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Review

Calcium cooperativity of exocytosis as a measure of Ca²⁺ channel domain overlap

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ABSTRACT

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neurotransmitter release rate to changes in the synaptic Ca^{2+} current, which is varied using appropriate voltage-clamp protocols or via pharmacological Ca^{2+} channel block under the condition of constant single-channel Ca^{2+} current. The slope of the resulting log–log plot of transmitter release rate versus presynaptic Ca^{2+} current is termed Ca^{2+} current cooperativity of exocytosis, and provides indirect information about the underlying presynaptic morphology. In this review, we discuss the relationship between the Ca^{2+} current cooperativity and the average number of Ca^{2+} channels participating in the exocvtosis of

exocytosis, and provides indirect information about the underlying presynaptic morphology. In this review, we discuss the relationship between the Ca^{2+} current cooperativity and the average number of Ca^{2+} channels participating in the exocytosis of a single vesicle, termed the Ca^{2+} channel cooperativity. We relate these quantities to the morphology of the presynaptic active zone. We also review experimental studies of Ca^{2+} current cooperativity and its modulation during development in different classes of synapses.

The number of Ca^{2+} channels contributing to the exocytosis of a single neurotransmitter

vesicle in a presynaptic terminal has been a question of significant interest and debate, and is important for a full understanding of localized Ca²⁺ signaling in general, and synaptic

physiology in particular. This is usually estimated by measuring the sensitivity of the

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62 1. Introduction

63 Synaptic neurotransmitter release and endocrine hormone 64 secretion are fundamental physiological processes, and there 65 has been sustained interest and active research aimed at 66 understanding better the steps leading from Ca²⁺ entry to exocytosis. Synaptic transmitter release occurs from active 67 zones, which contain Ca²⁺ channels and transmitter-filled 68 vesicles docked at release sites. The arrangement of channels 69 and vesicles is important in the release process, since 70 exocytosis is evoked by Ca²⁺ that enters the synaptic terminal 71 through voltage-dependent Ca²⁺ channels (Llinás et al., 1981; 72 Stanley, 1993) and remains highly localized to the channels' 73 Ca²⁺ domains (Augustine et al., 2003; Chad and Eckert, 1984; 74 Fogelson and Zucker, 1985; Neher, 1998a; Simon and Llinas, 751985). However, it is exceedingly difficult to determine active 76 77 zone morphology due to the small size of the active zone. Even in cases where such morphological information has been 78 79 determined in detail using freeze-fracture combined with 80 electron or atomic force microscopy, for instance at the frog 81 neuromuscular junction (Ceccarelli et al., 1979; Harlow et al., 82 2001; Heuser et al., 1979; Pumplin et al., 1981; Stanley et al., 83 2003), there remains a lack of complete knowledge of the 84 number of functional channels that open per action potential per vesicle, and the contribution of individual channels to 85 vesicle release. Given this limitation in the direct measure-86 ment of functional active zone morphology, indirect tech-87 niques are used to estimate the number of Ca²⁺ channels 88 contributing to an exocytotic event, which we will refer to 89 below as the Ca^{2+} channel cooperativity. These techniques 90 consist of varying the number of channels that open during 91 a stimulus while measuring both the presynaptic Ca²⁺ current 92and the release of transmitter through either presynaptic 93 capacitance measurements or postsynaptic measurements. 94 Typically, a log-log plot of the release variable and the 95 presynaptic Ca²⁺ current is made, and the slope of the plot is 96 determined (see e.g. Bucurenciu et al., 2010; Fedchyshyn and 97 98 Wang, 2005; Kochubey et al., 2009; Mintz et al., 1995; Quastel 99 et al., 1992; Wu et al., 1999). This slope, the Ca²⁺current cooperativity, provides indirect information about the mean 100 number of channels contributing to each exocytotic event and 101 the active zone morphology. A large Ca²⁺ current cooperativity 102suggests that many channels contribute to exocytotic events, 103 while a small Ca²⁺ current cooperativity is usually understood 104 to mean that release is gated by just a few proximal channels. 105 In particular, a Ca²⁺ current cooperativity near 1 is often taken 106 107 as an indication that each exocytotic event is gated by the 108 opening of a single channel (reviewed in Gentile and Stanley, 109 2005; Schneggenburger and Neher, 2005).

Measurements of the Ca²⁺ current cooperativity have been used to infer information about synaptic morphology in a wide variety of synapses, including the squid giant synapse (Augustine et al., 1991; Augustine and Charlton, 1986; Llinás et al., 1981), sensory ribbon synapses (Brandt et al., 2005; 114 Coggins and Zenisek, 2009; Jarsky et al., 2010; Johnson et al., 115 2008; Keen and Hudspeth, 2006; Thoreson et al., 2004), motor 116 nerve terminals (Quastel et al., 1992; Shahrezaei et al., 2006; 117 Yoshikami et al., 1989), the rodent calyx of Held (Borst and 118 Sakmann, 1996; Fedchyshyn and Wang, 2005; Kochubey et al., 119 2009; Sakaba and Neher, 2001; Wu et al., 1998; Wu et al., 1999) 120 and other central synapses (Bucurenciu et al., 2010; Gentile 121 and Stanley, 2005; Mintz et al., 1995). Theoretical studies have 122 also explored this experimental assay (Bertram et al., 1999; 123 Bucurenciu et al., 2010; Coggins and Zenisek, 2009; Matveev 124 et al., 2009; Meinrenken et al., 2002; Quastel et al., 1992; 125 Shahrezaei et al., 2006; Yoshikami et al., 1989; Zucker and 126 Fogelson, 1986). The first aim of this review is to clarify what 127 information is actually provided by the Ca²⁺ current cooperativity, 128 and to contrast this with the Ca²⁺ channel cooperativity, which is 129 only indirectly inferred from current cooperativity measure- 130 ments. The second aim is to review Ca²⁺ current cooperativity 131 studies and focus on several cases in which the current 132 cooperativity has been used to obtain important information on 133 active zone morphology or changes in morphology. 134

