Abstract. We have investigated the consequences of having multiple fusion complexes on exocytotic granules, and have identified a new principle for interpreting the calcium dependence of calcium-triggered exocytosis. Strikingly different physiological responses to calcium are expected when active fusion complexes are distributed between granules in a deterministic or probabilistic manner. We have modeled these differences, and compared them with the calcium dependence of sea urchin egg cortical granule exocytosis. From the calcium dependence of cortical granule exocytosis, and from the exposure time and concentration dependence of N-ethylmaleimide inhibition, we determined that cortical granules do have spare active fusion complexes that are randomly distributed as a Poisson process among the population of granules. At high calcium concentrations, docking sites have on average nine active fusion complexes.

Exocytosis involves many steps, including the biosynthesis of vesicles and their contents, vesicle transport to the plasma membrane, docking, the merging of membranes, and the release of vesicle contents. After exocytosis, there is often calcium-dependent granule mobilization (39) and endocytic membrane retrieval (48, 49). In sea urchin eggs, the cellular machinery that mediates membrane merger in calcium-triggered exocytosis, the fusion complex, resides on exocytotic granules (53) and can undergo conformational changes from inactive to active states (55). Transition between these two states is thought to be regulated by the binding of calcium to a "calcium sensor" (13, 38), but the identity and mechanism of the calcium sensor and the fusion complex is unknown. While binding of calcium to the sensor may be reversible, upon activation, a fusion complex can irreversibly commit to fuse (55). The sea urchin is a good preparation to study the calcium dependence of exocytosis because the fusion of preocked granules with the plasma membrane can be studied, in vitro, in the absence of reserve granule mobilization to docking sites, or in the absence of subsequent endocytic activity.

The simplicity of cortical granule exocytosis in vitro allows the investigation of a feature common to almost every fusing system (28): submaximal response to calcium. We have recently concluded that sea urchin cortical granules are heterogeneous in their response to calcium (Kaplan, D., P.S. Blank, M.S. Cho, I. Steinberg, and J. Zimmerberg. Biophys. J. 66:A284 [Abstr.]). Depending only on calcium concentration, a cortical granule will either fuse in time or will never fuse, regardless of how long or the number of times calcium is applied. This striking behavior occurs even though all granules are docked. In this in vitro system docking has been characterized by EM (8, 61), and is functionally evident by the ability of granules to remain attached to the plasma membrane despite extensive perfusion (61). In this fusion system, it is not known if docking and membrane merger are mediated by the same components. In terms of the above description, a fusion complex is irreversibly activated when a threshold calcium concentration is exceeded, i.e., all conformational changes associated with the calcium sensor and the fusion complex that are required for membrane fusion to ensue are complete. We define this state as an activated fusion complex.

Practically all docked cortical granules fuse with the plasma membrane of in vitro preparations (4, 15, 51, 61) at calcium concentrations in excess of 100 μM (45, 52). This suggests that virtually all cortical granules have at least one fusion complex that can be activated by calcium. The total surface area of the granule (1-μm diameter; 8, 61) is 400 times the surface area of a 50-nm synaptic vesicle, so there is ample area on a cortical granule for more than one fusion complex. Docked cortical granules are derived from much smaller vesicles that bud from the Golgi (14), and in other systems, such precursor vesicles fuse to each other before plasma membrane docking (16, 17). If each precursor vesicle contained at least one fusion complex, then docked cortical granules would contain multiple complexes, which may concentrate at their docking sites.

Since fusion for a single granule is all-or-none, fusion can be caused by one activated fusion complex. After fu-
sion, the rest of the fusion complexes will be redundant, and will not contribute to either the extent or the rate of fusion. Thus, if exocytotic granules have multiple activated fusion complexes before fusion, their number and distribution can affect both the rate and extent of triggered exocytosis. Attempts to model the extents and kinetics of calcium-dependent fusion must consider the consequences of having multiple activated complexes. These consequences are examined in this study.

