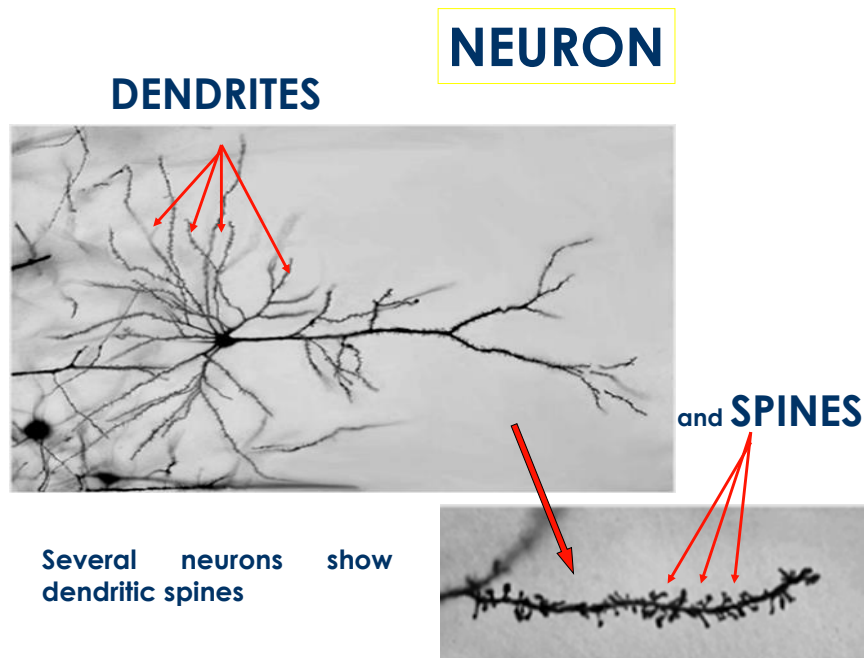
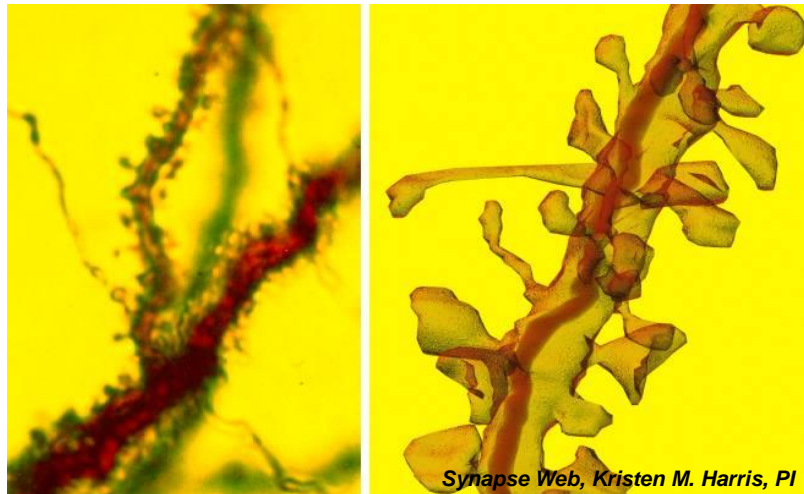


## Lesson (5)

### Inside the neuron I: Dendritic Spines

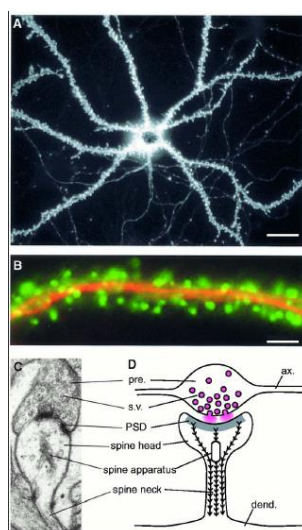


## DENDRITIC SPINES

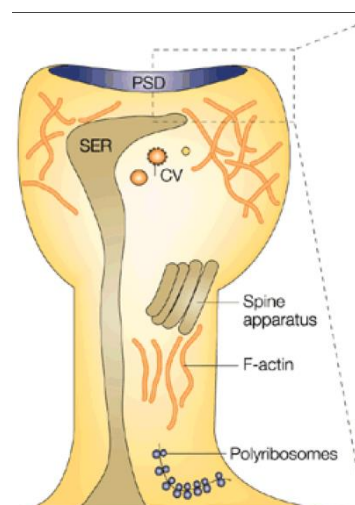


A tridimensional reconstruction from electron microscopy images

## DENDRITIC SPINES

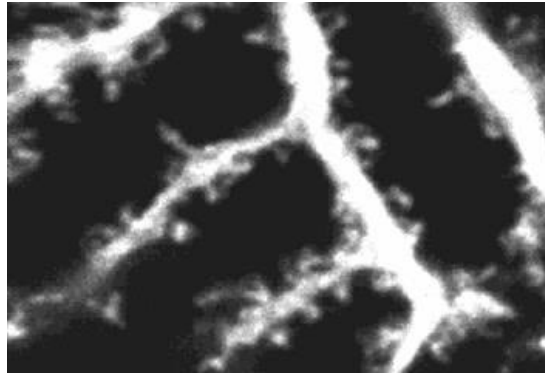


Sala et al. Neuron 2001



Hering and Sheng 2001

## DENDRITIC SPINES



movie

Dendritic spines have dynamic changes especially during early development, but also during learning