2. Biochemical Ca²⁺ cooperativity of exocytosis 136

Measurements of the Ca²⁺ current cooperativity that reflects 137 active zone morphology first arose in investigating the 138 biochemical (intrinsic) Ca²⁺cooperativity of release introduced 139 by Dodge and Rahamimoff(1967), which is independent of 140 synaptic morphology. The latter measure, which we denote by 141 *n*, provides a lower bound on the number of Ca²⁺ binding steps 142 required to evoke vesicle fusion (Dodge and Rahamimoff, 143 1967). The most direct biochemical cooperativity measure- 144 ment technique uses caged-Ca²⁺ compounds to raise the Ca²⁺ 145 concentration almost uniformly throughout the synaptic 146 terminal and Ca^{2+} imaging to measure the internal Ca^{2+} 147 concentration (Beutner et al., 2001; Bollmann et al., 2000; 148 Kochubey et al., 2009; Lando and Zucker, 1994; Schneggen- 149 burger and Neher, 2000). A less direct approach is to vary the 150 extracellular Ca²⁺ concentration [Ca²⁺]_{ext}, which will affect Ca²⁺ 151 influx through all open Ca2+ channels, and increase the 152 intracellular Ca²⁺ concentration (Augustine and Charlton, 153 1986; Borst and Sakmann, 1996; Dodge and Rahamimoff, 1967; 154 Katz and Miledi, 1970; Lester, 1970; Llinás et al., 1981; Mintz et al., 155 1995; Stanley, 1986). The biochemical cooperativity is then 156 obtained using the log–log slope of the Ca²-secretion curve: 157

$$n = \frac{d \log R}{d \log [Ca]} \tag{1}$$

where [Ca] represents the concentration of either intracellular or **159** extracellular Ca²⁺, varied in a non-saturating range. Alternative- 160 ly, some studies define *n* as the parameter of the Hill-function fit 161 to the entire saturating Ca²⁺-release curve (Jarsky et al., 2010; 162

Sakaba and Neher, 2001). However, in the absence of a biological 163 argument for such a functional relationship, the definition given 164 by Eq. (1) is preferable because it is model-independent (Quastel 165 et al., 1992). The measurement of n does not depend on the 166 number of Ca²⁺ channels that open during the stimulus, so it 167 provides no information on the active zone morphology. Instead, 168 it measures the intrinsic Ca²⁺ sensitivity of transmitter release, 169 and its value ranges from 1 to 5 across different preparations 170 171 (Augustine et al., 1985; Augustine and Charlton, 1986; Bollmann 172 et al., 2000; Borst and Sakmann, 1996; Brandt et al., 2005; Dodge and Rahamimoff, 1967; Duncan et al., 2010; Llinás et al., 1981; 173 Mintz et al., 1995; Reid et al., 1998; Schneggenburger and Neher, 174 2000; Stanley, 1986). Values of n greater than one obtained in 175 many synapses suggest that exocytosis requires the binding of 176 several Ca²⁺ ions to proteins gating release, possibly Synapto-177 tagmin, a well-described Ca2+ sensing protein that is also a 178 component of the SNARE protein complex, and plays a key role in 179 the gating of transmitter release (Fernandez-Chacon et al., 2001; 180 Geppert et al., 1994; Nagy et al., 2006; Pang et al., 2006; Stevens and 181 Sullivan, 2003; Xu et al., 2007). Isoforms 1, 2, and 9 all have five Ca² 182 ⁺ binding sites, with three on the C2A domain and two on the C2B 183 domain of the protein (see Rizo and Rosenmund, 2008 for review). 184 Note that several studies of sensory ribbon synapses suggest a 185 non-cooperative, linear relationship between release and [Ca²⁺], 186 and suggest an involvement of a different Ca²⁺ release sensor, 187 possibly otoferlin (Dulon et al., 2009; Keen and Hudspeth, 2006; 188 Roux et al., 2006; Thoreson et al., 2004) or non-neuronal 189 190 synaptotagmin IV (Johnson et al., 2010).

Since transmitter release becomes saturated at high con-191 centrations of internal Ca2+, measurements of biochemical 192cooperativity are made at non-saturating Ca²⁺ levels. It has been 193 shown recently (Lou et al., 2005) that the "intrinsic" biochemical 194cooperativity may vary with [Ca2+] even before saturation is 195reached, possibly due to an allosteric Ca²⁺ binding mechanism 196 that cannot be approximated as a simple serial or parallel 197 sequence of Ca²⁺ binding steps. It was suggested that this is due 198to the transition from asynchronous to synchronous release. 199Another study showed that the biochemical cooperativity for 200 asynchronous release, n=2, is considerably lower than that for 201 synchronous release, n=5, in the calyx of Held synapse (Sun 202 et al., 2007). Studies of asynchronous release in other synapses 203204have found the biochemical cooperativity to be the same as that for synchronous release, but with a lower Ca²⁺ affinity (Goda and 205Stevens, 1994; Ravin et al., 1997). There is evidence that 206biochemical cooperativity is not a constant property, but can 207be lowered by genetically reducing the expression level of the 208 SNARE proteins syntaxin 1A and synaptobrevin (Stewart et al., 2092000). It can also be lowered by pharmacologically cleaving 210SNAP-25 with Botulinum toxin (Cull-Candy et al., 1976) or 211 cleaving VAMP/synaptobrevin with tetanus toxin (Bevan and 212213Wendon, 1984). Finally, it has been proposed that biochemical cooperativity can be dynamically modulated by intracellular 214 kinases such as PKC (Yang et al., 2005). 215

216 3. Ca²⁺ channel cooperativity of exocytosis

An important functional characteristic of the active zone morphology is the mean number of channels that contribute ions to the triggering of a release event. This cannot be

measured experimentally because it is impossible to track the 221 paths of individual Ca²⁺ ions to determine their channel 222 source. However, it can and has been estimated using 223 computer simulations (Luo et al., 2008; Shahrezaei et al., 224 2006). The simulations by Shahrezaei et al., for the frog 225 neuromuscular junction, demonstrated that although there 226 were as many as six Ca²⁺ channel openings per vesicle per 227 action potential, only one or two proximal channels provided 228 the Ca²⁺ ions that evoke release from a nearby release site. 229 This computer simulation suggests that distal channels play 230 little role in gating release from this neuromuscular junction. 231 Since the number of channels contributing to a release event 232 can be no greater than the number of Ca²⁺ binding sites, this 233 measure of the Ca²⁺ channel cooperativity is bounded above 234 by the biochemical cooperativity, n. 235

The Ca²⁺ channel cooperativity was defined somewhat 236 differently in Matveev et al.(2009), where m_{CH} was quantified as 237 the number of channels contributing Ca^{2+} to the local Ca^{2+} 238 domain surrounding the vesicle release sensor. When defined 239 this way, many channels may contribute to the domain, although 240 ions from only a few (at most n) channels actually bind to proteins 241 at the release site. The channel cooperativity m_{CH} can exceed n but 242 is bounded by M, the number of channels in the vicinity of the 243 release site. The advantage of defining m_{CH} in this way is that 244 it provides useful information about the extent of overlap of the 245 Ca²⁺ nanodomains of individual open channels, and quantifies 246 the number of channels participating in release over several 247 exocytosis events. In fact, many studies tacitly assume this 248 second Ca²⁺ channel cooperativity definition; for instance, Borst 249 and Sakmann(1996) argue for a possibility of dozens of channels 250 being involved in the release of a vesicle in the rodent calyx of 251 Held synapse from immature animals. 252