Materials and Methods

Cell surface complex (CSC) was prepared and assayed for calcium-triggered exocytotic activity, as previously described (52). Sea urchin eggs were prepared from Strongylocentrotus purpuratus (Marinus, Long Beach, CA). Eggs in PKMEB (50 mM Pipes, 450 mM KCl, 10 mM MgCl₂, 5 mM EGTA, 1 mM BAPTA, 1 mM benzamidine, pH 6.7) at a ratio of 1:10 (vol/vol) were homogenized with 6–10 strokes of a Teflon-glass dounce homogenizer. The homogenate was centrifuged for 1 min at 200g in a tabletop centrifuge. The pellet of CSC was resuspended in HENPK buffer yielding a preparation consisting of exocytotic cortical granules docked on plasma membrane fragments attached to a glass coverslip. The coverslip was placed in a rapid perfusion chamber (25) and challenged with calcium buffers. In this chamber, buffer was completely replaced in <1 s. Light scattering was measured as previously described (61).

Fusion complexes were fractionally inactivated by mixing CSCs with HENPK containing various concentrations of NEM (up to 20 mM) on ice. After the appropriate incubation time (up to 2 h), unreacted NEM was neutralized by the addition of 20 mM DTT. The calcium dependence of cortical granule fusion in CSCs was determined at room temperature using a turbidimetric microtiter dish assay, as previously described (52), and HENPK buffer containing defined concentrations of free calcium. Free calcium concentrations were measured for these buffers using a calcium electrode (World Precision Instruments, Inc., Sarasota, FL) or with the fluorescent calcium indicator BTC (Molecular Probes, Inc., Eugene, OR), whose fluorescence was measured using a CytoFluor 2350 fluorimeter (Millipore, Bedford, MA) with a ratio of the light excited at 360 ± 40 and 485 ± 20 nm collected at 530 ± 25 nm. The free calcium concentration was calculated using the function [Ca²⁺] = K_d × Q × (R - R_min)/(R_max - R) and an apparent K_d value of 73 ± 1μM in HENPK buffer and a Q value of 66 ± 0.3.

Survival curves were measured by monitoring light scattering from planar isolated sea urchin egg cortices (61) that had been perfused with HENPK buffer containing defined concentrations of free calcium. Intact eggs were attached to polylysine-treated glass coverslips, and sheared with calcium-free HENPK buffer yielding a preparation consisting of exocytotic cortical granules docked on plasma membrane fragments attached to a glass coverslip. The coverslip was placed in a rapid perfusion chamber (25) and challenged with calcium buffers. In this chamber, buffer was completely replaced in <1 s. Light scattering was measured as previously described (61).

The fraction of refractile granules was measured using planar isolated sea urchin egg cortices (61). Granules were imaged using a Plan-Neofluor 63×1.25 NA objective and Nomarski optics on an upright microscope (Carl Zeiss, Inc., Thornwood, NY). Cortices were incubated in a buffer containing 50 mM Pipes, 450 mM KCl, 1 mM benzamidine, 5 mM DM-nitrophen, and 2.5 mM CaCl₂, pH 6.7. Caged calcium was released from the time constant for observing fusion.

In addition to the above assumptions, there are two assumptions that are unique to either the fixed or random model. The fixed model assumes that every granule in a population will have a fixed integral number of active fusion complexes at any specific calcium concentration. In contrast, the random model assumes that active fusion complexes are randomly distributed among granules in a population such that the total number of active fusion complexes is much greater than one, their average number per granule, and their number on any one granule.

Fixed Model

Triggered exocytosis of a population of granules can be treated as a survival process where the population of docked granules decreases by fusion. Let t = time in 1-s intervals, n = a fixed integer representing the number of active fusion complexes per granule, and p = the probability per second that an active fusion complex mediates a fusion event between a granule and the plasma membrane. The time dependence of exocytosis (survival) is:

S(t,n,p) = (1 - p)^nt

This formalism is the same as a coin toss problem. If granule fusion occurs on the first "heads" tossed, then p is equivalent to the probability per toss of getting a head, (1 - p) is the probability of not getting a head, t is the toss turn number, and n is the number of coins tossed per turn. When p is not very large (<0.1, S(t,n,p) decays exponentially with a time constant, np. If the parameters n or p are functions of [Ca²⁺], then increasing the value of either parameter would increase the rate of fusion. The rate of fusion is the derivative of 1 - S(t,n,p). A 0 value for either n

1. Abbreviations used in this paper: BAPTA, bisaminophenoxy ethane-N, N, N', N' tetraacetic acid; CSC, cell surface complex; HEDTA, N-hydroxyethylendiamine triacetic acid; NTA, nitritolriacetic acid.
or \( p \) would result in no fusion, while all other values for \( n \) result in every vesicle fusing (Fig. 1). For example, if each granule has one active fusion complex, \( n = 1 \), and every granule fuses.