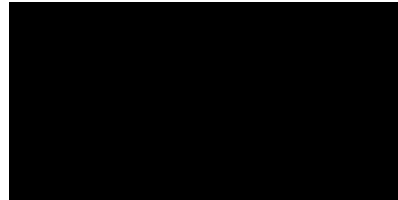
## Learning and Memory

*Neural activity plays an important role in the life history of dendritic spines by influencing the emergence of new spines, the stabilization of existing spines, and changes in spine morphology.*

Complex phenomena such as learning involve both the genesis and retraction of dendritic spines that are region-specific and temporally restricted.

## DENDRITIC SPINES

Changes in spine morphology have been observed following intense synaptic activity of the form that induces long-term potentiation (LTP), a persistent enhancement of synaptic strength that might underlie learning and memory.



movie

## DENDRITIC SPINES

Certain neurological and neuropsychiatric diseases are associated with alterations in dendritic spines.

The density and morphology of spines is abnormal in many types of mental retardation

In fragile X syndrome and Patau's syndrome, the spines have been described as long and tortuous

Down's syndrome the neurons are either completely depleted of spines or are covered with innumerable small spines with almost invisible pedicle

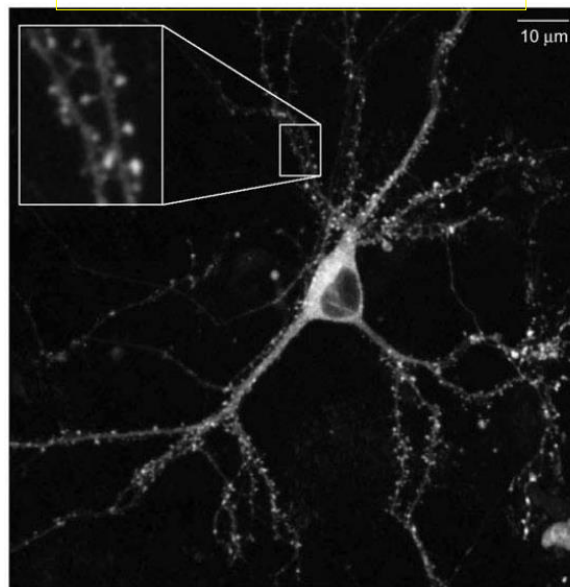
## DENDRITIC SPINES

Spine density decreases naturally during normal aging

Hormonal status also can have a substantial impact on dendritic spines.

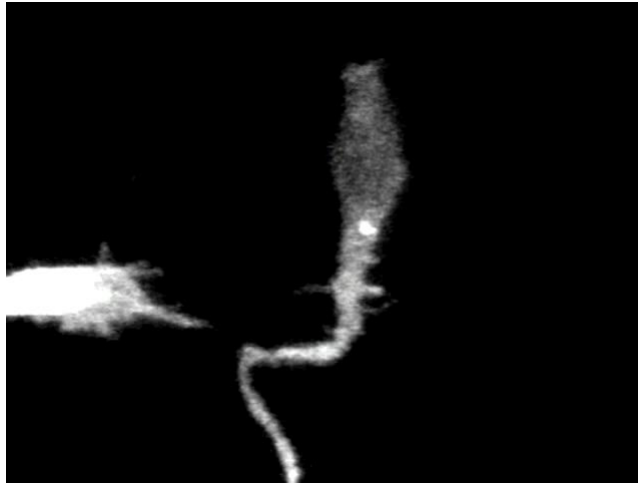
Among other factors, glutamate receptors and brain-derived neurotrophic factor (BDNF) have been implicated in activity-dependent synaptic remodeling