For the idealized case in which the channels are equidis- 253 tant from a release site, channel cooperativity equals the 254 average number of Ca^{2+} channels that open to trigger a release 255 event (i.e., the average number of open channels given that 256 exocytosis takes place). For example, if each release site is 257 surrounded by an average of a dozen equidistant channels, 258 each with open probability of 50% during a stimulus, and 259 assuming for simplicity that exocytosis occurs at every 260 depolarization event, then m_{CH} =6 since six channels open 261 and provide Ca^{2+} to the local domain at the release site. This 262 number is not limited by the biochemical cooperativity, which 263 would typically be less than six. 264

4. Ca²⁺ current cooperativity of exocytosis 266

Although knowledge of the number of Ca^{2+} channels involved 267 in a single exocytotic event is of significant interest for a full 268 understanding of localized Ca^{2+} signaling in general and 269 synaptic physiology in particular, as pointed out above such 270 a characteristic cannot be measured experimentally. This 271 limitation has led to the use of an indirect measure for the 272 number of channels participating in an exocytotic event. This 273 measure, the $Ca^{2+}current$ cooperativity of exocytosis, $m_{I_{ca}}$, was 274 originally introduced in the study of the biochemical Ca^{2+} 275 cooperativity (Augustine et al., 1985; Llinás et al., 1981), and 276 was soon hypothesized to depend on the localization of Ca^{2+} 277 influx (Augustine and Charlton, 1986; Chad and Eckert, 1984; 278

Yoshikami et al., 1989). Current cooperativity was first 279analyzed with the help of computational modeling by Zucker 280and Fogelson(1986) and analytically by Yoshikami et al.(1989) 281 and Quastel et al.(1992). Zucker and Fogelson (1986) showed 282 that $m_{L_{c}}$ would increase from 1 to *n* as the membrane potential 283increased, which changes the open probability and the driving 284 force. Here we review experiments in which the number of Ca²⁺ 285channel openings during a stimulus is varied without changing 286287the single-channel current, and both transmitter release and the presynaptic Ca^{2+} current (I_{Ca}) are measured. 288

There are two main methods for measuring $m_{I_{ca}}$, correspond-289ing to two different ways of modifying the number of open 290channels without changing the single-channel Ca²⁺ current. 291 One method involves voltage-clamping the pre-synaptic site 292and administering depolarizations of varying amplitude or 293duration (Augustine et al., 1985; Llinas et al., 1982). The range 294of depolarization amplitudes is kept sufficiently narrow to 295minimize the changes in the driving force for Ca²⁺, ensuring an 296approximately constant single-channel current (Quastel et al., 2971992). Avoiding this complication, a tail current protocol in-298volves a pre-depolarization to near the Ca²⁺ reversal potential, 299which activates the channels, and is followed by a step to a 300 hyperpolarized voltage, increasing the driving force so that Ca²⁺ 301 302floods into the cell through channels opened by the preceding 303 depolarization (Stanley, 1995; Stanley, 2005). In this case the 304 number of open channels is varied by changing the pre-305 depolarization duration. The second approach uses pharmaco-306 logical agents to block presynaptic Ca²⁺ channels (Mintz et al., 1995; Wu et al., 1999; Yoshikami et al., 1989). The agent can be 307 specific to a certain type of Ca²⁺ channel (for example, P/Q-type 308 channels can be blocked with ω-agatoxin IVA), or a non-specific 309 agents like Cd²⁺ could be applied to block Ca²⁺ channels of all 310 types. With this approach, the number of channels that open 311 during a stimulus is varied by applying different concentrations Q3,Q4 of the blocker. In the following we focus on non-specific channel 313 block except where indicated otherwise. 314

The Ca²⁺ current cooperativity can then be defined as the slope of the log–log plot of the release rate R versus the total Ca²⁺ current, I_{Ca} (Quastel et al., 1992)

$$n_{I_{Ca}} = \frac{d \log R}{d \log I_{Ca}} = \frac{d \log P(R)}{d \log p_o},$$
(2)

where P(R) is the probability of release, and p_o is the single-319 channel open probability. The second equality in Eq. (2) uses 320 the fact that the Ca²⁺ current is proportional to the single-321 322 channel open probability, and assumes that the influx is brief (which is true for the tail-current protocol), so that each 323 324 channel is either in an open or closed configuration (i.e., there is no flickering). As discussed below in more detail, the value 325 of $m_{I_{Ca}}$ is not constant, but initially increases with I_{Ca} (see Fig. 2) 326 and then decreases as I_{Ca} approaches saturating levels. 327

Finally, note that the biochemical cooperativity is known to be an upper bound for the Ca²⁺ current cooperativity (Mintz et al., 1995; Quastel et al., 1992; Wu et al., 1999; Zucker and Fogelson, 1986):

 $m_{I_{Ca}} \leq n.$

r

332 That is, $m_{I_{Ca}}$ initially increases with the number of open 334 channels but when there are many channels, the Ca²⁺ 335 concentration approximates a uniform rise owing to extensive domain overlap. For the case of sensory ribbon synapses, the 336 biochemical cooperativity is lower than in many other 337 synapses, with several studies reporting non-cooperative, 338 near-linear relationship between Ca^{2+} and release (Johnson 339 et al., 2008; Keen and Hudspeth, 2006; Thoreson et al., 2004). 340 This constrains the current–release relationship to be linear as 341 well, regardless of the degree of channel domain overlap. 342

5. Relationship between current and channel 343 cooperativities: equidistant channel model 345

What does $m_{I_{ca}}$ tell us about the number of channels contribut- 346 ing to release, m_{CH} ? If m_{CH} =1, then each release site has a single 347 proximal channel responsible for gating release, so $m_{I_{ca}}$ =1 348 (Augustine, 1990; Bucurenciu et al., 2008; Mintz et al., 1995; 349 Stanley, 1993; Wu et al., 1999; Yoshikami et al., 1989; Zucker and 350 Fogelson, 1986). Thus, if some fraction of presynaptic channels 351 is blocked, then the probability of release from the terminal will 352 be reduced by this same fraction since each release site 353 associated with a blocked channel will be inactive. The 354 relationship between $m_{I_{ca}}$ and m_{CH} is more subtle if an exocytotic 355 event is evoked by Ca²⁺ from more than one channel. To 356 illustrate, we examine the case of equidistant channels. While 357 this is a severe geometric constraint, it enables us to derive 358 formulas that provide insight into the relationship between 359 active zone morphology, the actions of Ca²⁺ buffers, and the Ca²⁺ 360 cooperativities we have defined, $m_{I_{Ca}}$, m_{CH} , and n. 361

Starting with the simplest case of two equidistant channels, $_{362}$ the probability of release can be expressed in terms of the $_{363}$ conditional probabilities of release given that a certain number $_{364}$ of channels are open, P(R|1) and P(R|2), and the probabilities that $_{365}$ either one or two channels open, P(1) and P(2): $_{366}$

$$P(R) = P(R|1)P(1) + P(R|2)P(2).$$
(3)

As illustrated in Fig. 1, $P(1)=2p_0(1-p_0)$ and $P(2)=p_0^2$, where p_0 is 368 the probability that a channel is open. We can then rewrite 369 Eq. (3) as 370

$$P(R) = P(R|1)(2p_o(1-p_o) + rp_o^2)$$
(4)

372

where

$$r = \frac{P(R|2)}{P(R|1)}$$
(5)



Fig. 1 – Probabilities of distinct configurations of the release site with two channels per vesicle. Denoting the open channel probability by p_0 , the probability that neither channel is open is $P(0)=(1-p_0)^2$, whereas the probability of one open channel is $P(1)=2p_0(1-p_0)$, and the probability of both channels being open is $P(2)=p_0^2$. Modified from Matveev et al. 2009.