**Random Model**

If active fusion complexes are randomly distributed among a population of granules, then different granules can have different numbers of active fusion complexes, with \( n \) and \( p \), as described above. \( S(t, <n>, p) \) is the survival curve for a population containing individual granules that may have 0, 1, 2, 3, . . . active fusion complexes. The relative abundance of granules with different numbers of active fusion complexes is set by the value of \( <n> \). Because we assume that large numbers of active fusion complexes are randomly distributed among large numbers of granules such that \( <n> \) is small relative to the total number of active fusion complexes, \( S(t, <n>, p) \) is the Poisson-weighted sum of individual, fixed model survival curves:

\[
S(t, <n>, p) = \sum_{n=0}^{\infty} \frac{(<n>)^n}{n!} e^{-<n>} (1 - p)^n e^{-<n> ((1 - p)^t - 1)}
\]

The maximal rate of fusion is the derivative of \( 1 - S(t, <n>, p) \) as \( t \) goes to 0:

\[
R_{\text{max}} = -<n> \ln (1 - p)
\]

The extent of fusion, \( E(<n>) \) is 1 - \( S(t, <n>, p) \) as \( t \) goes to infinity:

\[
E(<n>) = 1 - e^{-<n>}
\]

In the random model, increasing the value of either \( <n> \) or \( p \) as a function of \( [\text{Ca}^{2+}] \) would increase \( R_{\text{max}} \). In contrast to the fixed model, random model survival curves are not single exponentials (at low values of \( p \)), but are weighted sums of exponentials. More importantly, survival curves do not always extrapolate to 0 (100% fusion), but can asymptotically approach intermediate values as \( t \) approaches infinity (Fig. 1): the extent of fusion, \( 1 - S(t \to \infty, <n>, p) \), is a function of \( <n> \) alone.

**The Time Course of Exocytosis Can Be Fit by the Random Model**

If active complexes are randomly distributed, and an increase in \( [\text{Ca}^{2+}] \), corresponds to an increase in their number, then three predictions can be used to differentiate between these two models: (a) there should exist a range of calcium concentrations where submaximal extents of fusion are observed; (b) survival curves should be best fit by the random model; and (c) even at very high calcium concentrations (>1 mM), a small fraction of exocytotic granules should fail to fuse. In contrast, 100% should fuse if every granule has a fixed number of active fusion complexes (>0).

The extent of fusion was measured as a function of the calcium concentration in an in vitro preparation of sea urchin egg cortical granules that are predocked on isolated plasma membrane fragments. ATP, GTP, cytosol, and all other soluble factors were not included in the reaction mixture to obviate the possibility that calcium might, in addition to triggering fusion, also trigger fusion complex assembly. Consistent with the random model, submaximal extents of fusion were observed between 10 and 100 \( \mu \text{M} \) calcium (Fig. 2). To test the second prediction of the random model, survival curves were obtained from cortices that were exposed to either 16 or 350 \( \mu \text{M} \) calcium in a rapid perfusion chamber. At the lower calcium concentration, submaximal extents of fusion were observed, and the survival curve (Fig. 3A) was well fit by the random model. This curve is poorly described by the fixed model, even when the constraint of noninteger \( n \) is lifted (compare residuals of best fits in Fig. 3A and the error for the fitting parameters). At the higher calcium concentration, where the rate of fusion is much faster, the survival curve (Fig. 3B) appears to be a convolution of the delivery of calcium (45), a calcium activation process (55), and the underlying fusion process. These additional kinetic processes preclude our use of this technique for obtaining accurate values of \( <n> \) and \( p \). Despite the convolution, a better fit was obtained when this survival curve was fit to the random model (see Fig. 3B). The inability of the fixed model to fit either data set becomes even more apparent when the value of \( n \) is constrained to integer values. To test the last prediction of the random model, that even at high calcium concentrations a fraction of the granules (\( e^{-<n>} \)) fail to fuse, the fate of ~22,000 granules was determined after exposure to 2.5 mM calcium (see Fig. 4). Between 3–12 survivors were observed. This corresponded to a fusion efficiency >99.97% and to an \( <n> \) value of 8.4 ± 0.6 (mean ± SEM; \( n = 4 \)) in the random model. Since these predictions of the random model have been observed, it remains a viable description for the extent and rate of cortical granule exocytosis. In contrast, the fixed model is rejected.

