Study of synaptic vesicle cycling during exo- and endocytosis using optical approaches

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Declaration

I hereby declare that my doctoral thesis ‘study of synaptic vesicle cycling during exo- and endocytosis using optical approaches’ has been written independently with no other aids or sources than quoted.

Yunfeng Hua
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1. Introduction

In the central nervous system, communication between nerve cells mainly occurs at specialized junctions called synapses. At the chemical synapse, electrical signals trigger controlled secretion of neurotransmitter via exocytosis of synaptic vesicles (SV) at the presynaptic site. The fusion of SVs releases neurotransmitters that diffuse across the synaptic cleft and activate postsynaptic receptor channels. That generates a new wave of electric signal in the postsynaptic cell (Katz and Miledi, 1965, 1967). Given the limited number of SVs in a presynaptic terminal, complete fusion of many SVs with the plasma membrane would result in fast depletion of fusion-competent SVs and thereby the loss of the ability to transmit signals across synapses. Therefore, to support rapid and repeated rounds of release, a fast trafficking cycle of SVs is required to maintain synaptic function during sustained synaptic activities (Harata et al., 2001; Schweizer and Ryan, 2006; Sudhof, 2004). Although intense research over three decades has led to a deep understanding of the molecular mechanisms behind exocytic membrane fusion (Jahn and Scheller, 2006; Martens and McMahon, 2008) and endocytic retrieval of SV (Dittman and Ryan, 2009; Doherty and McMahon, 2009), comparatively little is known about the coupling between exo- and endocytosis. Hence, understanding the underlying mechanisms of the interplay between exo- and endocytosis still remains one of the central challenges of presynaptic physiology.
1.1. Synaptic Vesicle Pools

SVs are unique secretory organelles with an average diameter of 42 nm. Each SV contains about 1500-2000 neurotransmitter molecules. Through proteomic and lipidomic analysis approximately 40 integral membrane proteins have been identified to be vesicle residents (Figure 1.1), some of which are present in surprisingly low copy numbers per vesicle (Takamori et al., 2006). At each small synaptic bouton (~1 µm in diameter) of hippocampal primary neurons, the total vesicle population comprises about 200 vesicles (Figure 1.1). These vesicles consist of almost the same components and are clustered in a phosphorylation-dependent manner at the site of neurotransmitter release (Greengard et al., 1993; Schikorski and Stevens, 1997).

According to their release kinetics, SVs can be assigned to three different subgroups (Harata et al., 2001; Li et al., 2005; Rizzoli and Betz, 2005; Schikorski and Stevens, 2001). Vesicles of the readily releasable pool (RRP, about 5% of total population) are those spatially docked to the site of release and fuse immediately upon stimulation. Those vesicles recruited to the release site upon moderate physiological stimulation comprise the recycling pool (about 45% of total population), which is refilled by newly endocytosed vesicles after stimulation. The rest of vesicles comprise the resting pool, as those vesicles do not seem to cycle even upon exhaustive stimulation. Potential physiological functions of this pool are proposed to control presynaptic homeostasis during neuronal development (Andreae et al., 2012) and maintain high concentration of soluble proteins in the nerve terminals (Denker et al., 2011). These three vesicle pools are not anatomically segregated and mobilization of vesicles between different pools is possible (Rizzoli and Betz, 2005). Moreover, a recent study suggested that the size of the recycling and resting vesicle pools might be dynamically regulated by cyclin-dependent kinase 5 (CDK5) and calcineurin

- 6 -
(CN) activity, providing a potent control system for tuning synaptic performance (Kim and Ryan, 2010).

**Figure 1.1 Synaptic Vesicle**

(A) Molecular model of an average SV. Taken from (Takamori et al., 2006)

(B) Electron microscopic image of a chemical synapse on rat hippocampal neurons.
1.2. **Neurotransmitter Release**

**SNARE proteins and neuroexocytosis.** Upon arrival of an electrical impulse (action potential, AP) at the presynapse, membrane depolarization induces the opening of voltage-gated calcium channels (VGCC) that are clustered at specialized regions, called active zones (AZs). In the vicinity of these open channels localized high calcium concentrations (so-called Ca$^{2+}$ nanodomains) are created and trigger the fusion of docked and primed SVs with plasma membrane, releasing their chemical content into the synaptic cleft (*Rizo and Rosenmund, 2008*).

SV fusion during neuroexocytosis is the fastest form of intracellular membrane fusion. A key element of this process is the formation of a complex between components in the vesicular and target plasma membranes, the so-called SNARE (soluble NSF (N-ethylmaleimide-sensitive factor) attachment protein receptor) proteins (Figure 1.2). Once the complex is formed, its conformational change provides the driving force required for the fusion of the two membranes (*Sudhof and Rothman, 2009*). As suggested in several recent studies only few SNARE complexes are sufficient to accomplish vesicle fusion (*Mohrmann et al., 2010; Sinha et al., 2011; van den Bogaart et al., 2010*). However, to release neurotransmitter in a spatial and temporally defined manner, fusion of vesicles needs to be tightly controlled by the development of Ca$^{2+}$ nanodomains that are sensed by the vesicle associated protein synaptotagmin 1 (Syt1) through its Ca$^{2+}$-binding C2 domains. Indeed, it was found that neurons lacking Syt1 display complete absence of fast synchronized release (*Geppert et al., 1994*). Despite its important role in Ca$^{2+}$-triggered exocytosis, it still remains controversial whether Syt1 acts on SNARE complexes physically (*Dai et al., 2007; Lai et al., 2011*) or just serves as a distance regulator of SNARE complex formation (*van den Bogaart et al., 2011*).
Figure 1.2 SNARE Complexes and Neuroexocytosis

(A) The zippering model for SNARE-catalyzed membrane fusion. Assembly proceeds progressively from the membrane-distal N termini toward the membrane-proximal C terminal of the SNAREs. This generates an inward force that pulls the bilayers together, forcing them to fuse.

(B) When fusion has occurred, complete zippering turns the SNAREs in the low-energy cis-conformation. Taken from (Sudhof and Rothman, 2009).

Multiple roles of calcium ions in the regulation of neurotransmitter release. The intracellular Ca$^{2+}$ concentration not only has an important role in the triggering of neurotransmitter release but also in the regulation of short-term synaptic plasticity (STP) (Neher and Sakaba, 2008). A short burst of neurotransmitter release is initiated by high Ca$^{2+}$ concentration within nanodomains near VGCCs and can be electrophysiologically registered as a single peak during recording of excitatory postsynaptic current (EPSC). During repetitive presynaptic activities, however, the measured EPSC amplitudes drop progressively and reach a steady-state level within 10-20 stimuli, indicating a reduced rate
of neurotransmitter release. This phenomenon is known as short-term synaptic depression (STD) and is expected to modulate information processing in neuronal networks of the central nervous system by influencing the synaptic output during repetitive activity. A candidate mechanism for the induction of STD is depletion of vesicles in the RRP and the following steady-state is characterized by a balance of SV consumption and repopulation in this pool from a more distant reserve pool. During sustained synaptic activity, however, the recruitment of release-ready vesicles can be accelerated with a global rise in calcium concentration (Wang and Kaczmarek, 1998) and controls the level of STD as well as the recovery of the vesicle pool in periods of rest in an activity-dependent manner (Neher and Sakaba, 2008). Several calcium-binding proteins including synaptotagmin 2 (Young and Neher, 2009) and calmodulin (Sakaba and Neher, 2001) have been identified as key players in calcium-dependent vesicle replenishment by affecting important steps like vesicle docking and priming near VGCCs. In addition, it was reported recently that the intracellular Ca\textsuperscript{2+} level during stimulation also regulates the recycling pool size in the presynaptic bouton of rat hippocampal neuron (Li et al., 2011), suggesting an important role of calcium ions in dynamic regulation of synaptic strength.

**Visualization of vesicle fusion using a pH-reporter.** Recently, pH-sensitive GFP (pHluorin) has been used to visualize secretion at presynaptic boutons (Miesenbock et al., 1998). When tagged to the luminal domain of SV associated proteins like SV SNARE synaptobrevin 2 (Syb 2) or vesicular glutamate transporter (vGlut), pHluorin can be successfully introduced to the SV lumen upon overexpression in neurons. Inside the SV lumen, with a typical pH of 5.6, the fluorescence of spH molecules is almost completely quenched. Upon exposure to the extracellular environment and a neutral pH of 7.4 during exocytosis, each spH molecule gains a ~20-fold increase in fluorescence (Figure 1.3).
Using these pH-dependent changes in fluorescence intensities, it is possible to measure stimulation-dependent exo- and endocytosis in real time in presynaptic boutons of neurons (Sankaranarayanan et al., 2000; Sankaranarayanan and Ryan, 2000). In the past ten years, these genetically encoded reporters have been extensively used to study kinetics of vesicle recycling (Balaji and Ryan, 2007; Royle et al., 2008).

