⁺, increased the subsequent entry of FM1–43. Although Cd²⁺ and FM1–43 interacted in a complex manner, the results imply that both cooperate to increase membrane permeability, most likely by opening a channel.

Experiments on vesicle cycling can be contaminated by spontaneous dye entry. The characteristic delay before rapid intracellular staining was variable. Rapid entry could be triggered by a brief exposure to Cd²⁺ (Fig. 16), and we have the impression that the transition may sometimes be initiated in the absence of Cd²⁺ by bright light or exposure to a hydrophobic drug. Once dye entry started, it was not reversible. Therefore, our strategy was (1) to use FM1–43 only for short exposures (<120 sec in Fig. 5 and <30 sec in all other experiments) and FM4–64, which appeared to permeate more slowly (Fig. 14B), for longer exposures, (2) to record images of the terminal in every experiment and to reject an experiment if a bright fluorescent spot formed in the center of the terminal (as in Fig. 14), and (3) to elicit multiple periods of vesicle cycling and to include controls to be certain that all measurements were repeatable.

![Figure 15](image-url)
REFERENCES


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**Figure 16.** The combination of Cd$^{2+}$ and FM1–43 increased membrane permeability to both Cd$^{2+}$ and FM1–43. Cells were loaded with fura-2. A, Cd$^{2+}$ does not normally enter a terminal. The cell was superfused alternately with an extracellular saline containing 2 mM Mg$^{2+}$ (no added Ca$^{2+}$) and a saline containing 2 mM Cd$^{2+}$. There was little change in the ratio of fura-2 fluorescence produced by 331 and 380 nm light, indicating that Cd$^{2+}$ did not enter. B, Cd$^{2+}$ enters a terminal stained with FM1–43. The cell was superfused with an extracellular saline containing 2 mM Mg$^{2+}$ (no added Ca$^{2+}$). FM1–43 (20 μM) was added to the saline. When Mg$^{2+}$ was replaced by Cd$^{2+}$, fura-2 fluorescence indicated an increase in intracellular, divalent ions. After returning to the Mg$^{2+}$-rich saline, the rate of FM1–43 entry increased.