**Docking Sites Can Have on Average Nine Active Fusion Complexes**

We reasoned that the maximum number of fusion complexes, on average, could be determined by randomly and irreversibly modifying the proteinaceous components of fusion complexes needed for their activity, and then measuring the extent of fusion in response to saturating concentrations of calcium. By definition, granules that fail to
Figure 2. The top panel shows the calcium dependence of exocytosis in isolated cell surface complexes consisting of plasma membrane fragments and docked, ready releasable cortical granules. Points are mean ± SD, n = 8. Between 10 and 100 μM calcium, submaximal extents of exocytosis were observed. This behavior is consistent with the random model. The bottom panel shows the transformation of survival (S) into <n> according to the relationship <n> = −ln (S), derived from the random model. As survival values approach 0, the error in <n> increases. This is caused by our assay’s inability to distinguish between 97 and 100% granule exocytosis. Below 10 μM calcium, granule exocytosis was not observed. Between 10 and 80 μM calcium, <n> was proportional to pCa. The calcium dependence of <n> above 80 μM was not measurable with this technique.

Figure 3. Survival curves for sea urchin egg cortices exposed to either 16 μM (A) or 350 μM (B) calcium. Cortices were perfused for 1 s with the calcium buffers beginning at t = 0. Survival curves were fit to either the fixed model or the random model, and residuals of the fits are plotted in the lower panels. The fitting parameters (±SE) for the fixed model in A were n = 0.31 ± 0.33, p = 0.006 ± 0.006, and the parameters for the random model were <n> = 0.38 ± 0.00 and p = 0.007 ± 0.000. (B) The fitting parameters for the fixed model were n = 0.06 ± 0.80, p = 0.129 ± 0.549, and the parameters for the random model were <n> = 6.29 ± 0.53 and p = 0.169 ± 0.022. Survival data in A was from a single experiment, while the survival curve in B is the mean (±SEM) from five experiments illustrating that both single and averaged data can be fit. The fits for average data in B were weighted by the sample variance.

Figure 4. Four examples of exocytosis in response to 2.5 mM calcium from 60 individual experiments. Isolated sea urchin cortices were exposed to calcium before (left) and 1 min after (right). Occasionally, a granule failed to fuse in the presence of high calcium (see top video pair). Bar, 10 μm.
The Poisson distribution describes many biological processes, such as the release of quanta at synaptic terminals (26). A population of granules with a Poisson distribution of active fusion complexes will have a fraction of granules (e<sup>-<i>n</i>MAX</sup>) with no active fusion complexes. By measuring the fraction, <i>S</i>, of granules in a population that survives after exposure to calcium, we can calculate the value of <i><i>n</i></i> using the formula: <i><i>n</i></i> = -ln(1 - <i>S</i>) (see Fig. 2). <i><i>n</i></i> as a function of time of exposure to an inhibitor, or as a function of inhibitor concentration, enables extrapolation to the value of <i>n</i> in an untreated cortex. The maximum number of active fusion complexes per granule, on average (<i>n</i>MAX), can be measured at saturating calcium concentrations.

To measure the value of <i>n</i>MAX, the calcium and NEM sensitivity of cortical granule exocytosis was characterized in isolated cell surface complexes consisting of plasma membrane fragments and docked, ready releasable cortical granules. Submaximal extents of exocytosis were observed between 10 and 100 μM calcium (Fig. 5). Granule exocytosis was not observed below 10 μM calcium. Mild NEM treatment shifted the survival activation curve, while more extensive treatment decreased the fraction of fusible granules. The decrease in the fraction of fusion competent granules could not be overcome with higher calcium concentrations (>1 mM). The plateau region observed above 1 mM calcium is consistent with the saturation of calcium sensor-binding sites: above 1 mM calcium additional active fusion complexes cannot be recruited. This calcium-NEM dose-response behavior is expected if calcium increases the value of <i>n</i> while NEM treatment decreases <i>n</i>. This behavior is not unique to NEM, since treatment with proteases has the same effect (23). This classic pharmacological behavior, progressive shifting of a dose-response curve using mild treatments of irreversible inhibitors and the lowering of the extent of a reaction after harsher treatment, has been observed in tissue dose-

