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Bourgeois-Jaarsma, Q.

2020

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citation for published version (APA)

Bourgeois-Jaarsma, Q. (2020). Regulation of neurotransmitter release by C-domain Ca²-sensors. [PhD-Thesis - Research and graduation internal, Vrije Universiteit Amsterdam].

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Chapter 6

General discussion

1. Scope

As introduced in **chapter 1**, release of SVs can be evoked by action potentials or they can be released in their absence. Cytosolic Ca^{2+} plays a role in both forms of release. The Double C_2 proteins (Doc2a/b) contribute to neurotransmitter secretion by acting as a Ca^{2+} -dependent regulator of asynchronous^{1,2} and spontaneous vesicle fusion (mPSCs or minis)^{3,4}. AP-independent Ca^{2+} -elevations occur in the presynaptic cytoplasm. We refer to them as spontaneous Ca^{2+} elevations (SCEs) because no triggering mechanism has been identified. The intracellular Ca^{2+} elevation may originate from extracellular⁵, intracellular Ca^{2+} -stores^{6,7}. In the latter case, they might also be triggered by GPCR activation (see review⁸). These observations prompted the hypothesis that SCEs may activate Ca^{2+} sensor proteins such as Doc2 and trigger spontaneous release. Yet, experimental proof for this hypothesis is lacking and the molecular mechanism driving spontaneous vesicle fusion remains unknown.

This thesis deals with the regulation of synaptic transmission on the two distinct levels: first by fast, spontaneous presynaptic Ca^{2+} -signaling and second by various presynaptic Ca^{2+} -sensing C_2 domain proteins. The overall aim of this work described was to gain insight in the presynaptic mechanisms underlying regulation of spontaneous release, synaptic strength and plasticity.

1.1. Main findings

In **chapter 2**, we investigated the frequency and kinetic characteristics of SCEs in resting neurons using genetically encoded Ca^{2+} -indicators. Recent advances in live Ca^{2+} imaging offer unprecedented spatial and temporal resolution, which facilitates the detection of SCEs but also engenders an unmet need for data processing. The local and short-lived nature of SCEs, together with their low amplitude and frequency make the detection of these events challenging and their analysis by existing open source programs^{9,10} unsuitable for this purpose. Therefore, a MATLAB based analysis method was developed to automatically identify rapid Ca^{2+} rises in without predetermined regions of interest. The algorithm detected low-amplitude signals with various waveforms in noisy time lapse imaging data. Using a fast variant of GCaMP6 (GCaMP6f), the most common class of Ca^{2+} elevations, named SCTs (for spontaneous Ca^{2+} transients), exhibited fast kinetics with a mean Full Width at Half Maximum (FWHM) of 0.278 and an SD of 0.146 s. The signal amplitude was faint (mean $\Delta F/F_0 = 0.064$; SD = 0.011) and the SCTs occurred with variable frequencies of 0.174 Hz (mean; SD=0.322 Hz).

The reliability of the method was confirmed using both simulated and experimental data by testing its detection sensitivity using experimental data and simulated experiments with various SNRs. The availability of this method is highly valuable to speed up future investigations studying the role of SCEs in neuronal cells or circuits.

The method was employed in **chapter 3** to investigate if stochastic Ca^{2+} fluctuations could potentially cause spontaneous neurotransmitter release events. Spontaneous release probability is regulated by $[\text{Ca}^{2+}]_e$ ^{3-5,11} and intracellular Ca^{2+} ^{12,13}, but it is unclear if these effects are primarily attributable to global $[\text{Ca}^{2+}]_i$, SCEs, or both. We thus explored Ca^{2+} transients in glutamatergic hippocampal neurons by simultaneous electrophysiology and high-speed Ca^{2+} imaging using different Ca^{2+} probes. Among the Ca^{2+} reporters tested, SyGCaMP6f appeared the most efficient for the detection of presynaptic SCEs. Besides the fast SCTs, we also observed a variety of other SCE waveforms. An increase in extracellular Ca^{2+} caused an increase in global $[\text{Ca}^{2+}]_i$ as expected, but also increased the frequency of dynamic Ca^{2+} elevations. This primarily affected complex Ca^{2+} events. mEPSCs detected by patch-clamp recording were not time-locked to SCTs and SCEs. This suggests that quantal glutamate release is either regulated by global $[\text{Ca}^{2+}]_i$ or that SCEs induce SV secretion with a slow (>1s) or variable delay.

In **chapter 4**, we addressed the mechanism of Doc2b function which was postulated to act as a Ca^{2+} sensor³ or as a structural element supporting Ca^{2+} -dependent secretion⁴. Both studies relied on the effects of Ca^{2+} binding site mutants, named Doc2b^{DN} and Doc2b^{6A}. We therefore explored spontaneous and evoked release modification in a side-by-side comparison of Doc2b^{WT}, Doc2b^{DN} and Doc2b^{6A}. We found that the neutralization of critical aspartates had similar gain of function effects on the spontaneous release frequency at rest, which paralleled *in vitro* plasma membrane binding characteristics, reinforcing the idea of its action as a Ca^{2+} sensor. In addition, our results suggest that both mutants enhance the synaptic release probability (P_s) upon single AP stimulations, thereby affecting short-term plasticity and EPSC rundown during prolonged stimulation. This chapter also supports the idea that Doc2b contributes to synaptic recovery and potentiation in a manner that requires Ca^{2+} binding.

In **chapter 5**, we further investigated the synaptic mechanism responsible for the residual spontaneous secretion in absence of the main sensors for quantal release. As candidates we selected three C_2 domain proteins based on their high similarity with Doc2a and b: Doc2c, rabphilin-3a and Synaptotagmin-7 (Syt-7). Despite their high resemblance with Doc2a & b, gene inactivation of Doc2c and Syt-7 did not influence spontaneous release in excitatory hippocampal neurons. Rabphilin seemed to inhibit spontaneous release frequency in networks but this activity did not appear in autaptic culture. Our findings narrow down the repertoire of potential synaptic Ca^{2+} sensors for spontaneous release and redirect our interest to other C_2 domain proteins.

2. Ca²⁺ signaling of spontaneous release

2.1. Temporal characterization of Ca²⁺ transients

In presynaptic terminals, the intracellular Ca²⁺ concentration profile during AP-induced VGCC activation is characterized by highly local peaks immediately adjacent to the Ca²⁺ channels. The local nature of these Ca²⁺ signals matches a remarkably well organized subcellular architecture¹⁴ (for review^{15,16}). The nanoscale topography of Ca²⁺-channels relative to synaptic vesicles ensures rapid neurotransmission upon AP arrival. Thus, the delay time between Ca²⁺-entry and synaptic vesicle fusion is extremely short. A number of investigations reported a synaptic delay of ~1 ms (presynaptic AP to postsynaptic AP) and Ca²⁺ triggers vesicle fusion and neurotransmitter release in a few hundred of milliseconds¹⁷⁻²¹. Pioneering works studied the process at the NMJ²⁰, rat cerebellar fiber synapse²² and the Calyx of Held synapse^{23,24}. Simultaneous pre- and postsynaptic electrophysiological recordings in the squid giant synapse and mathematic isolation of the Ca²⁺-current estimated a latency between the inward Ca²⁺ current and transmitter release of ~200 μs¹⁷. In chapter 3 (Chapter 3, Figure 7A-B), we measured the latency between the presynaptic Ca²⁺-influx monitored by presynaptically targeted GECI sy-GCaMP6f and the AP evoked EPSC peak signal. The delay of the reported Ca²⁺ entry relative to the postsynaptic activation in our hands appeared to be +50 to +150 ms, meaning that the detection appeared after the EPSC, which is clearly a technical issue. Yet, electrophysiology and imaging acquisitions were precisely synchronized (Chapter 3, Figure S4) using internal controls which confirm accurate timestamping in the EMCD camera (an accuracy of ~10 ns is specified by the manufacturer). The delay in fluorescent peak detection is longer than described by purely electrophysiological approaches^{17,25,26}. The mismatch could be explained by the association kinetics of Ca²⁺ ions to the fluorescent indicators. Other studies with fluorescent Ca²⁺ indicators also report longer latencies than electrophysiological measurements. Previously reported measurements using OGB-1 exhibited a latency around 320 ms¹⁰. The Ca²⁺ peak FWHM triggered by a single AP with OGB-1 in our study was 176 ms (Chapter 3, Table 1). For GCaMP6f, the fastest GECI available, previous studies showed a FWHM of 360 ms and a time to peak latency from 50 to 75 ms²⁷ for a single AP. The association kinetics importantly depend on the [Ca²⁺]. For GCaMP1, the Ca²⁺ association time constant ranges broadly from 230 ms at 0.2 μM Ca²⁺ to 2.5 ms at 1 μM Ca²⁺²⁸, while the dissociation constant was not affected by [Ca²⁺].

