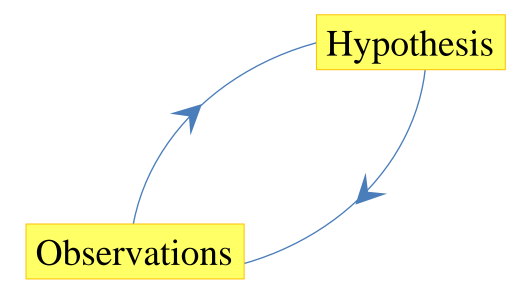
Introduction to Systems Neuroscience

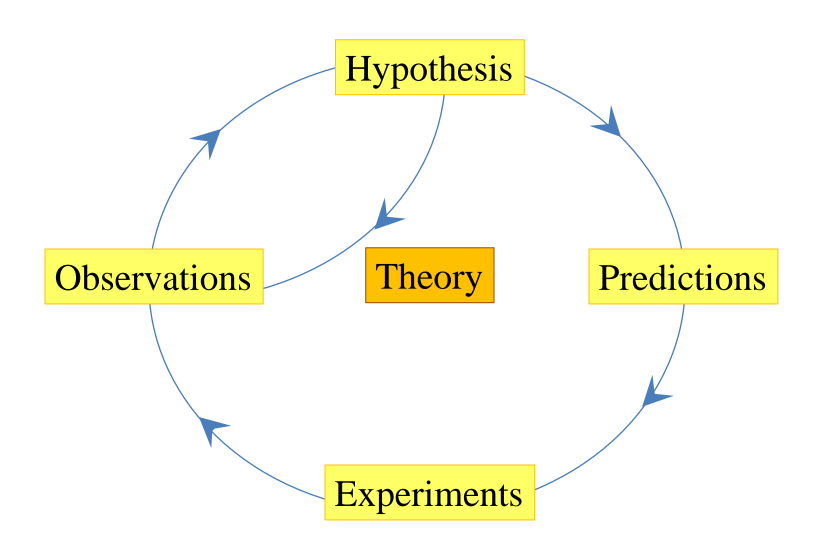
Methodologies used to study brain systems

Ehud Ahissar
Department of Neurobiology

- In natural sciences, unlike in mathematics, there is **no set** of **pre-defined axioms** from which other definitions can be derived and proved.
- What can be proven in natural sciences are only consistencies; hypotheses are considered valid as long as they are consistent with actual observations (e.g., gravity is considered valid as long as the movement patterns it entails are consistent with actual movements of bodies).
- The methods are thus developed to test consistencies of observations with assumptions



The scientific method



Research approaches

- Observational: we collect enough evidence to compose a hypothesis
- Correlational: we compare 2 variables the values of which have been collected without direct intervention. No causal relationships can be directly concluded.
- Experimental: an "independent" variable is systematically manipulated and the effects of this on a "dependent" variable are measured. Considered as the best approach for revealing causal relationships, although...

General guiding assumptions

- All mental functions are carried out in the brain
- All brain components are relevant
- Brain processes obey the rules of physics

Common working hypotheses

- The basic component of processing is a single neuron
- Neural processing is mediated by neurotransmitters, brain states are controlled by neuromodulators
- Memories are stored in synapses

Major difficulties

- Complexity
 - many levels (ions, ..., neurons, ..., systems)
 - many components (e.g., 10¹¹ neurons, 10¹⁵ synapses)
 - many variables (physical, chemical, mechanical, electrical, physiological, psychological)
- Small sizes (neurons ~ 10 μm, synapse <1 μm)
- Closed loops
- Self organization
- plasticity

Cannot repeat an experiment twice

Major difficulties - Methods to address them

- Complexity
 - many levels (ions, ..., ne many components (e.g
 many components (e.g
 rons, 10¹⁵ synapses)

 - many variables (physical, chemical, mechanical, electrical, physiological, psychological)
- Closed loops
- Self organization
- plasticity

• Small sizes (neurons Small tools, magnifications, genetic tools

Opening the loops by anesthesia, flashed stimuli, cuts

Cannot repeat an experiment twice

Using statistics

About this lecture

- Review of popular methods for experiments in systems neuroscience
- Focus on resolutions and potential power. Less on limitations
- Not all methods will be covered. You are invited to complete the picture through the web or text books
- Slide numbers appear in MOST of the slides, not in all...

		Spatial re	solution		Te	emporal i	resoluti	on
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	<u>device</u>	<u>signal</u>	<u>spikes</u>	heuristic
behavior	brain	brain	> 10 ¹¹	station	1 ms	10 ms	10	
2DG, c-fos	10 μm	10 μm	1		NA	30 min	> 10 ⁶	
fMRI	> 1 mm	> 200 µm	> 10 ⁵		1 s	> 10 s	1000	10 ms
EEG	10 mm	100 mm	> 10 ⁹	station	< 1 ms	50 ms	50	1 ms
MEG	10 mm	100 mm	> 10 ⁹		1 ms	50 ms	50	
ECoG	10 mm	1-10 mm	> 10 ⁴⁻⁶		< 1 ms	10 ms	10	
Intr. Signal	10 μm	200 μm	> 1000		< 1 ms	1 s	1000	10 ms
VSD Imaging	10 μm	30 μm	< 100		< 1 ms	10 ms	< 1	
μElec	50 μm	10 μm	1	10 μm	< 1 ms	1 ms	1	
μDialysis	< 1 mm	100 μm	> 10 ⁴		> 1 min	100 ms	> 10 ⁶	
Intracell elec	10 μm	10 μm	< 1		< 1 ms	< 1 ms	< 1	
Ca imaging	1 μm	1 μm	< 1		1 ms	> 100 ms	> 100	

Stimulating / perturbing neural activity

		Spatial resolution				Temporal resolution				
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	device	signal	<u>spikes</u>	heuristic		
sensory	modality	modality	> 10 ⁹	station	< 1 ms	< 1 ms	< 1			
TMS	100 mm	10 mm	> 10 ⁹	station	< 1 ms	100 ms	100			
μStim	< 10 µm	> 100 µm	> 50		< 1 ms	10 ms	10			
μPharmac	< 10 µm	> 100 µm	> 50		1 ms	> 10 s	10			
single cell	< 10 µm	< 10 µm	1		< 1 ms	< 1 ms	< 1			
sub-cell	>100 μm	< 1 µm	> 50		< 1 ms	< 1 ms	< 1			

Measuring structure (anatomy)