**Figure 6.** (A and B) The time and concentration dependence of the NEM inactivation of cortical granule survival after transformation into <i>n</i>. The decrease in <i>n</i> is an exponential function of both the time of NEM inactivation or NEM concentration at the indicated calcium concentration. All experimental points are mean ± SD, n = 3. Dotted lines are weighted averages of all individual single exponential fits using the equation: <i>y = a e^{(-b x)}</i>, where <i>a</i> is a scaling parameter, <i>b</i> is either the time or concentration decay constant for NEM inactivation, and <i>x</i> is either the time or NEM concentration. The exponential extrapolation to either 0 NEM concentration or time 0 is mean ± SEM derived from individual fits and displayed on a log-linear plot to explicitly show the exponential nature of this process.
vation at calcium concentrations >1 mM. The decrease in \(<n>\) is an exponential function of both the time of NEM treatment and the NEM concentration during a fixed time period (Fig. 6). A single exponential decay would be observed if fusion complex inactivation was both random and required either a single-site modification or modification of multiple identical sites. At all calcium concentrations measured, above 1 mM calcium \(<n>_{\text{MAX}} = 9.0 \pm 0.2\) (mean \(\pm 95\%\) confidence) when extrapolated to a NEM concentration of 0, and \(<n>_{\text{MAX}} = 8.1 \pm 2.2\) (mean \(\pm 95\%\) confidence) when extrapolated to time 0. These values of \(<n>\) at high calcium concentrations are in agreement with the value of 8.4 \(\pm 0.6\) derived from the fraction of granules that failed to fuse at 2.5 mM calcium (Fig. 4). At lower concentrations of calcium (<1 mM), the extrapolated values of \(<n>\) were lower. For example, at 809 \(\mu\)M calcium, \(<n>\) had an extrapolated value of 6.0 \(\pm 2.2\) (mean \(\pm SD\)).

We have demonstrated that at >1 mM calcium, \(<n>\) reaches a maximum value of 9.0 \(\pm 0.1\) (mean \(\pm SEM, n = 10\)). Below 1 mM calcium, the extent of fusion can be interpreted as the fractional activation of fusion complexes in response to calcium. Fig. 7 depicts the cumulative results of four techniques used to measure the calcium dependence of \(<n>\) from sea urchin exocytotic preparations in vitro. No fusion was observed below 10 \(\mu\)M calcium, so \(<n> = 0\). Between 10 and 800 \(\mu\)M calcium, \(<n>\) increased with pCa. It is important to realize that 50\% of granules fuse between 18 and 22 \(\mu\)M calcium, while in contrast, 50\% of the fusion complexes, which can be activated, were active at \(~100\ \mu\)M calcium. These different half maximal calcium concentrations are derived, in part, from the same data set, and they illustrate the model dependence of assigning calcium-binding affinities.

To examine the applicability of the random model to describe the time course of exocytosis in intact cells, we visualized a population of cortical granules in *L. pictus* eggs during fertilization and measured the time course of exocytosis (Fig. 8, A–C). The exocytotic survival curve for this egg is shown in Fig. 8 D. After perfusion with sperm, there was a lag time during which sperm must find the egg, fuse with the oolemma, and release calcium from the egg’s intracellular stores. Presumably, when the calcium concentration at docking sites reaches a threshold, granule fusion complexes are activated and mediate fusion with the plasma membrane. This survival curve can be parameterized by the random model (Fig. 8 D). In the egg, as observed in the isolated sea urchin egg cortex (Fig. 3 B), the initial portion of the survival curve (Fig. 8 D) may be convolved with the temporal and spatial characteristics of the calcium delivery and a calcium activation process. During fertilization, the subcortical space that was occupied by cortical granules before fusion (Fig. 8 A) is not invaded by cellular organelles in the cytoplasm after exocytosis (see Fig. 8 C).