## DENDRITIC SPINES



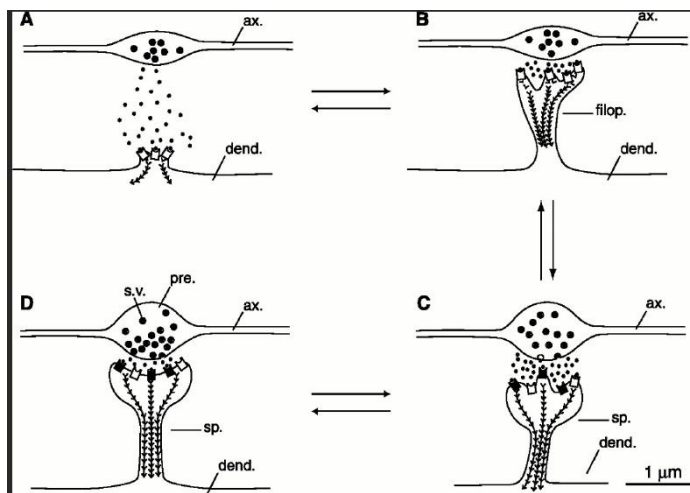
Sala et al. Neuron 2001

## Spine and filopodia formation during development



Sala et al. Neuron 2001

## DEVELOPMENT OF DENDRITIC SPINES



Sala et al. Neuron 2001

# DENDRITIC SPINES

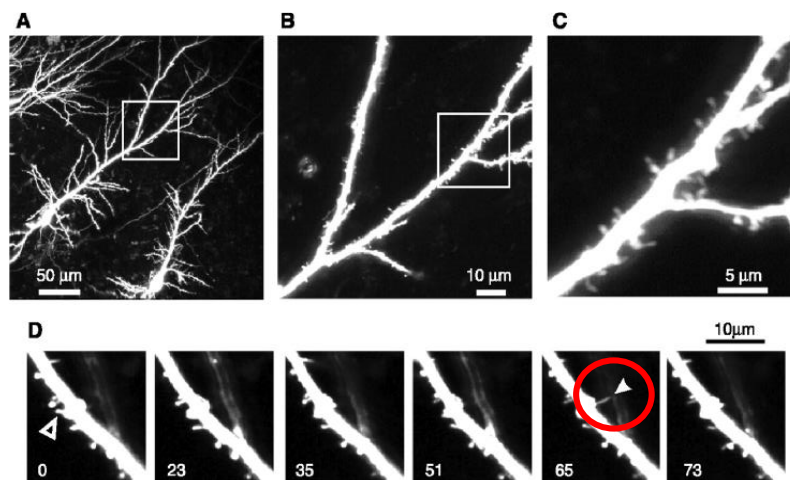
## Spine Maintenance and Pruning

Once created, spines appear to require signals from their afferent neurons to remain intact.

In many brain regions spines initially are overproduced, and are then gradually and selectively eliminated in an activity-dependent manner during later development.

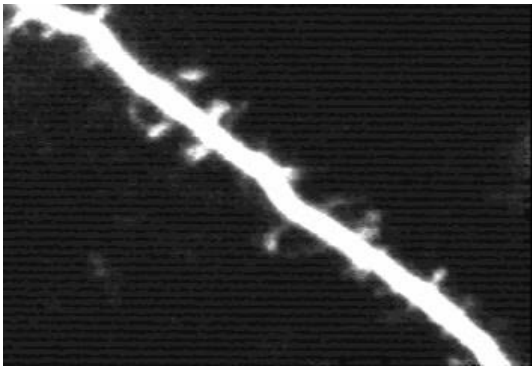
Potential functions of spine motility and dynamic actin include regulated protein scaffolding, retrograde signaling and synapse stabilization.

## Spine and filopodia formation during development



Sala et al. Neuron 2001

# Spine and filopodia formation during development



## The Disappearance of Filopodia Coincides with Appearance of Persistent Spines

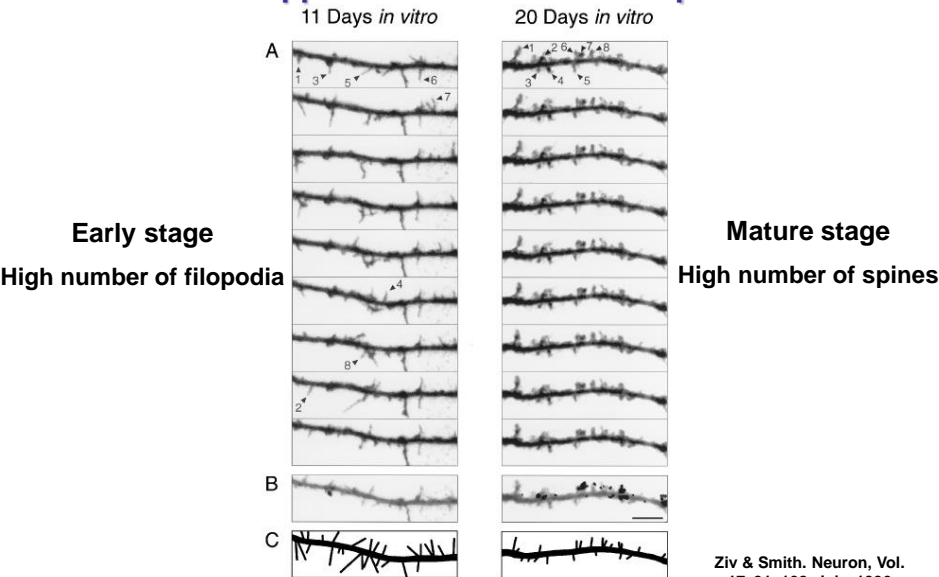


Figure 3. Time-Lapse Sequences of Dendritic Filopodia and Dendritic Spines



## DENDRITIC SPINES TYPES

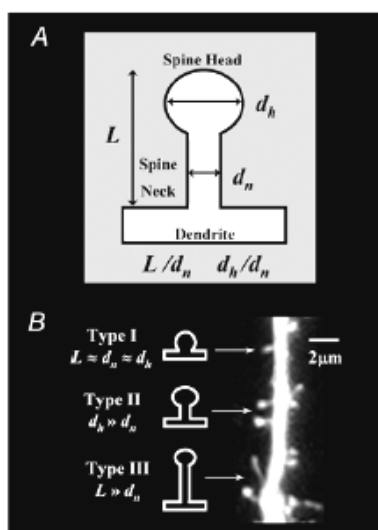
Ultrastructural studies have traditionally classified spines into major categories based on their distinct morphologies:

- stubby,
- thin,
- mushroom

The number and shape of dendritic spines are highly mutable on time scales ranging from seconds to days, suggesting that several different intrinsic mechanisms control spine dynamics.