Another indication that fluorescent indicators markedly delay the Ca²⁺ peak signal can be deduced from a direct comparison of data obtained with GCaMP6f versus OGB-1. For low [Ca²⁺] amplitude SCTs, OGB-1 signals had a FWHM of ~49 ms (Table 1) compared to ~260 ms for GCaMP6f (Chapter 3, Stat. table 6 & Figure 6). The ratio of these FWHM values indicates that OGB-1 is a ~5.75 fold faster reporter than GCaMP6f. Thus, in the full chain of events from Ca²⁺ channel opening to optical measurement of Ca²⁺ induced fluorescence signals and dynamics, the delay from binding of Ca²⁺ onto the indicator to the emergence of fluorescence forms a profound rate-limiting step. In contrast, imaging framerates were not rate-limiting in our experiments, while channel dynamics and Ca²⁺ diffusion have minor effects. Altogether, before performing Ca²⁺-imaging, it must be taken into consideration that

Ca^{2+} signals are optically reported with a delay which is an inherent property of the Ca^{2+} indicator used. The delay also varies with the nature of the $[\text{Ca}^{2+}]$ signal. It would be helpful if a next generation GECIs could be improved to achieve shorter latencies close to that of chemical Ca^{2+} -probes, allowing to detect Ca^{2+} events at a higher temporal resolution while retaining possibilities for subcellular targeting.

The image acquisition speed is also an important parameter. In our experiments the framerate of ~ 30 Hz was more than sufficient for accurate detection of Ca^{2+} -events, which had a mean peak width (FWHM) of ~ 8 frames (FWHM = 278 ms and acquisition every 33 ms).

2.2. Spatial characterization of Ca^{2+} transients in relation to spontaneous release

The ultrastructure and topology of presynaptic protein architectures have started to elucidate how scaffold proteins establish nanodomains that connect VGCCs with release-ready SVs in a radius of approximately 30 nm²⁹ (Chapter 1, see Figure 1). Scaffold proteins seem to operate as molecular linkers, limiting SV-VGCC distances to tens of nanometers, thereby regulating the probability and plasticity of SV release³⁰. Measurements of quantal secretion have shown that the vesicular release probability decreases threefold with a doubling of the distance between the Ca^{2+} channel and the synaptic vesicle from 25 to 50 nm³¹.

In chapter 3, Ca^{2+} -transients emerged all over the neuron which led to the idea that they might appear in an unpredictable spatial pattern. Deeper analysis of GCaMP6f fluorescence revealed that 70% of the SCTs co-localize with synapsin-eGFP. In line with this, Sy-GCaMP6f detected 2.5-fold more SCT events compared to global GCaMP6f (Chapter 3; Figure 5) which suggests a spatial enrichment of fast transients in the presynaptic element.

Various Ca^{2+} sources have been observed to affect the frequency of mPSCs: 1. Intracellular stores, from which Ca^{2+} can be liberated through IP_3R and/or RyR channels^{7,32}, 2. The spontaneous opening of VGCC at the resting potential, also referred to as VGCC flickering^{5,33}. Those two mechanisms can generate global $[\text{Ca}^{2+}]_i$ fluctuations depending on the equilibrium between Ca^{2+} liberation and Ca^{2+} buffering and clearance. Caffeine application^{32,34} and $[\text{Ca}^{2+}]_e$ increase³⁵ are well described and commonly increase spontaneous release. Application of an agonistic caffeine concentration significantly increased the SCT frequency from 0.23 Hz to 0.36 Hz (Chapter 2, Figure 6). These data support the implication of intracellular Ca^{2+} -stores in the generation of presynaptic Ca^{2+} -transients but do not rule out the possible involvement of extracellular sources. Prior works revealed the involvement of the Ca^{2+} -induced Ca^{2+} -release mechanism in spontaneous release⁶. This mechanism was shown to be critical for the amplification of faint Ca^{2+} -events in the postsynaptic element³⁶. Extracellular $[\text{Ca}^{2+}]_e$ rise from 0 mM to 10 mM promoted the frequency of both mEPSCs and SCEs (Chapter 3, Figure 4 & 6). The same treatment also caused an increase in global $[\text{Ca}^{2+}]_i$ as confirmed by Fura-2 imaging (Chapter 3, Figure 4). The concomitant increase in global $[\text{Ca}^{2+}]_i$ may be a direct consequence of the increased $[\text{Ca}^{2+}]_e$, implying a passive Ca^{2+} leakage following the concentration gradient, or an indirect result of the higher SCE frequency.

In both cases, the limited Ca^{2+} clearance capacity likely yields a progressive cytosolic build-up, coercing the cell to a higher $[\text{Ca}^{2+}]_i$ steady state (Chapter 3, Figure 4B). In Chapter 2 and 3, complex Ca^{2+} transients were described as all events lasting longer than 280 ms, usually covering larger areas than SCTs and forming very different waveforms. This may suggest the existence of multiple triggering mechanisms. In the GCaMP6f experiments, complex events represented 43% of all events (Chapter 2). They were frequent in the presynaptic compartment (0.082 Hz vs 0.13 Hz for GCaMP6f vs SyGCaMP6f respectively; Chapter 3, Figure 5) and their frequency was affected more by extracellular $[\text{Ca}^{2+}]_e$ than that of SCTs (Chapter 3, Figure 6). SCEs could either arise from the summation of SCTs by simultaneous channel unit opening or via the CICR amplification mechanism. Further investigations are necessary to disclose the pathway driving SCEs.

Altogether, our data support the preferential involvement of long lasting, complex Ca^{2+} fluctuations rather than fast SCTs in SV fusion in resting glutamatergic hippocampal neurons. The observation that caffeine increases the frequency of TTX-insensitive SCTs, together with the preferential involvement of SCEs in spontaneous release suggests that the CICR mechanism is also functional in the presynapse for minis.

2.3. Ca^{2+} dose-dependence of spontaneous release

Many investigations attempted to precisely measure the Ca^{2+} dependence of neurotransmitter release. A wide range in Ca^{2+} affinities of evoked and spontaneous release were reported, likely because of the variety in methodologies used and the different brain structures investigated. Early studies investigated the relationship between intraterminal $[\text{Ca}^{2+}]_i$ using a combination of Ca^{2+} -indicators during photolysis of caged Ca^{2+} , ionomycin perfusion and either intracellular electrophysiology or capacitance measurements. Ratiometric Fura-2 imaging and intracellular recording of miniature end plate potentials (MEPPs) at the frog neuromuscular junction reported a resting $[\text{Ca}^{2+}]_i$ of around ~ 100 nM together with marked increase in the frequency of MEPPs at 500 nM³⁷. Caged Ca^{2+} photolysis³⁸ and ionomycin application³⁹ methods yielded lower Ca^{2+} -sensitivities for synchronous and asynchronous release than for spontaneous release. In the Calyx of Held, a combination of ratiometric and non-ratiometric Ca^{2+} probes indicated a resting $[\text{Ca}^{2+}]_i$ of 40 nM and a peak amplitude of Ca^{2+} transients elicited by single APs around 400 nM to 500 nM⁴⁰. In apical dendrites of hippocampal CA1 neurons, resting $[\text{Ca}^{2+}]_i$ was evaluated by OGB-1 imaging to be 32 nM to 59 nM and monitoring evoked Ca^{2+} transients yielded estimates of peak $[\text{Ca}^{2+}]_i$ of 178 nM to 312 nM⁴¹. As the availability of the Ca^{2+} -indicators diversified, the use of Fluo-4 together with OGB-1 highlighted the importance of endogenous Ca^{2+} buffering capacity and subcellular volume which influence the $[\text{Ca}^{2+}]_i$ peak during single APs (respectively 528 ± 251 nM in the spine and 401 ± 139 nM in the dendrite)⁴². The $[\text{Ca}^{2+}]_{\text{rest}}$ in individual giant mossy fiber boutons was evaluated to 116 ± 20 nM with Fluo-4 and 103 ± 14 nM with OGB-1⁴³. A later study estimated a much higher peak Ca^{2+} concentration at the VGCC mouth of approximately 100 μM during evoked neurotransmitter release²⁹, likely due to the advancement of temporal and spatial resolution. It stands out that reported Ca^{2+} -event properties differ depending on the spatial (subcellular compartment or whole cell) and temporal focus of the study.

Altogether, the Ca^{2+} rise following a single action potential is approximately 5- to 10-fold higher than resting Ca^{2+} concentration. In our hands, OGB-1 reported a 4-fold increase in the $\Delta F/F_0$ peak signal during a single AP compared to the amplitude of spontaneous events (Chapter 3, Table 2).

The contrast between the low-threshold Ca^{2+} requirement for spontaneous release and the high Ca^{2+} threshold of fast evoked release is striking. The different phases of release, namely spontaneous, asynchronous and synchronous release arise along a wide range of $[\text{Ca}^{2+}]$ and show a Ca^{2+} -sensitivity from 50 nM to 50 μM in the Calyx of Held, however with different Ca^{2+} -cooperativity⁴⁴⁻⁴⁶. Taken together, the $[\text{Ca}^{2+}]_i$ window in which a Ca^{2+} -sensor could exert its function for spontaneous SVs fusion can be situated above the resting Ca^{2+} concentration of 40-50 nM and below the activation threshold of evoked release around 400-500 nM. Considering concordant reported values of absolute peak $[\text{Ca}^{2+}]$ during a single AP (in average ~ 387 nM), together with our comparison with the $\Delta F/F_0$ obtained from OGB-1 imaging (Chapter 3, Table 1&2), one can estimate an absolute peak $[\text{Ca}^{2+}]$ during SCTs of ~ 121 nM. This estimate is coherent with the high Ca^{2+} -affinities of membrane binding activity by identified Ca^{2+} sensors for spontaneous release Doc2a, b (EC_{50} of respectively 450 nM and 175 nM, Chapter 4)⁴⁷.