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Fig. 2 – (A) Log-log plot of release probability versus open channel probability (computed using Eq. (4)), assuming two equidistant channels, and no saturation of release $(r=2^5=32)$. The value of the slope yields Ca²⁺ current cooperativity of exocytosis, $m_{I_{ca}}$, and is different along different parts of the curve. The black curve is a regression line, the slope of which would be the experimentally determined $m_{I_{ca}}$. (B) The dependence of both $m_{I_{ca}}$ and m_{CH} on p_o . Colors of points correspond to colors of line segments in (A). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

is the ratio of release given that two channels open over that when one channel opens. The release ratio, r, describes the relative contribution that a second open channel makes to release. The advantage of expressing cooperativity measures in terms of r is that it fully quantifies the sensitivity of release to the number of open channels, without need to describe explicitly the details of the underlying Ca²⁺ binding process or [Ca²⁺] diffusion.

To fully analyze a particular synaptic release model it is 381 necessary to calculate r, as we do below to examine the effect 382 of channel distribution and buffering, but some simple cases 383 can be understood without calculation. For example, if r=1384 then the release site is saturated by the Ca²⁺ from one open 385 channel, and the opening of a second channel makes no 386 contribution. At the other extreme of no saturation, the 387 opening of a second channel doubles the probability that 388 each Ca²⁺ binding site will be occupied. Since there are *n* such 389 sites (the biochemical cooperativity), $r=2^n$ in this case of no 390 saturation. The Ca2+ buffering and the distance of the 391 channels from the release sites also affect the level of 392 saturation. If the channels are close to the vesicle, or buffering 393 is weak, then r is near 1. Conversely, large channel-vesicle 394separation and/or strong buffering results in a release ratio 395 that is near 2^n . Apart from the release ratio, the only other 396 parameter affecting current and channel cooperativities is the 397 probability that a channel is open, p_0 , or equivalently, the 398 fraction of open channels in the terminal. 399

400 Eqs. (2)–(5) can be used to derive relationships between the 401 Ga²⁺ current and channel cooperativities and the two inde-402 pendent parameters r and p_o (see Matveev et al., 2009 for 403 details). In the case of two equidistant channels, these 404 relationships are:

 $m_{CH} = \frac{1 + (r-1)p_o}{1 + (r-2)\frac{p_o}{2}}$

$$m_{I_{Ca}} = \frac{1 + (r-2)p_0}{1 + (r-2)\frac{p_0}{2}}$$
(7)

where $m_{l_{ca}}$ is obtained by taking the logarithmic derivative of 408 Eq. (4) with respect to p_{o} , while m_{CH} is the average number of 409 open channel given that release has occurred. 410

While these expressions are similar, there is nevertheless a 411 difference. One can see that, in this case of two equidistant 412 channels, 413

$$m_{I_{Ca}} \leq m_{CH} \leq 2.$$

Other results can be deduced from Eqs. (6) and (7) by 415 considering some limiting cases. When r=2, release scales 416 linearly with the number of open channels (there is twice as 417 much release with 2 open channels compared to 1 open 418 channel), so $m_{I_{Ca}}$ is identically 1. In contrast, in this case m_{CH} 419 depends on the open channel probability, $m_{CH}=1+p_o$, linearly 420 increasing from 1 to 2 as p_o is increased from 0 to 1. This is 421 reasonable, since m_{CH} is conditioned on the fact that release 422 occurs (so at least one channel must open to evoke the release) 423 and in the linear regime (r=2) m_{CH} reflects the fraction of 424 instances in which the second channel opens. Another 425 instructive case is large r. In this case both channels have to 426 open to trigger release, so m_{L_2} and m_{CH} approach their upper 427 bound of 2 (the number of available channels) regardless of the 428 fraction of open channels during the stimulus. 429

Fig. 2 illustrates the definition of $m_{I_{ca}}$ for two equidistant 430 channels (Eq. (7)), showing its relationship to the experimen- 431 tally measured value of $m_{I_{ca}}$. The experimental value is 432 typically calculated as the slope of the log-log plot of some 433 measure of release versus I_{Ca} , using linear regression through 434 several data points to construct the curve. The corresponding 435 theoretical definition of $m_{I_{ca}}$ is demonstrated in Fig. 2A, where 436 P(R) is given by Eq. (4). Here we vary the channel open 437 probability, as would be the case in an experiment using either 438 the tail current or non-specific channel block protocol, and we 439 fix the model active zone morphology. A release mechanism 440 with 5 Ca²⁺ binding sites is assumed, and we consider the case 441 where there is no saturation, so $r=2^5=32$. The slope of the 442

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black regression line through the points is the Ca^{2+} current 443 cooperativity that would be calculated experimentally if all of 444 the data points in the figure were measured. However, the 445 slope of the log(P(R)) versus $log(I_{Ca})$ curve given by Eq. (4) that 446 defines the points in this figure varies with po. This is 447 illustrated with the three colored line segments in Fig. 2A. 448 The slopes of these line segments are given by Eq. (7), with 449 r=32 and three different values of p_o . These slopes, $m_{I_{ca}}$, are 450451 plotted in Fig. 2B as correspondingly colored points. They lie on a curve that is the slope of the log(P(R)) versus log(I_{Ca}) curve 452for p_o ranging from 0 to 1. This $m_{I_{co}}$ curve is just the plot of 453Eq. (7) with r = 32 and p_o ranging from 0 to 1. 454

