

Lesson (8)

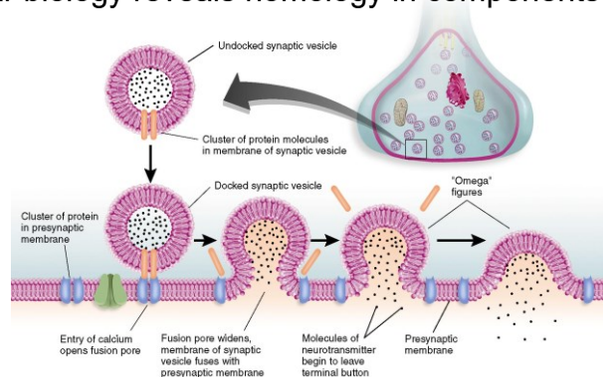
Inside the neuron IV:

Presynaptic secretion

What is the molecular machinery of synaptic vesicles fusion ?

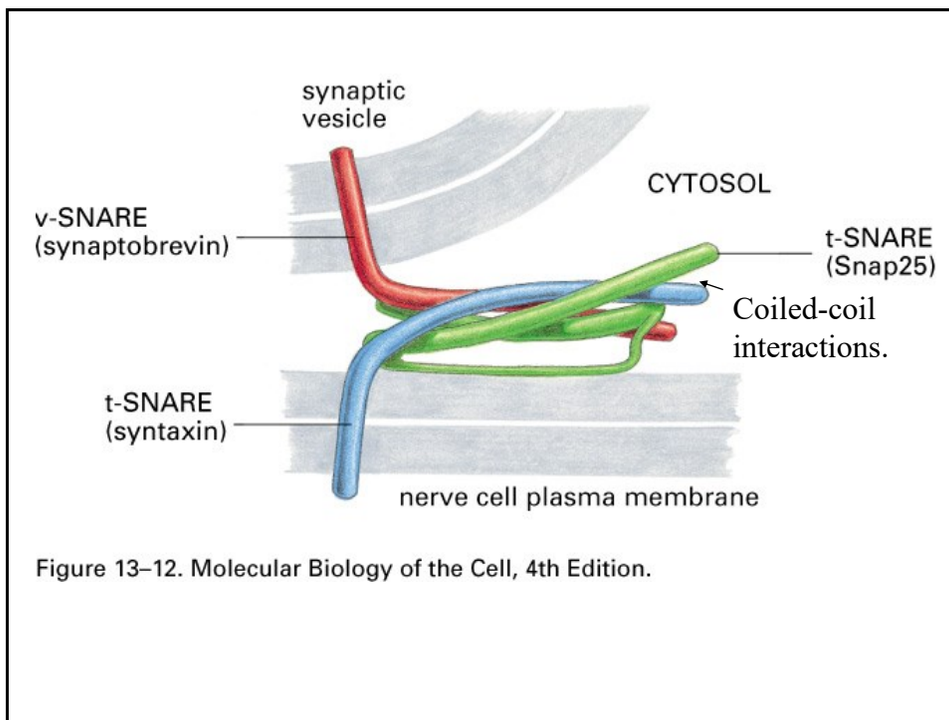
Some observations:

- Synaptic transmission, a specialized form of normal exocytosis.
- Specific proteins on vesicles and targets guide the process
- Molecular biology reveals homology in components from yeast to man

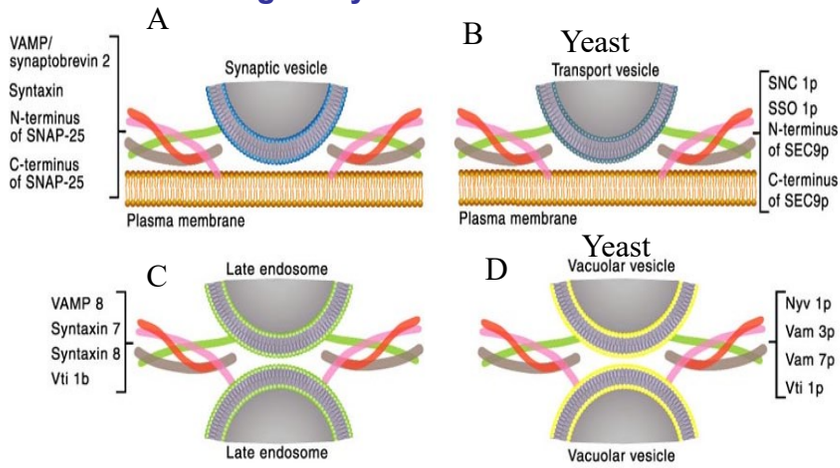


The SNARE hypothesis

- Synaptic vesicles have a specific protein that directs them to a receptor on the plasma membrane (with the aid of other proteins SNAPS etc.).
- The vesicle associated SNAP receptor (v-SNARE) in neurons is believed to be synaptobrevin (VAMP).
- Target sites such as the plasma membrane (nerve terminal) would have a corresponding t-SNARE (syntaxin and SNAP-25).
- At sites other than the nerve terminal specific vesicle and target SNAREs would function to target vesicles to specific compartments.

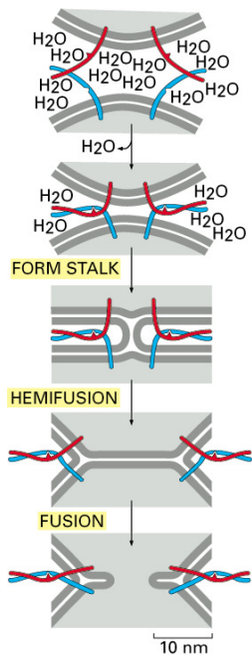


The SNARE mechanisms is shared among many membrane fusion events



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FIGURE 7 Neurotransmitter release shares a core mechanism with many membrane fusion events within eukaryotic cells. The fusion of synaptic vesicles (A) is driven by a particular complex of four coiled-coil domains contributed by three different proteins. Exocytosis in yeast (B), the fusion of late endosomes in mammalian cells (C), and the fusion of vacuolar vesicles in yeast (D) exemplify the closely related four-stranded coiled-coil complexes required to drive fusion in other membrane-trafficking steps.



Generic fusion mechanism involves a tightening of the SNAREs to push out H₂O

Figure 13-15. Molecular Biology of the Cell, 4th Edition.

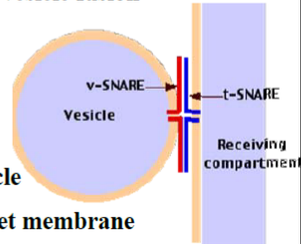
Specificity of vesicle fusion

Need mechanism for selective vesicle trafficking -controlled by **SNAREs** and **Rab** proteins

SNARE hypothesis proposes specific interactions between **v- SNAREs** and **t- SNAREs** govern vesicle docking and fusion

Each organelle has specific SNAREs leading to specific vesicle fusion

Vesicle docking controlled by Rab proteins



Monomeric GTPases attach to surface of budding vesicle

Rab-GTP on vesicle interacts with Rabeffector on target membrane

After vesicle fusion GTP hydrolysed, triggering release of Rab-GDP

Different Rab proteins found associated with different membrane-bound organelles

Comparison between intra-organelles and synaptic vesicle trafficking

Different Rab **GTPases** & Rab effectors control docking of different vesicles on target membranes: vesicle docking controlled by Rab protein.