	Spatial resolution				Temporal resolution			
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	device	signal	<u>spikes</u>	heuristic
cell density								
receptor dens	receptor density							
transmitter de	ensity							
tract tracing								
single-cell								

Manipulating structure

		Spatial resolution					Temporal resolution			
	<u>device</u>	levice signal neurons heuristic				<u>device</u>	signal	<u>spikes</u>	heuristic	
Neuropsychology		> 10 mm	> 10 ⁷				months			
lesions	> 100 µm	> 100 μm	> 50			>1s	> 1 min			

		Spatial re	solution		Temporal resolution				
	device	signal	neurons	heuristic	device	signal	spikes	heuristic	
behavior	brain	brain	> 10 ¹¹	station	1 ms	10 ms	10		
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EEG	10 mm	100 mm	> 10 ⁹	station	< 1 ms	50 ms	50	1 ms	
MEG	10 mm	100 mm	> 10 ⁹		1 ms	50 ms	50		
ECoG	10 mm	1-10 mm	> 10 ⁴⁻⁶		< 1 ms	10 ms	10		
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VSD Imaging	10 μm	30 μm	< 100		< 1 ms	10 ms	< 1		
μElec	50 μm	10 μm	1	10 μm	< 1 ms	1 ms	1		
μDialysis	< 1 mm	100 μm	> 10 ⁴		> 1 min	100 ms	> 10 ⁶		
Intracell elec	10 μm	10 μm	< 1		< 1 ms	< 1 ms	< 1		
Ca imaging	1 μm	1 μm	< 1		1 ms	> 100 ms	> 100		

Measuring neural activity

Behavior

- Present sensory stimuli
- Measure response accuracy, threshold, speed
- Infer related neural activity

Example: odor sensitivity

- Present mixture of 2 odors
- Ask if the mixture is more similar to A or B (2AFC)
- Response accuracy: randomize stimuli and compute a psychometric curve (% correct as a function of ΔO)
- Discrimination threshold: use staircase paradigm and wait for stabilization
- Perceptual speed: measure reaction time
- Infer relevant neuronal pathways and stations by known constrains

heuristics → Station resolution by known functional anatomy

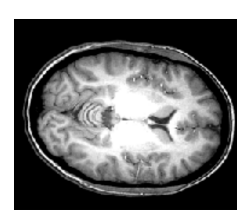
		Spatial re	solution		Te	emporal i	resoluti	on
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	<u>device</u>	<u>signal</u>	<u>spikes</u>	heuristic
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EEG	10 mm	100 mm	> 109	station	< 1 ms	50 ms	50	1 ms
MEG	10 mm	100 mm	> 10 ⁹		1 ms	50 ms	50	
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μDialysis	< 1 mm	100 μm	> 10 ⁴		> 1 min	100 ms	> 10 ⁶	
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		Spatial re	solution		Te	emporal i	resoluti	on
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	<u>device</u>	<u>signal</u>	<u>spikes</u>	heuristic
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Ca imaging	1μm	1 μm	< 1		1 ms	> 100 ms	> 100	

MRI vs. fMRI

MRI

high resolution (1 mm)

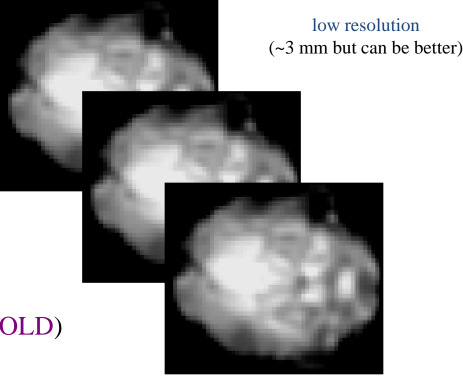


one image

fMRI

Blood Oxygenation Level Dependent (BOLD) signal indirect measure of neural activity

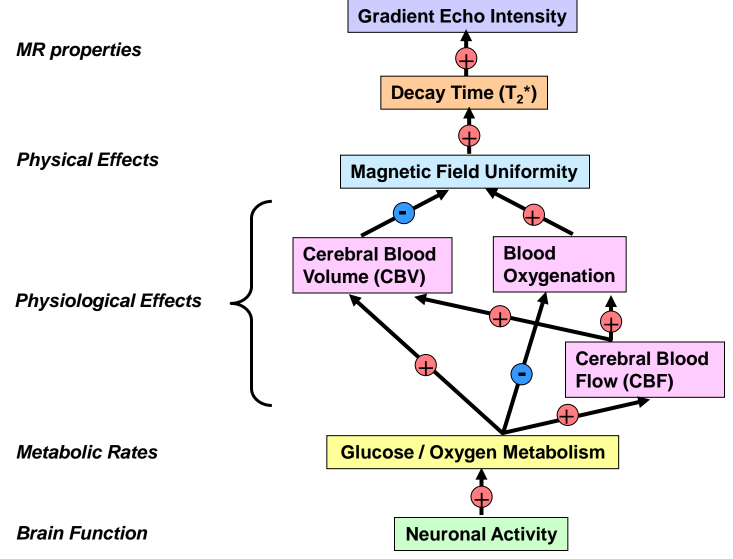
fMRI

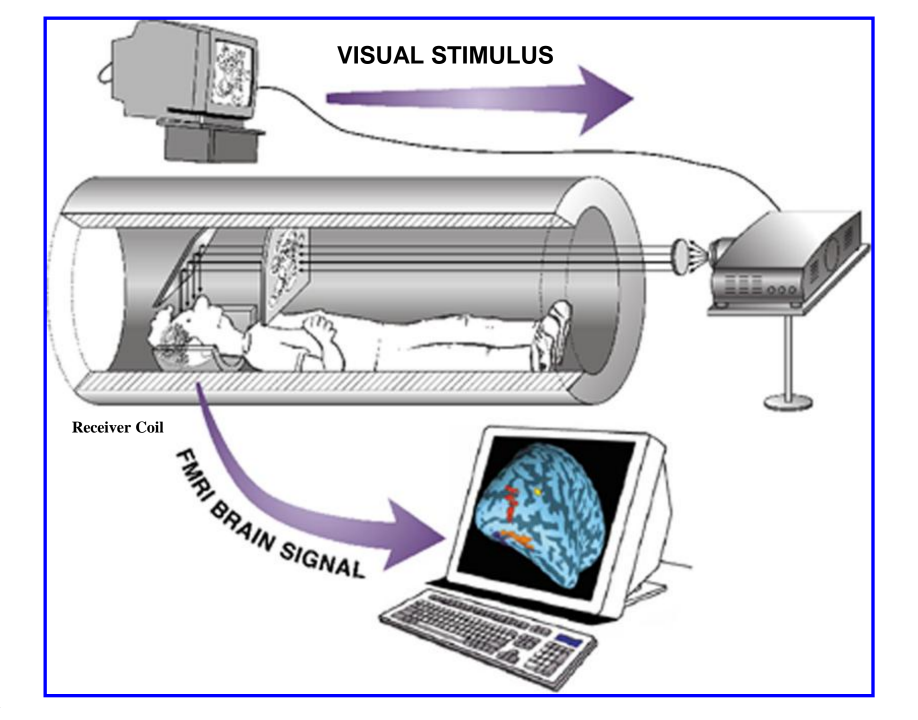


many images (e.g., every 2 sec for 5 mins)