Given that $m_{I_{ca}}$ varies with p_o , which range of p_o values is 455most realistic physiologically? Studies of central synapses 456 found a very high open probability during an action potential, 457typically between 50% and 70% (Borst and Sakmann, 1998; Li 458et al., 2007; Sabatini and Regehr, 1997) (but see Jarsky et al., 2010). 459Thus, a value of $p_0 = 0.5$ would be reasonable during an impulse. 460 When $m_{I_{Ca}}$ is determined through the Ca²⁺ channel block 461 procedure, p_o may be decreased over 1–2 orders of magnitude, 462from about 0.5 to about 0.01. Over this range of p_0 values, the 463 slope of the regression line to the points on the dashed 464 theoretical curve in Fig. 2A equals $m_{I_{ca}} \approx 1.51$, which is close to 465 466 $m_{\rm L_{c}} \approx 1.55$ corresponding to the midpoint of this range (the slope 467 of the blue line segment). (The slopes would differ more if the 468 theoretical curve were not nearly linear.) In contrast, in the tail 469 current protocol, po would be varied from near 1 (functional 470 channels fully activated) to about 0.01. The corresponding fullrange $m_{I_{c_{\alpha}}}$ slope value would be well matched by the slope of the 471 solid black regression line in Fig. 2A. Fig. 2B summarizes how 472both $m_{I_{Ca}}$ and m_{CH} vary with the channel opening probability. 473 The two cooperativity measures are both close to one at low p_{o} 474 values, since in this case there is usually a single channel 475opening per each release event. However, m_{CH} and $m_{I_{Ca}}$ move 476 apart for higher p_0 . Note that they approximate the number of 477 available channels per release site only in the limit of p_o 478approaching 1. This implies that the log-log slope of release 479versus I_{Ca} should preferably be measured over the range of p_o 480 values corresponding to low channel block fraction to determine 481 the number of channels in close proximity to release sites. 482

In Fig. 3 the theoretical results given above are generalized
to M=5 equidistant channels. In general, we find that

$$m_{I_{Ca}} \le m_{CH} \le M \tag{8}$$

485 regardless of the value of r when the channels are equidistant from the release site. Note that for M = 5 equidistant channels, 487 the separation between $m_{I_{Ca}}$ and m_{CH} is even greater than with 488 two channels, and $m_{I_{ca}}$ is far less than the number of 489 equidistant channels. In summary, then, these calculations 490 show that the Ca²⁺ current cooperativity provides only a lower 491 bound on the channel cooperativity and the number of 492 channels proximal to the release site. Thus, whereas low 493 494 channel cooperativity implies low current cooperativity, low 495 current cooperativity does not imply low channel cooperativity. In general, the estimate of m_{CH} provided by $m_{I_{Ca}}$ de-496 teriorates as M increases. However, even if M is large, $m_{L_{ca}}$ is 497 close to m_{CH} if single-channel opening probability p_0 is low and 498 saturation of release is also low (i.e. release ratio r is high). The 499 fact that $m_{I_{Ca}}$ is a strong underestimate of m_{CH} means that a 500

modest value of $m_{I_{Ca}}$ of about 2–3 may imply a fairly large m_{CH} , 501 at least in the equidistant channel case (see Fig. 4; Coggins and 502 Zenisek, 2009). 503

6. Effects of Ca²⁺ channel distance and buffers 505 on channel and current cooperativity 506

Endogenous Ca²⁺ buffers, which may be stationary or mobile 507 (Neher, 1998b), trap Ca²⁺ ions that enter through open 508 channels and change their diffusion characteristics. It is 509 therefore expected that buffers may significantly affect the 510 Ca²⁺ channel cooperativity, as confirmed by simulation results 511 illustrated in Fig. 4, which assume 5 equidistant channels per 512 release site. (Ca²⁺ diffusion simulations in the presence of a 513 mobile buffer are performed using standard equations de- 514 scribed in Matveev et al., 2009). Fig. 4A shows that $m_{L_{c}}$ 515 increases as the channels are moved away from the release 516 site, since now more channels must open to gate the release. 517 There is a saturation of m_{I_c} , and even a slight decrease, far 518 below the upper bound of M=5. At the large distances where 519 this decrease occurs the Ca²⁺ reaching the vesicle location is 520 sufficiently small to become comparable to background Ca²⁺, 521 which will thus play a larger role in gating the release, thereby 522 reducing the current cooperativity. 523

Fig. 4B shows how $m_{I_{ca}}$ varies with the mobile buffer 524 concentration, with a fixed channel distance of 30 nm. For 525 lower buffer concentrations the current cooperativity in- 526 creases with increases in the buffer concentration. This is 527 because the buffer's main effect is to limit the extent of a 528 single channel Ca²⁺ nanodomain, reducing the probability of 529 release when a small number of channels are open, and thus 530 requiring more open channels to supply Ca2+ in the local 531 domain at the release site. When the buffer concentration is 532 very large, however, so much of the Ca²⁺ entering through 533 open channels is buffered that the background release rate 534 becomes comparable to the evoked release rate. The primary 535 effect of increasing the buffer concentration under these 536 conditions is to further reduce the contribution of evoked 537 release to the total release, and thus there is a small reduction 538 in $m_{I_{cc}}$. Hence, $m_{I_{cc}}$ first increases, and then decreases slightly 539 as the buffer concentration is increased. 540

The observation that both channel distance and buffer 541 concentration affect the current cooperativity suggests that 542 the values of $m_{I_{ca}}$ that can be achieved by changing channel 543 distance can also be achieved by changing the buffer 544 concentration. In fact, Fig. 4D shows that there is indeed an 545 exact correspondence between distance and buffering, so that 546 the same change in release rate and cooperativity is achieved 547 by increasing either the buffering or the distance. The five 548 colored points in Fig. 4D correspond to the five colored points 549 in Figs. 4A and B. For example, the black point has $m_{L_{ca}}$ =2.34. 550 To increase $m_{I_{ca}}$ to 2.87 (blue point) one could either increase 551 the buffer concentration to 1000 μ M while keeping channel 552 distance at 30 nm (x-axis of graph in Fig. 4D), or increase the 553 channel distance to about 50 nm while keeping the buffer ${\rm 554}$ concentration constant at 200 µM (y-axis). Each point on the 555 curve in Fig. 4D, then, tells what channel distance (with buffer 556 fixed at 200 μ M) provides the same release probability and 557 current cooperativity as would be achieved with the buffer 558

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Fig. 3 – (A) Log–log plot of the probability of release versus the open channel probability with M=5 equidistant channels and no saturation of release (release is proportional to 5th power of the number of open channels). (B) Ca²⁺ current and channel cooperativities as a function of p_0 . Equations for all quantities are given in Matveev et al.(2009). Colors of points correspond to colors of line segments in (A). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

concentration on the x-axis (with channel distance fixed at 30 nm). Panel E shows the effect of distance on $m_{I_{Ca}}$. At short distances, saturation is significant and $m_{I_{Ca}}$ severely underestimates *M*, whereas at longer distances the approximation improves somewhat. Similarly, $m_{I_{Ca}}$ gives a better approximation when buffer concentration is large.