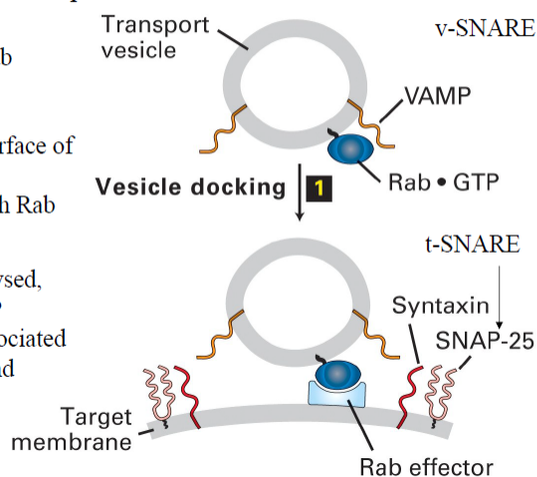
Vesicle docking controlled by Rab proteins

Monomeric GTPases attach to surface of budding vesicle

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After vesicle fusion GTP hydrolysed, triggering release of Rab-GDP

Different Rab proteins found associated with different membrane-bound organelles



Paired sets of SNARE proteins mediates fusion of vesicles with target membranes.

Analysis of yeast *sec* mutants defective in each of the >20 SNARE genes.

In vitro liposome fusion assay.

SNARE-mediated fusion → exocytosis → secretory protein

In this case, v-SNARE as VAMP (vesicle associated membrane protein)

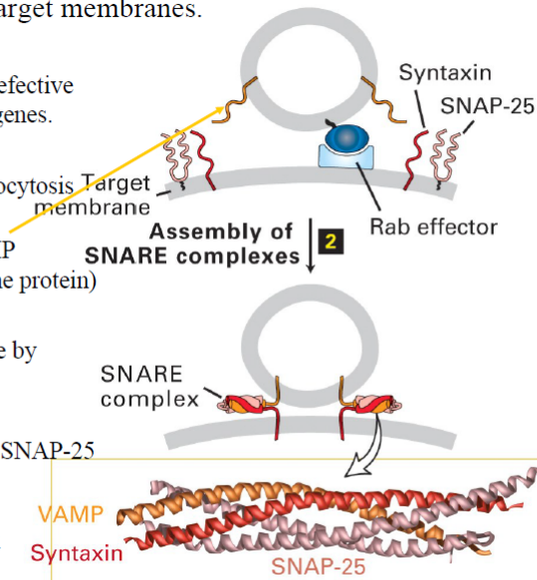
t-SNAREs are syntaxin

SNAP-25 attached to membrane by hydrophobic anchor.

Formation of four-helix bundle:

VAMP (1), Syntaxin (1) and SNAP-25 (2)

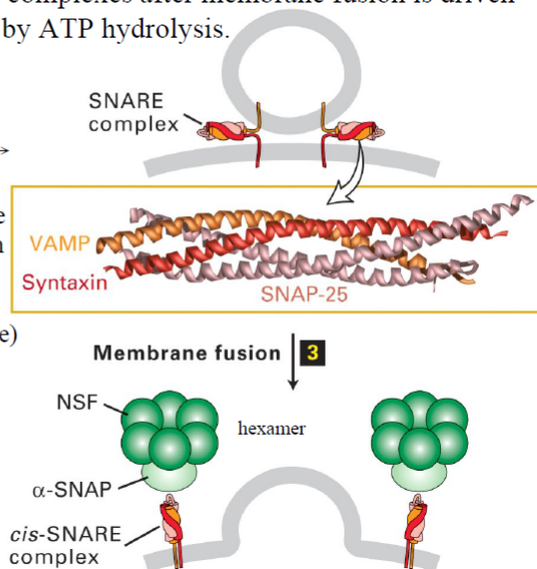
But, in COPII with cis, each SNARE has provide one helix
SNARE complex had specificity



Dissociation of SNARE complexes after membrane fusion is driven by ATP hydrolysis.

SNARE complex formation by non-covalent interaction.
Dissociate → free SNARE → can fuse next time

Two protein play important role of dissociation or fusion with a target membrane: NSF (NEM-sensitive factor, blocked by N-ethylmaleimide) & α-SNAP (soluble NSF attachment protein).



Soluble (i.e. cytoplasmic) Factors

NSF or n-ethylmaleimide (NEM) Sensitive Factor

SNAP- Soluble NSF Attachment Proteins

NSF + SNAP bind to target membranes (synaptic vesicle & plasma membrane)

Receptors for NSF and SNAP are synaptobrevin (vesicle), SNAP-25 (plasma membrane) and syntaxin (plasma membrane)

Membrane targets are called **SNAREs** (v- and t-) Soluble NSF Attachment protein **RE**ceptors

SNAP-25- Synaptosome Associated Protein of 25 kDa

- Over-expression of truncated SNAP-25 blocks release
- Syntaxin, 15 kDa protein
- Sensitive to botulinum toxin A cleavage - release prevented

Dissociation of SNARE complexes after membrane fusion is driven by ATP hydrolysis.