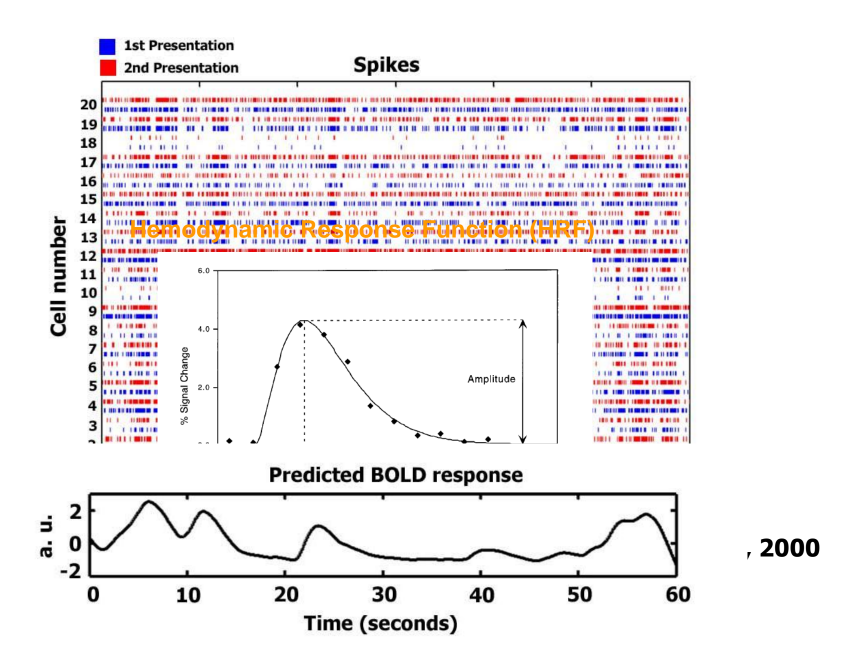
↑ neural activity → ↑ blood oxygen → ↑ fMRI signal

fMRI using BOLD





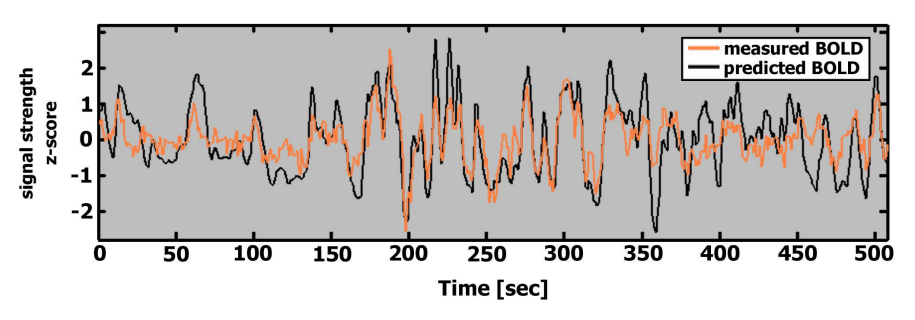
From spike trains to fMRI BOLD predictor:



Predicting fMRI BOLD signal in one subject from spike activity in another subject during the same movie

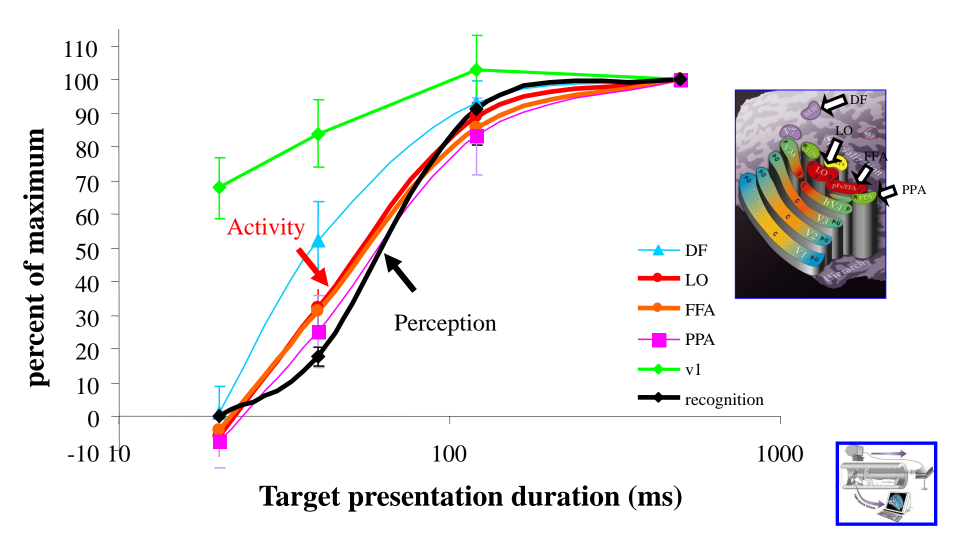






Correlation = 0.73, p<0.001

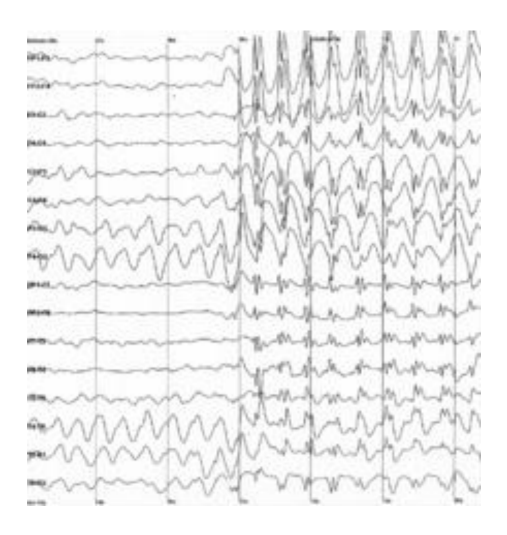
Non-linear amplification revealed in fMRI signals