2.4. The Ca^{2+} trigger of Doc2

Doc2 is the main sensor involved in quantal neurotransmitter release, yet the relation of Doc2b driven spontaneous release with the Ca^{2+} -source is not resolved. Based on its high Ca^{2+} -sensitivity⁴⁷ and considering that Syt-1 activation requires higher $[\text{Ca}^{2+}]$ ($\text{EC}_{50} \sim 3.5 \mu\text{M}$)^{48,49} it is very likely that Doc2b is specifically activated by Ca^{2+} levels that do not activate Syt-1. As suggested by our investigation in chapter 3, both global cytosolic Ca^{2+} and SCEs potentially control quantal release but the portion of Doc2b-driven release events in these assays isn't known. Unlike Syt-1, Doc2 is a cytosolic protein with no physical association to Ca^{2+} microdomains, as far as known. This feature may render Doc2 activatable by Ca^{2+} events that occur at distinct locations relative to Syt-1. Being associated to vesicles, Syt-1 is preferentially activated by Ca^{2+} -influx through VGCCs. The localization of Doc2b is likely more dynamic: it has several interactors that can target it to release sites. The N-terminal MID enables its co-translocation with Munc13 to the plasma membrane⁵⁰⁻⁵². This activation pathway is stimulated by phospholipase C-dependent production of diacylglycerol, which can be mimicked experimentally by phorbol ester administration⁵¹. The Doc2b-Munc13 association mechanism promotes vesicle release and synaptic augmentation⁵³. Doc2b also associates with the SNARE-accessory protein Munc18^{54,55} and the SNARE proteins syntaxin-1 and SNAP25³, providing another pathway for Doc2b release site targeting. Therefore, global cytosolic Ca^{2+} increases might enhance Doc2b-driven quantal release via the PLC / DAG / IP_3 / Munc13 association pathway (Chapter 1, Figure 5), whereas Ca^{2+} transients caused by VGCC flickering could directly trigger Doc2b molecules already present near the exocytotic apparatus. The latter idea may not be relevant for mouse hippocampal GABAergic neurons, because mIPSC events sensitive to Cd^{2+} blockade of VGCCs are Doc2b-independent⁵⁶. Yet, this pathway might occur in other cell types. Further investigations are necessary to clarify the specific Ca^{2+} trigger for Doc2 driven spontaneous release.

2.5. Limitations to our study of SCE – mEPSC correlation

In chapter 3 of this thesis, we investigated the relation between spontaneous neurotransmitter release and presynaptic Ca^{2+} -events. The temporal correlation between SCEs and mEPSCs appeared poor at high $[\text{Ca}^{2+}]_e$ (Chapter 3, Figure 4 & 6). Refined analysis of the temporal correlation of mEPSCs relative to single or complex events at 4 mM $[\text{Ca}^{2+}]_e$ (Chapter 3, Figure 7 & 8) led to a similar conclusion. Yet, a parallel increase of cytosolic Ca^{2+} and minis was reported multiple times^{7,37}. Several technical and physiological limitations could cause a hypothetical coupling between SCEs and mEPSC to remain undetected. The first technical limitation relies on the Ca^{2+} indicator performance. The latency from AP-induced Ca^{2+} peak and the EPSC was mentioned earlier to be in order of 50 to 150 ms with the GCaMP6f. We took this delay into account in our temporal analysis, by inspecting minis in a time window of 250 ms and 1 s before and after Ca^{2+} transients. However, the slow Ca^{2+} -association kinetics of the GECl, particularly for low peak $[\text{Ca}^{2+}]_i$, could then leave fast SCTs undetected²⁸. Indeed, low Ca^{2+} binding affinity and slow kinetic can give enough time for endogenous Ca^{2+} buffering, extrusion and diffusion mechanisms to clear intracellular Ca^{2+} before it can bind the fluorescent reporter. The second restriction could arise from the imaging methodology. The imaging field of view was roughly ~70% smaller than the entire autaptic neuron, which is expected to lead to a proportional reduction of detected SCEs (but not mEPSCs).

Moreover, GCaMP6f associates with Ca^{2+} with a K_d around 375 nM²⁷ which means that small Ca^{2+} elevations below this threshold could remain undetected. Yet, quantal release is at least partially driven by high affinity sensors as Doc2b which show an EC_{50} around 175 nM. Therefore, one can envision that low-amplitude SCEs escape detection, but trigger spontaneous SV fusion.

The last restriction could relate to different mechanisms for spontaneous release in excitatory vs. inhibitory synapses^{33,56–58}. Spontaneous fusion events appear preferentially triggered by local Ca^{2+} micro-domains generated via stochastic openings of VGCCs only at GABAergic, but not glutamatergic terminals³³. Selective and unselective VGCC blockers affected mIPSC but not mEPSC frequency in neocortex neurons⁵⁸. A recent investigation reported the preferential involvement of Syt-1 in spontaneous GABA release⁵⁶, consistent with the strategic location of the fast sensor nearby VGCCs in support of quantal GABA release. In contrast, spontaneous release frequency was altered by $[\text{Ca}^{2+}]_e$ and both EGTA and Cd^{2+} at cholinergic and GABAergic synapses at *C. Elegans* NMJ⁵⁷. Both mEPSCs and mIPSCs were shown to rely on CaV_1 and CaV_2 channel opening. mIPSCs exhibited a higher proportion of $[\text{Ca}^{2+}]_e$ -independent events than mEPSCs as suggested by the effect of EGTA and Cd^{2+} ⁵⁷. Thus, spontaneous release seems to be, at least in part, differently regulated at excitatory and inhibitory synapses. In Chapter 3 where we observed no tight coupling between SCEs and mEPSCs, our investigations were performed on excitatory hippocampal neurons, while only a minority of GABAergic neurons was present in the culture. It is thus important to note that the lack of tight coupling cannot be extrapolated to inhibitory neurons.

In conclusion, an accurate time correlation between Ca^{2+} -transients and vesicle fusion at rest may possibly have remained undetected if the triggering signal would rely on an extremely rapid, low amplitude Ca^{2+} -peak, considering the technical limits of available tools. To acquire more certainty, this quest would be facilitated by the use of improved indicators with faster Ca^{2+} association kinetics that simultaneously report Ca^{2+} and fusion events.

2.6. Spatial correlation of synaptic vesicle fusion and SCTs

The all-optical alternative

As an alternative approach, an all optical technique can be employed to assess presynaptic Ca^{2+} -signalling together with synaptic release events. This method requires indicators with appropriate excitation and emission wavelengths to simultaneously detect synaptic fusion and Ca^{2+} events. Optical methods for synaptic vesicles trafficking at the active zone have been developed using a pH-sensitive GFP fluorophore termed pHluorin⁵⁹. pHluorin can be targeted to the synaptic vesicle lumen by fusion to synaptophysin (sypHy)⁶⁰ or other vesicle associated proteins. It is quenched by the low acidic pH in the vesicles but the fluorescence increases as the vesicles fuse with the PM, neutralizing the pH due to content mixing with the extracellular environment. Fusion events are identified as a sudden brightness increase in sypHy fluorescence which declines upon vesicle re-acidification following endocytosis.

Dual channel detection of Ca^{2+} and SV secretion has been achieved using GCaMP3 as Ca^{2+} indicator in the green emission channel, combined with red-shifted variants of sypHy as indicators for vesicle fusion. Examples of the latter are VGLUT-mOr2⁶¹ or SypHyTomato⁶². An alternative approach uses the red shifted calcium indicator R-GECO1⁶³ and sypHy in a unique probe sypHy-RGECO⁶⁴ respectively targeted to the intracellular space and the vesicular lumen. This enables to simultaneously track Ca^{2+} -events and concomitant vesicular fusion with a single-molecule indicator. Our laboratory recently developed a new imaging fluorescent probe, sypHyJREx, made of sypHy and JREx, a red-shifted Ca^{2+} indicator developed from REX-GECO1⁶⁵. pHluorin located in the vesicle lumen and JREx on the cytoplasmic outer surface are excited with the same wavelength (480 nm) but their emission peaks occur respectively at 510 and 585 nm. This allows to report Ca^{2+} -secretion coupling by simultaneous dual channel imaging. The slow Ca^{2+} -association kinetics of the GECIs will still cause a delay, but on the other hand, optical detection offers the advantage of providing spatial information. The invasiveness of neuronal stimulation can also be overcome by optical stimulation methods. For example, the red-shifted channelrhodopsin Chrimson⁶⁶ gives promising perspectives for photostimulation in combination with the detection of Ca^{2+} and vesicle fusion events.

To summarize all the above, our investigation in hippocampal excitatory neurons yielded new insights in the various types of spontaneous Ca^{2+} fluctuations that occur during rest in the presynaptic compartment. Although our methods do have some limitations, we did not detect a tight temporal coupling of quantal fusion events to SCEs. Several mechanistic models, summarized in Figure 1 (Chapter 6), can explain how Ca^{2+} may regulate spontaneous release. The contributions of each of these mechanisms remain to be investigated in the future.