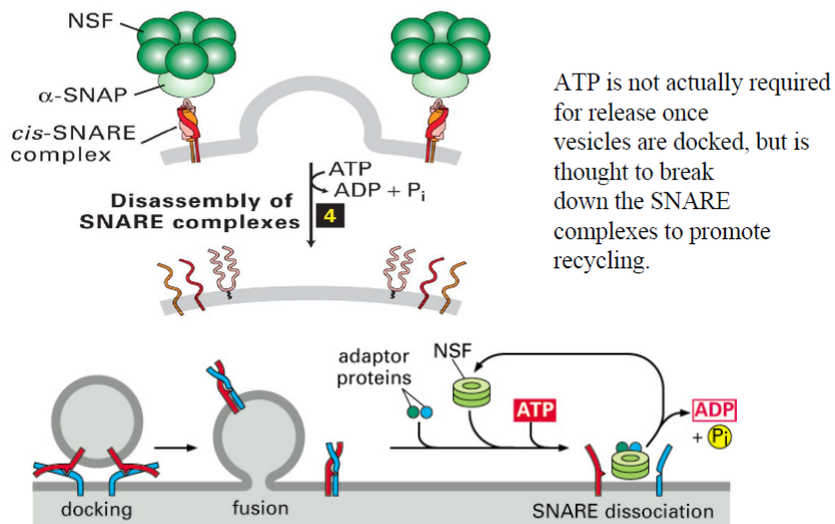
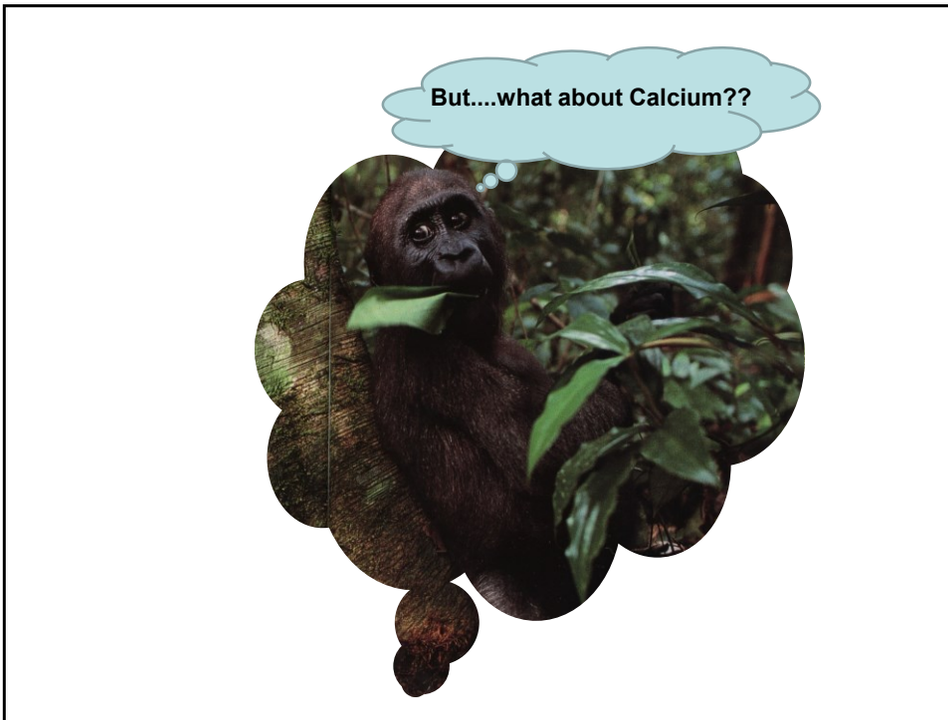
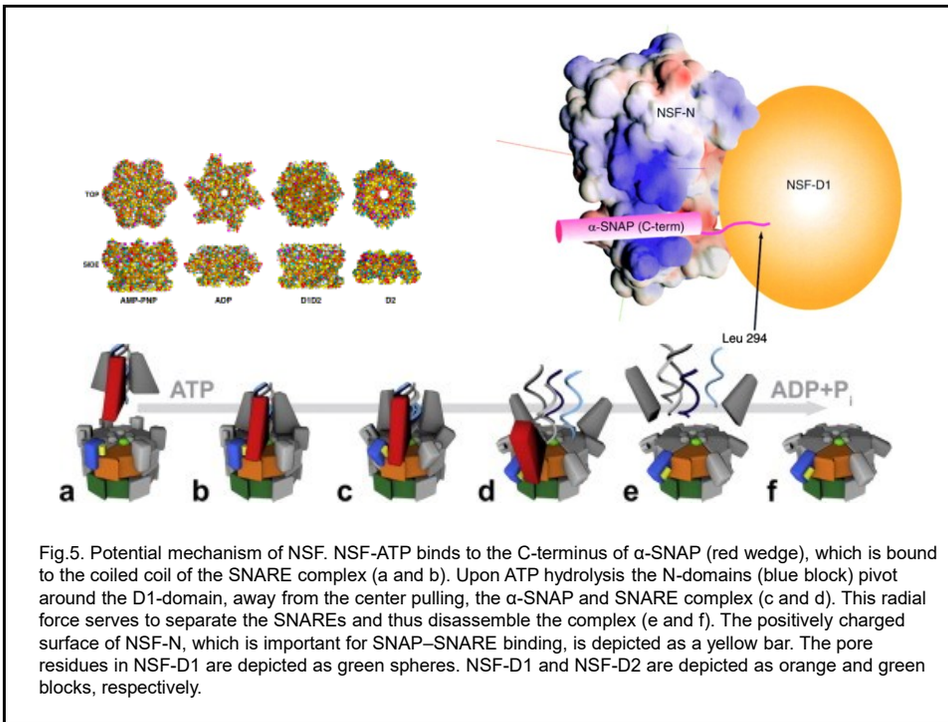
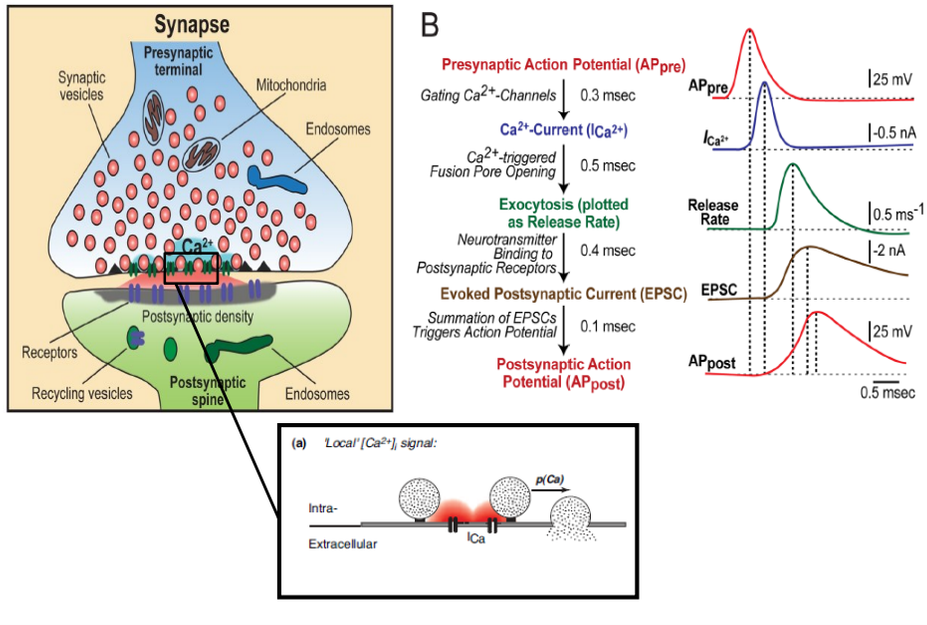


Figure 13-13. Molecular Biology of the Cell, 4th Edition.