Backward masking

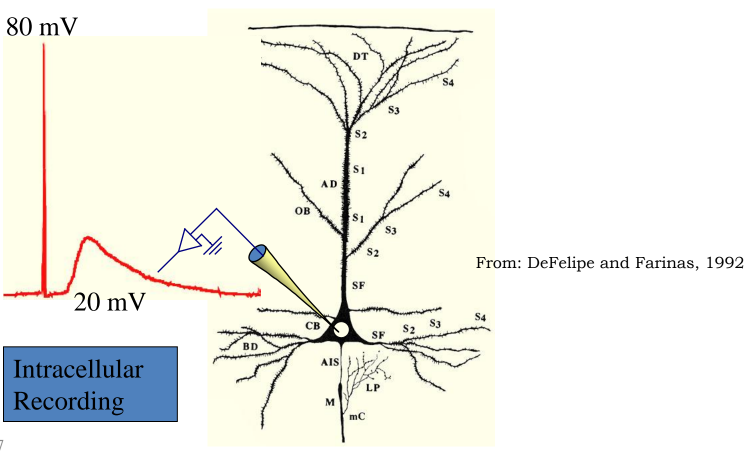
		Spatial re	solution		Te	emporal i	resoluti	on
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	<u>device</u>	signal	spikes	heuristic
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EEG	10 mm	100 mm	> 10 ⁹	station	< 1 ms	50 ms	50	1 ms
MEG	10 mm	100 mm	> 10 ⁹		1 ms	50 ms	50	
ECoG	10 mm	1-10 mm	> 10 ⁴⁻⁶		< 1 ms	10 ms	10	
Intr. Signal	10 μm	200 μm	> 1000		< 1 ms	1 s	1000	10 ms
VSD Imaging	10 μm	30 μm	< 100		< 1 ms	10 ms	< 1	
μElec	50 μm	10 μm	1	10 μm	< 1 ms	1 ms	1	
μDialysis	< 1 mm	100 μm	> 10 ⁴		> 1 min	100 ms	> 10 ⁶	
Intracell elec	10 μm	10 μm	< 1		< 1 ms	< 1 ms	< 1	
Ca imaging	1 μm	1 μm	< 1		1 ms	> 100 ms	> 100	





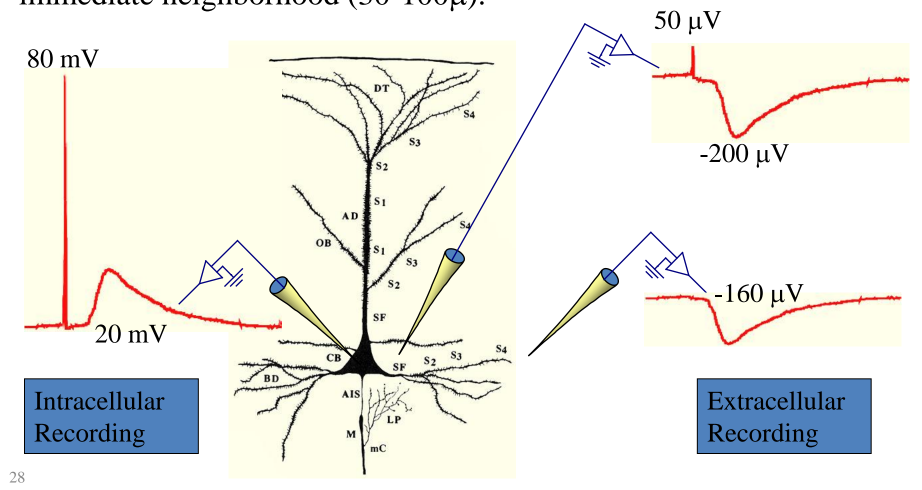
Epileptic spike and wave discharges monitored with EEG.

Dipole Formation in EEG Potentials

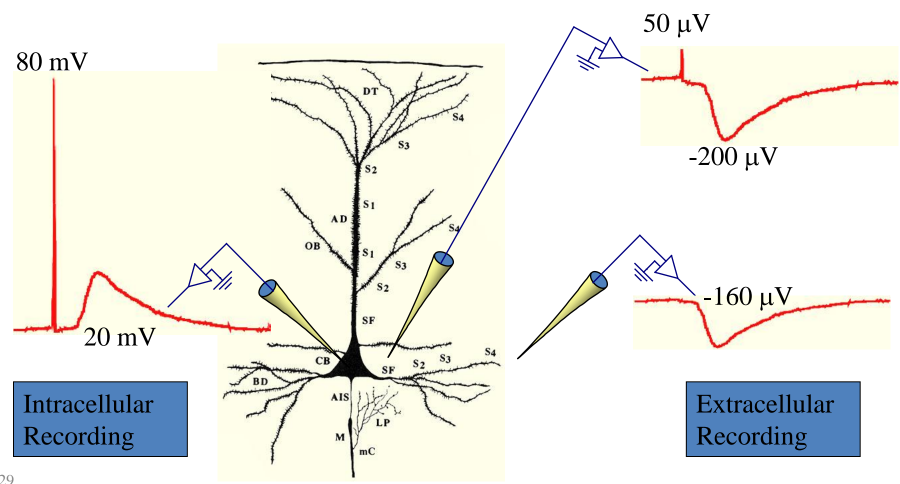


PRINCIPLES of VOLUME CONDUCTION

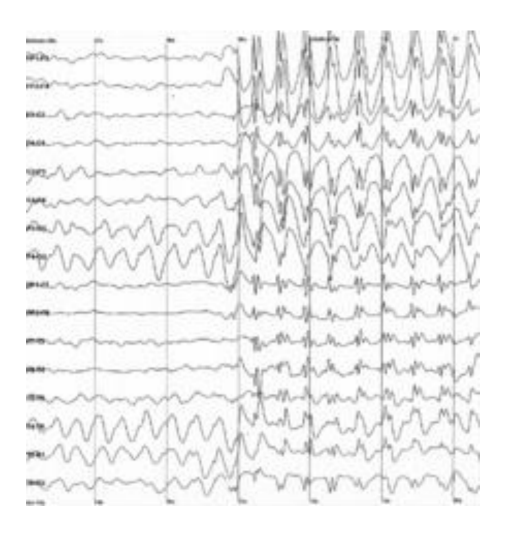
The amplitude of spikes strongly decay, and therefore, they do not summate to levels that is detectable by electrodes not located in their immediate neighborhood (30-100 μ).