3. Mechanism(s) of spontaneous release

3.1. Functional diversity of C₂ domains

In this research project different proteins containing similar C₂ domain architectures were investigated using the same methodology (see Chapter 5). C₂ domains typically form independent Ca²⁺-binding modules⁶⁷ that endow Ca²⁺-dependent lipid binding properties to the protein. However, as an exception to this typical pattern, some C₂ domains have diverged evolutionarily into Ca²⁺-independent forms⁶⁷ suggesting that C₂ domains also have other functions. Ca²⁺-independent isoforms of protein kinase C such as PKC δ and θ lack Ca²⁺- and phospholipid binding, but instead bind phosphotyrosine instead^{68,69}, providing an alternative activation mechanism. As a second example, the Syt-12 isoform which colocalizes with Syt-1 on synaptic vesicles does not bind Ca²⁺, but its overexpression still specifically increases spontaneous release in cultured neurons⁷⁰. This suggests that C₂ modules provide Ca²⁺-independent secretory functions. The third member of the Doc2 protein family, Doc2c, despite its high amino acid sequence identities with Doc2a and b (45.6 % and 43.2% respectively), does not contain Ca²⁺-dependent phospholipid binding activity⁷¹ nor associates with the PM in response to Ca²⁺ influx^{47,71}. Doc2c contains a unique highly basic sequence^{177RLRRRRR183}, situated in the CBL3 of the C₂A domain⁷², that could theoretically support Ca²⁺-independent interactions with negatively charged membranes. In chapter 5, we show that Doc2c is not involved in spontaneous (Chapter 5, Figure 2) or evoked (Chapter 5, Figure 3) neurotransmitter release in absence of the main sensors for spontaneous release in hippocampal neurons. This is in line with the lack of conservation of critical aspartate residues in its C₂A domain. Most proteins with C₂ domains function in signal transduction or membrane traffic, e.g. for the generation of lipid second messengers or exocytosis. Whereas Doc2c does not function in glutamate exocytosis, we cannot rule out a role in other secretory processes.

The conserved amino acid sequence of the C₂B domain suggests it might still bind Ca²⁺ whereas the C₂A domain may act as Ca²⁺-independent targeting module for the protein (see review)⁷³. Quantitative single cell RT-PCR revealed substantial Doc2c co-expression together with Doc2a,b in cultured hippocampal neurons⁷⁴. The MID domain sequence is also fully conserved meaning that the DAG-dependent Munc13 association is functional. It could be hypothesized that Doc2c acts on vesicle trafficking through the regulation of Munc13. The basic amino acid sequence in the Doc2c's C₂A domain has been proposed to function as a nuclear localization signal⁷² which targets Doc2c to the nucleus^{47,72}. Mouse transcriptome data indicates that it is highly expressed in the heart⁷⁵, unlike other Doc2s. Cardiac myocytes present cytosolic ([Ca²⁺]cyt) and nuclear [Ca²⁺] ([Ca²⁺]nuc) oscillations which respectively control the beat-to-beat contractile activation and gene expression⁷⁶. Both activities converge to an excitation-transcription feedback loop that integrates contractile activity for adequate cardiomyocyte reprogramming. Thus, in the heart, Doc2c activation via C₂B could conceivably contribute to nuclear envelope dynamics or gene expression reprogramming. In Chapter 5, we reported the first knock-out mice model for Doc2c, which are viable and display no sign of major impairment in neural functions or survival. Therefore, its hypothesized activity cannot be essential for vital aspects of brain and heart function.

Another difference observed in different C₂ domains is the nature of the change induced by Ca²⁺-binding. Electrostatic modifications are considered to be the major contributor to C₂ domain activation of Syts^{77–79} and Doc2b⁸⁰. In addition, hydrophobic interactions support membrane penetration of loops extending out of the Ca²⁺-binding site. Yet, a long variant of piccolo's C₂A domain^{81,82} and the CBL1 in the C₂A domain of rabphilin-3A⁸³ undergo significant conformational changes upon Ca²⁺-binding. The Piccolo long variant shows low Ca²⁺ affinity, which is increased in the short C₂A variant⁸², suggesting that a conformational difference directly affects the Ca²⁺ affinity. In both PCLO and rabphilin, the backbone and side chains of the Ca²⁺-binding aspartates adopt conformational modifications that have been observed in structural studies of the Ca²⁺-free/bound structures by NMR and X-ray crystallography.

Thus, the range of C₂ domain activation mechanisms seems more broad than previously thought. Many C₂ domains show limited conformational variations upon Ca²⁺ binding, likely due to the compact arrangement of C₂ domains together with other fusogenic proteins, which may not permit conformational changes. Another conclusion is that slight structural modifications from one C₂ domain to another may account for important functional changes and biophysical properties. Given the significant differences in their mode of action, it is important to specifically consider each protein in the context of structural models for their activity.

3.2. C₂A and C₂B domains: role in phospholipid binding and membrane fusion

There is an important distinction to make between single C₂ domain proteins (e.g. PKC, PLC) and multiple tandem C₂ domains (Syts, Doc2). Single C₂ domains regulate protein membrane targeting and activity⁸⁴ whereas tandem C₂ domains have an additional activity in inducing membrane deformation, which contributes to the regulation of membrane fusion probability together with the SNARE-dependent activities^{85–89}. It has been reported that tandem C₂ domains in close proximity are necessary to bridge opposing membranes in contrast to individual C₂ domains that fail to trigger membrane cross-linking^{77,90}. Accordingly, the activity of Syt-1 C₂ domains to induce membrane curvature and fusion requires that the C₂A and C₂B domains are linked to each other in tandem⁸⁷. The presence of two or more C₂ domains in proteins enhance the membrane- and Ca²⁺-binding affinity beyond that of individual C₂ domains. Syt-1 tandem C₂ domains were shown to Ca²⁺-dependently penetrate deeper into the membrane bilayer compared to individual C₂A and C₂B, suggesting cooperativity^{85,91} and Syt-7 tandem C₂AB binds liposomes with greater Ca²⁺-sensitivity than individual domains⁹². Additionally, Syt-1 C₂AB dissociates from PIP₂-free liposomes much more slowly than either of its individual C₂ domains and Syt-7. Therefore, tandem C₂A and C₂B domains do not act independently but influence their mutual membrane penetration by cooperative energetic effects.

Isolated and tandem C₂ domains of Syts and Doc2s present different Ca²⁺- and phospholipid-binding properties. Quantitative measurements of the Ca²⁺ dose-dependency of phospholipid binding by Syt-1 C₂ domains have yielded various estimates, which strongly depend on the

lipid composition. The Ca^{2+} -sensitivity of the C_2B domain appeared higher than that of the C_2A . PIP_2 binding to the C_2B domain increases the Ca^{2+} -sensitivity and strongly influences the sensitivity of full length Syt-1, while C_2A binding is predominantly governed by negatively charged phospholipids⁴⁹. Consistently, Ca^{2+} association to the C_2A domain of Syt-1 appears important but not essential for the Ca^{2+} -dependent function of the protein⁹³, because abrogation of Ca^{2+} binding to this domain by mutations (D232N and D238N) did not induce major changes in synaptic transmission. Loss of function mutations in the C_2B domain of Syt-1 have more prominent effects on synaptic transmission than similar mutations in the C_2A domain^{85,93-95}. Thus, the C_2B domain is the main determinant for Syt-1-driven Ca^{2+} -secretion coupling. Yet, a recent publication reported the absolute requirement of a functional C_2A domain for neurotransmitter release⁹⁶. Mutation of two hydrophobic residues (M224E and F286E) present at the tip of each CBL and adjacent to aspartate residues, abolished release more severely than a Syt-1 null mutation in *Drosophila*⁹⁶. A plethora of mutants was generated in the fast sensor C_2 domains and their effect on neurotransmitter release was scrutinized. Substitutions in Syt-1, homologous to those in Doc2b, produce different phenotypes^{97,98}. D230, 232N neutralization suppressed the Ca^{2+} binding activity of the protein⁹⁹ and SNARE binding which can be interpreted as a loss-of-function. Neutralization of positively charged residues R233 and K366 present at the membrane binding interface also impaired Ca^{2+} -sensitivity for liposome interaction and reduced exocytosis¹⁰⁰. Similar to mutagenesis in Doc2b, mutations in Syt-1 demonstrated that changing its apparent Ca^{2+} affinity for either its phospholipid interactions, or its SNARE binding, altered the apparent Ca^{2+} affinity of release⁸⁵.

In Syt-7, the C_2A domain has a much higher Ca^{2+} -sensitivity of membrane binding compared to Syt-1 C_2A ¹⁰¹. This high Ca^{2+} -sensitivity dominates the phospholipid binding property of full length Syt-7¹⁰². Quantitative fluorescence measurements revealed a slower dissociation of Syt-7 C_2A from lipid bilayers¹⁰¹. The crystal structure of Syt-7 Ca^{2+} -bound C_2 domains was found very similar to that of Syt-1 C_2 domains^{102,103}. Isothermal titration calorimetry (ITC) data also indicate that the intrinsic Ca^{2+} -binding properties of Syt-7 C_2 domains are comparable to those of the Syt-1 C_2 domains^{102,103}. Altogether, those data do not provide an explanation for the differences in the functional importance of C_2 domains between Syt-1 and Syt-7. The different behavior of Syt-7 and Syt-1 is ascribed to subtle amino acid substitutions in the Ca^{2+} binding interface. In Syt-7, Ca^{2+} association to the C_2A domain is the major mediator in asynchronous release¹⁰².