Ca²⁺ signalling at presynaptic sites



Ca²⁺ signalling at presynaptic sites

Review

Trends in Neurosciences May 2011, Vol. 34, No. 5

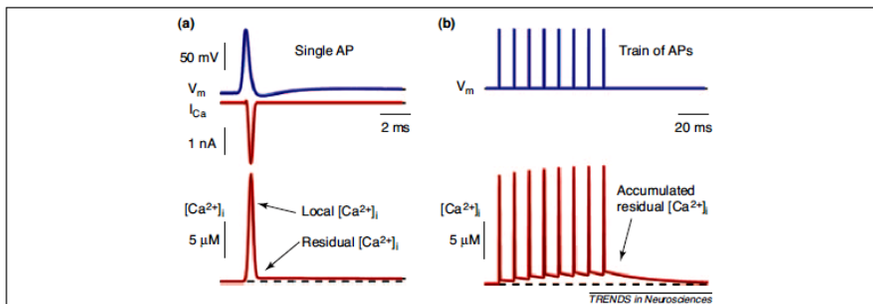
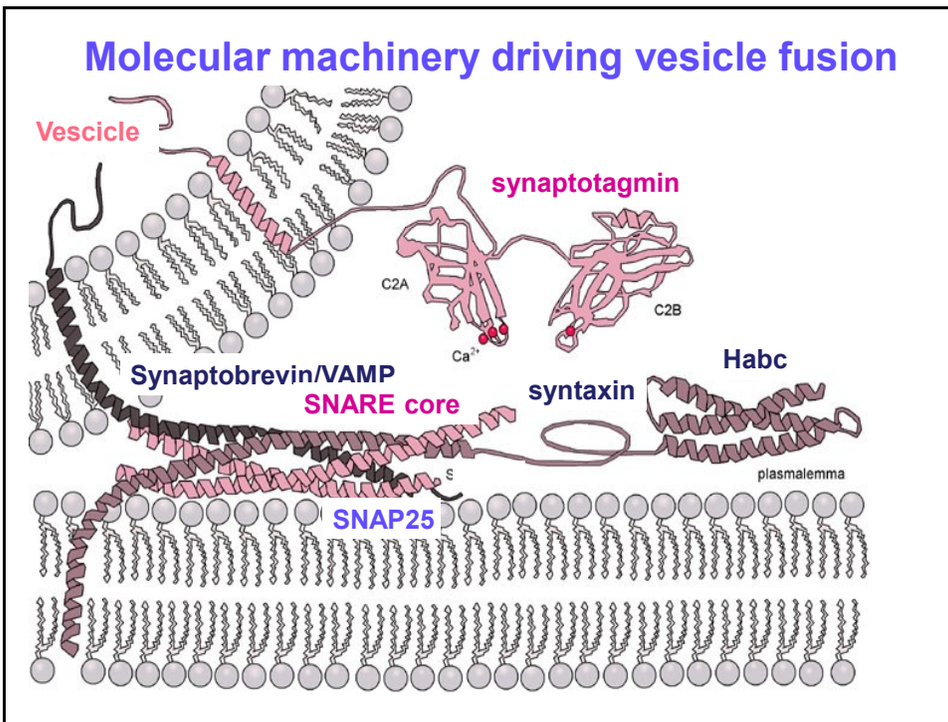
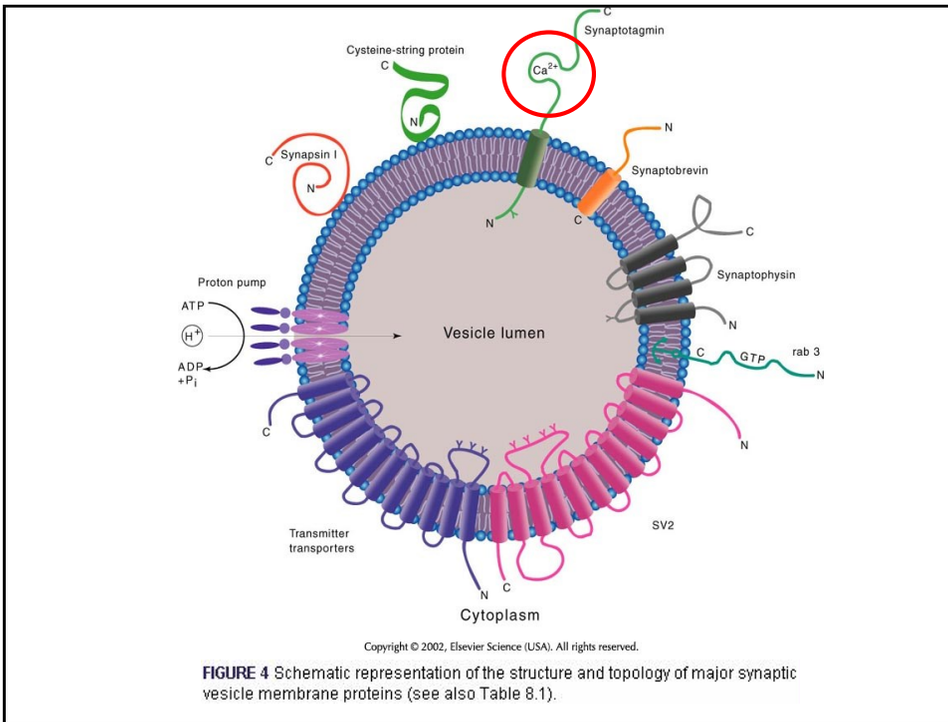
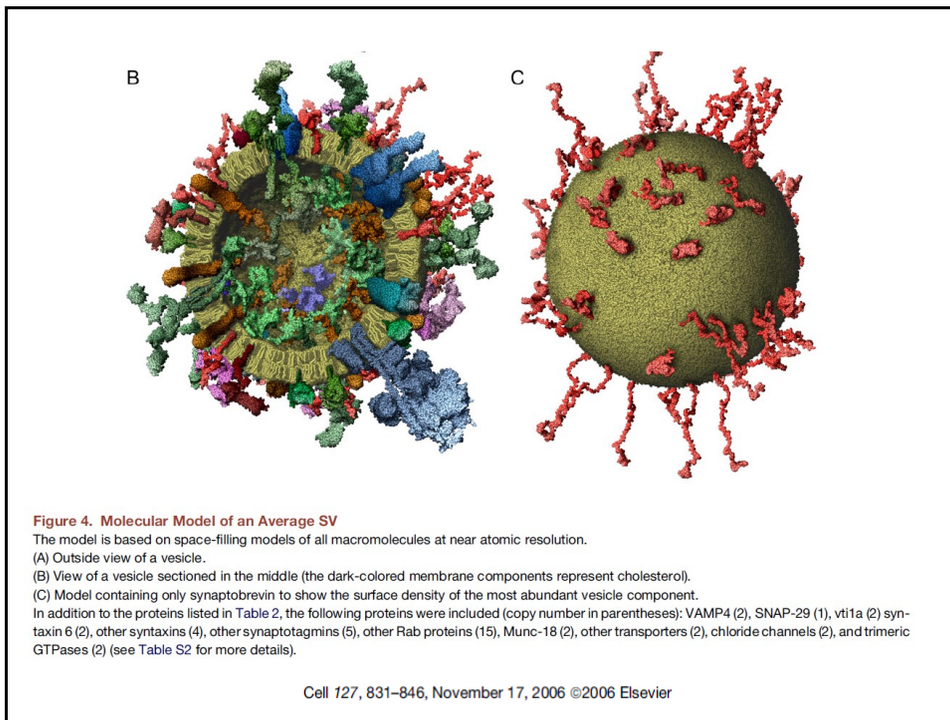
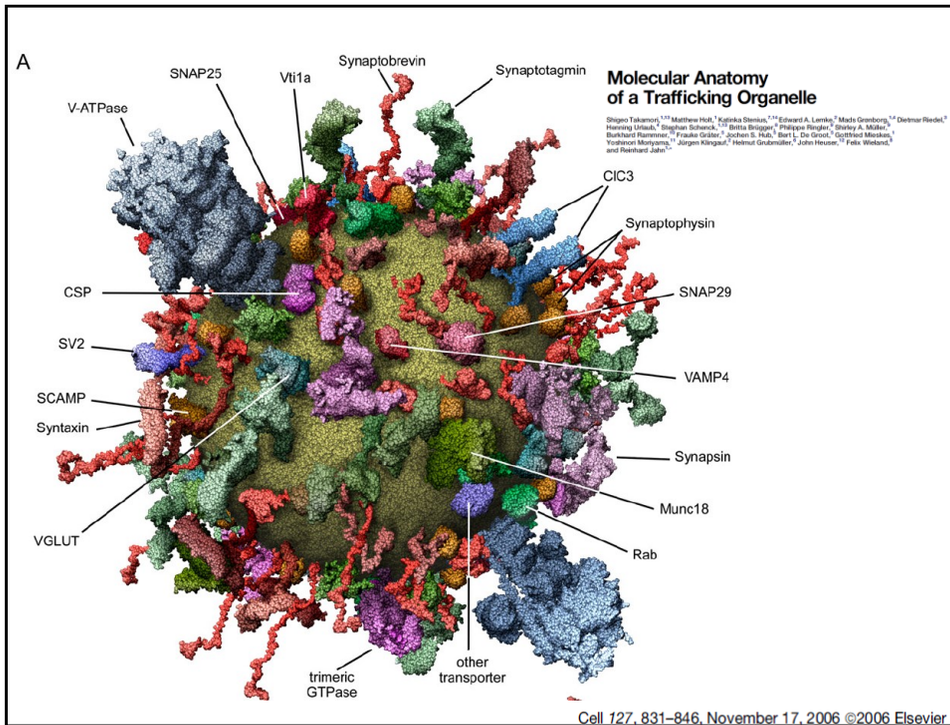


Figure 1. Schematic illustration of AP-evoked presynaptic [Ca²⁺]_i signaling at the release site. (a) A single AP (top panel) causes a short (~0.4 ms half-width) Ca²⁺ current (I_{Ca}), which in turn creates a brief [Ca²⁺]_i transient at the release site ('local' [Ca²⁺]_i). The local [Ca²⁺]_i signal is followed by a longer-lasting (τ_{1/2} ~30–50 ms) and widespread residual [Ca²⁺]_i signal of smaller amplitude (< ~0.5 μM). Residual [Ca²⁺]_i can cause a slower and asynchronous phase of release. (b) During a high-frequency train of APs (top panel), the residual [Ca²⁺]_i signal (bottom panel) can readily sum up to values in the low μM range, leading to increased rates of asynchronous release. Thus, phasic transmitter release during the presynaptic AP and the synchronous release phase are driven by local [Ca²⁺]_i signals with profoundly different amplitudes and kinetics.