**Figure 1.3 Imaging SV Recycling with pHluorin-based Probes**

The pHluorin is quenched by the low pH inside the vesicle lumen. Upon stimulation (black arrow), vesicle fuses with the membrane and deprotonation switches the pHluorin in to its fluorescent form (upward deflection, in the trace, black). The fluorescence drops to the base line following endocytosis and vesicle reacidification. Taken from (Royle et al., 2008).
1.3. Synaptic Vesicle Recycling

**Multiple vesicle retrieval pathways.** After membrane fusion, compensatory endocytosis is required for cell to maintain constant surface area by specifically retrieving membrane and proteins inserted by calcium-triggered exocytosis (Dittman and Ryan, 2009; Murthy and De Camilli, 2003). The mechanism of the retrieval is comparatively more controversial than that of the fusion, although it occurs at longer time scales and can be more easily studied. Evidence from electron micrographs reveal that newly formed vesicles are mainly found at specialized region called endocytic site, which surrounds the presynaptic AZ (Heuser and Reese, 1973; Miller and Heuser, 1984). After having been refilled with neurotransmitter, those vesicles are ready for a next round of exocytosis. Multiple vesicle retrieval pathways might coexist with different characteristic time constants (Gandhi and Stevens, 2003; Smith et al., 2008). In total, three different recycling pathways have been proposed (Figure 1.4). A fast mode called ‘kiss and run’ describes neurotransmitter release via a narrow fusion pore during a brief connection between the SV and the plasma membrane. Rapid closure of the pore enables the vesicle being retrieved intact within one second (Aravanis et al., 2003; Klingauf et al., 1998; Pyle and Pelletier, 2003; Zhang et al., 2009). The more classical model proposed by Heuser and Reese involves the full fusion and collapse of the SV followed by clathrin-mediated endocytosis (CME) and is thought to be the major pathway for SV recycling. These coated vesicles are derived from invaginations of the plasma membrane which later bud off, forming a nascent SV for successive rounds of exocytosis (Granseth et al., 2006; Heuser and Reese, 1973). According to this mechanism, SV collapse and retrieval occur with a time constant of about tens of seconds. A third pathway, called bulk endocytosis, is observed during heavy stimulation when large membrane infoldings appear which bud off vesicles via CME (Koenig and Ikeda, 1996; Takei et al., 1996). Although having been intensively studied
for many years, some aspects of this topic still remain unclear, including whether different pathways produce functionally different pools of vesicles and what role the different modes may have on neurotransmission (Kavalali, 2007; Voglmaier and Edwards, 2007). Furthermore, a recent study in the mouse motor nerve terminals suggests a change in the location of preferred endocytic sites upon different stimulation strengths, suggesting a possible stimulus-dependent modulation of retrieval pathways (Gaffield et al., 2009).

Figure 1.4 Different Modes of SV Recycling

Schematic illustration of different modes of SV recycling. A fast 'kiss-and-run' mechanism, where the vesicle connects only briefly to the plasma membrane without full collapse ("kiss & run"), a slow pathway via large infoldings and endosomes ("bulk endocytosis"), and recovery of vesicle membrane by clathrin-coated pits (clathrin-mediated endocytosis, "CME"). Taken from (Haucke and Klingauf, 2006).
Modulation of endocytosis: calcium and exocytic load. Quantitative estimates of membrane retrieval after exocytosis provide information about roles of modulated endocytic pathways. Two different techniques are widely used. Whole-cell capacitance measurements, in which real-time changes of cell surface area are monitored by measuring the membrane capacitance with great temporal resolution through patch-clamp pipettes, are well established in studies of large nerve terminals like Calyx of Held (Augustine and Neher, 1992; Sun and Wu, 2001; von Gersdorff and Matthews, 1994). The second technique involves fluorescent optical tracers at small synapses and enables us mapping in a large scale the key feature of membrane or vesicle protein uptake (Ryan et al., 1993; Sankaranarayanan and Ryan, 2000). The appearance of distinct kinetic components observed from both techniques has been interpreted as participation of different retrieval modes. Interestingly, aside from the importance in signaling, the cytosolic calcium level at nerve terminal seems to have a role in regulation of endocytosis as well. In most types of synapses in the central nervous system, intracellular calcium elevation leads to accelerated retrieval of vesicular proteins (Sankaranarayanan and Ryan, 2001; Wu et al., 2005). However, an inhibitory effect of calcium on endocytosis has also been uncovered in the retinal bipolar cell terminal (von Gersdorff and Matthews, 1994). However the functional role of such inhibition under physiological conditions is still unclear. In addition, the progressive decline of the rate of endocytosis has been consistently observed after prolonged electrical stimulation in hippocampal synaptic boutons, despite the fact that endocytosis was initially accelerated under conditions that led to elevated intracellular calcium concentration. Thus, it has been suggested that this slowing of endocytosis with the accumulation of vesicle components may result from saturation of the total endocytic capacity of the synapse (Balaji et al., 2008; Sankaranarayanan and Ryan, 2001).
**Coupling of exo- and endocytosis.** The spatial and temporal coupling of exocytic membrane fusion and compensatory retrieval is a crucial and central element in the maintenance of synaptic transmission in the central nervous system during sustained synaptic activity. Without compensatory retrieval, continuous fusion of SVs with the plasma membrane would lead to a fast depletion of the SV pool and expansion of the presynaptic membrane. Evidence for such functional coupling includes the dual function of many SV proteins (e.g. Syb 2, Syt 1 or synaptophysin) that were originally considered to play a role exclusively in exocytosis, but have been found to have an additional role in compensatory endocytosis (Daly et al., 2000; Deak et al., 2004; Kwon and Chapman, 2011; Nicholson-Tomishima and Ryan, 2004; Poskanzer et al., 2003). Additionally, interfering with the function of endocytic proteins (e.g. dynamin) leads to a strong depression of exocytosis (Hosoi et al., 2009; Kawasaki et al., 2000). The local intracellular calcium concentration, which rises and falls rapidly during an AP, seems to serve as a central regulatory factor in these processes. During stimulation, high concentrations of calcium not only trigger exocytosis, but also facilitate endocytosis (Sankaranarayanan and Ryan, 2001). Recently, the SV associated transmembrane protein known as ‘flower’ (a proposed calcium channel) was shown to regulate SV endocytosis, and thus, potentially controls exo-endocytic coupling (Yao et al., 2009). These data imply a crucial role of vesicular proteins in regulating synaptic plasticity by shifting the balance between exo- and endocytosis during sustained synaptic activities. However, the underlying mechanisms remain controversial, probably due to the complexity emerging from the involvement of the multiple recycling pathways discussed above.
1.4. **Readily Retrievable Surface Pool**

An unexpected finding from studies using the spH assay (see above) was that up to 30% of overexpressed probes are localized on the plasma membrane even under resting conditions (Fernandez-Alfonso et al., 2006; Wienisch and Klingauf, 2006). A minor surface expression of synaptophysin-pHluorin (~ 8%) (Granseth et al., 2006) and vGlut1-pHluorin (~ 2%) (Balaji and Ryan, 2007) was also reported recently. This implies the existence of a pool of vesicular proteins stranded on the membrane surface and raises questions about the reason why these vesicular proteins are excluded during compensatory endocytosis. To explore the function of these surface-stranded SV proteins, their pHluorin-tags were selectively silenced through either proteolysis or photo-bleaching, while those in the vesicle lumens remained intact. Both manipulations resulted in a normal exocytosis-driven fluorescence increase followed by an impaired fluorescence decay, suggesting that newly exocytosed vesicular proteins exchange with surface-stranded ones during brief stimulation (Wienisch and Klingauf, 2006). Based on these observations, it was speculated that these membrane-stranded vesicular proteins may support the first wave of compensatory endocytosis and thereby participate actively in exo-endocytic cycling. Further evidence for a functional pool of preassembled SV components at the presynaptic membrane came from experiments using botulinum toxin (BoNT/A), which completely abolishes exocytosis by cleaving the membrane SNARE SNAP-25, but did not block endocytosis completely (Neale et al., 1999). This clearly suggests an exocytosis-independent retrieval of SV components from a pre-existing pool on the presynaptic membrane. If such a ‘readily retrieval pool’ (RRetP) of vesicle components exists, it may serve as a coupling factor between exo- and endocytosis through dynamic control of the pool size. Thus, it is crucial to optically resolve this functional pool and understand how it may couple exo- and endocytosis in a temporal and spatial defined manner.
1.5. **Release Site Clearance**

It is well established that in many types of synapses the level of neurotransmitter release during ongoing activity is mainly limited by the availability of release-ready vesicles and, therefore, depends on the speed of vesicle recruitment. Surprisingly, many synapses in the central nervous system can sustain synaptic activity even upon high frequency stimulation (Kopp-Scheinpflug et al., 2008; Lorteije et al., 2009; Rancz et al., 2007). It is evident that vesicles are required to undergo fast recycling to prevent depletion of the vesicle pool (Fernandez-Alfonso and Ryan, 2004; Sudhof, 2004). It has been reported in several recent studies that interfering with the function of endocytic proteins causes a fast, stimulus-frequency-dependent depression of vesicle exocytosis (Hosoi et al., 2009; Kawasaki et al., 2000). Some of these inhibitory effects developed so rapidly that they could not simply be explained by depletion of recycled vesicles. Therefore, it has been postulated that the respective manipulations affect a step in between exo- and endocytosis (Haucke et al., 2011; Neher, 2010). Translocation of SV components from sites of release to the endocytic sites may be a most likely candidate for this step. Acute accumulation of newly exocytosed vesicular proteins may cause a functional block of the previously used release site by interfering with the docking of subsequent vesicles. Knowing how these newly exocytosed vesicular proteins are removed from the release site may allow us to better understand the interplay between exo- and endocytosis and to gain a more accurate picture of neurotransmitter release.
1.6. **Aims of This Study**