Also for Doc2b, the apparent Ca^{2+} affinities of the C_2 domains differ from each other and are influenced by the presence of membranes and their phospholipid composition. ITC titrations of isolated C_2 domains in absence of lipids indicated that only C_2B binds Ca^{2+} ¹⁰⁴ and was solely responsible for Ca^{2+} binding of the C_2AB tandem. In the same study, in presence of lipids, C_2A appeared as the main actuator in the Ca^{2+} -dependent liposome association of the C_2 tandem¹⁰⁴. Crystal structures of the C_2A and C_2B domains indicated that they accommodate two Ca^{2+} ions each with minor conformational changes⁸⁰. In tandem C_2AB , the C_2B domain appeared to be the primary Ca^{2+} sensing unit whereas C_2A had an additive effect in enhancing the interaction of C_2AB with the PM⁸⁰.

Similarly, the isolated C₂A domain binds to liposomes in a Ca²⁺ dependent manner while binding of the C₂B domain is limited³. In chapter 4, we measured the apparent Ca²⁺-affinities of phospholipid binding by the C₂A and C₂AB domains of Doc2b. The C₂AB domain showed a drastically higher Ca²⁺-affinity compared to C₂A (176 nM and 435 nM respectively; Chapter 4, Figure 3) which illustrates the synergistic effect. Aspartate substitutions in the C₂A domain of Doc2b drive its constitutive PM association^{2,104–106}, whereas similar mutations in the C₂B domain do not^{2,104}. Taken together, it is clear that the C₂A and C₂B have complementary functions where Ca²⁺ activation of soluble Doc2b is likely driven by the C₂B domain and membrane binding is mostly governed by C₂A domain interactions.

3.3. Multiple sensors for spontaneous release

Many neurons co-express a fast sensor (Syt-1, 2 or 9) together with a sensor for slower asynchronous release (Syt-7) and for spontaneous (AP)-independent release (Doc2a,b), together encoding the complex stimulus-responsiveness of synaptic secretion^{107–109}. The fact that co-expressed sensors operate together becomes apparent upon deletion of the fast Ca²⁺-sensor, leaving the remaining component still Ca²⁺-sensitive^{110,111} yet showing altered Ca²⁺-dependency. In many systems the loss of fast release is accompanied by increased asynchronous and spontaneous release^{110–116}. Given the co-existence of these multiple Ca²⁺ sensors in synapses and other secretory systems, it remains the question how these sensors interact together to give rise to the overall stimulus-secretion coupling properties of the secretory apparatus¹⁰⁹.

The competition model

One model hypothesizes that several Ca²⁺ sensors with different Ca²⁺/membrane binding affinity and kinetics compete for vesicular release. This scenario is commonly used to explain why the deletion of one Ca²⁺ sensor causes changes the Ca²⁺-sensitivity and cooperativity of release (for review)¹⁰⁷. The occupancy of partially assembled SNARE complexes could be the biochemical basis for such competitive activity of Ca²⁺ sensors. Subcellular targeting of fast sensors could provide a competitive advantage due to local enrichment at these sites, for example encoded by the N-terminal domain of Syt-1 and additional interactions of its cytosolic part with N-, P/Q-, L-type Ca²⁺ channels irrespective of the presence of Ca²⁺^{117–119}. This model provides a plausible explanation for the increase in the apparent Ca²⁺ sensitivity and higher release probability at resting Ca²⁺ concentrations in absence of Syt-1 where Doc2 is able to occupy vacant Syt-1 driven release site (Chapter 6, see Figure 1).

Null mutations in Doc2a/b induces a decrease in spontaneous release rate by 50-75% compared to wildtype neurons^{3,4}. In chapter 4, we investigated the relation between Doc2b Ca²⁺-binding site mutants and spontaneous release regulation. We suggest that resting stochastic release parallels Doc2b's Ca²⁺-dependent membrane binding (Chapter 4, Figure 2, 3, 5). A recent investigation revealed that removal of Doc2a/b in Syt-1 KO hippocampal neuron culture did not reduce the high mEPSCs rate, meaning that Doc2 proteins do not account for the increased minis upon Syt-1 ablation¹²⁰. This outcome is supported by a prior publication⁴ where they used an shRNA-dependent quadruple knockdown (KD) of all four

Ca²⁺-binding proteins of the Doc2 family in cortical neurons. Knockdown synapses exhibited a marked decrease in the spontaneous event frequency in Syt-1-expressing neurons, but the knockdown did not suppress the high spontaneous release frequency in Syt-1-deficient neurons. The remaining spontaneous release in Syt-1, Doc2a,b deficient neurons could be supported by a still undiscovered other class of Ca²⁺-sensor. In Chapter 5, we investigated several candidate C₂ domains proteins but did not identify the one responsible for the residual spontaneous release.

In addition to Syt-1 and Doc2, the high affinity Ca²⁺-sensor Syt-7 was shown to modulate the spontaneous component^{74,108}. Electrophysiological investigations revealed no phenotype upon Syt-7 ablation in wildtype neurons¹²¹ but its overexpression in Syt-1 KO neurons⁷⁴ and removal in Syt-2 KO neurons¹⁰⁸ respectively decreased and increased spontaneous exocytosis. Comparably to Syt-1, this suggests a supplementary clamping function for this isoform in absence of the fast sensor. This is not in line with evidence presented in chapter 5, as Syt-7 ablation in Doc2a, b, c, Raphilin knock-out genetic background had no effect on spontaneous release rate (Chapter 5, Figure 6), confirming that the effect of Syt-7 on this component only becomes apparent upon Syt-1, -2 removal. This hypothesis is supported by the fact that fast Ca²⁺ sensors mask the role of Syt-7 in short-term plasticity¹²² and asynchronous release¹⁰⁸. Thus, a pattern is emerging where faster Ca²⁺ sensors contribute more dominantly to overall synaptic release, masking the activities of the slower ones.

The dual fusion / clamping model

An alternative explanation for the mixed effects of Syt-1/2 deletion on different phases of release states that fast sensors synchronize Ca²⁺-evoked release by a dual mechanism, which combines a positive role in fusion enhancement with an inhibitory role under resting conditions. This inhibitory role is often referred to as ‘clamping’. Clamping could result from competitive inhibition of a more sensitive 2nd sensor as predicted by the competition model. Paradoxically, Syt-1 point mutations that moderately (D238N) or severely (R233Q) alter the apparent Ca²⁺-affinity for phospholipid binding decrease spontaneous release¹¹⁴. If clamping was mediated by the Ca²⁺-dependent membrane binding capacity of Syt-1, those mutations would have unclamped Doc2 (or other sensors) and increase spontaneous release. Instead, spontaneous release is reduced, meaning that spontaneous fusion inhibition isn’t mediated by a membrane attachment mechanism.

A mutational analysis of Syt-2 identified a role of the poly-lysine motif in the C₂B domain in spontaneous release inhibition¹¹¹. In this study, the triple substitution K327Q, K328Q, K332Q blocked t-SNARE binding. The resulting Syt-2^{3K} mutant fully rescued evoked release but lost its activity to inhibit spontaneous release at rest, demonstrating that clamping requires association with t-SNAREs or PIP₂. In Syt-1, the clamping function was selectively affected in a variant termed Syt-1^{9Pro}, in which proline substitutions in the flexible linker between C₂A and C₂B constrains the orientation of both domains relative to each other¹²³. These observations suggest that the inhibitory mechanism is distinct from the fusogenic one, with differences pertaining to the relative disposition of the C₂ domains and the mode of target membrane association via t-SNAREs or PIP₂.

The dual effect of Syt-1 is paralleled by complexins which clamp spontaneous release at rest by arresting SNARE complex zippering via its central helix and an accessory helical domain^{124,125}. This Ca^{2+} -sensitive inhibition synchronizes Ca^{2+} triggered SVs fusion¹²⁶. The collaborative action of Syt-1 and complexins in synchronizing Ca^{2+} -triggered membrane fusion is reinforced by a reconstituted single vesicle-vesicle fusion assay where Syt-1 and complexins individually inhibit spontaneous fusion and show a further negative effect when both incorporated¹²⁶. A recent crystal structure of the SNARE proteins together with complexins and Syt-1 provides a structural basis for their synergistic function¹²⁷. Altogether, this structure seems to constitute the elementary Ca^{2+} -sensitive platform in the primed vesicle state.

Díez-Arazola et al. reported an unexpected inhibitory effect of Doc2b on spontaneous release upon its overexpression in Syt-1, Doc2a,b TKO hippocampal neurons¹²⁰. A plausible explanation is that this inhibition is caused by increased spontaneous GABA release in mixed cultures. Long-term spontaneous release of GABA can regulate the mEPSC frequency of glutamatergic cells in a mixed culture¹²⁸. In addition, Doc2b is predominantly expressed in GABAergic neurons⁵⁶. Thus, the inhibitory effect of Doc2b in excitatory neurons may be the consequence of an increased spontaneous GABA release specifically. This idea is consistent with the positive endogenous role of Doc2b in spontaneous release³. In chapter 4, neither Doc2 overexpression in wild type neurons (Chapter 4, Figure S3) nor its rescue in Doc2a,b KO neurons (Chapter 4, Figure 5) lower the mEPSC rate, indicating that endogenous Doc2b isn't limiting spontaneous release rate.