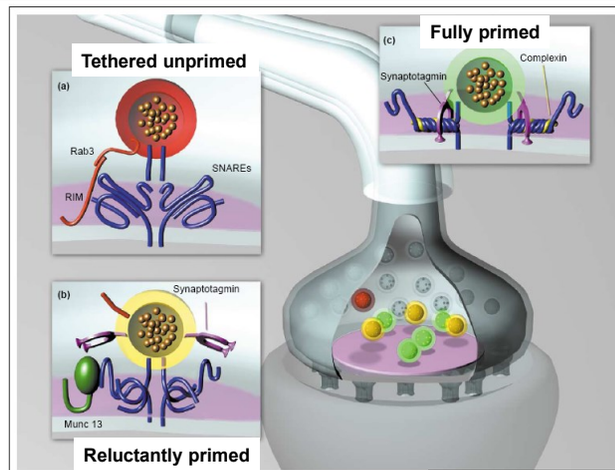
Olexiy Kochubey¹, Xuelin Lou² and Ralf Schneggenburger¹





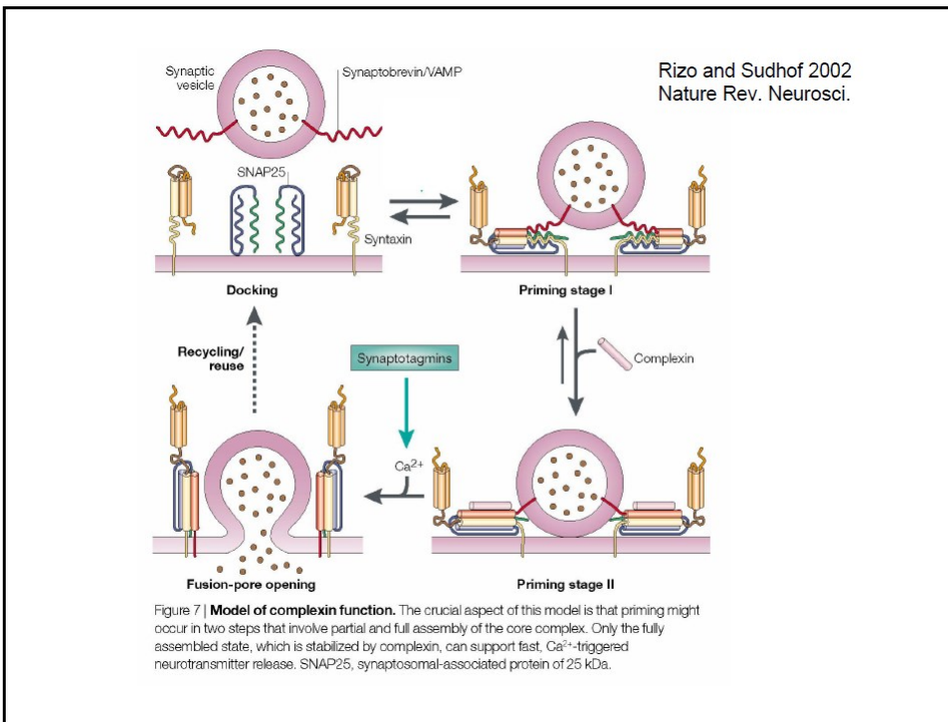
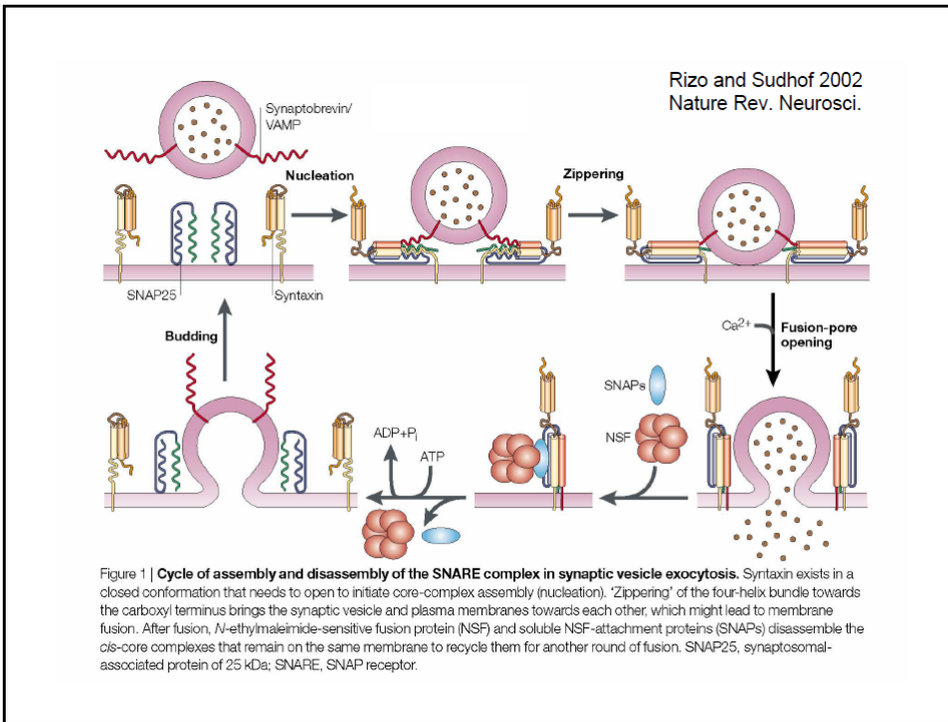
Synaptotagmin (P65) Ca²⁺ binding.

- Single polypeptide, one transmembrane spanning region and large cytoplasmic domain.
- Protein kinase C like homology in carboxyl terminal (C2 domain), involved in calcium and phospholipid binding. Injection of C2 peptide can block release after the vesicle is docked (squid; Augustine lab).
- Forms tetramers with each binding calcium (possible) cooperatively consistent with 3-4 power of Ca dependence of release (Sudhof and others).
- In vitro studies it binds Ca cooperatively at concentrations in the physiological range (10-100 μ M) for release.
- In the absence of Ca²⁺ synaptotagmin may serve as a brake for release. Ca²⁺ removes the brake. May explain how release in non-neuronal cells occurs without synaptotagmin.



Current Opinion in Neurobiology 2003, 13:509-519

Different stages of vesicle priming in central nerve terminals. The presynaptic terminal contains various neurotransmitter-filled vesicles, which are functionally defined as reserve vesicles (grey opaque), tethered unprimed vesicles (red), reluctantly primed (yellow), and fully primed (green). The reluctant and fully primed vesicles constitute the readily releasable vesicle pool. At each active zone (ilac circle), this pool consists of about 5-10 vesicles [24]. The inserts document the current view of how presynaptic proteins regulate the priming and maturation status of vesicles and how this affects their properties. (a) The interaction of active zone specific RIM and the vesicular Rab3 is thought to be involved in vesicle docking and the initiation of vesicle priming. (b) During priming, a 'loose' form of the SNARE complex, consisting of synaptobrevin, SNAP-25 and Syntaxin 1 (SNAREs are the blue rods on the diagram) is formed with the aid of Munc13. The vesicles mature to this stage relatively quickly, but they exocytose only reluctantly in response to single action potentials. Their release probability is in the range of 1-2% and they are released rather asynchronously (time course 20 msec). The release probability of these vesicles is potentiated two to five-fold by Munc13s interacting with DAG/phorbol esters [31**]. (c) Fully primed vesicles require additional interactions with complexin and synaptotagmin 1 to increase vesicle release probability (to 12-14%) and to speed up the release time course (2 msec). It is thought that fully primed vesicles obtain their release properties either through the formation of a tightened SNARE complex structure as shown here, a defined interaction of synaptotagmin 1 with the plasma membrane and the SNARE complex, and/or their proximity to voltage-activated Ca²⁺ channels.