## DENDRITIC SPINES

Classification of spine categories



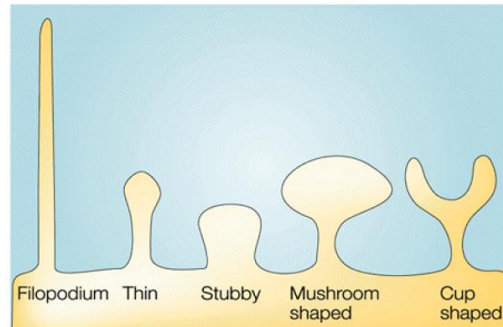
Dendritic spines were classified using the length ( $L$ ), neck diameter ( $d_n$ ) and head diameter ( $d_h$ ) of individual spines

• TYPE I : stubby spines,

• TYPE II : mushroom spines

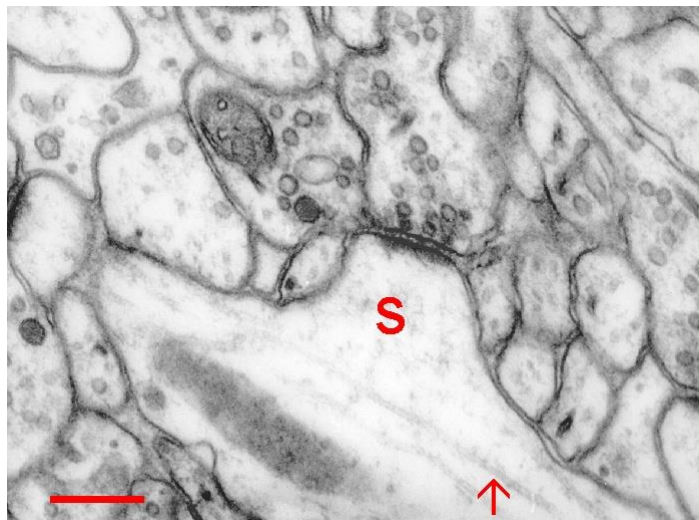
• TYPE III : thin spines,

# DENDRITIC SPINES



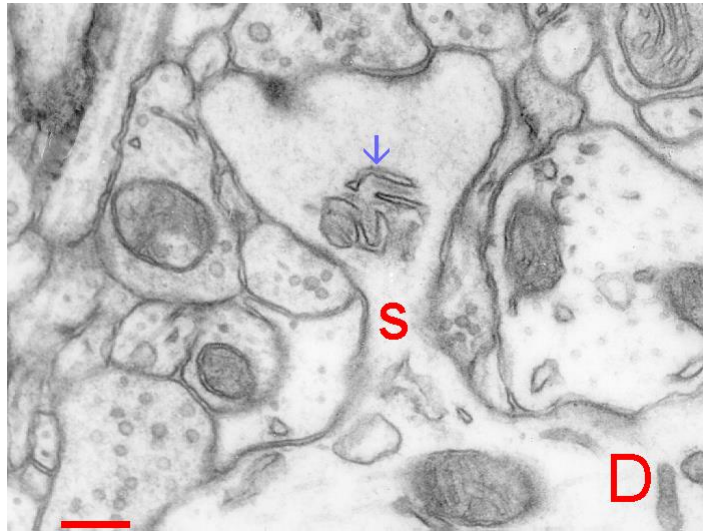
Hering and Sheng 2001

Nature Reviews | Neuroscience



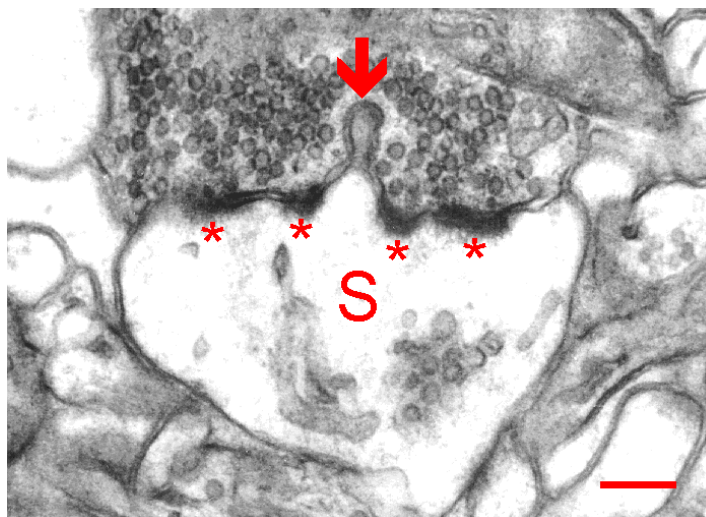
**Stubby dendritic spine (S) with axo-spinous synapse. Microtubule labeled by arrow**

from "[Atlas of Ultrastructural Neurocytology](#)"