EEG signals reflect synchronous waves of dendritic activities

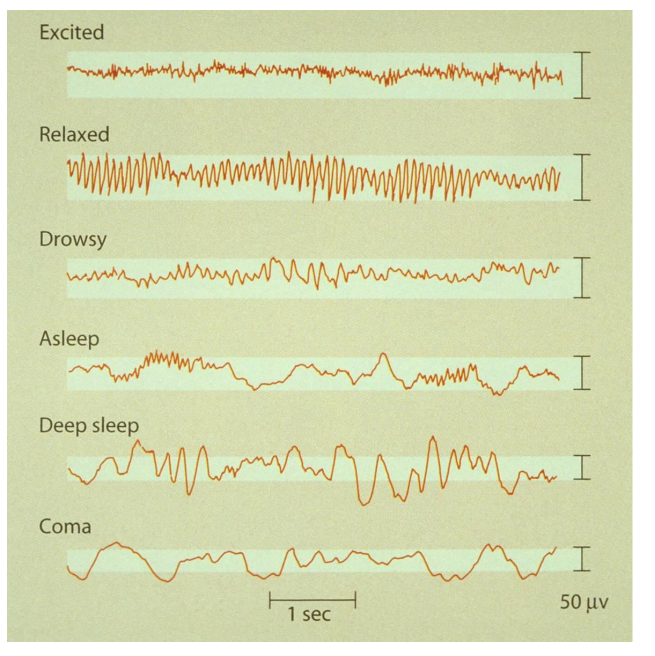






Epileptic spike and wave discharges monitored with EEG.

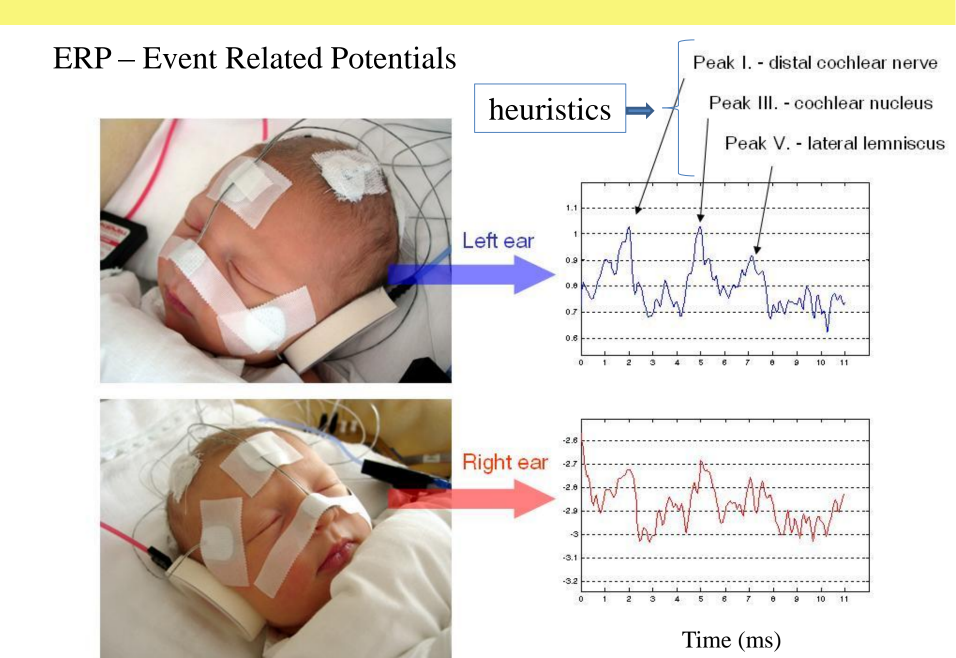
EEG & brain states



From: Jasper, 1956

ERP – Event Related Potentials

When an event is repeated tens or hundreds of times, and the time-related EEG signals are averaged, the resulting signal is considered as the average potential evoked by the event



		Spatial re	solution		Temporal resolution				
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	<u>device</u>	<u>signal</u>	<u>spikes</u>	heuristic	
behavior	brain	brain	> 10 ¹¹	station	1 ms	10 ms	10		
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μElec	50 μm	10 μm	1	10 μm	< 1 ms	1 ms	1		
μDialysis	< 1 mm	100 μm	> 10 ⁴		> 1 min	100 ms	> 10 ⁶		
Intracell elec	10 μm	10 μm	< 1		< 1 ms	< 1 ms	< 1		
Ca imaging	1 μm	1 μm	< 1		1 ms	> 100 ms	> 100		

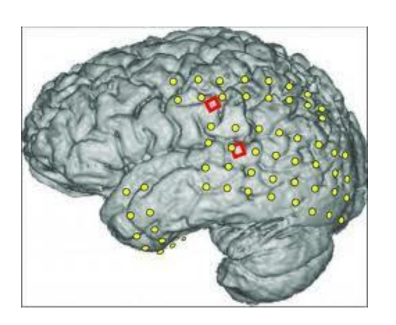
- MEG picks up magnetic fields generated by ionic currents in the brain
- Like EEG, it will show meaningful signals only for synchronized and coordinated currents.
- Like EEG, the major source of these sugnals are synaptic currents in cortical pyramidal neurons
- Unlike EEG, MEG is sensitive to the direction of the summed current
- As a result, comparison of MEG and EEG can increase the resolution of source localization
- So far, source localization is very limited.

		Spatial re	solution		Te	emporal i	resoluti	on
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	<u>device</u>	<u>signal</u>	<u>spikes</u>	heuristic
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μElec	50 μm	10 μm	1	10 μm	< 1 ms	1 ms	1	
μDialysis	< 1 mm	100 μm	> 10 ⁴		> 1 min	100 ms	> 10 ⁶	
Intracell elec	10 μm	10 μm	< 1		< 1 ms	< 1 ms	< 1	
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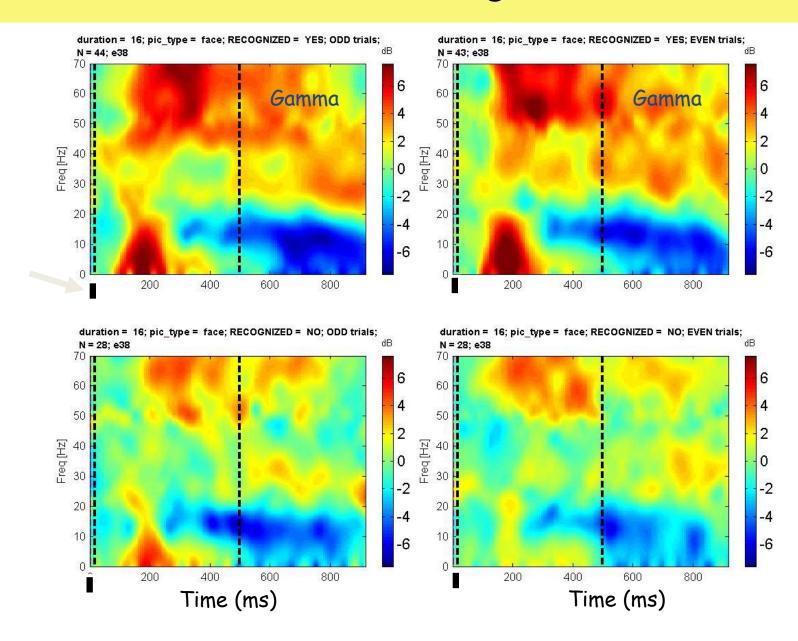
ECoG recording