Synchronous and Asynchronous release of synaptic vesicles

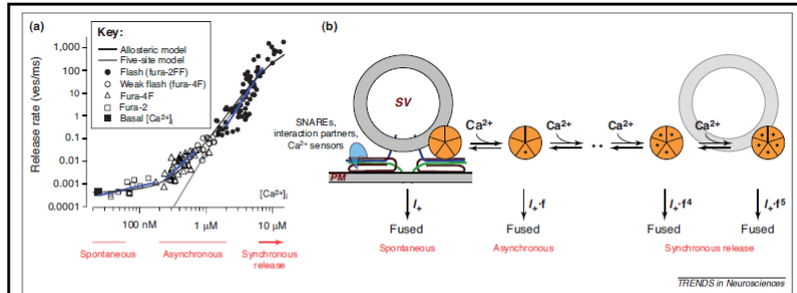


Figure 2. The intracellular Ca^{2+} -sensitivity of transmitter release over a wide range of $[\text{Ca}^{2+}]_i$, and the allosteric model of Ca^{2+} binding and vesicle fusion. (a) Dose-response curve of the intracellular Ca^{2+} -sensitivity of release determined over a wide range of $[\text{Ca}^{2+}]_i$ in young rat calyces of Held (postnatal days P8-P11) [18]. For the different concentration ranges, $[\text{Ca}^{2+}]_i$ was either controlled by presynaptic Ca^{2+} -buffer loading together with the indicated Ca^{2+} indicators (i.e. fura-4F or fura-2), or instead by using weak or strong flashes in Ca^{2+} uncaging experiments as indicated. The blue fit lines show linear fits in double-logarithmic coordinates (slope values 0.66 at $[\text{Ca}^{2+}]_i < 0.2 \mu\text{M}$, 2.77 at intermediate $[\text{Ca}^{2+}]_i < 1 \mu\text{M}$, and 4.2 at high $[\text{Ca}^{2+}]_i > 2 \mu\text{M}$). Note that the five-site model (grey line) failed to predict release rates at low $[\text{Ca}^{2+}]_i < 2 \mu\text{M}$, whereas the allosteric model could fit the data well (black line). The $[\text{Ca}^{2+}]_i$ ranges relevant for spontaneous, asynchronous and phasic Ca^{2+} -evoked release are highlighted. Reproduced, with permission, from [18]. (b) Graphical representation of the allosteric model of Ca^{2+} binding and synaptic vesicle (SV) fusion [18]. The model allows low rates of vesicle fusion from the Ca^{2+} -free state (spontaneous fusion; rate constant J) and from partially Ca^{2+} -bound states of the vesicle fusion machinery. Fusion rates rise in geometric progression upon further Ca^{2+} -binding to a general Ca^{2+} sensor complex with maximally five Ca^{2+} ion-binding sites (shown in orange). The states with four and five bound Ca^{2+} ions largely carry the synchronous release component evoked by $[\text{Ca}^{2+}]_i > 2 \mu\text{M}$ (the complete state diagram of the allosteric model is presented in Figure S1b in the supplementary material online).

The temporal resolution between Ca^{2+} influx induced by action potentials and SV fusion coupled to Ca^{2+} influx differs in synchronous and asynchronous release. The temporal precision largely depends on the distance between SV and VGCC, which matters for the timing of SV fusion. Fast synchronous neurotransmission is mediated by SV coupled to VGCC, while recruitment of new vesicles to the proximity of VGCC seems to contribute to the delayed asynchronous release.

Structure and Ca^{2+} properties of synaptotagmins

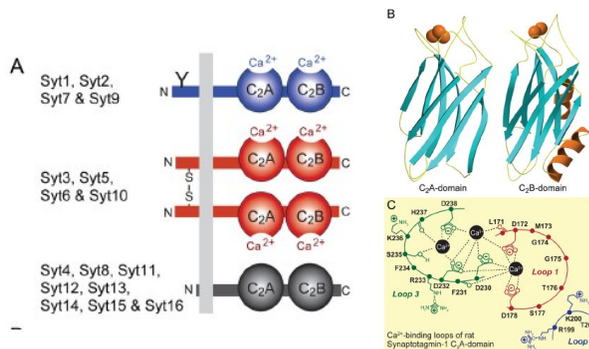
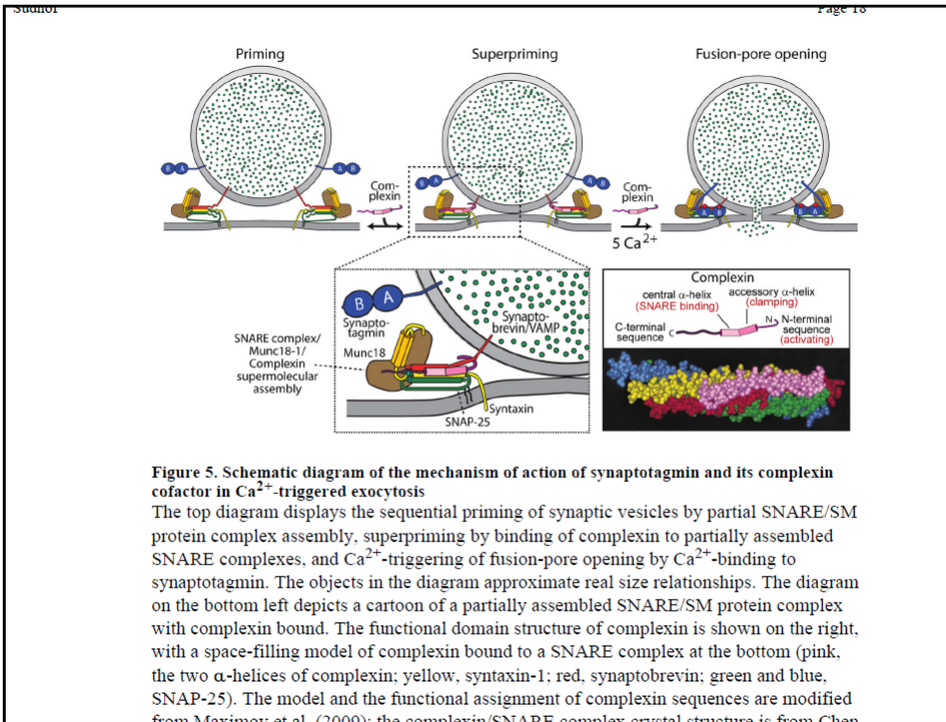


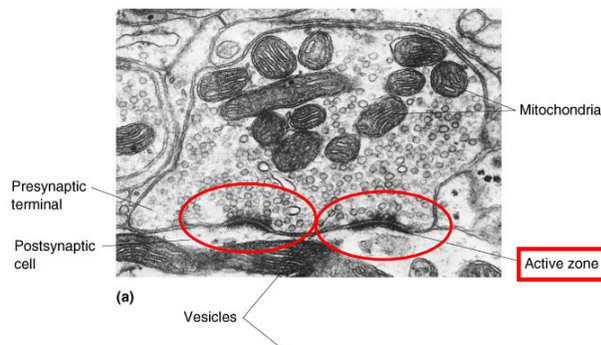
Figure 2. Structures and Ca^{2+} -binding properties of synaptotagmins

A. Canonical domain structures and classification of synaptotagmins. Mammals express 16 synaptotagmins composed of an N-terminal transmembrane region preceded by a short non-cytoplasmic sequence and followed by a variable linker sequence and two C2-domains; in addition, a 17th related protein called B/K-protein contains the same domain structure but an N-terminal lipid anchor instead of the transmembrane region (see Pang and Südhof, 2010, for a discussion of additional synaptotagmin-related proteins). Eight synaptotagmins bind Ca^{2+} (Syt1, 2, 3, 5, 6, 7, 9, and 10; blue and red); the remaining synaptotagmins do not (black). The eight Ca^{2+} -binding synaptotagmins fall into two broad classes that differ in the absence (Syt1, 2, 7 and 9; blue) or presence (Syt3, 5, 6, and 10; red) of disulfide-bonded cysteine residues in their N-terminal sequences. Note that Syt1 and 2 include an N-glycosylated sequence at the N-terminus (indicated by a 'Y'), and that Syt7 is extensively alternatively spliced in the linker sequence (Han et al., 2004).