**Mushroom-shaped dendritic spine (s) containing a spine apparatus (arrow) in its head. D**

from "[Atlas of Ultrastructural Neurocytology](#)"



**The head of a *mushroom-shaped dendritic spine* (S) with a *spinule* (arrow) protruding into the presynaptic axon terminal**

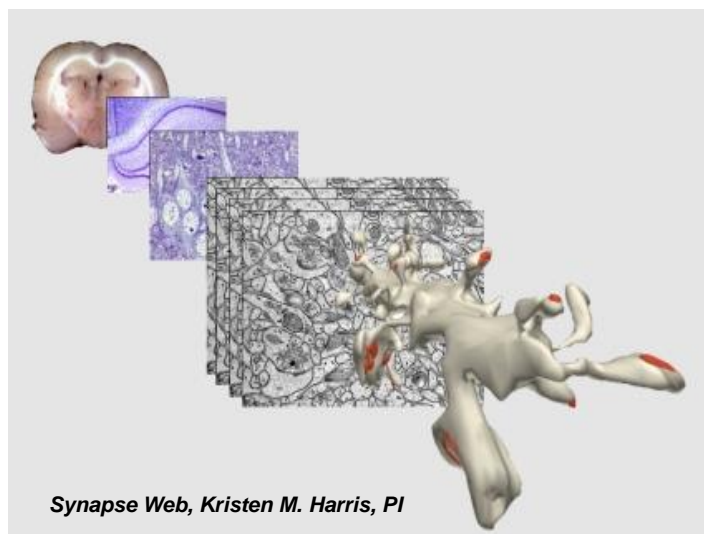
from "[Atlas of Ultrastructural Neurocytology](#)"



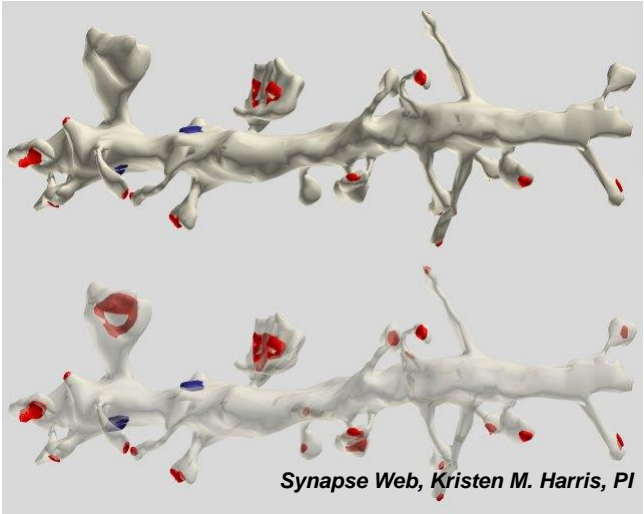
*Thin dendritic spine (S) with axo-spinous synapse.  
D - dendritic stem.*

from "[Atlas of Ultrastructural Neurocytology](#)"

### 3D reconstructions of EM pictures of spines



*Synapse Web, Kristen M. Harris, PI*



A segment of pyramidal cell dendrite from stratum radiatum (CA1) with thin, stubby, and mushroom-shaped spines.  
<http://synapses.clm.utexas.edu/>

OTHER DENDRITIC SPINES TYPES

Table 1.3 Synaptic specializations of dendrites		
Pattern	Characteristics	Examples
Varicosity	An enlargement in a thinner dendrite associated with synaptic contacts	Retinal amacrine cells
Filopodium	A long, thin protrusion with a dense actin matrix and few internal organelles	All neurons during developmental synaptogenesis
Simple spine	Sessile	Sessile spine
	Stubby spine	Cerebral pyramidal cells
	Crook thorn	Neurons of dentate nucleus
Pedunculated	Bulbous enlargement at tip	Cerebral pyramidal cells
	Thin spine	Cerebral pyramidal cells
	Mushroom spine	Cerebral pyramidal cells
Branched spine	Gemmule	Olfactory bulb granule cell
	Each branch has a unique presynaptic partner and each branch has the shape characteristics of a simple spine	CA1 pyramidal cells Granule cells of dentate gyrus Cerebellar Purkinje cells
Synaptic crest	Crest-like protrusion with a synapse on either side of a thin lamellar neck region	Cerebral pyramidal cells Neurons of habenula, subfornical organ, and interpeduncular nucleus

Table 1.3 (continued) Synaptic specializations of dendrites		
Pattern	Characteristics	Examples
Claw ending	Synaptic protrusions at the tip of the dendrite associated with one or more glomeruli	Granule cells of cerebellar cortex and dorsal cochlear nucleus
Brush ending	Spray of complex dendritic protrusions at the end of dendrite that extends into glomerulus and contains presynaptic elements	Unipolar brush cells of cerebellar cortex and dorsal cochlear nucleus
Thorny excrescence	Densely lobed dendritic protrusion into a glomerulus	Proximal dendrites of CA3 pyramidal cells and dentate gyrus mossy cells Proximal dendrites of thalamocortical relay cells
Racemose appendage	Twig-like branched dendritic appendages that contain synaptic varicosities and bulbous tips	Inferior olivary neurons Relay cells of lateral geniculate nucleus
Coralline excrescence	Dendritic varicosity extending numerous thin protrusions, velamentous expansions and tendrils	Neurons of dentate nucleus and lateral vestibular nucleus