3.4. Working model: how Doc2 proteins drive spontaneous release

Fast Ca^{2+} sensors prevent spontaneous fusion at rest. This helps to preserve fusogenic vesicles for rapid fusion upon Ca^{2+} activation. The spontaneous component is strongly affected by the presence or absence of different Syts^{129,130}. This property is likely to arise from the privileged location of Syts on the vesicle membrane (Syt-1) or the opposed PM (Syt-7). At rest the soluble Doc2 protein is localized in the cytosol (Chapter 6, Figure 1B) but can be recruited into the vicinity of fusion competent vesicles by various actors. The high Ca^{2+} -sensitivity and phospholipid affinity offer a near-resting activation capacity that is highly responsive to spontaneous Ca^{2+} fluctuations (Chapter 6, Figure 1C) or residual $[\text{Ca}^{2+}]_i$ increases induced by synaptic activity. These Ca^{2+} fluctuations are too small to activate synaptotagmins (Chapter 6, see Figure 1D). This hypothesis is supported by previous finding where Doc2b dependent quantal release rate is strongly increased following train stimulation and extracellular Ca^{2+} rise³ both of which conditions increase intracellular Ca^{2+} events (Chapter 3, Figure 4 and 6). Our model conceptualizes mechanisms of action of Doc2 in four different contexts (Chapter 6, Figure 1). During AP-dependent release, VGCC opening induces a spatially restrained Ca^{2+} -influx with a high peak concentration (Chapter 6, Figure 1A). In that scenario, Doc2 is mainly inactive in the cytoplasm. Few Doc2 copies might be present at the PM, by chance or recruited to it by a Munc13-, Munc18- or SNARE-dependent mechanisms. Synchronous neurotransmission is triggered by Syt-1 activation, located at the release site.

Syt-1 may inhibit Doc2 or Syt-7, competitively by release site occupancy (e.g steric hindrance at the primed proteolipid complex) or by active clamping together with complexins. In absence of AP depolarization, represented by the second scenario, spontaneous release is hypothetically driven by fast Ca^{2+} transients (SCTs) of short duration caused by activation of metabotropic receptors or VGCC flickering (Chapter 6, Figure 1B). Doc2 membrane association is determined by the basal resting $[\text{Ca}^{2+}]_i$ and complementary recruitment pathways (Munc13, -18, SNAREs). In this context, spontaneous release may be triggered by Syt-1 or Doc2, depending on Ca^{2+} amplitude and the proximity of the sensors from the VGCC. This theory might be more relevant for GABAergic neurons because Syt-1 and Doc2b are both required for spontaneous release in inhibitory neurons⁵⁶. The third scenario postulates that spontaneous release might be triggered by complex Ca^{2+} events (SCEs) generated by activation of metabotropic receptors or single or several VGCCs, amplified by the CICR process (Chapter 6, Figure 1C). The Ca^{2+} wave could enable a significant Ca^{2+} /Munc13-dependent Doc2 translocation and concomitantly induce Doc2-driven spontaneous release. In the fourth scenario, spontaneous events are generated by a global Ca^{2+} (Chapter 6, Figure 1D), as after a repetitive synaptic activity where residual Ca^{2+} gradually decreases. Syt-1 and Syt-7 are then inactive, but Doc2 could translocate to the PM and elicit spontaneous release. Massive activation of Doc2 could override the inhibitory effect of Syt-1 or alternatively, disassembly of Syt-1 oligomers after repeated synchronous release could enable Doc2 mediated fusion. Global $[\text{Ca}^{2+}]_i$ could enable Doc2 translocation via simple Ca^{2+} -association or via DAG-dependent Munc13 binding.

In Chapter 4, we confirmed previously reported data on high Ca^{2+} -affinity properties of wildtype Doc2b (Chapter 4, Figure 3) but also showed a novel inhibitory effect of high $[\text{Ca}^{2+}]_i$ on lipid binding behavior. At high Ca^{2+} concentration, the C_2A and C_2AB fragments (Chapter 4, Figure 3) showed a decreased liposome binding, suggesting that each sensor might only function in a limited $[\text{Ca}^{2+}]_i$ window and are inhibited by higher $[\text{Ca}^{2+}]_i$. A striking parallel was observed between the Ca^{2+} -dependent lipid binding properties of Doc2b and its effect on exocytosis at rest or during activity (Chapter 4, Figure 5). Finally, we revealed a potential function in synaptic recovery (Chapter 4, Figure 7) which is likely regulated by residual Ca^{2+} , because the effect was abrogated in mutant forms. The "site-occupancy" model mentioned above fits well with our experimental results. Altogether, multiple Ca^{2+} sensors seem to be organized in hierarchy, dominated by Syt-1 (2 and 9; fast Syt) mediating the synchronous phase, followed by Syt-7 (slow Syt) involved in asynchronous release and finally Doc2 for spontaneous fusion. This hierarchy is likely based on their respective localization relative to the release machinery and their biophysical activation properties¹³¹.

3.5. Spontaneous release: Ca^{2+} sensor or not?

An ongoing point of debate was whether Doc2b acts as a direct Ca^{2+} -sensitive protein for exocytosis³ or as a structural element⁴ (see chapter 4). Neutralization of two critical aspartates D218 and D220 in the C_2A domain of Doc2b appeared to mimic a Ca^{2+} -bound state which caused constitutive membrane enrichment^{2,105} associated with an increased spontaneous release frequency³ supporting its role as a Ca^{2+} -sensor^{3,47}.

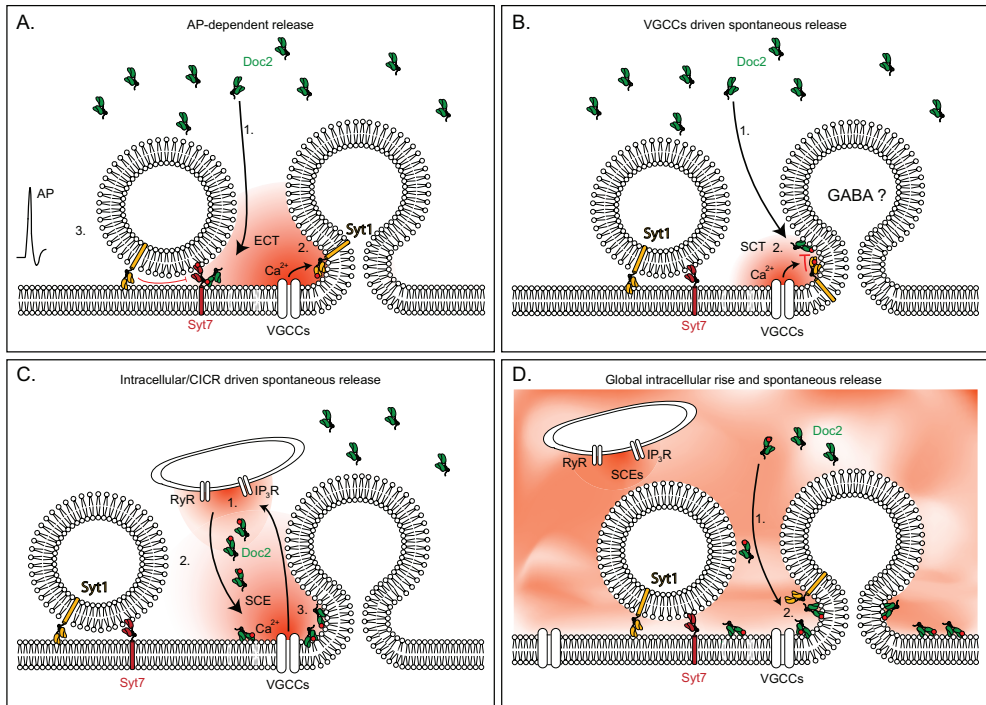


Figure 1: Model for the interplay of Ca^{2+} sensors and the Ca^{2+} -sources which may trigger evoked (A) and spontaneous (B-D) release. A. In the scenario of neuronal activity, VGCC opening induces a spatially restrained Ca^{2+} -influx with a high peak concentration. 1. Prior to the evoked Ca^{2+} -transient, only few Doc2 copies might be recruited to the PM by a Munc13-, Munc18- or SNARE-dependent mechanism (these proteins are not depicted for simplicity). 2. Synchronous release is elicited by Syt-1 activation following the ECT. The Ca^{2+} -sensor activation following Ca^{2+} association is represented by a red dot on the C_2 domain. In parallel, due to the preferential location of Syt-1 to the release site, its occupancy or sterical hindrance at the fusogenic proteolipid complex, Syt-1 may competitively inhibit Doc2 or Syt-7 (red arrows). 3. In a second phase or further away from Ca^{2+} nanodomains, Syt-1 is not activated and inhibits fusion, leaving the vesicle in a primed state where spontaneous fusion is unlikely. B. In the scenario of spontaneous release driven by VGCC flickering, fast SCTs occur which cause low-amplitude Ca^{2+} transients of short duration. 1. The proportion of membrane-bound Doc2 is determined by resting $[\text{Ca}^{2+}]_i$ and complementary recruitment pathways (Munc13, -18, SNAREs). 2. Spontaneous release may be triggered by Syt-1 or other sensors depending on Ca^{2+} amplitude, sensor affinity and proximity. This theory might be more relevant for GABAergic neurons because Syt-1 and Doc2b are both required for spontaneous release in inhibitory neurons, while Syt-1 is not necessary for mEPSCs which solely depend on Doc2a⁵⁶. C. In the scenario of spontaneous release elicited by SCEs, Ca^{2+} -influx through single or several VGCCs could be amplified by RyR/IP₃R opening (1.) (CICR). 2. This Ca^{2+} -fluctuation would enable a significant Ca^{2+} /Munc13-dependent Doc2 translocation and concomitantly induce a (3.) SCE-dependent Doc2-driven spontaneous fusion event. D. In the last scenario of a global Ca^{2+} wave e.g. following a stimulation train, Syt-1 and Syt-7 are inactivated but Doc2 translocates to the PM (1.) and elicits spontaneous release. Under these conditions, massive activation of Doc2 could override the inhibitory effect of Syt-1 (2.). Moreover, global $[\text{Ca}^{2+}]_i$ could enable Doc2 recruitment via simple Ca^{2+} -association but also via DAG-dependent Munc13 binding. In this state, VGCCs are closed but RyR and IP₃R opening could nourish/maintain the Ca^{2+} wave.