Does synaptic vesicles fusion occur at specific places ?

Figure 5.3
Chemical synapses, as seen with the electron microscope. (a) Fast excitatory synapse in the CNS. (Source: Adapted from Heuser and Reese, 1977, p. 262.) (b) A synapse in the PNS, with numerous dense-core vesicles. (Source: Adapted from Heuser and Reese, 1977, p. 278.)



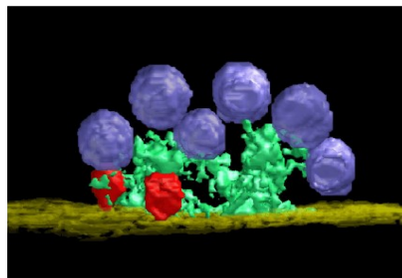
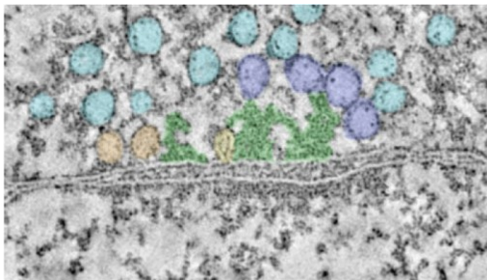
The active zone: definition and structure

Morphologically, active zones are defined as sites of synaptic vesicle docking and fusion, and physiologically they are defined as sites of neurotransmitter release.

Active zone have three morphologically and functionally distinct components:

1. the plasma membrane juxtaposed to the PSD where synaptic vesicle fusion occurs,
2. the cytomatrix immediately internal to the plasma membrane where synaptic vesicles dock, and
3. the electron-dense projections extending from the cytomatrix into the cytoplasm on which synaptic vesicles are tethered

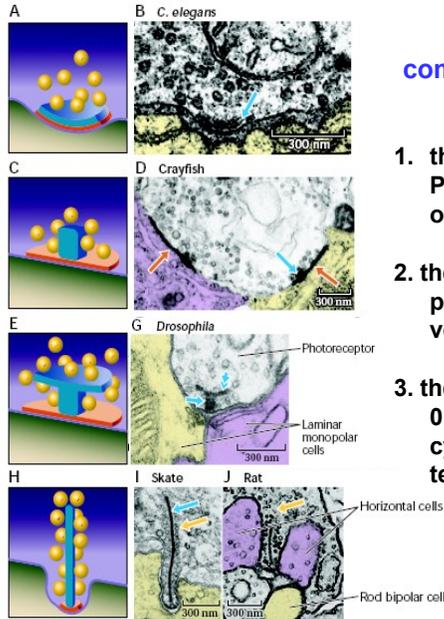
Architecture of the presynaptic Active Zone



Reconstruction of two dense projections (green) showing their relationship with six synaptic vesicles (blue) and two fusing vesicles (red).

Burette et al., *Microsc Microanal* 13(Suppl 2), 2007

Architecture of the presynaptic Active Zone



All active zones have three components, with variable size and shape of dense projections:

1. the plasma membrane juxtaposed to the PSD where synaptic vesicle fusion occurs,
2. the cytomatrix immediately internal to the plasma membrane where synaptic vesicles dock,
3. the electron-dense projections extending 0.5-1 μm from the cytomatrix into the cytoplasm on which synaptic vesicles are tethered (SYNAPTIC RIBBONS)

Zhai & Bellen *PHYSIOLOGY* 19: 262-270, 2004

Sikou et al. • Architecture of Presynaptic Terminals

J. Neurosci., June 27, 2007 • 27(26):6868 – 6877 • 6871

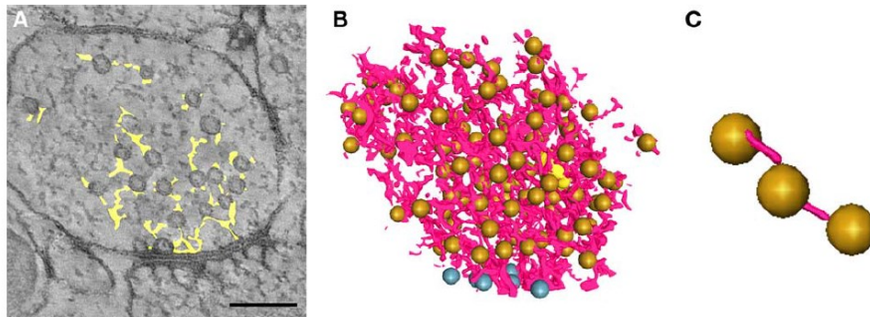


Figure 2. Network of filaments linking SVs in the presynaptic terminal. *A*, The electron-dense materials surrounding the SVs (yellow) are superimposed on a virtual section. *B*, 3D reconstruction emphasizing that docked (blue) and other (gold) SVs are embedded in a tight meshwork of filaments (pink). *C*, Higher magnification of a partial reconstruction exemplifying the connections (pink) between neighboring SVs. Scale bar, 200 nm.

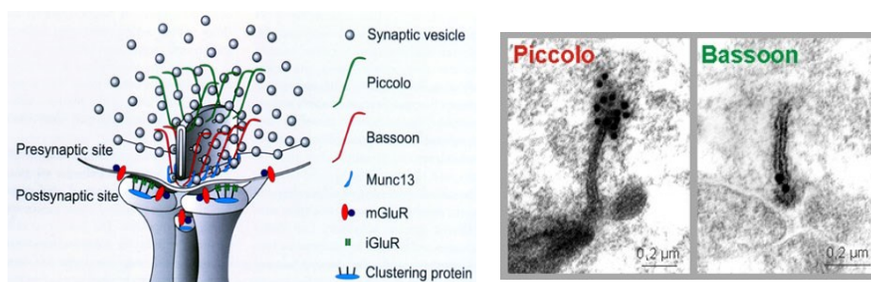
Proteins identified in the active zone cytomatrix can be classified into 3 categories.

- **First**, the classical cytoskeletal proteins corresponding to actin, tubulin, myosin, spectrin-chain and -chain, and -catenin are the fundamental elements of the framework of active zone cytomatrix.