566 7. Case of non-equidistant channels

While the simplifying assumption of equidistant channels 567helps in developing intuition about the connection between 568 current and channel cooperativities, in many cases of interest 569cooperativity depends on the relative contributions of both 570proximal and distal Ca²⁺ channels. There are a number of 571experimental studies focused on the effect of channel distance 572on Ca2+ current cooperativity of exocytosis (Fedchyshyn and 573Wang, 2005; Meinrenken et al., 2002; Shahrezaei et al., 2006). 574Modeling is useful in analyzing and interpreting the results of 575such experimental studies, and may make it possible to infer 576the channel cooperativity values from the measured current 577 cooperativity. This in turn requires a generalization of the 578channel cooperativity, since this is not a straightforward 579extension of the definition for equidistant channels. With 580equidistant channels, each of the channels contributes 581equally when open, while now some open channels contribute 582more Ca²⁺ to the local domain of a release site than others. If 583there are two channels, one distal and one proximal, and both 584585channels open, then both channels contribute to the local domain, but since the proximal channel is closer it will 586 contribute more ions. To determine the contribution made 587by each, it is therefore reasonable to determine the Ca^{2+} 588 concentration at the release site when either channel opens 589 separately (Ca10 and Ca01, where the first subscript corre-590sponds to the proximal channel and the second to the distal 591 channel and 1 means open). The cooperativity when both 592 channels open can then be defined as $m_{CH} = \frac{Ca_{10} + Ca_{01}}{Ca_{20}}$. This 593makes intuitive sense, since if the channels are at the same 594

distance then $Ca_{01} = Ca_{10}$ and $m_{CH} \approx 2$. If the distal channel is 595 much further from a release site than the proximal channel, 596 then $Ca_{01} \ll Ca_{10}$ and so $m_{CH} \approx 1$. If the proximal channel 597 contributes 4 times as much Ca^{2+} to the local domain as the 598 distal channel, then $m_{CH} = \frac{4+1}{4} = 1.25$, which is closer to 1 599 than to 2, as one would expect. This definition can be 600 generalized naturally to the case of an arbitrary number of 601 non-equidistant channels:

$$m_{CH} = \frac{\sum Ca_i}{\max Ca_i}.$$
(9)

where Ca_i is the Ca²⁺ concentration at the release site when 603 the ith channel is open and others are closed. 605

In Fig. 5 we consider a scenario in which there are two 606 proximal Ca²⁺ channels per release site and six distal 607 channels. The distal channels are situated at a distance of 608 90 nm from a release site, while the location of the proximal 609 channels is varied. When the proximal channels are quite 610 close, 30 nm, they dominate the release. This is reflected in a 611 low current cooperativity of about 2 and a channel coopera- 612 tivity between 2 and 3. Both forms of cooperativity increase as 613 the proximal channels are moved further from the release site. 614 Note that for a small range of distances $m_{CH} < m_{L_c}$, which 615 would not be possible if channels were equidistant. However, 616 the m_{CH} curve has upward concavity while the $m_{I_{Ca}}$ curve has 617 downward concavity, so for larger distances the current 618 cooperativity again provides only a lower estimate of the 619 channel cooperativity. When all eight channels are equidis- 620 tant we know that $m_{I_{Ca}}$ must be less than m_{CH} , and at 90 nm the 621 current cooperativity is indeed far less than the eight channels 622 available for release. 623

8. Selective channel block

Varying the number of open channels using pharmacological 626 channel block rather than a tail current protocol enables one 627 to examine the preferential coupling to exocytosis of specific 628

623



Fig. 4 – Dependence of $m_{l_{ca}}$ on the channel-vesicle distance (A) and buffer concentration, B_{total} (B), for a ring of 5 equidistant channels (C), with a single-channel current of 0.05 pA and 1 ms duration, with channel open probability of p_o =0.6. In (A), buffer concentration is B_{total} =200 μ M; in (B), channel distance is 30 nm. (D) Correspondence between increasing channel distance and increasing buffering. On x-axis, B_{total} is varied while keeping channel distance fixed at 30 nm. On y-axis, the channel distance is varied while keeping B_{total} fixed at 200 μ M. (E) Current cooperativity as a function of channel number M, for different channel-vesicle distances, with 200 μ M of mobile buffer (D=50 μ m²/ms) of 1 μ M affinity. Ca²⁺ dynamics is simulated in a 1 μ m³ box; a square Ca²⁺ current pulse arrives through each open channel. Open channel probability is varied around the value of p_o =0.6. Release is modeled using the Ca²⁺ binding scheme in Felmy et al.(2003). Each data point represents a weighted average over all possible open channel configurations. Calcium Calculator modeling software is used for all simulations (Matveev, 2008). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

subtypes of Ca²⁺ channels (Mintz et al., 1995). In this case, 629 larger current cooperativity values suggest closer coupling of 630 631 the pharmacologically manipulated channel type to exocytosis. Since part of the Ca²⁺ current arrives through channels not 632 affected by a particular blocker, in the case of selective block 633 the relationship between synaptic release and Ca²⁺ current 634 can be much steeper than in the case of non-specific channel 635 block. In fact, the bounds on $m_{I_{ca}}$ given by Eq. (8) would not 636 hold, and in particular, current cooperativity can in this case 637 easily exceed the number of available channels (Bertram et al., 638 1999; Matveev et al., 2009; Wu et al., 1999). For example, if a 639 640 channel blocker preferentially targets channels that are tightly coupled with exocytosis, while the majority of total presyn-641 aptic Ca²⁺ influx arrives through channels remote to the 642 vesicles, then the blocker would strongly reduce release with 643 only a minor decrease of presynaptic Ca²⁺ current. This would 644 lead to very large values of current cooperativity. Conversely, 645

if the blocker affects the channels that are remote to the 646 release site and loosely coupled to release, the decrease in I_{Ca} 647 would greatly exceed the concomitant decrease in release 648 rate, resulting in small values of current cooperativity. 649

For example, the study of Wu et al.(1999) found differences 650 in current cooperativity at rat calyx synapses with selective 651 block of three distinct subtypes of Ca²⁺ channels: P/Q (Cav2.1, 652 ω -agatoxin IVA-sensitive), N- (Cav2.2, ω -conotoxin GVIA- 653 sensitive) and R-type (Cav2.3) channels. The Ca²⁺ current 654 cooperativity of P/Q channels was higher than the other two 655 subtypes, indicating that a fraction of N- and R-channels were 656 located further away from release sites. Similar studies 657 revealed lower current cooperativity for N-type channels 658 than for P/Q type channels in rat and mouse hippocampal 659 synapses (Qian and Delaney, 1997; Qian and Noebels, 2001), 660 and cerebellar parallel fiber synapses (Mintz et al., 1995), 661 although no difference in $m_{I_{Ca}}$ between channel subtypes was 662