Another Ca^{2+} -ligand mutant Doc2b^{6A}, in which six critical aspartates were neutralized by alanine substitution, still supported spontaneous fusion while it did not bind Ca^{2+} , supporting its involvement as a structural element⁴. In chapter 4, we addressed the apparent discrepancy of those results and revealed that both mutants share very similar characteristics. Recently the 6A substitution was clearly identified as mimicking a Ca^{2+} -bound state⁵⁶. In this experiment set, the mutant caused an increased mIPSC frequency that was largely Ca^{2+} -independent. These data, in agreement with the absence of translocation behavior and the Ca^{2+} -insensitive membrane binding properties consistently point to a Ca^{2+} -mimicking effect, causing gain-of-function behavior at rest. Recently, a genuine loss-of-function substitution was identified (D303N) which completely abolishes Doc2b translocation behavior^{2,56} and its capacity to support asynchronous² or spontaneous release⁵⁶. That draws a clear correlation between Doc2b Ca^{2+} -dependent membrane association and its ability to enhance exocytosis. The loss-of-function mutant is particularly interesting as it can be used as a negative control and provide clarification on Doc2b's function in exocytosis. Finally, the finding that the C₂AB domain of Doc2b enhances hemifusion in a Ca^{2+} -dependent manner^{132,133} further consolidates the Ca^{2+} -sensor theory.

Intracellular Ca^{2+} -chelation by BAPTA-AM provokes the loss of most (75% to >95%) spontaneous release events^{3,111,114} and $[\text{Ca}^{2+}]_e$ rise increases them^{3,4,114} (Chapter 3, Figure 4 & 6). The Ca^{2+} -dependence could possibly reflect a pool recruitment mechanism. Vesicle recruitment is Ca^{2+} -dependent¹³⁴. It accelerates during accumulation of Ca^{2+} within the presynaptic terminal¹³⁵ and diminishes after application of the fast Ca^{2+} buffer BAPTA, but not the slower buffer EGTA¹³⁶. The process possibly involves synapsins, Munc13 alone or together with calmodulin^{137–139}. In line with the positive effect of $[\text{Ca}^{2+}]_e$ on global $[\text{Ca}^{2+}]_i$ (Chapter 3, Figure 4A-C) and considering the effect of $[\text{Ca}^{2+}]_i$ rise on the RRP size, the dependence of spontaneous release to extracellular Ca^{2+} might partially depend on the accumulation of release-ready vesicles, independently of Ca^{2+} sensors. The enhancement of synaptic recovery by Doc2b overexpression after train stimulations (Chapter 4, Figure 7) could imply a Ca^{2+} -sensitive function for Doc2b in RRP maintenance. Yet, no change in RRP size was found in Doc2 deficient neurons^{3,4}. An evaluation of the RRP size by sucrose application upon various $[\text{Ca}^{2+}]_e$ perfusion in either Doc2 removal or overexpression would clarify our questioning whether Doc2 speeds-up RRP refilling in a Ca^{2+} -dependent manner.

Another question is whether spontaneous release absolutely requires Ca^{2+} sensors or can be sensor-less, only needing SNARE zippering for spontaneous fusion of synaptic vesicles. Residual spontaneous events remain upon Ca^{2+} -chelation by EGTA and BAPTA^{3,34,37,111,114}. This might be explained by the inability of the Ca^{2+} -chelators to completely absorb free Ca^{2+} ions, as the K_d for EGTA and BAPTA is ~100 nM and spontaneous release could occur by $[\text{Ca}^{2+}]_i$ below this value. Alternatively, a portion of spontaneous release could be Ca^{2+} -independent. *In vitro* reconstitution experiments with synaptobrevin and syntaxin-1–SNAP-25 incorporated into separate populations of liposomes¹⁴⁰ indicated that the SNAREs alone can induce vesicle fusion¹⁴¹. However, spontaneous events *in vivo* are tightly regulated and rendered Ca^{2+} -dependent by the complexin-Syt-SNARE complex, preventing stochastic fusion.

Still, one can imagine that a temporary lack of a strong fusion clamp would allow spontaneous SV fusion with the PM. Syt-1 was recently reported to self-assemble into a ring-like oligomers¹⁴² which would be essential for the clamping function of Syt-1¹⁴³. The latter investigation showed that destabilization of ring-shaped Syt-1 oligomers releases its clamping function, resulting in uncontrolled spontaneous fusion in absence of Ca²⁺¹⁴⁴. In this context, SVs would diffuse freely on the bilayer until they fuse spontaneously even in absence of Ca²⁺ sensors. This hypothesis provides an explanation for the spontaneous release phenotype in Syt-1 KO neurons and supports Ca²⁺-independent spontaneous release.

3.5.1. The impact of mutation in C₂ domains

C₂ domains are centrally involved in Ca²⁺-sensor function by interacting with SNAREs and membranes and they enhance membrane fusion (for review)^{145,146}. Selective mutagenesis of the polybasic motif inhibited the SNARE interaction in Doc2b^{K237,319E}³, but did not abrogate liposome binding. Vice versa, substitution of the conserved amino acids H158, F222 and I360 to alanine in the Ca²⁺-binding loops termed 4A mutation produced a lipid-binding-incompetent protein without blocking the SNARE interaction³. Thus, SNARE versus phospholipid association can happen independently. In chapter 4, we compared Doc2b^{DN} and Doc2b^{6A} mutations. Both versions of Doc2b mimic a Ca²⁺-bound state (constitutively active) which enhance spontaneous release. The Ca²⁺ ligand site mutation D218,220,357,359N, termed C2A_{CLM}B_{CLM}^{2,104} abolishes Ca²⁺ association to C₂ domains but greatly increases spontaneous fusion rate. Extensive mutagenesis work on Doc2b where the five critical aspartate residues in each Ca²⁺-binding loop were mutated by paired or single substitutions showed that the D220N substitution within the C₂A Ca²⁺-binding-pocket is solely responsible for the constitutive membrane binding² and the mEPSC frequency increase. The behavior of the DN, 6A and CLM/CLM substitution is therefore explained by the single D220 neutralization which thus governs Doc2b lipid binding.

3.6. Distinct spontaneous release mechanisms in excitatory and inhibitory neurons

It was suggested that glutamatergic and GABAergic events are driven by the expression of Doc2a and -b respectively, although both isoforms are redundant in their ability to rescue spontaneous release⁵⁶. In this study, both Syt-1 and Doc2 were confirmed to regulate minis. The contribution of Doc2a vs Doc2b on minis depends on whether one are dealing with glutamate or GABA secretion⁵⁶. Single cell mRNA levels did not systematically corroborate specific expressions of Doc2a and Doc2b in pyramidal cells and interneurons respectively¹⁴⁷. Yet, recent studies reported a predominant expression of Doc2a in CA3 pyramidal cells and of Doc2b in GABAergic striatum neurons^{56,148}. The specific expression and function of Doc2a and b in excitatory and inhibitory neurons is still debated. It appears that mechanisms for Ca²⁺ signal that trigger spontaneous release at inhibitory but not excitatory synapses mostly rely on VGCCs³³. Divergent outcomes were reported about the effect of [Ca²⁺]_e on mEPSCs⁵. In our experiments (Chapter 3 and 5), the mEPSC frequency release rate correlated with global [Ca²⁺]_i changes but the events were not time-locked to SCTs. This result from excitatory neurons does not exclude such coupling in GABAergic synapses (Chapter 3). It would thus be valuable to extend our investigation on the correlation between SCEs and spontaneous release to inhibitory neurons.

The high spontaneous release frequency in neurons lacking Syt-1 is more apparent in mixed glutamatergic/GABAergic pairs than in autaptic culture^{112,128}. This is apparently the consequence of the GABAergic innervation of glutamatergic neurons. We can relate this finding to results we obtained in chapter 5 (Chapter 5, Figure 4) where spontaneous release increased upon rabphilin ablation in neuronal networks, but not autapses. Rabphilin might specifically promote spontaneous events in inhibitory neurons. Its ablation would then lower stochastic GABA release rates, indirectly causing a mEPSC frequency rise in networks. In light of all those findings, it can be recommended that investigations evaluating the implication of fusogenic proteins in spontaneous release take place in mixed glutamatergic/GABAergic cultures.