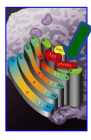
- To increase resolution one has to invade the brain
- The first step inside is with Electro-cortico-graphy (ECoG) – using electrodes that are placed on the surface of the brain, above or below the dura mater.
- The method is used mainly in the treatment of epilepsy, but also used to collect experimental data





ECoG recording





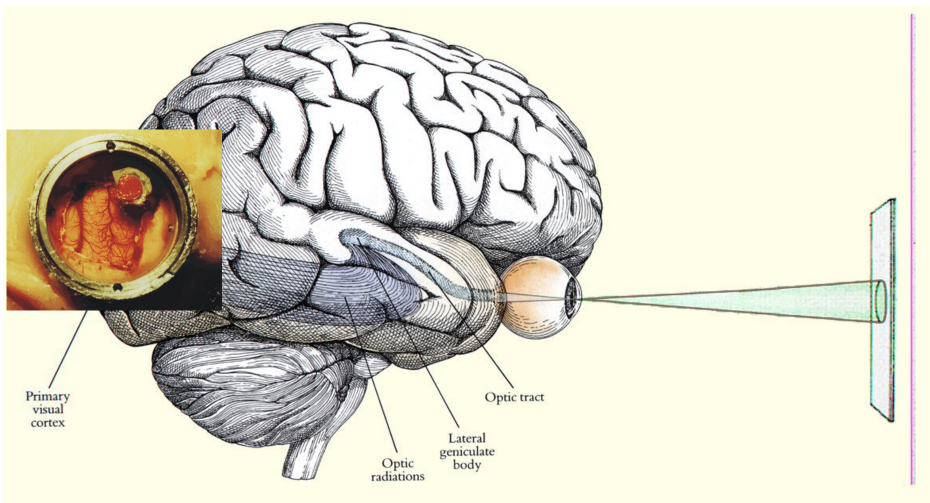
Methods table

Measuring neural activity

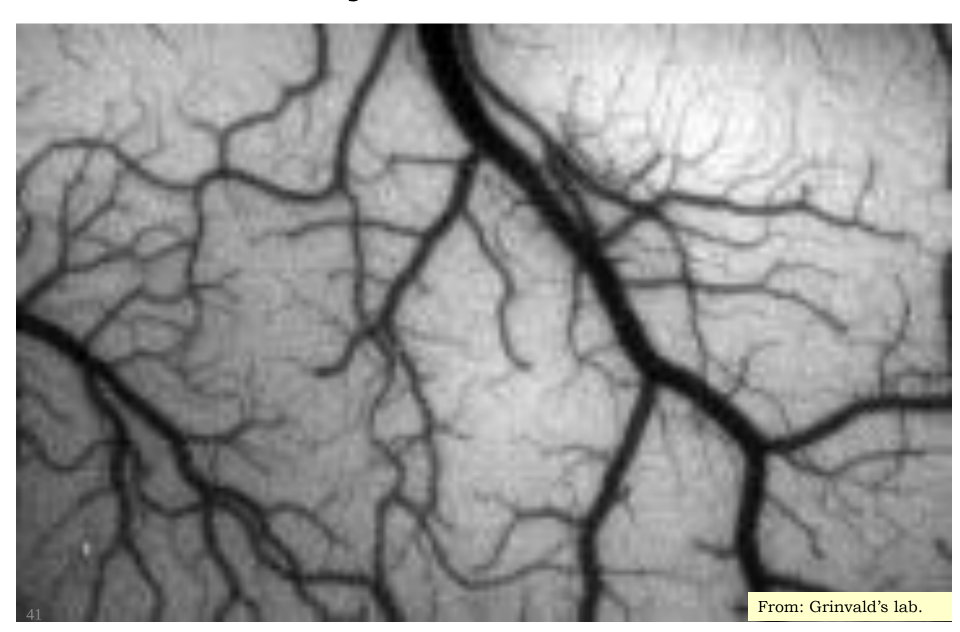
	Spatial resolution				Temporal resolution				
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	<u>device</u>	<u>signal</u>	<u>spikes</u>	heuristic	
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Intracell elec	10 μm	10 μm	< 1		< 1 ms	< 1 ms	< 1		
Ca imaging	1 μm	1 μm	< 1		1 ms	> 100 ms	> 100		