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Fig. 5 – Cooperativity and reliability with 2 proximal and 6 distal channels per release site. (A) The distal channels are located at a distance of 90 nm from a release site, while the proximal channel distance is varied. The mobile buffer concentration is 200 μ M and the channel opening probability is $p_0 = 0.5$. (B) The Ca²⁺ current and channel cooperativities increase as the proximal channels are moved away from the release site.

observed in experiments on guinea pig hippocampal cells (Wu 663 and Saggau, 1994) and rat hippocampal autapses (Reid et al., 664 1998). Further, a study of Reid et al.(1997) used selective 665 channel block to show that the distribution of specific channel 666 types is not uniform across distinct synaptic terminals, even 667 668 those efferent from the same cell (see also Reuter, 1995). Differences in the exocytotic coupling of distinct channel 669 subtypes using pharmacological channel block were also 670 found in chromaffin cells by Artalejo et al.(1994) 671

679 9. Functional implications

What are the functional implications of having some channels 674 close to release sites and some further away? It has been argued 675 that low values of $m_{I_{co}}$ observed for instance in several sensory 676 ribbon synapses contribute to linearity, fidelity and the dynamic 677 range of the sensory response (Brandt et al., 2005; Goutman and 678 Glowatzki, 2007; Jarsky et al., 2010; Johnson et al., 2008; Keen and 679 Hudspeth, 2006; Thoreson et al., 2004). In addition, studies that 680 infer tight vesicle-channel coupling suggest that low current 681 cooperativity is associated with faster and more Ca²⁺ efficient 682 synaptic response (Fedchyshyn and Wang, 2005; Kochubey 683 et al., 2009). In contrast, higher $m_{I_{ca}}$ values could be required for 684 response specificity, leading to a more all-or-none synaptic 685 transmission profile with lower noise due to fewer false 686 positives (Coggins and Zenisek, 2009). These hypotheses are 687 summarized in Fig. 7. As illustrated in this figure, small channel 688 cooperativity can be achieved via two distinct morphological 689 mechanisms: a coupling of a single channel to each vesicle, or 690 691 the action of several channels with very low release probability, a scenario inferred for instance in mouse retinal ribbon bipolar 692 cell synapses (Jarsky et al., 2010). In turn, higher current 693 cooperativity could either signify a lose association between 694 the spatial vesicle and channel distributions, or a coupling of 695 each vesicle to an array of several proximal channels, each 696 allowing only a non-saturating Ca²⁺ current. 697

A meta-analysis of several current cooperativity measure- 698 ments served as the basis for a hypothesis that Ca²⁺ channel 699 domain overlap is adapted to the physiological extracellular 700 Ca²⁺ concentration, so that at lower external Ca²⁺ concentra- 701 tion the synaptic morphology compensates through higher 702 channel clustering (Gentile and Stanley, 2005). 703

10. Dynamic and spatial modulation of current 704 cooperativity 706

The Ca²⁺ current cooperativity need not be fixed. In fact, there is 707 evidence for developmental changes in $m_{I_{Ca}}$ in synapses. For 708 example, in the rat calyx of Held $m_{I_{Ca}}$ in immature (pre-hearing) 709 calyces was determined to be 4.6 using tail currents (Kochubey 710



Fig. 6 – Hypothesis for the developmental change in vesicle-channel coupling at the calyx of Held. (A) Channels are poorly coupled to release in an immature synapse.
(B) With development, a subset of Ca²⁺ channels moves closer to the vesicle location.

9

et al., 2009), and 4.8 when presynaptic I_{Ca} was varied by 711 systematically increasing the duration of an action potential 712 waveform that provided the input (Fedchyshyn and Wang, 2005). 713 In mature (post-hearing) calyces the Ca²⁺ current cooperativity 714 was reduced to 3.7 when measured with tail currents and to 2.6 715 when measured with action potential waveforms. The reduction 716 in $m_{L_{c}}$ during development is consistent with a scenario in which 717 there is a tightening of the release site/Ca²⁺ channel complex, so 718 that in mature calyces there are a few proximal channels assisted 719 720 by more distal channels. The proximal channels could have moved in from more distant locations (Fig. 6), which would reduce 721 $m_{\rm L_{c}}$ as in Fig. 5. To test this hypothesis, Fedchyshyn and Wang 722 sealed a patch electrode containing the slow Ca²⁺ chelator EGTA 723 onto mature and immature calyces. Because of its slow binding 724 rate, EGTA can buffer Ca²⁺ from distal channels before it reaches 725 release sites more effectively than Ca²⁺ from proximal channels. 726 In immature calyces exposed to EGTA the quantal output was 727 reduced by ~70%, while in mature calyces the EGTA-mediated 728 reduction was much smaller, only ~20%. In contrast, the fast 729 buffer BAPTA was equally effective in reducing release in both 730 mature and immature calyces. These findings are consistent with 731 the hypothesis that channels are closer to vesicles in the mature 732 case. Distance may not be the only effect; Fedchyshyn and Wang 733 734 also found that action potentials were wider in the immature calyces, which would further enhance the effectiveness of EGTA 735 736 relative to mature calyces. In a subsequent study, a synaptic 737 protein, Septin 5, was identified that appears to act as a physical 738 barrier to vesicle docking to release sites (Yang et al., 2010). The data suggest that in the immature calyx, septin 5 proteins block 739 vesicle docking to release sites in the active zone, where most of 740 the Ca²⁺ channels are located. This results in a large distance 741 between channels and vesicles, leading to a relatively large 742 current cooperativity reported in Fedchyshyn and Wang(2005). 743

During maturation, the septin 5 proteins move to the active zone 744 periphery, removing the barrier to vesicle docking to active zone 745 release sites. Consequently, docked vesicles are closer to the Ca²⁺ 746 channels, resulting in a lower current cooperativity. 747

Studies using selective blockers of different channel types 748 have shown that N-, P/Q-, and R-type Ca²⁺ channels contribute 749 to transmitter release in calyces from 8 to 10-day-old rats (Wu 750 et al., 1998), but that P/Q-type channels contribute more to 751 release than the other two types (Wu et al., 1999). This is 752 consistent with imaging using subtype-specific antibodies, 753 which showed that a sizable fraction of the N- and R-type 754 channels are more distant from vesicles than are P/Q-type 755 channels (Wu et al., 1999). A complementary study, again 756 using channel type-specific blockers, found that the contribu-757 tion to release of P/Q-type channels greatly increased from age 758 4 d to 10 d, so that by age 10 d release was gated entirely by 759 P/Q-type channels (Iwasaki and Takahashi, 1998). Thus, it 760 seems likely that in the experiments of Fedchyshyn and Wang 761 (2005), application of EGTA blocked the contribution of the 762 more distant N- and R-type channels, while the contribution 763 from the closer P/Q-type channels was preserved. 764