In this study I explored the usability of the genetically encoded pH-sensitive fluorescence probe spH ([Miesenbock et al., 1998; Sankaranarayanan et al., 2000](#)) as well as a newly developed exogenous pH-indicator ([Hua et al., 2010; Hua et al., 2011](#)) to visualize exo- and endocytosis in cultured rat hippocampal neurons. Using these probes, I tried to address several unresolved questions concerning SV recycling. First: Is there a pre-existing pool (so-called RRetP) of endogenous vesicle components on the membrane surface and what is its location relative to the presynaptic AZ? Second: Are endogenous vesicular proteins also preferentially retrieved from this surface pool during compensatory endocytosis as implied by studies using overexpressed probes and is the retrieval from such putative RRetP kinetically faster? If the RRetP has only a limited capacity for fast endocytosis, exhaustion of this pool will lead to a reduction of the endocytic rate and thereby should cause massive protein accumulation on the membrane surface. Will this eventually result in functional block of the previously used release site and become the rate-limiting step during sustained synaptic activities?
2. Material and Methods

Cell Culture. Primary neuronal cultures (~ 5000 – 7500 cells per cover slip) were prepared according to the revised protocol established by C. Rosenmund with minor modifications (Rosenmund and Stevens, 1996). Hippocampal CA1-CA3 regions were dissected from one-day-old Wistar rats under the microscope and collected in a tissue dish filled with Hank’s Balanced Salt Solution (HBSS, Invitrogen) on ice. This tissue was then transferred into a falcon tube and the following procedures in preparation were performed under sterile conditions. After being bubbled with carbogen gas for 10-30 minutes, 1 ml digestion solution containing 20-25 units Papain was added to the tissue culture. After 1-hour incubation at 37 °C, the tissue was then transferred to inhibition solution and incubated for 5 minutes. The dispersion of the tissue into individual neurons was completed in required amount of serum-free Neurobasal A Medium (NBA, Invitrogen) by triturating with Pasterur pipette. The suspension was diluted with primary medium of required volume. Finally the neurons were plated onto 18 mm cover slips (Assistant, REF 1001/18; 0.13-0.16 mm) coated with Matrigel (Beckton-Dickinson, Heidelberg, Germany) and placed in a 12 well plate (BD 35 30 43). Neuronal cultures were kept in an incubator (Heraeus BBD 6620) at 37 °C and 5 % CO₂ before use (DIV 14-28).

Transfection of spH. Cultured neurons were transfected at 3 DIV by a modified calcium phosphate transfection procedure (Wienisch and Klingauf, 2006). First NBA solution was prewarmed 30-minute prior to the transfection. Calcium phosphate/DNA precipitates were then formed by mixing 5–30 μg/ml plasmid DNA, diluted in sterile double distilled water to a final concentration of 0.2 mg/ml, with 5 μl of 2.5 M CaCl₂ stock solution and 50 μl of 2× BBS-buffered saline (50 mM BBS (N, N-bis[2-hydroxyethyl]-2-aminoethanesulfonic
acid), 280 mM NaCl, 1.5 mM Na₂HPO₄, pH 7.05) kept at room temperature in darkness for 20 minutes. Mixture containing the precipitates was added drop-wise to 900 μl NBA solution while gently vortexing. Neuronal cultures were taken out and moved to new wells, to each of which 500 μl of the diluted calcium phosphate/DNA precipitate solution was added. After 20-minute incubation at 37°C and 5 % CO₂, a fine sandy precipitate covering the cells was observed. The cultures were then washed with prewarmed HBSS and finally replaced back to the original growth media. The efficiency was typically between 5 and 20 %, and there was no apparent toxicity to the cells. Imaging was performed at about 21 DIV.

*Live Cell Immunostaining.* In this study both cypHer5E-conjugated antibodies (mouse monoclonal αSyt1-cypHer5E: 105 311cpH; rabbit polyclonal αVGAT-cypHer5E: 131 103cpH) were purchased from Synaptic Systems (Göttingen, Germany). Neurons were incubated in cell culture incubator for 3 hours with the staining buffer (1:200) containing 120 mM NaCl, 5 mM KCl, 2 mM MgCl₂, 2 mM CaCl₂, 10 mM glucose and 18 mM NaHCO₃. Before use, the stained neuronal culture was washed twice with antibody-free medium. For pH titration measurements of cypHer, αSyt1-cypHer5E-stained neuronal culture was fixed with 4 % paraformaldehyde (PFA) in phosphate buffered saline (PBS) at room temperature. 0.4 % Saponin was used to permeabilize the cell membranes, making the internalized antibodies accessible to perfused buffers of different pH. The selective staining of the surface-residual SV proteins was performed on the microscope stage by incubating the neurons with αSyt1-cypHer5E (1:50) in the presence of 1 μM Tetrodotoxin (TTX) at room temperature for 5 minutes. The neurons within the region of interest were then constantly washed with Tyrode solution through perfusion capillary during the recording.
**Optical Measurement, Microscopy and Analysis.** Coverslips were placed in a perfusion chamber containing a modified Tyrode solution (in mM: 120 NaCl, 2.5 KCl, 2 CaCl$_2$, 2 MgCl$_2$, 10 Glucose, 10 HEPES, pH was adjusted to 7.4 by NaOH). To stimulate neurons, an electric field stimulation were delivered via two platinum electrodes spaced at 10 mm with 1 ms pulses of 50 mA and alternating polarity applied by a constant current stimulus isolator (WPI A 385, World Precision Instruments); 10 µM 6-cyano-7-nitroquinoxaline-2,3-dione (CNQX) and 50 µM DL-2-amino-5-phosphonovaleric acid (AP5) were added to prevent recurrent activity. Fast solution exchanges were achieved through a two-barrel glass tubing perfusion system controlled by a piezo-controlled stepper device (SF778, Warner Instruments). Fluorophores were excited at 488 nm (for spH) or 640 nm (for cypHer) with a computer-controlled monochromator (Polychrom IV, Till Photonics) and time lapse images were acquired using an electron-multiplying CCD camera (Andor iXon+ DU-897E-BV camera; Andor Technology) which was mounted on an inverted Nikon TE2000 microscope equipped with a ×60, 1.2 numerical aperture water-immersion objective and an FITC/Cy5 dual-band high performance filter set (AHF analysentechnik AG, Tübingen). Images were analyzed using self-written program in Matlab (MathWorks). Active boutons were localized using an automated routine in which a difference image is generated by subtracting an average of five frames before from after stimulation. Square regions of interest (1.5 × 1.5 µm$^2$) were centered on these spots and average fluorescence profiles were then plotted. Fluorescence traces from individual synapses were normalized before calculating the average response.

**Theoretical Basis for pH-based Optical Signals.** By given pH, indicators are in the equilibrium with free protons. Therefore instantaneous fluorescence intensity reflects the proportion of protonated molecules, which can be calculated by the following equation,
where pK is the negative logarithm of the equilibrium constant for protonation and [$\text{Ind}^-$] for the fraction of protonated molecules. During the exo-endocytic cycling, the change in the fraction of protonated molecules can be quantified by the following expression, due to the exposure of intra-vesicular molecules to extracellular environment or the reacidification of freshly retrieved vesicles.

$$
\Delta[\text{Ind}^-] = \frac{1}{1+10^{7.4-pK}} - \frac{1}{1+10^{5.6-pK}}
$$

**Photobleaching Correction for cypHer Signals.** As cypHer molecule displays much less photo-stability compared to its prototype (Cyanine 5 dye), it is required to correct the intensity of time-lapse images properly for photo-bleaching. Considering that the cypHer molecules would be quenched and temporarily protected from photo-bleaching, while being exposed to extracellular medium, the kinetics of exo-endocytic cycling could be successfully isolated using a memorial deconvolution algorithm. It is assumed that the intensity decline due to photo-bleaching can be described as an exponential decay (see below), where $I(t)$ is the instantaneous observation of intensity. Practically, the decay constant $\lambda$ is able to be obtained from observations of intensity-time courses in blank experiments, using the same exposure time and acquisition frequency, which indeed showed monoexponential bleaching characteristics.

$$
\frac{dI(t)}{dt} = -\lambda I(t)
$$

Therefore, the instantaneous intensity change caused by exo-endocytosis can be determined iteratively by the following equation, where is the instantaneous intensity observed at interval $t-1$. An example of the correction is shown in Figure 2.1.
Deconvolution Analysis. The normalized fluorescence decay of calibration response was used as the template for the deconvolution. To minimize the noise, the template was simplified to a single exponential decay function obtained from fitting. The deconvolution was then performed by dividing the template from the normalized fluorescence time course in the Fourier space using Matlab built-in function.

\[
I(t) = I(0) - \sum_{i=0}^{t-1} \lambda I(i) + \Delta I(t)
\]
Figure 2.1 Photobleaching Correction for cypHer Signals

(A) During exo-endocytosis the fluorescence of cypHer is quenched and dequenched by pH-change of the surrounding, which led to different photobleaching rates of different phases. An iterative correction is required to isolate the pure pH-dependent fluorescence change. Average fluorescence time course of neurons stimulated by 600 APs was obtained by averaging the original data after normalization (blue). Standard bleaching curve (green circles) can be fit well to an exponential decay function (red line). The red dotted line indicates the working curve for the correction.

(B) Average cypHer fluorescence time course in (A) is plotted after correcting for photobleaching. Taken from (Hua et al., 2011).