3.8. Candidate sensors: how to find them

Although our findings revealed no effect in favour of the regulation of minis by Doc2c, Syt-7 or rabphilin, many unidentified Ca^{2+} sensors could exist. Proteins containing C_2 domains are potential candidates. Among the plethora of presynaptic proteins, Syt isoforms, Munc13s, ferlins and copines¹⁴⁹ are potential regulators of spontaneous fusion in glutamatergic hippocampal neurons. A Ca^{2+} sensor promoting spontaneous vesicle fusion is expected to meet several requirements. It should be active near resting Ca^{2+} concentrations and increase the release probability of fusion in the absence of the fast sensors Syt-1, -2^{109,110}. It should display high Ca^{2+} -sensitivity of membrane binding, higher than Syt-1 and Syt-7 and be present at least temporarily in the vicinity of fusion competent vesicles. Additional bioinformatics analysis, based on sequence profile searches, phylogenetic and phyletic-pattern and structure-prediction, have implemented the long list of proteins containing C_2 domains and greatly facilitate the quest. Related C_2 domain proteins have been included in the large collection of proteins families of the PFAM database (<http://pfam.sanger.ac.uk>). Proteins showing a high degree of amino acid sequence homology with Syts and Doc2s should then be favoured.

4. Spontaneous release in health and pathologies

Miniature synaptic potentials have been considered for long as physiologically meaningless events and only considered to define the size of a quantum, quantal release evoked by an AP and modeling of random release¹⁵⁰. Yet, consideration towards spontaneous release changed as cortical and hippocampal neurons were shown to sometimes communicate via a single synapse which release only one neurotransmitter quantum (mini) upon minimal stimulation¹⁵¹⁻¹⁵³. It suggests that single quantum release must have a physiological importance. Interest in spontaneous release grew further as investigations exposed that spontaneous release can produce a significant inhibitory tone in postsynaptic neurons¹⁵⁴ and influence the firing of small interneurons¹⁵⁵. It can then directly shape neuronal excitability as inhibitory quanta could rapidly terminate neuronal firing whereas small numbers of coincident excitatory quanta can trigger firing¹⁵⁵. Alternatively, spontaneous release modulates postsynaptic excitability by regulating the tonic activity of high-affinity receptors^{154,156}. Low concentrations of ambient GABA can persistently activate certain synaptic and extrasynaptic GABA receptors to generate a tonic conductance.

Spontaneous release also participates in postsynaptic homeostasis in mature synapses. Synaptic homeostasis is a process based on feedback regulation which enables the tuning of neurotransmission efficiency to maintain neuron excitability balance. It was suggested to be independent of evoked neurotransmission at the *Drosophila* NMJ¹⁵⁷ as it appeared in presence of TTX. An increase in evoked synaptic strength, resulting from an increase in the efficacy of transmission was reported after blockade of postsynaptic receptors for several minutes. The mechanism implicated presynaptic modulation of VGCCs (CaV_{2.1}) by retrograde signaling. AP-independent homeostatic plasticity could be partially regulated by minis. Synaptic scaling can be engaged by a change in AMPA receptor density. Initial experiments in *in vitro* models revealed that pharmacological blockade of activity (TTX) scales up the AMPA-receptor mediated mEPSCs amplitude¹⁵⁸. The effect of spontaneous neurotransmitter release on synaptic structure conservation and postsynaptic scaling is likely mediated by the immobilization and density maintenance of postsynaptic receptors¹⁵⁹⁻¹⁶¹.

Spontaneous release may also play a role during development. It was suggested to modulate postsynaptic enzymatic activity and protein synthesis^{162,163}. Minis might keep resting protein synthesis processes in check and sensitive to stimuli for synapse strengthening. Rat hippocampal neurons show very high levels of exclusively spontaneous release in young neurons, which progressively decrease during synapse formation and maturation¹⁶⁴. Cortical layer 4 neurons also display an important increase in spontaneous events from development stages P12 to P23¹⁶⁵. Neurotransmitters are released spontaneously from the growth cones of embryonic neurons¹⁶⁶, suggesting a role in nerve growth and axon guidance. One model proposes that spontaneous glutamate release from developing axons organizes axodendritic contact by signaling over long ranges to NMDA receptors on developing dendrites¹⁶⁷, thereby shaping neuronal arborization. Once synaptic contact is established, postsynaptic protein synthesis required for continued growth cone guidance is stopped^{163,168}. Therefore, spontaneous release of neurotransmitter might be important in the wiring of the brain during development¹⁶⁹. Spontaneous release is also involved in the maintenance of dendritic spines via AMPAR receptor activation¹⁷⁰. CA1 pyramidal cell spines decreased in density and length after application of an AMPAR antagonist but not TTX, suggesting that spontaneous input is sufficient for synaptic maintenance.

Precise control of synaptic development, connectivity and maintenance is critical for brain functions. The function of spontaneous release detailed above raises the question which neuropathologies may be triggered in case of dysregulation. One can easily imagine that aberrant development of neuronal circuitry could lead to drug-resistant epilepsy^{171,172}, encephalopathy or autism where mEPSC increases in a critical time window were correlated to the risk of Autism Spectrum Disorders¹⁷³. Disorders associated with faulty neuronal circuits also include schizophrenia¹⁷⁴ or fragile X Syndrome¹⁷⁵. This is corroborated by findings that point to an increased expression of DOC2A (involved in spontaneous release regulation) has been observed in human and rat models of Temporal Lobe Epilepsy¹⁷⁶, while copy number variants of DOC2A have been associated with Schizophrenia¹⁷⁷.

5. Conclusion, perspectives and future directions

The work described in this thesis explored neurotransmitter release regulation by spontaneous presynaptic Ca^{2+} fluctuations and several C_2 domain proteins at excitatory hippocampal synapses. Prior studies have revealed the complexity of the Ca^{2+} -dependent neurotransmitter release in neurons. Despite the physiological significance of spontaneous release, a direct evidence for a relationship between cytosolic Ca^{2+} -signals and resting vesicle fusion is still missing. The first part of this thesis examined presynaptic SCEs and SCTs. The SICT method developed for Ca^{2+} -imaging data analysis significantly accelerated the analysis and appeared sensitive enough to detect changes in the spontaneous release frequency induced by caffeine. $[\text{Ca}^{2+}]_c$ increased both bulk/global cytosolic $[\text{Ca}^{2+}]$ and SCEs in parallel with the mEPSC frequency rise. Although increases in $[\text{Ca}^{2+}]_c$ clearly enhanced minis, a tight coupling between SCEs and mEPSCs was not detected. Calcium transients triggered by intracellular Ca^{2+} sources typically give rise to slower Ca^{2+} waveforms. Additional studies are necessary to further unravel the Ca^{2+} -dependency of spontaneous neurotransmission. New optical tools, such as sypHyJReX are good opportunities to investigate any temporal relation between Ca^{2+} and spontaneous SV fusion. Using this dual indicator, efforts should be made to evaluate the role of different Ca^{2+} influx pathways by examining a temporal coupling at high Ca^{2+} , upon Cd^{2+} blockade of VGCCs, in presence of selective VGCC blockers (ω -Agatoxin for P,Q-type channels; ω -Conotoxin for N-type; SNX-482 for R-type and TTA-P2 for T-type channels) ⁵ in parallel with Ca^{2+} -store depletion by thapsigargin both in glutamatergic and GABAergic neurons. Additional insight could be gained by the development and use of Ca^{2+} probes offering higher Ca^{2+} detection sensitivity and speed (or faster Ca^{2+} association). Altogether, extensive studies of the different types of Ca^{2+} events and their sources are necessary to apprehend their physiological role.

The second part of this thesis focused on the regulation of evoked and spontaneous neurotransmitter release by various C_2 domain proteins. We established that two Doc2b mutants, Doc2b^{DN} and ^{6A} previously categorized respectively as gain and loss of function both constitutively bind to the plasma membrane. The modification of Doc2b Ca^{2+} -dependent properties altered spontaneous release rate, supporting its role as a Ca^{2+} sensor in the release process. Furthermore, potential other Ca^{2+} sensors were investigated regarding their regulation of residual spontaneous release in absence of Doc2a,b. Rabphilin ablation altered mEPSC frequency selectively in neuronal networks, suggesting a positive role on quantal GABA release. Yet, no apparent effect on spontaneous release became prominent upon removal of Doc2c or Syt-7.

Additional investigations are necessary to definitely determine the function of the above-mentioned proteins in spontaneous fusion. Because fast sensors can mask the effects of slower ones, one suggestion would be to examine the effect of Rabphilin, Doc2c and Syt-7 deletion in absence of Syt-1 in excitatory and inhibitory neurons.

Overall, the findings presented in this thesis provide new insight on the regulation of quantal release by Ca^{2+} signaling and C_2 proteins. Despite a collective effort made to decipher the

regulation of spontaneous neurotransmission by multiple sensors, several questions remain: What are the molecular mechanisms and interactor molecules necessary for the inhibition of second sensors by fast sensors? Also, does this inhibitory activity require activation of fast sensors or is it a function that happens at rest? Is the dual capacity of Syt-1 to promote and clamp SVs fusion a common feature for all C₂ domains implicated in neurotransmission? Are the fine-tuned expression levels of fast sensors relative to other sensors important for the Ca²⁺-dependency of neurotransmitter release and short-term plasticity? Answers will considerably increase our knowledge of synaptic physiology, with essential implications for overall brain function.

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