Although the studies above point to a developmental 765 tightening of the release site complex, they do not rule out a 766 parallel developmental change in the biochemical Ca^{2+} 767 cooperativity, *n*. This possibility was investigated by Kochubey 768 et al.(2009). They raised the Ca^{2+} concentration throughout the 769 calyx using laser-induced flash photolysis to uncage Ca^{2+} from 770 exogenous chelator. Different flash intensities were used to 771 raise Ca_i to different levels (measured with fura-2), while 772 simultaneously measuring EPSCs in the postsynaptic cells. 773 This approach was used to determine *n* in immature calyces 774 (from 8 d to 9 d rats) and more mature calyces (from 12 d to 775 15 d rats). It was found that *n* was approximately 3.6 in both 776



I_{Ca} (# open channels per release event)

Fig. 7 – Possible functional consequences of different Ca²⁺ current cooperativity values. A linear relationship between Ca²⁺ influx and release leads to greater sensitivity and dynamic range of the response, as well as Ca²⁺-efficiency and speed of response due to tighter vesicle-channel coupling. In contrast, high current cooperativity may lead to a more specific, all-or-none response with a lower false-positive rate. High current cooperativity requires an overlap of multiple channel domains (upper-right inset), whereas low current cooperativity can be achieved through either the tight coupling of a single channel to each vesicle (middle-right inset) or low probability of channel opening, whereby only one proximal channel is likely to open with each pulse (lower-right inset; black filled ovals indicate closed channels).

cases, and thus there was no developmental change in the biochemical cooperativity over the period of time when the current cooperativity declines.

Another example of a developmental change in the Ca²⁺ 780 current cooperativity is the ribbon synapse of the mammalian 781 inner hair cell. These ribbon synapses are specialized to allow 782 sustained release of a large number of vesicles. Cells on the 783 sensory neuroepithelium are tuned to different characteristic 784 785 frequencies; those in the apical region respond primarily to lowfrequency sound, while those in the basal region respond 786 primarily to high-frequency sound (Fettiplace and Fuchs, 787 1999). Johnson and colleagues studied developmental changes 788 in *m*_L of ribbon synapses in mouse inner hair cells, and in apical 789 and basal cells of the gerbil (Johnson et al., 2008). Using a voltage 790 clamp protocol to vary the number of open Ca²⁺ channels, in the 791 latter study they found $m_{I_{Ca}} \approx 3.8$ in immature synapses from 792 both apical and basal cells. In cells from mature gerbils, $m_{L_{c}}$ was 793 significantly lower, again showing a developmental reduction in 794 the current cooperativity as in the calyx of Held. However, this 795 reduction was not uniform across the neuroepithelium; synap-796 ses from apical (low-frequency) cells had $m_{L_{c}} \approx 2.2$, while those 797 from basal (high-frequency) cells had $m_{I_{ca}} \approx 1$. 798

To investigate the origin of the different current cooperativ-799 800 ities in the two cells, Johnson et al. (2008) examined the biochemical cooperativity of synapses from each cell type. 801 802 They determined that n=2.8 for mature apical cells, and n=1.2803 for mature basal cell synapses. Hence, the origin of the different 804 current cooperativities in the low-frequency and high-frequency cells of mature animals appears to be at least partially due to 805 differences in the Ca²⁺ trigger for release in the two cells, and 806 may indicate involvement of a distinct Ca²⁺ sensitive proteins, 807 such as otoferlin or non-neuronal synaptotagmins (Duncan et 808 al., 2010; Johnson et al., 2010; Keen and Hudspeth, 2006; 809 Mirghomizadeh et al., 2002). Taken together with studies done 810 in the calyx of Held, it appears that developmental changes in 811 both the structure of the channel/release site complex and the 812 vesicle release mechanism itself can occur so as to reduce $m_{\rm L}$ 813 and increase reliability as the animal matures. 814

816 **11.** Conclusions

In the absence of a direct way to observe the functional coupling 817 of individual Ca²⁺ channels to neurotransmitter release, mea-818 surements of current cooperativity have proved very useful in 819 elucidating the degree of Ca²⁺ channel nanodomain overlap in 820 synaptic vesicle release in a variety of synaptic terminals, from 821 invertebrate neuromuscular junctions to central mammalian 822 synapses. However, care must be taken in interpreting $m_{\rm L}$ 823 measurements, since a given value of current cooperativity does 824 not allow one to directly infer the functional Ca²⁺channel 825 cooperativity, i.e. the number of Ca²⁺ channels participating 826 on average in the release of a single vesicle. In particular, a small 827 828 value of m_{I_c} is not necessarily an indication that a small number 829 of Ca²⁺ channels participates in release. As reviewed above, current cooperativity can provide a good approximation of the 830 underlying channel cooperativity only when the number of 831 channels is very small (2-3), and under further restrictive 832 conditions such as low release saturation or low channel 833 opening probability. Therefore, current cooperativity is not a 834

very sensitive characteristic of synaptic morphology. Still, 835 combined use of experimental and modeling techniques in 836 interpreting $m_{I_{Ca}}$ measurements provides greater insight into 837 functional synaptic morphology (Bucurenciu et al., 2010; 838 Coggins and Zenisek, 2009; Jarsky et al., 2010; Meinrenken 839 et al., 2002; Quastel et al., 1992; Shahrezaei et al., 2006; 840 Yoshikami et al., 1989; Zucker and Fogelson, 1986). 841

Current cooperativity measurements served a particularly 842 important role in the long-standing argument whether 843 synaptic neurotransmitter release is controlled by the opening 844 of a few channels, or whether an overlap of Ca²⁺ nanodomains 845 of many channels is required to trigger exocytosis (Gentile and 846 Stanley, 2005; Schneggenburger and Neher, 2005). However, 847 recent data strongly suggest that the Ca²⁺ influx varies 848 between different types of synaptic terminals, and can indeed 849 vary with development at each particular synapse (Fedchy- 850 shyn and Wang, 2005; Johnson et al., 2005; Johnson et al., 2008; 851 Kochubey et al., 2009; Yang et al., 2010). Further, possible 852 values of current cooperativity are directly dependent on the 853 intrinsic (biochemical) cooperativity, which can also change 854 developmentally, either through modulation of the synaptic 855 machinery or through the presence of heterogeneous pools of 856 vesicles, which may in turn have distinct biochemical 857 cooperativity values under developmental regulation. 858

It is interesting to speculate on the functional significance 859 of different $m_{I_{Ca}}$ values. It has been suggested that the small 860 values of $m_{I_{Ca}}$ observed in many sensory ribbon synapses are 861 connected to the requirement of linear coding of the sensory 862 signal over a wide dynamic range, while the higher channel 863 cooperativity found in other central synapses may indicate a 864 requirement for an all-or-none response with lower noise due 865 to false positive, spontaneous activity.

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