Dual-color STED Microscopy. Dual-color super-resolution images were recorded with a custom-built confocal microscope using laser stimulated emission depletion (STED) technique. The setup combines two pairs of excitation and depleting laser beams all derived from a single supercontinuum laser source as described in (Liu et al., 2011). STED images were processed using a linear deconvolution function integrated into Imspector Software. The point spread function (PSF) for deconvolution was generated by using a 2D Lorentz function with its half-width and half-length fitted to the half-width and
half-length of each individual image. Both Syt1 antibodies (mouse monoclonal, 604.2, Synaptic Systems, Göttingen, Germany) were directly labeled with either ATTO 647N or ATTO 590 and diluted (1:200) with Tyrode solution before use. For selective surface pool staining, a pre-incubation of neuron culture with 1 μM TTX was required to effectively suppress spontaneous network activities. A detailed procedure of staining was described in the main text. After having been stained, sample was washed twice and fixed with 4 % PFA at room temperature for 15 minutes and then mounted on a glass slice and transferred to the microscope stage for image acquisition. The colocalization of two Syt1 patches was quantitatively analyzed using Pearson correlation coefficients between signals from two channels after background subtraction.
3. Results

3.1. Surface Retrieval of SV Proteins

To provide direct evidence for the stimulation-locked retrieval of surface-stranded vesicular proteins, a new assay was developed in our lab where the luminal domains of vesicular proteins are tagged with an exogenous pH-indicator cypHer 5E (referred to here as cypHer) through antibodies (Adie et al., 2003; Martens et al., 2008). Protonation of this fluorophore can cause a five-fold increase in its quantum yield, internalization of labeled vesicular proteins will thereby lead to an increase in the fluorescence during reacidification inside the vesicle. To test this idea, antibodies against the luminal domain of synaptotagmin 1 (Syt1), and vesicular GABA transporter (VGAT) were labeled with cypHer and each separately applied to the neurons in culture. In both cases large accumulation of cypHer molecules in synaptic boutons were confirmed by a punctate staining due to dequenching of the fluorophores in the SV lumen (Figure 3.1).

Next, we attempted to monitor dynamics of surface residual vesicular proteins (This work was a collaboration with Raunak Sinha and has been published (Hua et al., 2011)). Brief incubation of the neuron culture in a saturating concentration of αSyt1-cypHer was conduct to selectively label the surface pool of vesicular proteins with limited dye internalization. 1 µM tetrodotoxin (TTX) was added to the staining buffer to further reduce intermixing between the surface residual protein population and vesicular population through spontaneous activities by suppressing neural network activity (Groemer and Klingauf, 2007; Hua et al., 2010). When stimulated with 50 APs at 20 Hz after washout of TTX, the stained synaptic boutons responded with an increased fluorescence which gradually reached a plateau with a characteristic time constant, clearly
Figure 3.1 cypHer-coupled Antibodies Are a Reliable Tool to Measure Exo-endocytic Cycling.

(A) Average normalized (norm.) fluorescence of anti-Syt1-cypHer-stained hippocampal boutons from multiple trials (each trial consisted of > 30 boutons) as a function of pH. The solid curve is a fit to a Henderson-Hasselbalch equation, yielding a pKa of 7.05. Error bars represent s.d.

(B) Schematic depicting fluorescence changes of cypHer during exo-endocytic cycling. The red dots show cypHer-coupled antibodies, which bind to the extracellular domain of SV proteins and remain quenched owing to the neutral extracellular pH. During endocytosis and reacidification, however, the cypHer fluorescence is dequenched. Re-exocytosis upon stimulations results in quenching of the cypHer fluorescence at the extracellular neutral pH.

(C) Representative fluorescence image of hippocampal neurons labeled with anti-VGAT-cypHer. The label forms a punctate pattern with puncta typically representing individual boutons. Taken from (Hua et al., 2011)

indicating that stimulation triggers the retrieval of the RRetP to form fresh SVs (Figure 3.2 A). To further examine whether those SV recruited from the RRetP are functional and participate in vesicle cycling during repeated stimulation, a second round of stimulation was applied to surface Syt1-labeled neurons 60 seconds later. The fluorescence responses indeed now showed a fast drop in intensity due to partial exocytosis of previously recycled RRetP components again followed by an increase indicating endocytosis. The
released fractions of SVs originating from the RRetP were 27.6 ± 5.1% for 50 APs and 49.7 ± 9.2% for 200 APs (Figure 3.2 B), which is in close agreement with similar experiments using FM styryl dyes as tracers (Vanden Berghe and Klingauf, 2006). However, unexpectedly, the fluorescence increase due to endocytosis after the second stimulus exceeded that of the exocytic quenching. This could be due to incomplete depletion of the RRetP during the first stimulus of 50 APs, but is more likely to be due to partial replenishment of the RRetP after the first stimulus by exocytoosed vesicles that had taken up cypHer by spontaneous recycling during the initial labeling phase (Groemer and Klingauf, 2007; Hua et al., 2010).

Figure 3.2 Readily Retrievable Surface Pool of SV Components.

(A) Average fluorescence profile of surface Syt1-labeled bouton (n = 3 experiments with > 50 boutons for each) stimulated first with 50 APs at 20 Hz followed by a second stimulation with either 50 (black) or 200 (red) APs at 20 Hz. Error bar represent s.e.m.

(B) In response to the second stimulus, a marked fraction of SV recycled from the RRetP upon the first stimulation is released (27.6 ± 5.1% for 50 APs and 49.7 ± 9.2% for 200 Aps). Taken from (Hua et al., 2011)
3.2. **Exo-endocytosis Cycling of Endogenous SV Proteins**

Next, to clarify the question of whether fluorescence-labeled antibodies report same recycling pathway as spH, neuronal cultures were incubated with the cypHer-conjugated antibodies for three hours to label recycling vesicle during spontaneous neural activity so that recycling kinetics could be measured as fluorescence changes of cypHer upon electrical stimulation. For comparison, the staining was repeated in neurons transfected with spH. The wide spectral separation enables dual-color imaging of spH and cypHer without bleed-through (Figure 3.3 A). As the spH transfection efficiency is only ~ 10 %, not all αSyt1-cypHer-stained boutons express spH. Therefore, automatic detection of active boutons was performed using spH responses and the created mask was then applied to cypHer channel. After having examined the fluorescence changes in response to 200 APs at 20 Hz in both channels, it was found that spH and cypHer responses of the same boutons showed mirror-image signals with near-identical endocytic time constants for monoexponential fits (~ 23 seconds versus ~ 22 seconds). This indicates that overexpressed fusion proteins reliably report exo-endocytosis and do not artificially affect SV recycling.

In order to investigate the recycling kinetics of the inhibitory synapses, spH-transfected boutons were stained with αVGAT-cypHer (Figure 3.3 B). A ~ 32 % fraction of all spH-expressing boutons could be identified with αVGAT-cypHer, representing the fraction of inhibitory synapses in the cultures. Stimulation with 200 APs at 20 Hz induced similar fluorescence transients (~ 22 seconds endocytic time constant) in both channels. Therefore, the kinetics of endocytosis and reacidification in inhibitory synapses are indistinguishable from the whole population response as probed with spH and cypHer antibodies.
Figure 3.3 Comparison of Vesicle Recycling Kinetics Probed with spH and cypHer-coupled Antibodies.

(A) Top: staining of spH-transfected hippocampal neurons with anti-Syt1-cypHer. Fluorescence images show colocalization of individual boutons expressing spH (green) with endogenous Syt1 (red) labeled with anti-Syt1-cypHer. Bottom: average fluorescence transients of double-labeled boutons in response to 200 APs at 20 Hz (n = 5 experiments with >50 boutons for each). Traces were corrected for photobleaching. Error bars represent s.e.m.

(B) Top: staining of spH-transfected hippocampal neurons with anti-VGAT-cypHer. Fluorescence images show colocalization of individual boutons expressing spH (green) with the inhibitory synapse marker VGAT (red). Bottom: average fluorescence transients of double-labeled boutons in response to 200 APs at 20 Hz (n = 5 experiments with >50 boutons for each). Traces were corrected for photobleaching. Error bars represent s.e.m. Scale bars represent 5 µm. Taken from (Hua et al., 2011)
3.3. **Spatial Organization of the Readily Retrievable Pool**

To address the question how is the functional surface pool spatially organized at the hippocampal synaptic bouton, we used the same antibody, αSyt1, to specifically label the surface pool, at 4 °C to minimize spontaneous activities (This work was a collaboration with Cora Thiel & Roman Schmidt and has been published (Hua et al., 2011)). We fixed the cells immediately and identified synapses by colabeling with antibodies to bona fide cytoplasmic matrix of AZ (CAZ) or postsynaptic density (PSD) proteins. The spatial organization of the Syt1 surface pool was investigated using dual-color isoSTED nanoscopy (Schmidt et al., 2008). In the three-dimensional reconstruction of hippocampal en passant boutons from isoSTED image stacks (Figure 3.4), distinct Syt1 patches were found in the vicinity of the AZ.

From a large list of antibodies to CAZ elements, we obtained good signal-to-noise ratio only with antibodies for RIM1 and RIM2. The observed antibody fluorescence distribution was broad, extending up to 1 μm along its long axis. AZ sizes of up to 700 nm have been documented in hippocampal synapses in electron microscopy of single sections (Schikorski and Stevens, 2001), as well as in recent full three-dimensional reconstructions (Staras et al., 2010). The larger patches measured in our experiment indicate that AZ sizes are indeed overestimated by labeling the large RIM proteins at the N-terminal zinc-finger domain.