# DENDRITIC SPINES

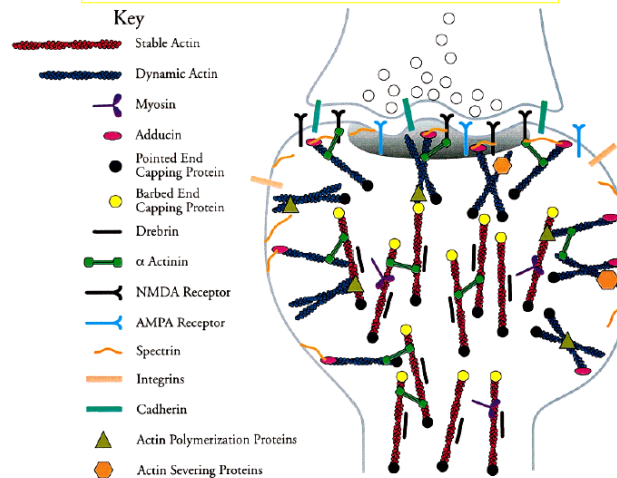
The spine cytoskeleton is based on actin filaments. Intermediate filaments and microtubules, the other two main filament systems of the cytoskeleton, are rare or nonexistent in spines.

Actin filaments are extremely enriched in spines relative to other parts of the mature neuron

# DENDRITIC SPINES

Protein	Concentration	Evidence
β-actin	Enriched	Light microscopy
γ-actin	Enriched	Light microscopy
α-actinin	Enriched	Light microscopy
α-adducin	Present	Electron microscopy
ADF	Present	Electron microscopy
Arp2/3 complex	Predicted	
Calcineurin	Present	Electron microscopy
Calmodulin	Present	Electron microscopy
Calpain	Present	Electron microscopy
CaMKII	Present	Light microscopy
Capping protein	Predicted	
Cofilin	Predicted	
Drebrin	Enriched	Electron microscopy
Ena/VASP family members	Predicted	
Fyn	Present	Subcellular fractionation
Gelsolin	Predicted	
α internexin	Present	Electron microscopy
Myosin IIB	Present	Light microscopy
NAP22	Present	Electron microscopy
NF-L subunit of neurofilament	Present	Subcellular fractionation
PKA	Present	Subcellular fractionation
PKC-γ	Present	Electron microscopy
Profilin	Predicted	
PP1 α	Enriched	Electron microscopy
PP1 γ1	Enriched	Electron microscopy
RC3/neurogranin	Present	Electron microscopy
Synaptopodin	Enriched	Electron microscopy
Spectrin	Present	Light microscopy
Spinophilin/neurabin II	Enriched	Electron microscopy
Tropomodulin	Predicted	
Tropomyosin	Predicted	
WASP family members	Predicted	

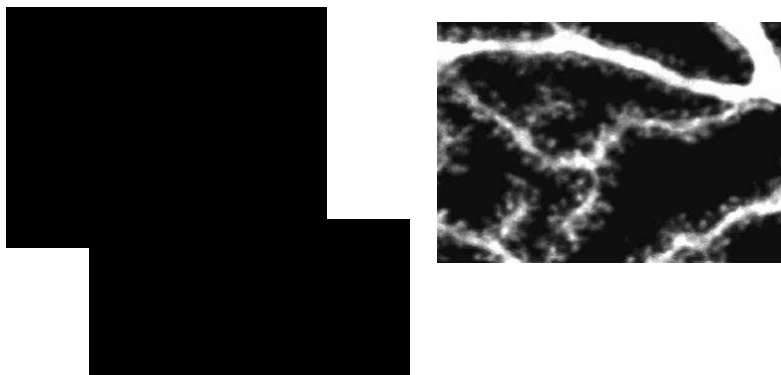
# DENDRITIC SPINES



Spines contain two distinct pools of actin filaments (one stable, the other unstable) that provide the spine with both a stable core structure and a dynamic, complex shape

## Regulation of dendritic spines motility

Motility is inhibited by blockers of synaptic transmission. AMPA receptor activation causes the influx of  $\text{Na}^+$  ions through receptor-associated channels, postsynaptic membrane depolarization and spine motility.