**Discussion**

**The Random Model**

We envision cortical granule docking sites as having, depending upon [Ca\(^{2+}\)], an average of between 0–9 active fusion complexes. Using the random model, we can rationalize much of the calcium-dependent exocytotic behavior observed in this preparation. These include (a) the sigmoidal dependence of survival on pCa due to the existence of submaximal responses (4, 60); (b) the increase in fusion rate with increase in [Ca\(^{2+}\)] (10, 54); (c) the discrepancy between the calcium concentration required for maximal extents and rates (61); (d) shifting of the calcium activation curve with inhibitors of exocytosis (23, 34, 58); and (e) aging (45, 61), where a progressive decrease in calcium sensitivity is observed with time in the isolated planar cortex. The random model succeeds in describing all of these behaviors in the sea urchin with the simple hypothesis that an increase in [Ca\(^{2+}\)] corresponds to an increase in the average number \((<n>)\) and perhaps the efficacy \((p)\) of active fusion complexes at docking sites. According to this model, the number of active fusion complexes per granule is a Poisson-distributed random variable, with a maximal average value of \(<n> = 9\). In contrast, deterministic models such as the fixed model are not compatible with the data.

The random model links the rate and extent of degranulation, and can be used to fit the time dependence of exocytosis (Fig. 3). The fraction of granules that fuse at any calcium concentration is solely a function of one of these parameters, \(<n>\), the average number of active fusion complexes at a docking site. Submaximal extents of fusion are observed because a fraction of granules containing no active fusion complexes will always exist when active fu-
sion complexes are randomly distributed. In the random model, the maximal rate of exocytosis is directly proportional to the average number of active fusion complexes: as $[Ca^{2+}]$, increases $<n>$, the maximal rate of exocytosis will increase. Since the extent of exocytosis is an asymptotic function of $<n>$ and the maximal rate is a linear function of $<n>$, they need not plateau at the same value of $<n>$. This differential dependence on $<n>$ can explain the difference in the calcium dependence of rate and extent observed in sea urchin eggs (61) and chromaffin cells (3). Since $<n>$ does plateau at $>1$ mM calcium, the maximal rate should also plateau at these concentrations. A plateau in the apparent maximal rate has been observed at $>1$ mM calcium in preliminary experiments (data not shown).

The random model does have limitations. In essence, the random model shifts the burden of explaining incomplete exocytosis to that of explaining incomplete conversion of inactive to active complexes. The nature of the coupling between calcium sensors and fusion complexes is not intrinsic to the model, nor is the assembly, activation, and other unknown aspects of producing a Poisson distribution of active fusion complexes. We can only speculate on issues such as the number and nature of calcium-binding sites, cooperativity between these sites, and the molecular mechanism for producing and maintaining Poisson-generated fusion complex heterogeneity. The origin of this be-

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure8.png}
\caption{Cortical granule exocytosis observed in intact sea urchin eggs. Sea urchin eggs were perfused with a 1:1,000 dilution of sperm in artificial seawater at $t = 0$ s. (A–C) The subcortical space, normally occupied by cortical granules in the unfertilized egg, was imaged using differential interference video contrast microscopy at 40, 70, and 100 s after perfusion. The bar in C is 5 \mu m.

The Kaplan-Meier survival data (C; mean $\pm$ SD) for sperm-triggered cortical granule exocytosis is shown in D. The solid line in D is a survival curve based on the random model ($S = e^{-\lambda(t - \theta)/(t - \lambda)}$) with $<n> = 4.5$, $p = 0.029$, and $\Delta t = -48.6$ s to allow for nonzero starting times.
}
\end{figure}
behavior is under investigation. Fusion complexes which do not contact a target membrane may not be functional (55), and the model yields only the average number of functional complexes per granule. Thus, the number of 'functional' complexes may differ from the total number of complexes. Changes in the probability that a fusion complex initiates the fusion process have not been incorporated in this version of the model, nor have potential rate limiting steps, such as time dependent delivery of calcium or time dependent activation of fusion complexes. Indeed, there is evidence that fusion complexes are not "instantaneously" activated (61). This can be seen in the discrepancy between the initial time course of exocytosis and the initial time course generated using the random model (see Figs. 3 and 8). Finally, it is known that in vitro, granules adjacent to granules that have already fused are more likely to fuse themselves (45). This behavior might result from a local modification of n and p values on neighboring granules. This version of the model has not been modified to account for this.