To provide an additional fingerprint of synapses, we modified the staining by targeting the PSD opposing the AZ with antibodies to Homer1. This introduced a characteristic spatial separation between the two channels. We analyzed the distribution of the Syt1 patches with respect to Homer1 in each of the three bouton regions, as depicted in the histograms. The mean distance between Syt1 and Homer1 increased as the azimuth angle α between
the estimated orientation of the axon and the Syt1–Homer1 connecting vectors approaches 0 or ± π, owing to a higher probability of Syt1 patches localized adjacent to the axolemma but not associated with Homer1. At other angles, the observed distances are smaller and reach minimum distances of about 100 – 300 nm for a bouton. This finding agrees well with a localization analysis of a large number of synapses in 4Pi microscopy (Huve et al., 2008; Kano et al., 2001). Considering this result together with our live cell imaging data, we therefore conclude for that the associated Syt1 patches constitute the preassembled RRetP.
Figure 3.4 Spatial Organization of the RRetP

(A) Z-projection of a three-dimensional image stack of Syt1 at synapses labeled by the CAZ markers RIM1 and RIM2.

(B) Z-projection of Syt1 at synapses labeled by the PSD marker Homer1, with synapses at regions of interest numbered 1 and 3. Arrows mark the local orientation of the axon.

(C) Perspective views of volume-rendered data. The Syt1 labeling reveals preassembled patches that are localized in the periphery of the AZ, typically within 500 nm distance from the postsynaptic marker.

(D) Spatial distribution of Syt1 with respect to Homer1 for each region of interest. Histograms show Syt1-Homer1 cross correlation, binned according to Syt1-Homer1 distance $d$ and relative azimuth angle $\alpha$, which was measured with respect to the orientation of the axon as marked by arrows in B. Distances between Syt1 patches and the postsynaptic marker accumulate at about 100-300 nm at angles where the probability of finding Syt1 patches that are not associated with Homer1 is reduced. Taken from (Hua et al., 2011)
3.4. Quantitative Measurement of Cumulative Release

Spatial separation of the AZ and the endocytic site implies lateral diffusion of newly exocytosed patches of vesicular proteins from the release site towards the rim of the AZ where they are eventually retrieved. Therefore, sustained synaptic activities may cause an acute accumulation of vesicular proteins around the release site. This may eventually lead to a functional blockade of the previously used AZ slot, thereby causing STD as recently postulated (Neher, 2010). However, no study, to my knowledge, has been conduct to clarify whether unavailability of the release sites can become the rate limiting step during sustained synaptic activities.

To reliably measure the level of depression, attempts were made to quantify the amount of released vesicles upon different stimulation strengths using well established pHluorin assay. Under the same experimental conditions the absolute amplitude of fluorescence change is proportional to the amount of spH molecules in exocytosed vesicles and can differ from cell to cell due to inhomogeneous expression of the probe and variation in release probability of the synaptic bouton. Therefore, normalization of the recorded responses is required for quantitative measurements of release. However, in previous studies normalization of the fluorescence responses were performed either to the vesicle pool size or simply to the base-line fluorescence, neither of which correlates strongly to the factors discussed above (Burrone et al., 2006; Hua et al., 2010; Newton et al., 2006; Sankaranarayanan et al., 2000). Even after such normalization, large variation in characteristic parameters of response remains, especially for small sample populations. To address this problem, an internal normalization reference was introduced by using a paired-stimulation protocol based on the fact that responses to brief stimuli remain stable over many trials when an interval of about 60 seconds is given for recovery after each stimulus (Burrone et al., 2006). In practice, a calibration stimulus which consisted of 50
APs fired at 20 Hz was employed, since such a stimulus is believed to trigger the fusion of only docked vesicles without being affected by any downstream processes after exocytosis (Murthy and Stevens, 1999; Schikorski and Stevens, 1997). After a 60 second recovery interval, synapses were then stimulated with different test protocols. Each fluorescence transient from a given region of interest was normalized to the amplitude of the calibration stimulus response. Since the release probability and the average number of spH molecules per vesicle are conserved at the given synapse, their contribution to the variation between responses is negligible.

Using this method, I examined the effect of increasing stimulation frequencies on vesicle release and found that a higher level of normalized peak fluorescence could be reached for a given number of stimuli, if stimulation firing rate was increased (Figure 3.5). An obvious explanation for this is ongoing endocytosis during stimulation, which counteracts protein accumulation at the plasma membrane and therefore reduces the peak value of spH fluorescence. In order to compensate the effect of endocytosis and further characterize the role of stimulation frequency on release rate, attempts were made to quantify release rate and cumulative release from normalized fluorescence responses. First I extracted information about the kinetics of endocytosis and reacidification by fitting an exponential function to the decay phase of each calibration response. The vesicle retrieval time constant has been reported recently to be a cell-wide property and remains constant upon multiple rounds of trials for individual cells (Armbruster and Ryan, 2011). Therefore, assuming that the measured signal is a linear sum of elementary events, the relative release rate could be calculated by deconvolution of the decay from the normalized fluorescence transient. An increased endocytic time constant was constantly observed, whenever the cell was stimulated by 200 APs at more than 10 Hz (Figure 3.5), in line with previous work (Balaji et al., 2008). A potential explanation for such a progressive decline
in endocytic rate is the exocytosis-driven protein accumulation on the surface that exceeds the buffering capacity of the RRetP. To test this idea, I explored the relationship between the stimulation strength versus the resulted endocytic time constant and found that at given stimulation frequency the endocytic rate remains constant until a given number of stimuli, beyond which the rate decreases rapidly (Figure 3.6). During low frequency stimulation, recovery of the RRetP may be fast enough to relieve the exocytic load and thereby ensure maximum speed of endocytosis for prolonged stimulation. This result confirms the existence of a limited surface capacity for fast membrane retrieval from the RRetP and suggests that the repopulation of this pool limits the rate of the endocytosis.

**Figure 3.5 Normalization of spH Response Using a Paired-stimulation Protocol**

(A) Average spH responses to 50 Aps at 20 Hz followed by 200 Aps at 5 (blue), 10 (cyan), 20 (green) or 40 Hz (red) with 60 s of recovery in between (n = 5 experiments with > 20 boutons for each). Each curve was normalized to its peak value of the first response, decay of which was fit with an exponential function. The time constant was 16.1 s. Error bars represent s.e.m.

(B) The second responses of fluorescence transients from (A) were fit with a single exponential decay function, respectively. The estimated endocytic time constant increases with stimulation frequency: 16.6 s for 5 Hz, 18.3 s for 10 Hz, 20.5s for 20 Hz and 23.1s for 40 Hz.
Figure 3.6 Color-coded Endocytic Time Constants of Synapses Challenged with Different Stimulation Protocols.

The fluorescence decay was fit by a single exponential function and obtained time constants were normalized to that of the calibration response evoked by 50 APs at 20 Hz. High stimulation rate and prolonged stimulation lead to increased endocytic time constants.

When the endocytic load does not exceed the buffering capacity of the RRetP, the endocytic rate reaches its maximum and remains invariant, ensuring reliability of the outcome from the deconvolution method. This analysis revealed stepwise increases in release extending throughout the period of stimulation, indicating a linear release rate increase with mild stimulation up to 200 APs at 5 Hz, in agreement with several previous studies using alkaline trapping (Ariel and Ryan, 2010; Li et al., 2005). Integrating the release rate time course provided a good estimation of cumulative release for quantitative comparison (Figure 3.7). The resulting cumulative release was proportional to the number of APs up to 200, implying the absence of release depression during mild stimulation.
Figure 3.7 Quantitative Measurement of Cumulative Release

(A) Representative spH responses to 50 APs at 20 Hz followed by 200 APs at 5 (blue), 100 at 10 Hz (cyan), 50 at 20 Hz (orange) or 50 Hz at 40 Hz (red) with 60 s of recovery in between. Each fluorescence transient from a given region of interest was normalized to the amplitude of the calibration stimulus response.

(B) Time courses of cumulative release calculated from (A). In normalized fluorescence transients of (A), the decay phase of each calibration response was fitted with an exponential function. The relative release rate could be calculated by deconvolution of the decay from the fluorescence transient. Integrating the release rate time course provided a good estimation of cumulative release for quantitative comparison.
3.5. **Limited Vesicle Reuse during Mild Stimulation**

Despite its narrow range of applicability, the deconvolution method has an advantage over other methods that either block compensatory endocytosis or prevent vesicular reacidification, in fact it directly measures the rate of exocytosis without any perturbations. When applicable, it provides a better estimate for exocytosis, since it takes into account the contributions of fast vesicle reuse. In fact, comparing the results of deconvolution with that of using e.g. Folimycin (see below) should allow one to estimate the contribution of vesicle reuse.

I next compared the deconvolved-integrated signal with that from Folimycin (V-ATPase inhibitor) and Dynasore (dynamin GTPase inhibitor) experiments and found that during 200 AP stimulation firing at 5 Hz only the signal amplitude under Dynasore treatment was ~ 9.2 % smaller than that obtained from deconvolution (Figure 3.8). This difference suggests a minor contribution of fast reused vesicles to synaptic transmission or incomplete block by the agents. In agreement with several previous studies (Betz and Bewick, 1993; Li et al., 2005; Wu and Wu, 2009), this finding argues strongly against a major contribution of reused vesicles to synaptic transmission within a 40 second stimulation period. We cannot exclude the possibility of a kiss-and-run type of recycling pathway acting in addition, since fast opening and closing of fusion pores may only cause a relatively small pH-fluctuation and therefore will not be fully reported by deconvolution (Zhang et al., 2009).
Figure 3.8 Same Amount of Vesicles Participating in Exocytosis during Mild Stimulation at 5 Hz

(A) The basic experimental design is schematically illustrated.