## Signalling cascade that regulates spine shape and motility

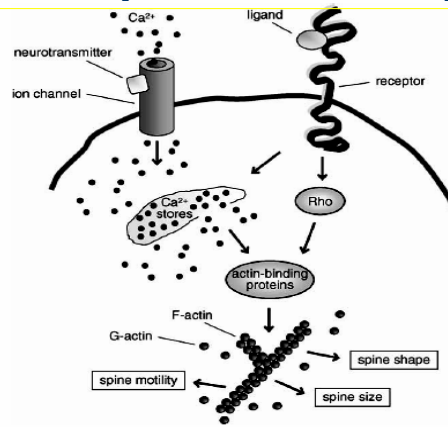
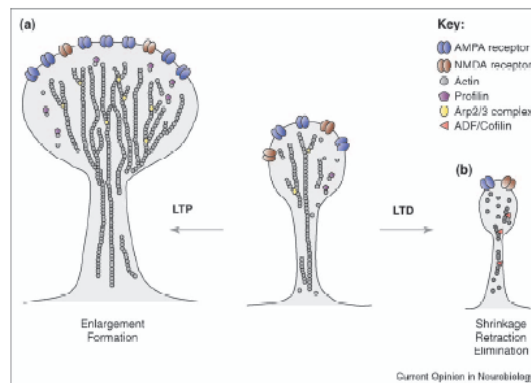


Fig. 3. The signaling cascades that regulate dendritic spine shape and motility involve cell surface receptors and ion channels, which activate signaling cascades controlling the activity of Rho GTPases and the  $\text{Ca}^{2+}$  concentration within the spine.  $\text{Ca}^{2+}$  permeable ion channels include neurotransmitter-gated ion channels, as shown, and voltage-gated  $\text{Ca}^{2+}$  channels (not shown). Cell surface receptors include receptor tyrosine kinases and cell adhesion and recognition molecules. These receptors can bind soluble ligands, as shown, or ligands anchored to adjacent cell surfaces or to the extracellular matrix (not shown). *LM. Ethell, E.B. Pasquale/Progress in Neurobiology 75 (2005) 161–205*

## Changes in actin polymerization with LTP and LTD leads to changes in spine shape



Changes in actin polymerization and spine morphology with LTP and LTD. (a) LTP is associated with a shift of actin equilibrium toward F-actin (F-actin is depicted as linear chains of monomeric G-actin [single circle]) in spines, enlargement of the spine head, and recruitment of more AMPA receptors to the postsynaptic membrane. Profilin promotes actin filament assembly by increasing the availability of actin-ATP for polymerization. The Arp2/3 complex stimulates nucleation of new actin filaments and formation of branches. (b) By contrast, LTD stimulation shifts the equilibrium toward actin depolymerization, resulting in shrinkage or loss of spines. The actin severing protein ADF/cofilin might be involved in spine shrinkage.

Tada & Sheng. Current Opinion in Neurobiology 2006, 16:95–101



## SUMMARY 1

Dendritic spines undergo several types of transformations, ranging from growth to collapse, and from elongation to shortening, and they experience dynamic morphological activity on a rapid time scale.

Changes in spine number and morphology occur under pathological conditions like excitotoxicity, but also during normal central nervous system development, during hormonal fluctuations, and in response to neural activity under physiological circumstances.

## PROPOSED FUNCTIONS OF SPINES

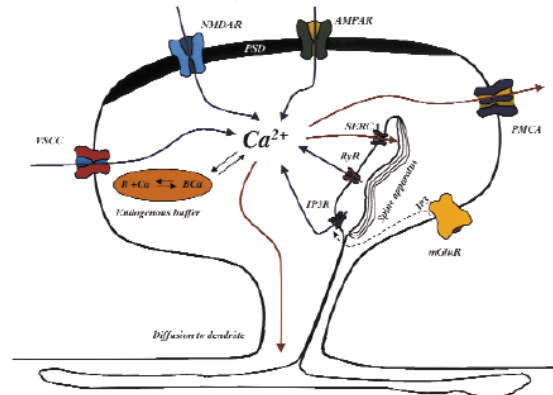
Dendritic spines are targets of most excitatory inputs in the central nervous system (CNS) and are morphologically heterogeneous

The spine is suggested to act as a biochemical compartment which isolates subsynaptic activity-driven calcium ion flux from the rest of the dendrite and therefore also from other nearby synapses .

The restriction of the calcium pulse to the spine is believed to allow localized signaling cascades, which act on spine constituents to induce local modifications of spine structure and composition.

## Ca<sup>2+</sup> entry in dendritic spines

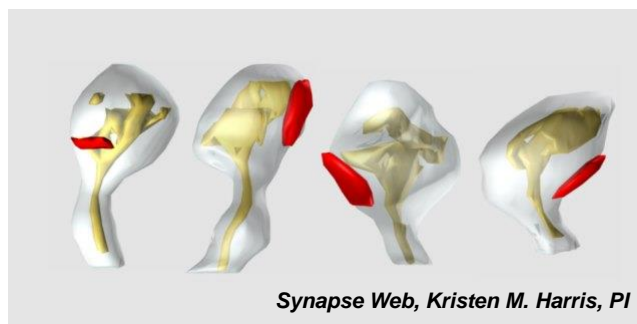
Fig. 4. Model of a spine. Cartoon illustrating the rich diversity of mechanisms that control calcium kinetics in spines. Blue arrows depict the influx of calcium into the spine; red arrows show calcium efflux. Influx can occur through NMDA channels, voltage-sensitive calcium channels (VSCCs) and calcium-permeable AMPA receptors. Calcium can also enter the cytoplasm from internal stores through a ryanodine receptor-dependent mechanism or through an IP<sub>3</sub>R after activation of metabotropic glutamate receptors. In the cytoplasm, calcium can bind to calcium buffers (orange, B) whose kinetics and affinities alter the shape of the calcium transient. Calcium then leaves the spine by extrusion mechanisms such as the plasma membrane calcium ATPase (PMCA), which pumps calcium into the extracellular space, or the SERCA pump, which sequesters calcium into intracellular stores. Calcium can also dissipate by diffusing through the thin spine neck into the adjacent dendrite.