**Alternative Hypotheses**

One might speculate that a granule-docking site contains a single fusion complex, but that the NEM-sensitive site resides on a factor required to assemble that complex in response to an increase in the calcium concentration. Because fusion complex assembly is thought to require ATP and other soluble proteins and factors (5, 44), and our assay was conducted in the absence of ATP, GTP, and all other soluble factors and proteins, this seems unlikely. Furthermore, cortical granule exocytosis in vitro has been observed within tens of milliseconds after calcium delivery (54); as it is thought in other systems (2), it is likely that only preassembled fusion complexes are functional.

By invoking schemes involving a single preassembled fusion complex at a docking site, it may be possible to explain some of the individual calcium-dependent behaviors associated with exocytosis. First, it is known that calcium increases the rate of fusion (10, 54) and NEM decreases the fusion rate (45). By invoking an intrinsic, time-dependent inactivation process, one can explain both submaximal extents of fusion and the shift in calcium dose–response observed with NEM treatment. However, there is no evidence for time-dependent inactivation in this system (45; Kaplan, D., P.S. Blank, M.S. Cho, I. Steinberg, and J. Zimmerman. *Biophys. J.* 66:A284 [Abstr.]). Second, if progressive NEM treatment on a single fusion complex progressively shifts its calcium-binding constant until the complex is destroyed, one can explain why mild treatment with inhibitors shifts the calcium dependence of exocytosis and extensive treatment with inhibitors decreases the fraction of granules that fuse. However, in this scheme, the calcium dependence of rate and extent should be identical, contrary to our data (extent plateaus at ~100 μM, while rate plateaus at >1 mM calcium). Finally, any scheme having a single fusion machine per granule would require a cellular mechanism to enforce a one-granule—one-fusion complex distribution. No such mechanism has been described, and seems improbable when one considers the ontogeny of these large cortical granules.

**In Vivo Application**

The primary physiological role of cortical granule exocytosis is to prevent polyspermy (24), thus it is important that most of an egg's 15,000 cortical granules fuse within the first minute after sperm entry. An egg takes 9–10 s for 50% of the granules to fuse (35) in response to a calcium concentration estimated to reach between 150 and 170 μM (36). If we set random model parameters <n> = 4.5 and p = 0.03, 98.7% granule fusion will occur by 3 min with an apparent half-time of ~10 seconds, and at a maximum rate of ~2,000 granules fusing per second. An <n> value of 4.5 requires that the calcium concentration at docking site reach ~100 μM, which is comparable to values reached at synapses (1, 33, 43). A maximum rate of 2,000 granules fusing per second is comparable to exocytotic rates observed in neuronal cells using capacitance techniques (3, 18). In the sea urchin egg, as in neuronal cells (6, 20, 42), calcium-triggered exocytosis is followed by endocytosis (59). Interestingly, docking sites that were occupied by cortical granules before fusion are not invaded by other exocytotic organelles located deeper in the cytoplasm when observed by differential interference contrast microscopy during the first few minutes after fertilization (see Fig. 8). This suggests that organelle access to the subcortical space is limited, consistent with the recent finding that newly formed endosomes occupy this space (59).

**Spare Receptor System**

It is well known that fractional occupation of receptors by ligand need not correspond directly to fractional biological response (41, 46), as in cells having spare receptors (29). This behavior has been reconstituted and studied in vitro (40). Activation of a single surface receptor may invoke maximal response despite the fact that most surface receptors are free to bind ligand. Apparently, this behavior can also be invoked in intracellular receptor-mediated systems, such as calcium-triggered exocytosis. Spare receptor mechanisms allow low affinity sites to mediate responses to low concentrations of ligand. More importantly, spare receptor systems may bring about a sharper dose response transition, comparable to cooperative ligand-binding transitions (30, 37). Thus, a biological switch with a sharp transition may be implemented by spare receptor system or strong positive cooperativity.

The calcium dependence of exocytosis has been studied in different cell types, as models for synaptic release (3, 9, 11, 18, 19, 21, 22, 27, 50, 57). We suggest that the rate and extent of calcium-triggered exocytosis at synapses may also be a function of the number, distribution, and efficacy of fusion complexes at docking sites, in addition to the speed, amplitude, and duration of calcium delivery to the exocytotic calcium sensor. It would be remarkable if Poisson analysis, which was essential for constructing the vesicular hypothesis for synaptic release (26), would now allow for the interpretation of its calcium dependence.

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