(B) Comparison of the estimated cumulative release using three different methods. Same experiment like in Figure 3.7 was repeated in the presence of either Dynasore (gray) or Folimycin (black). Comparison of the deconvolved-integrated signal with that from Folimycin and Dynasore experiments revealed that during 200 APs firing at 5 Hz the signal amplitude from Dynasore-treatment was only ~ 9.2% smaller than that obtained from deconvolution.
3.6. **Blocking Endocytosis Causes Rapid Release Depression**

As demonstrated above, synapses do not preferentially reuse freshly exocytosed vesicles during brief stimulation with low frequency (200 APs at 5 Hz). When stimulating at frequencies of 10-100 Hz, a reduction in synaptic response, known as STD, is observed in many types of glutamatergic synapses during electrophysiological recording. Upon sustained stimulation the rate of neurotransmitter release drops rapidly and reaches a steady state within 10-20 stimuli, reflecting a balance between vesicle usage and recruitment of new vesicles. To further examine the dynamics of vesicle cycling during stimulation with higher frequencies, I first repeated experiments using stimulation protocols similar to those of Figure 3.7 B, but with increased frequency and in the presence of 65 nM Folimycin (Fig 3.8 A). Fluorescence amplitudes were found to be similar except for a slight drop for the highest stimulation frequency at 40 Hz. The similarity of responses for 5, 10 and 20 Hz may suggest the absence of significant release depression and fast vesicle reuse. As sustained high release rate requires faster replenishment of vesicles in the RRP than that at rest, this result provides strong evidence for an activity-dependent replenishment mechanism to meet the needs of high release rate within a short period. This is not only in line with several previous studies (Dittman and Regehr, 1998; Stevens and Wesseling, 1998; Wang and Kaczmarek, 1998), but has also been further confirmed by the fact that a high-rate depleting pulse or increased extracellular Ca$^{2+}$ concentration can drive more vesicles to participate in the exocytosis by enlarging the recycling pool size (Suppl. 1).
Figure 3.9 Frequency-dependent Release Depression in the Presence of Dynasore.

(A) Average spH responses to 50 APs at 20 Hz followed by 200 APs at 5 (blue), 10 (red), 20 (green) or 40 Hz (cyan) in the presence of Folimycin. Each fluorescence transient from a given region of interest was normalized to the amplitude of the calibration stimulus response. (n = 6 experiments with each > 20 boutons for 5 and 40 Hz; n = 6 experiments for 10 and 20 Hz). Fluorescence increases were insensitive to the stimulus frequency. Error bars represent s.e.m.

(B) Same experiments like in (A) were repeated in the presence of Dynasore (n = 5 experiments with > 20 boutons for each). The amplitude of fluorescence responses dropped monotonically with increased AP-firing frequency. Error bars represent s.e.m.

Based on this finding I then tested whether STD is also enhanced after acute block of dynamin activity in primary neurons, as reported in the Calyx of Held (Hosoi et al., 2009). Indeed, when similar experiments were performed in the presence of 100 μM Dynasore, the amplitude of fluorescence responses dropped monotonically with increased AP-firing frequency (Figure 3.9). This suggests that release depression during high frequency stimulation after block of dynamin activity is not a unique feature of large nerve terminal like Calyx of Held, but maybe a common property of excitatory synapses. The same experiment was repeated using cypHer as the pH-indicator (Hua et al., 2011) and similar results confirmed that the observed effect is not due to genetic perturbation by using spH (Suppl. 2). To further confirm that the reduction in fluorescence amplitude was Dynasore-
specific, I examined spH response upon 200 APs at 20 Hz in the presence of both Dynasore and Folimycin (Figure 3.10) and found that addition of Folimycin could neither rescue nor intensify the release depression caused by Dynasore.

![Figure 3.10. Reacidification has no Effect on Dynasore-enhanced Release Depression.](image)

Average spH responses to 50 APs at 20 Hz followed by 200 APs at 40 Hz in the presence of Dynasore (red), Folimycin (gray) or both (black broken line). Between stimuli a 60 s interval was given for recovery. Each fluorescence transient was normalized to the amplitude of the calibration stimulus response. In the presence of both Folimycin and Dynasore resulted in same signal amplitude compared to that of only using Dynasore and about 38 % smaller than that of using Folimycin.
3.7. **Vesicle Recruitment during Multiple Spaced Stimuli**

When vesicles of the recycling pool are trapped in the alkaline state, mixing of these vesicles with the RRP will lead to a progressive reduction in the evoked fluorescence response, since fusion of alkaline-trapped vesicles does not contribute to the fluorescence increase. Such an effect was consistently observed during multiple rounds of short stimuli in the presence of a V-ATPase inhibitor (Li et al., 2005). A similar phenomenon was also reported in hippocampal cultures treated with Dynasore (Newton et al., 2006), which was believed to be the consequence of vesicle pool depletion during sustained activity in the absence of dynamin-dependent endocytosis. To test and verify this idea, I analyzed responses of synapses expressing spH to multiple rounds of a brief stimulus (50 APs at 20 Hz) at 60 second interval in the presence of either Dynasore, Folimycin or the delivery vehicle DMSO. Deconvolution was performed on control responses recorded with DMSO (Figure 3.11) and showed stepwise increases with identical amplitudes in cumulative release, ensuring the absence of major changes in synaptic function for at least 10 trials. When Folimycin was applied, fluorescence responses decreased gradually to ~ 43% of the initial response, indicating an increased percentage of alkaline-trapped vesicles in the RRP. In the presence of Dynasore, however, there are no vesicles to be reused due to block of endocytosis and all release sites are occupied by fusion-competent vesicles recruited from a larger recycling pool. Similar response amplitudes were thereby expected at least during the initial few trials. Surprisingly, an accelerated reduction in responses was observed compared to that in the presence of Folimycin. This finding presented us with a puzzle about the mechanism behind the strong release depression caused by impaired dynamin activity even during the initial few trials. It is not expected that a small consumption of vesicles by exocytosis after few trials would affect the total number of recycling vesicles and cause strong release depression due to lack of release-ready
vesicles. We therefore conclude that blocking of endocytosis by Dynasore must cause the enhanced STD by a mechanism other than insufficient vesicle supply.

**Figure 3.11 Recruitment of Reserve Vesicles during Multiple Spaced Stimuli**

(A) Representative spH responses to consecutive runs of stimulation (50 APs at 20 Hz) with 60 second interval for recovery (red solid line) plotted together with the calculated release rate (black) and cumulative release (read broken line). By comparison, same experiment was repeated in the presence of Folimycin (blue) or Dynasore (green). All fluorescence transients were normalized to the initial response, respectively.

(B) Plot of changes in fluorescence amplitudes upon each trial. In control experiment, evoked responses (red) remain almost constant over at least ten trials, while that of Dynasore (green) or Folimycin (blue) decreases with different rates (n = 3 experiments with > 20 boutons for each). Error bars represent s.e.m.
3.8. Surface Accumulation of SV Components Caused by Dynasore

The major phenotype of impaired dynamin activity has been shown to be a reduction in the vesicle pool size and the appearance of coated pits and invaginations at the stimulated synapses (Ferguson et al., 2007; Newton et al., 2006). An acute block of endocytosis or the loss of dynamin activity has been postulated to directly perturb the clearance of vesicle components from release sites, thereby interfering with release probability (Haucke et al., 2011; Kawasaki et al., 2000; Neher, 2010). Here I took the advantage of the newly developed super resolution STED microscopy technique to follow the fate of newly exocytosed vesicular proteins on the plasma membrane. Since Syt1, one of the vesicle associated proteins, is enriched at the periphery (potential endocytic site) of synapses at rest (see above), immunostaining of those surface residual proteins should indicate the location where exocytosed vesicle components will finally be retrieved (Hua et al., 2011).

I first performed such surface staining using antibody coupled with red-colored fluorephore (ATTO 647N) on live hippocampal neuron culture, to which 1 µM TTX was applied to suppress the network activities. After washing with normal saline, the culture was changed to another staining buffer containing the same Syt1 antibody coupled with green-shifted fluorephore (ATTO 590). The binding of ATTO 590 labeled antibody to the cells was completely blocked due to the absence of free epitopes on the surface, until the culture was stimulated with a train of 200 AP at 20 Hz (Suppl. 3), which exposed unlabeled vesicular Syt1 to the surface. The detailed staining protocol is illustrated in Figure 3.12 A. Note that during the whole process Dynasore was present to block compensatory endocytosis, while control experiments were performed with DMSO. Two populations of Syt1 could thereby be well distinguished from each other by their colors using super resolution STED microscopy. Images revealed that Dynasore prevented newly exocytosed Syt1 patches from intermingling with the preexisting ones on the membrane.
(Fig 3.12 B). For a quantitative analysis the overlap of the two Syt1 populations was calculated using Pearson correlation coefficients between signals from two channels after background subtraction (Fig 3.12 C). Statistical significance in the difference between Dynasore and control groups argues strongly for a disruption in translocation of vesicular components towards endocytic sites. Together with reduced release probability observed under the same stimulation condition, this finding suggests a role of dynamin activity in the fast clearance of vesicular components from release sites, which is crucial for synapses to maintain high release rate during prolonged stimulation.
Figure 3.12 Accumulation of Vesicle Components in the Presence of Dynasore

(A) Experimental protocol for labeling surface-stranded and newly exocytosed Syt1 patches.