Ca<sup>2+</sup> entering through NMDA receptor channels could initiate a feedback loop in which actin-driven changes in spine shape influence the efficacy of transmission at individual synapses.

Calcium-sensitive interactions between the actin-binding protein  $\alpha$ -actinin

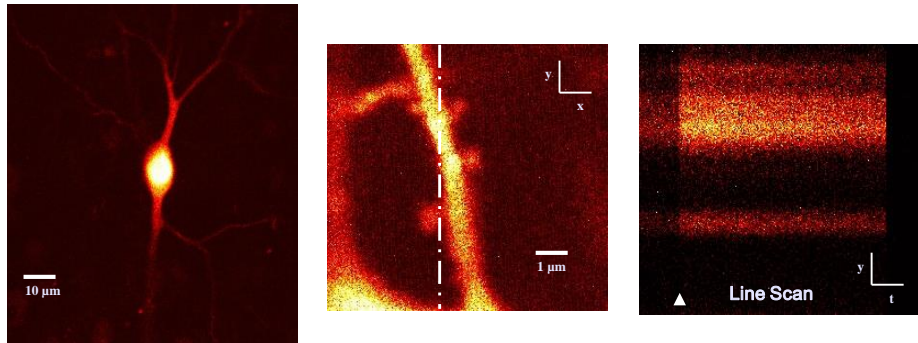
## Ca<sup>2+</sup> stores in dendritic spines



Dendritic spines are rich in smooth endoplasmic reticulum (known as the [spine apparatus](#)). This is found in mushroom spines of hippocampus, neocortex and cerebellum.

The spine apparatus plays a major role as Ca<sup>2+</sup> store in spines

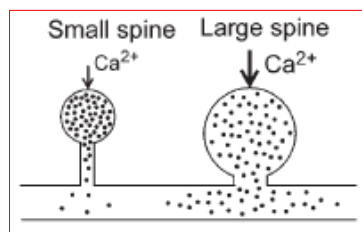
## Confocal Laser Scanning Microscopy of CA1 hippocampal dendritic spines in rat organotypic slices



Pablo d'Alcantara, PhD  
Department of Neurophysiology  
National Institute for Medical Research  
London, United Kingdom

## Spine necks regulate biochemical and electrical signals in large and small spines

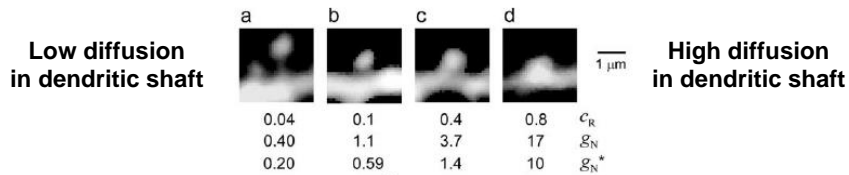
Calcium influx into spines is mediated by calcium channels and by NMDA and AMPA receptors and is followed by fast diffusional equilibration within the spine head



Compartmentalization of  $\text{Ca}^{2+}$  within the spine head is controlled by spine neck dimensions in both mushroom and thin spines of CA1 pyramidal cells: spines that have narrower or longer necks appear to retain more  $\text{Ca}^{2+}$  in their heads following synaptic activation than do wider shorter spines.

## Relation between spine type and $\text{Ca}^{2+}$

- Long and thin spines (type-III) spines may isolate  $\text{Ca}^{2+}$  transients from the parent dendrite and other spines



- Shorter and stubbier spines (type-I) are thought to promote more coordinated and widespread  $\text{Ca}^{2+}$  transients in the parent dendrite, as well as to coordinate  $\text{Ca}^{2+}$  signalling among adjacent spines,

## SUMMARY

Dendritic spines are biochemical microcompartments that are isolated from their parent dendrites and neighboring spines. Spines compartmentalize  $\text{Ca}^{2+}$  and perhaps other messengers, such as  $\text{IP}_3$  and  $\text{Na}^+$ .

Imaging studies of spine  $[\text{Ca}^{2+}]$  dynamics have revealed that  $\text{Ca}^{2+}$  can enter spines through voltage-sensitive and ligand-activated channels, as well as through  $\text{Ca}^{2+}$  release from intracellular stores.

In thin spines  $\text{Ca}^{2+}$  is more strictly compartmentalized

In large, short stubby spines  $\text{Ca}^{2+}$  is less strictly compartmentalized and can diffuse more easily in the parental dendrite

## CURRENT RESEARCH

- Neuronal dendritic arbor development and dendritic integration into functional circuits is a critical step in brain development and plasticity
- The molecular mechanisms control of this process remain unknown.