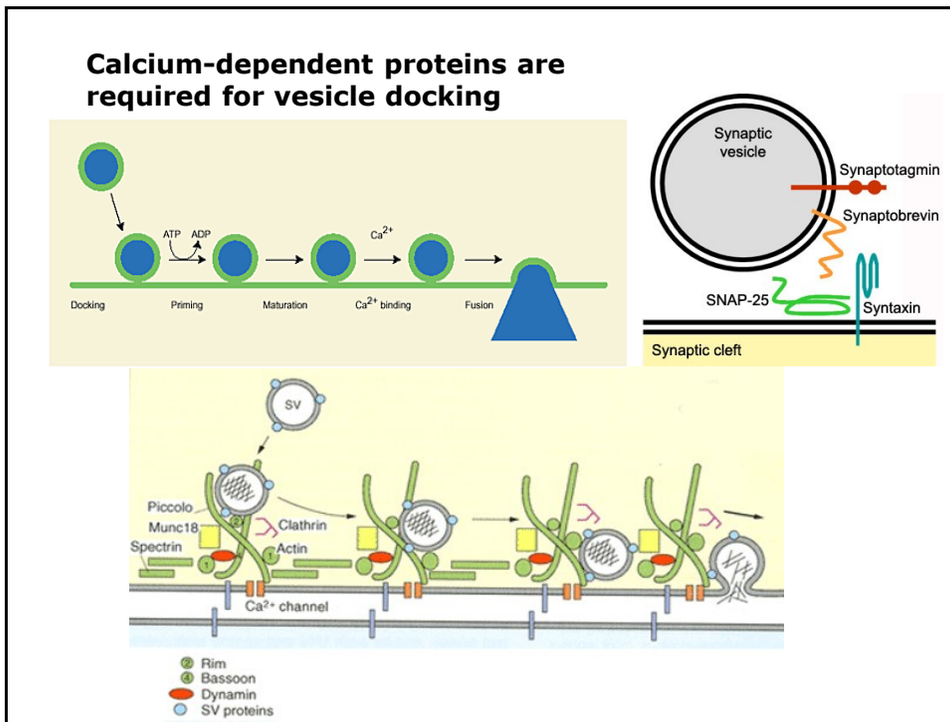
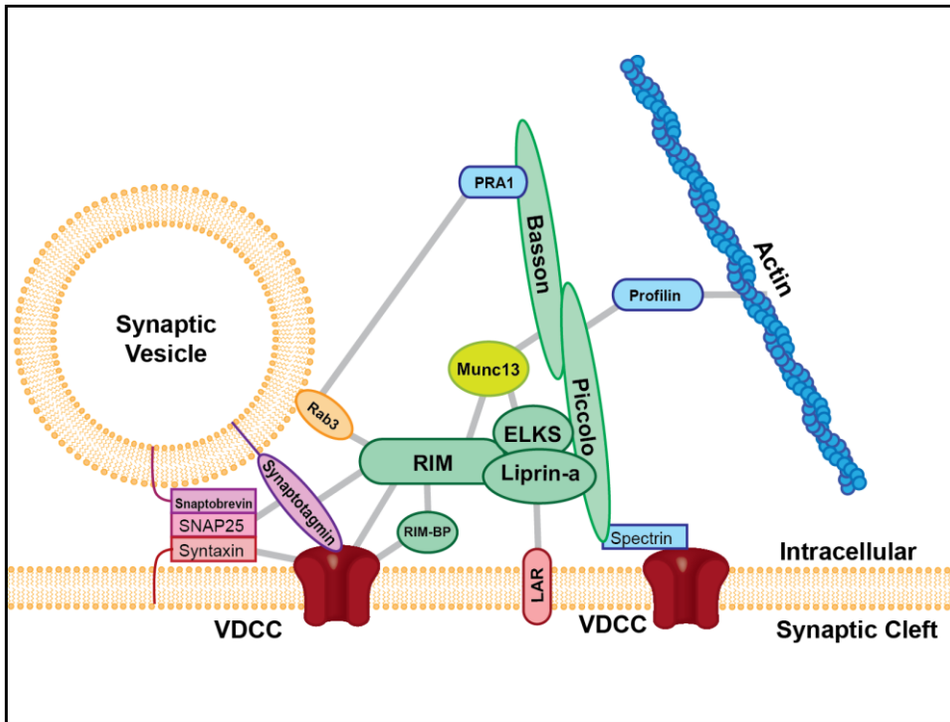
- **Second** are the known scaffolding proteins, including SAP97, and CASK/LIN-2

- **Third**, there are the active zone-specific proteins including RIM1, Munc13/unc13, Bassoon, Piccolo/Aczonin, and CAST/ERCs

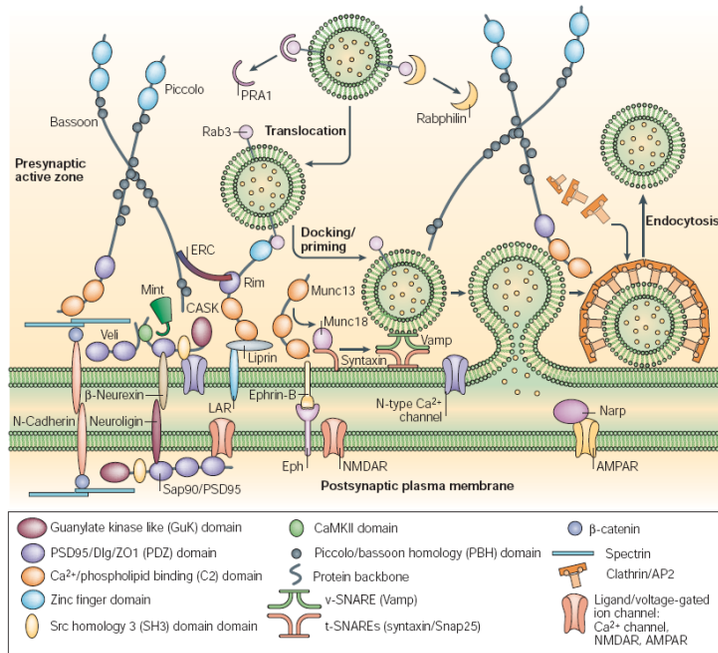
Molecular structure of a ribbon synapse



Structural molecules piccolo e bassoon anchor the synaptic ribbon on the membrane



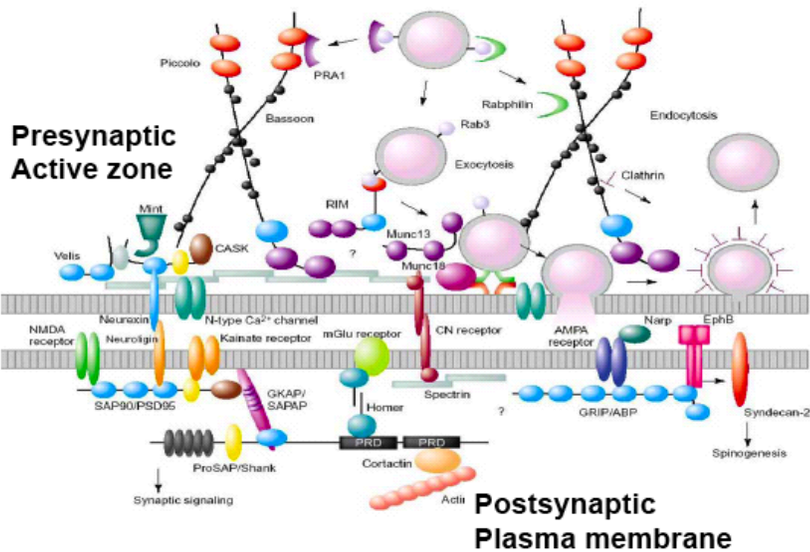
Box 1 | Molecular structure of CNS glutamatergic synapses



Current studies indicate that these distinct complexes help to define the active zone (see TABLE 1 for a list of key active

active zone cytomatrix

Proteins identified in the active zone cytomatrix can be classified into 3 categories. cytoskeleton, scaffolding, active zone proteins



Proteins identified in the active zone cytomatrix can be classified into 3 categories.

1. **cytoskeletal proteins :**
actin, tubulin, myosin, spectrin, catenin
2. **scaffolding proteins :**
SAP90/PSD95/Dlg, SAP97, CASK/LIN-2
3. **active zone-specific proteins :**
RIM1, Munc13/ unc13, Bassoon,
Piccolo/Aczonin, and CAST/ERCs

Trafficking mediated by Dynein, Kinesin, microtubule associated protein (MAP)