(B) Representative images of surface-stranded Syt1 (red) and newly exocytosed Syt1 (green). 100 µM Dynasore was added to block compensatory endocytosis. Overview was captured in the confocal mode (B1 and B2). Scale bar represents 2 µm. Images of 80 nm resolution were obtained by scanning synaptic boutons in the STED mode (B1a, B2a and B2b). Clear spatial separation between Syt1 patches was observed in STED images from experiments performed in the presence of Dynasore, indicating that Dynasore disrupts the translocation of newly exocytosed Syt1 patches towards RRetP. Scale bar represents 400 nm.

(C) Statistic analysis of protein colocalization. Pearson coefficients were calculated for images from experiments performed either in the presence of Dynasore (test) or DMSO (control) after background subtraction. Significant reduction in Pearson coefficient was observed in Dynasore treated group compared to control group (p < 0.001, n = 15 experiments each).
4. Discussion

During my PhD, I have devised a new technique to measure synaptic activity in hippocampal synaptic boutons. Using cypHer-conjugated antibodies against luminal domains of Syt1 or VGAT, monitoring endogenous vesicle proteins during exo-endocytic cycling has achieved. Action-potential-driven fluorescence transients of αSyt1-cypHer and αVGAT-cypHer-stained synapses showed transients that were the mirror image of spH transients owing to their inverse pH dependence of cypHer fluorescence. Kinetics of endocytosis and reacidification assayed by both methods were nearly identical, showing that this cypHer-based approach reliably monitors recycling of endogenous synaptic vesicle proteins. The good spectral separation between cypHer and spH allowed us to use the two pH-dependent optical reporters in tandem, resulting in two independent readouts of presynaptic activity.

As previously shown, a significant percentage of SV proteins reside on the presynaptic membrane surface and SV interchange their membrane proteins with such surface reservoir during recycling (Fernandez-Alfonso et al., 2006). By bleaching or cleaving the pHluorin-tags of these surface SV proteins, it has been demonstrated that upon brief stimulation (40 APs) the vesicle proteins exo- and endocytosed lost their identity completely (Wienisch and Klingauf, 2006). To provide direct evidence for the active participation of RRetP-derived SVs in exo-endocytosis, I selectively labeled the RRetP with cypHer-conjugated antibodies and depleted the RRP by eliciting 50 Aps at 20 Hz. The observed increase in cypHer fluorescence demonstrated preferential endocytosis of cypHer-stained Syt1 and VGAT residing on the presynaptic membrane rather than freshly exocytosed unlabeled ones. The existence of this functional pool of surface-stranded SV proteins is also corroborated by the nanoscopy of surface-stranded Syt1 at the synaptic periphery, which is generally identified as the site of endocytosis. Images with a lateral
resolution of 80 nm revealed several preassembled Syt1 patches, from which new synaptic vesicles could bud. Note that even for the low-copy-number protein VGAT, there was a functional surface pool that was preferentially internalized upon stimulated exo-endocytosis, raising the question of whether this transporter might be active in the plasma membrane.

These findings demonstrate the existence of a functional RRetP of native vesicle constituents such as Syt1 and VGAT at the synaptic periphery. The preferential uptake of those vesicle constituents from the surface pool explains why an endocytosed vesicles is not composed of the most recently exocytosed vesicular proteins (Wienisch and Klingauf, 2006). Although overexpression of spH-tagged SV proteins may lead to a substantial increase of the total surface pool, the cypHer experiments clearly demonstrate the existence of an endogenous vesicular protein surface pool participating actively in SV cycling (Fernandez-Alfonso et al., 2006; Wienisch and Klingauf, 2006). Existence of such RRetP can explain the first wave of CME observed in the first reconstruction of the endocytic time course from electron micrographs of frog neuromuscular junctions (Miller and Heuser, 1984). Also, it provides an explanation for the reason why the initial endocytic rate remains constant irrespective of the stimulation strength, as suggested both in this and other studies (Balaji et al., 2008; Fernandez-Alfonso and Ryan, 2004; Sankaranarayanan and Ryan, 2000). However, after this RRetP is depleted, the rate of endocytosis drops because the endocytic machinery has to invest more energy and time to recapture and re-sort freshly exocytosed vesicle components to the sites of retrieval (Figure 3.7). Because endocytosis of the RRetP itself, however, takes several seconds, the main advantage of an RRetP might not be so much a gain in speed but rather a gain in precision of re-sorting and reclustering to ensure maximum fusion competence of newly endocytosed vesicles. The counterintuitive finding that knockdown or knockout of major
sorting factors such as adaptor protein 1 (AP1) and AP2 subunits results in similar or even faster endocytic kinetics of a large fraction of retrieved SV proteins (Glyvuk et al., 2010; Kim and Ryan, 2009), can be brought in line with present investigations under the hypothesis that this fast endocytosis represents bulk endocytosis. The availability of endocytosed SVs is slowed and their fusion competence is diminished (Glyvuk et al., 2010) due to lack of a maturation step to an RRetP. In three recent studies, the importance of Ca\textsuperscript{2+} for coupling of exo- and endocytosis was emphasized (Hosoi et al., 2009; Wu et al., 2009; Yao et al., 2009). Work in Drosophila suggested that there might be a vesicular Ca\textsuperscript{2+} channel that, once incorporated into the plasma membrane by fusion, would allow Ca\textsuperscript{2+} entry at the site of endocytosis and thus might trigger the retrieval from the RRetP (Yao et al., 2009). However, the exact molecular mechanism that couples exocytosis from the RRP with compensatory endocytosis from the RRetP and the regulatory factor of such pool still remains elusive.

So far this study demonstrated that exocytosed and endocytosed vesicles are driven in parallel from RRP and RRetP, respectively. In order to release neurotransmitter steadily at high rate, however, a mechanism is required to efficiently translocate newly exocytosed patches of vesicular proteins from the release site towards the rim of the AZ where the endocytic sites are located. Through examining the Dynsore-enhanced release depression using quantitative spH assay, I provide direct evidence for a novel mechanism that enhances STD, by functionally blocking the previously used AZ slot due to a release site clearance defect.

From technical point of view, I improved pHluorin-based assays for quantitative measurements of neurotransmitter release by using a novel normalization strategy of the spH response. The new protocol greatly reduces the variation in responses between synapses by applying a sequence of a standardized calibration stimulus followed by
various test stimuli. With this method the inhomogeneity across synapses between responses to calibration and test stimulus is cancelled out. Furthermore, I combined this with a deconvolution method that provides an estimate of the cumulative release without any pharmacological perturbations. The deconvolution method distinguishes itself from alkaline-trapping by including the contribution of reused vesicles, which are not monitored by the latter method. Therefore, comparison between results from deconvolution with those from alkaline-trapping measurements serves as an excellent tool for quantifying vesicle reuse.

A surprising result in our data was the absence of preferential reuse of exocytosed vesicles under mild stimulation. I observed identical signal amplitudes for both deconvolution and alkaline-trapping in response to 200 APs at 5 Hz. Also, further experiments using Folimycin excluded the existence of additional vesicle reuse when firing frequency was increased up to 20 Hz, while a small reduction in the recorded amplitudes during 40 Hz stimulation indicated some amounts of vesicle reuse, which may account for the reduction (Figure 3.9). Additionally the amount of release upon 200 APs appeared to be insensitive to stimulation frequency up to 40 Hz, which may be a consequence of the activity-dependent replenishment of RRP to meet the high demand of release-ready vesicles at AZ (Dittman and Regehr, 1998; Stevens and Wesseling, 1998; Wang and Kaczmarek, 1998). Moreover, I demonstrated that a high-frequency depleting pulse or elevated extracellular calcium concentration results in an increased recycling pool size (Suppl. 1). This suggests an accelerated replenishment that recruits vesicles from a release-reluctant pool rather than the slowly endocytosed vesicles. Taking this evidence together, I conclude that fast RRP replenishment alone can guarantee a sufficient vesicle supply during short period of physiological stimulation without additional contribution of rapid vesicle reuse.
The measured cumulative release upon different stimulation conditions (Figure 3.10) provided quantitative information about how many vesicles can be eventually exocytosed in response to a seconds-long electrical stimulus. Based on that knowledge I was able to probe the release depression caused by the acute block of dynamin activity. Consistent with previous work performed at Calyx of Held using capacitance measurement (Hosoi et al., 2009), I found that in cultured rat hippocampal neurons the perturbation of dynamin function leads to a significant reduction in the cumulative release during sustained high frequency (40 Hz) stimulation. When the same number of stimuli was applied at low rate (5 Hz), release depression was almost undetectable. It has been postulated that insufficient vesicle supply accounts for the release depression in Dynasore-treated neurons (Newton et al., 2006), since depletion of fusion competent vesicles is a direct consequence of impaired endocytosis under dynamin blockade. However, two aspects of our results challenge this view. First, stimulation at 5 Hz in the presence of Dynasore does mobilize more vesicles for fusion than at 40 Hz, implying that the release depression upon high frequency stimulation occurs before vesicle pool depletion. Second, alkaline-trapping experiments show that even during high frequency stimulation for up to few seconds, vesicles are mainly recruited from the pre-existing vesicle pool rather than originating from recently endocytosed vesicles. Note that in these experiments the drug was applied to the cell culture only five minutes before recording to avoid any possible chronic changes like e.g. depletion of vesicles in the synaptic bouton, as reported in other studies where genetic knockout of the protein was employed (Lou et al., 2012).

So far, these results demonstrate that dynamin is essential for a process that occurs between exo- and endocytosis with a time constant shorter than that of vesicle recruitment. It manifests itself rapidly after dynamin activity is abolished. As proposed in recent works, translocation of synaptic components from sites of exocytosis to a
“periactive zone” may be a good candidate for this step (Haucke et al., 2011; Neher, 2010). This is supported by findings that perturbations of dynamin and several interacting proteins enhance short-term synaptic depression upon high frequency stimulation (Hosoi et al., 2009; Marie et al., 2004). These studies include intersectin which interact with the actin regulatory proteins nwASP (neural wiskott-Aldrich syndrome protein) and CDC42 (cell division control protein 42) (Pechstein et al., 2010). Therefore, perturbed dynamin function may lead to the accumulation of vesicle components around release sites, as I observed under the STED microscope (Figure 3.13). This may prevent previous used release sites from participating in the recruitment and the release of additional vesicles. From the physiological point of view, release site clearance as a hypothetical mechanism, which nicely explains the deep depression during sustained high frequency stimulation after interference with dynamin function. Free diffusion of the exocytosed vesicular proteins on the plasma membrane may account for the remaining slow restoration of the release sites, while the effect of CDC42 block may suggest that the cytoskeleton is involved in rapid site clearance. Moreover, it has also been reported by Gaffield and colleagues that during high-frequency stimulation of mouse motor nerve terminal, the preferred sites of endocytosis were found much closer to the release sites as compared to mild stimulation (Gaffield et al., 2009). If this is also the case for cultured hippocampal neurons, deeper depression may be expected for high-frequency stimulation because of protein crowding around the release sites. This may provide a reason why a 60 second interval between multiple stimuli failed to rescue the depression in the presence of Dynaosre, if reduced distances between sites for exo- and endocytosis further restrict the diffusion of the exocytosed vesicle components. These results broaden the physiological importance of intact dynamin function by showing it is essential for the proper function of the synapse during sustained activity (Hosoi et al., 2009).
In conclusion, our study demonstrates that impaired dynamin activity leads to a novel form of short-term depression, which is not related to insufficient vesicle supply at the synaptic bouton, but is the result of slow clearance of vesicular components from the release sites. This finding implies an important role of dynamin for sustained synaptic transmission at high rates beyond its well-known role in mediating a late step of endocytosis.
Summary

In the central nervous system, communication between nerve cells mainly occurs at synapses via chemical neurotransmitters. Upon stimulation the transmitters are released by Ca\textsuperscript{2+}-triggered exocytosis of SVs from presynaptic terminal to the synaptic cleft, where they activate postsynaptic receptors. In order to repopulate the pool of synaptic vesicle for further rounds of release, vesicle proteins need to be resorted and retrieved from the surface by compensatory endocytosis.

In this study, both exogenous cypHer-based and genetically encoded GFP-based (spH) pH-switchable reporters are used to investigate the vesicle cycling in cultured rat hippocampal neurons. Using cypHer-conjugated antibodies against luminal domains of vesicle proteins (e.g. Syt1 and VGAT), visualizing cycling of endogenous vesicle proteins has been achieved. In addition, by monitoring surface-stranded vesicle proteins, constituting the so-called readily retrievable pool (RRetP), using cypHer-antibody, it was clearly demonstrated that upon stimulation stained Syt1 residing on the presynaptic membrane rather than freshly exocytosed unlabeled ones are preferentially retrieved to form fresh vesicles, indicating an active participation of the RRetP pool in the vesicle cycling. Furthermore, the existence of this RRetP is also corroborated by the observation of surface-stranded Syt1 at the synaptic periphery, which is generally identified as the site of endocytosis. These evidences indicate that exocytosis and subsequent compensatory endocytosis of vesicles are driven in parallel from the RRP and the RRetP, respectively. Since the surface pool is repopulated by newly exocytosed vesicle components, fast translocation of newly exocytosed vesicle proteins from AZ towards the synaptic periphery is expected. To explore potential roles of this translocation in synaptic transmission, dynamin-inhibitor Dynasore was applied to induce protein crowding by
blocking endocytosis. Under influence of this drug a clear release depression was observed during high frequency stimulation, which was, however, absent under control conditions. Based on this observation, it is concluded that functional block of previously used release sites rather than insufficient vesicle supply accounts for the fast release depression after blocking endocytosis. This finding implies an important role of dynamin for sustained synaptic transmission at high rates beyond its well-known role in mediating fission, a late step of endocytosis.
5. Supplementary Figures

Suppl. 1 Calcium Level Regulates the Recycling Pool Size.

(A) Average spH response to two different depleting pulses in the presence of Folimycin. Vesicle pool was depleted either with a train of 1200 APs at 30 Hz followed by 1200 APs at 5 Hz (gray) or in an opposite order. A brief pulse of NH₄Cl was applied at the end of the recording to determine the total vesicle population. Traces were normalized to the maximum dynamic range of spH.

(B) Average spH response to 1200 APs at 5 Hz in the presence of Folimycin. Experiments were conducted in a buffer containing either 2 mM (gray) or 4 mM calcium (red). A brief pulse of NH₄Cl was applied at the end of the recording. Traces were normalized to the maximum dynamic range of spH.
Suppl. 2 Dynasore-enhanced Release Depression Probed by cypHer

(A) Average cypHer responses to 50 APs at 20 Hz followed by 200 APs stimulation at 5 (red) or 40 Hz (blue) in the presence of Folimycin. Each fluorescence transient from a given region of interest was normalized to the amplitude of the calibration stimulus response. (n = 3 experiments with > 20 boutons for 5 and 40 Hz). Error bars represent s.e.m.

(B) The same experiment as in (A), but was performed in the presence of Dynasore (n = 3 experiments with >20 boutons for each). Each fluorescence transient from a given region of interest was normalized to the amplitude of the calibration stimulus response. Error bars represent s.e.m.
Suppl. 3 Live Staining of Newly Exocytosed SV Proteins

(A) Representative images of neuron cultures stained with ATTO 647N labeled αSyt1-antibodies. The cultures were preincubated with unlabeled antibodies to block surface residual epitopes and then challenged with stimulations of different APs at 20 Hz to evoke exocytosis. Dynasore was always present to block compensatory endocytosis. Without stimulation no signals of ATTO 647N was observed (left), suggesting the absence of free epitopes on the surface, while fluorescence intensity increases with prolonged stimulation (middle, right), indicating that more epitopes become access to the antibodies due to exocytosis. Scale bar represents 10 µm.
6. Reference


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### 7. Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>AP</td>
<td>Action Potential</td>
</tr>
<tr>
<td>AP-1, AP-2</td>
<td>Adaptor Protein-1,2</td>
</tr>
<tr>
<td>AZ</td>
<td>Active Zone</td>
</tr>
<tr>
<td>CAZ</td>
<td>Cytoplasmic Matrix of Active Zone</td>
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<tr>
<td>CME</td>
<td>Clathrin-mediated Endocytosis</td>
</tr>
<tr>
<td>CNQX</td>
<td>6-cyano-7-nitroquinoxaline-2,3-dione</td>
</tr>
<tr>
<td>DIV</td>
<td>Day in Vitro</td>
</tr>
<tr>
<td>DL-AP5</td>
<td>D-amino-5-phosphonovaleric acid</td>
</tr>
<tr>
<td>EPSC</td>
<td>Excitatory Postsynaptic Current</td>
</tr>
<tr>
<td>HBSS</td>
<td>Hank’s Balanced Salt Solution</td>
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<tr>
<td>NBA</td>
<td>Neurobasal A Medium</td>
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<tr>
<td>NSF</td>
<td>N-Ethylmaleimide-Sensitive Factor</td>
</tr>
<tr>
<td>PFA</td>
<td>Paraformaldehyde</td>
</tr>
<tr>
<td>PSD</td>
<td>Postsynaptic Density</td>
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<tr>
<td>RIM 1/2</td>
<td>Rab3-interacting Molecule 1/2</td>
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<tr>
<td>RRP</td>
<td>Readily Releasable Pool</td>
</tr>
<tr>
<td>RRetP</td>
<td>Readily Retrieval Pool</td>
</tr>
<tr>
<td>SNARE</td>
<td>Soluble NSF Attachment Protein Receptor</td>
</tr>
<tr>
<td>spH</td>
<td>Synaptobrevin2-pHluorin</td>
</tr>
<tr>
<td>STED</td>
<td>Stimulated Emission Depletion</td>
</tr>
<tr>
<td>STD</td>
<td>Short-term Synaptic Depression</td>
</tr>
<tr>
<td>STP</td>
<td>Short-term Synaptic Plasticity</td>
</tr>
<tr>
<td>SV</td>
<td>Synaptic Vesicle</td>
</tr>
<tr>
<td>TTX</td>
<td>Tetrodotoxin</td>
</tr>
<tr>
<td>vGlut</td>
<td>Vesicular Glutamate Transporter</td>
</tr>
<tr>
<td>VGAT</td>
<td>Vesicular GABA Transporter</td>
</tr>
<tr>
<td>VGCC</td>
<td>Voltage-gated Calcium Channel</td>
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