Intrinsic signals recording

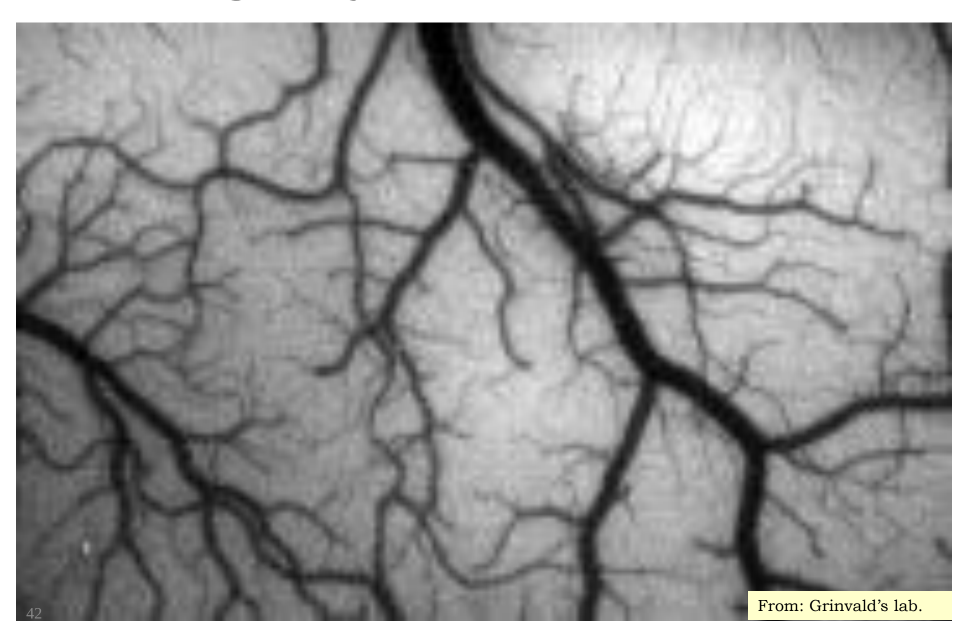
The Cranial Window



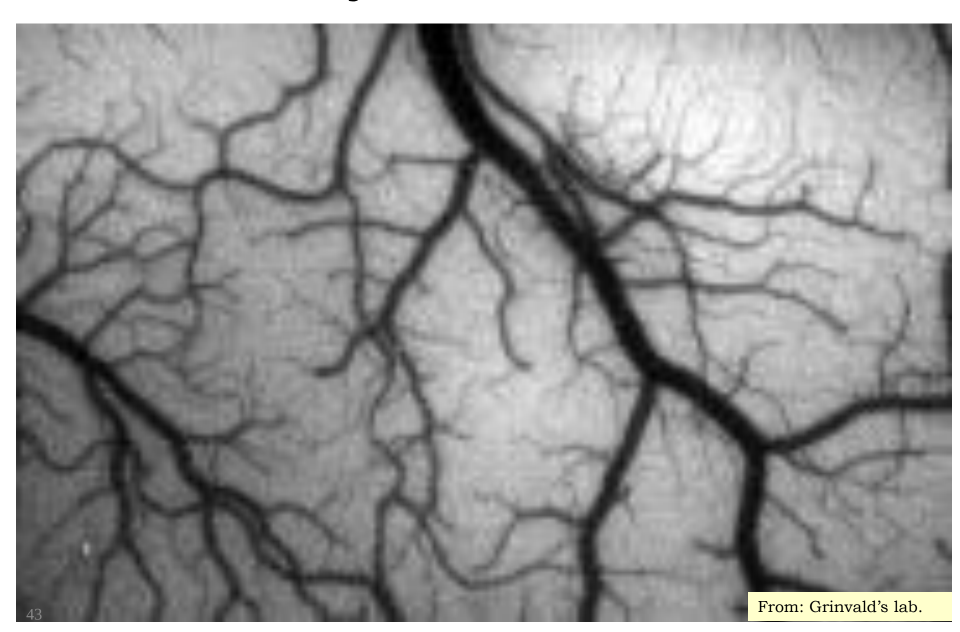
Left eyes was stimulated



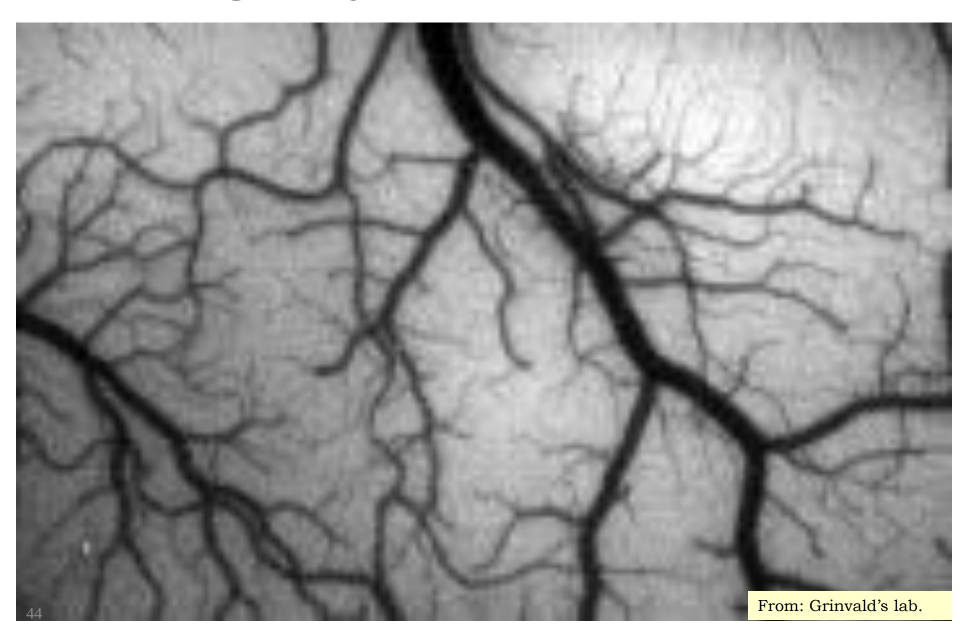
Right eyes was stimulated

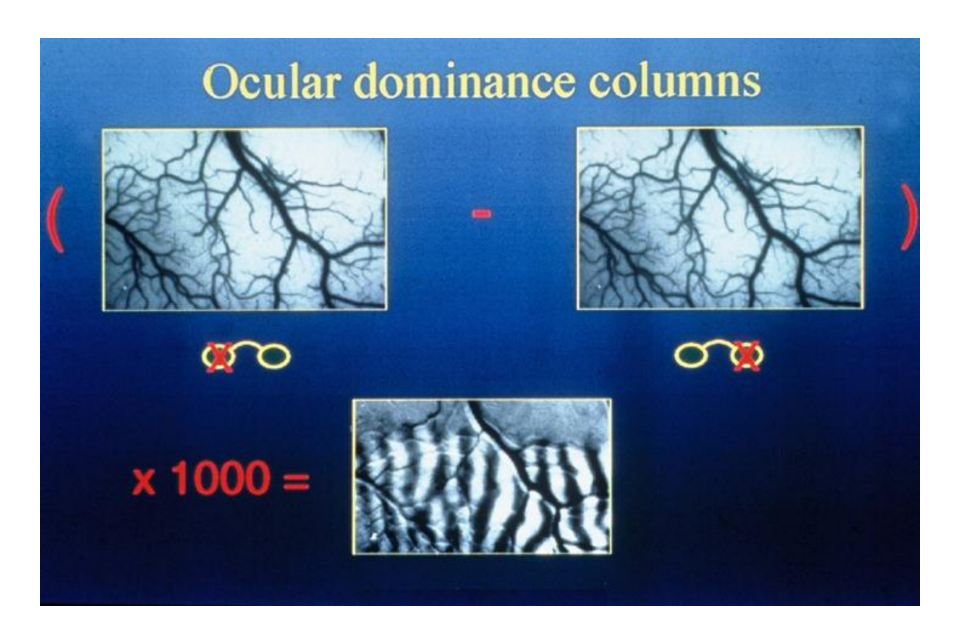


Left eyes was stimulated

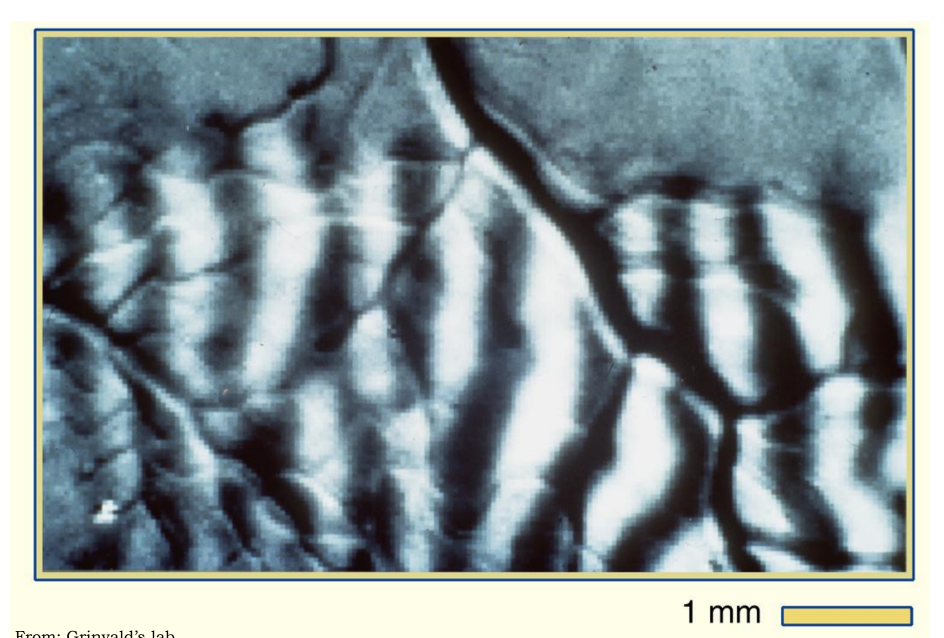


Right eyes was stimulated





From: Grinvald's lab.



SOURCES OF INTRINSIC SIGNALS

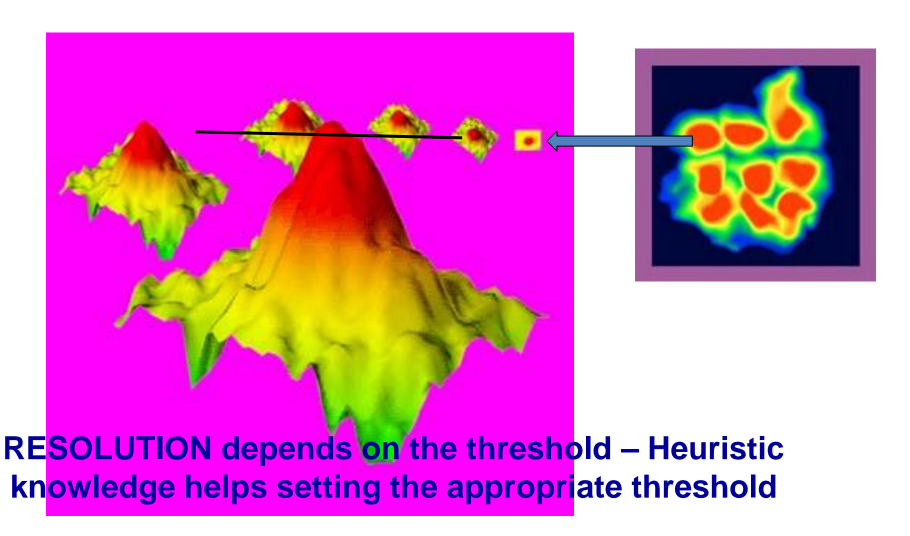
- Changes in absorption (similar to BOLD) due to:
 - Changes in oxygenation
 - Changes in blood volume
 - Changes in blood flow
- Changes in Light scattering due to:

Ion movement; water movement; shrinkage or expansion of the extracellular space; transmitter release; Volume changes due to capillaries dilation

From: Grinvald's lab.

RESOLUTION OF INTRINSIC (and BOLD) SIGNALS

Full 3-D view (no threshold) of a single WFR



Barrel area ~ 0.15 mm²; WFR area ~ 15 mm²

Methods table

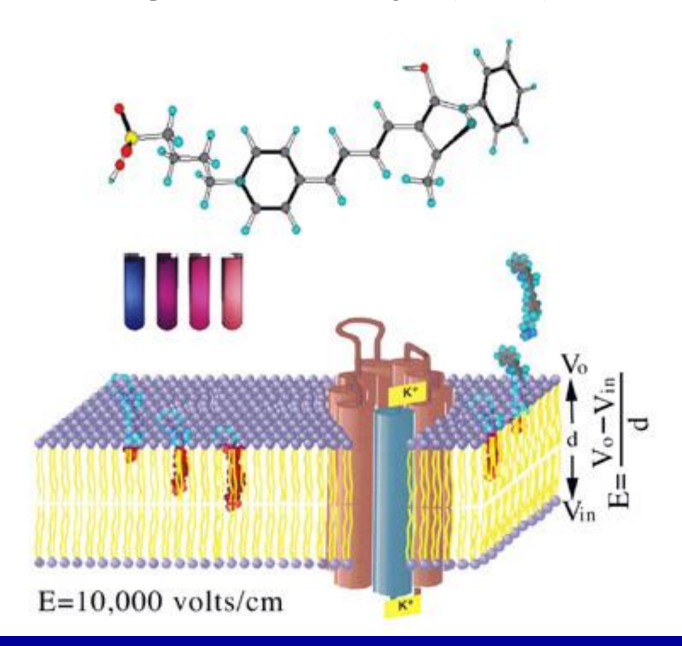
Measuring neural activity

		Spatial re		Temporal resolution				
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Intracell elec	10 μm	10 μm	< 1		< 1 ms	< 1 ms	< 1	
Ca imaging	1 μm	1 μm	< 1		1 ms	> 100 ms	> 100	

I

Voltage Sensitive Dye (VSD)

Merocyanine Dye RH-890



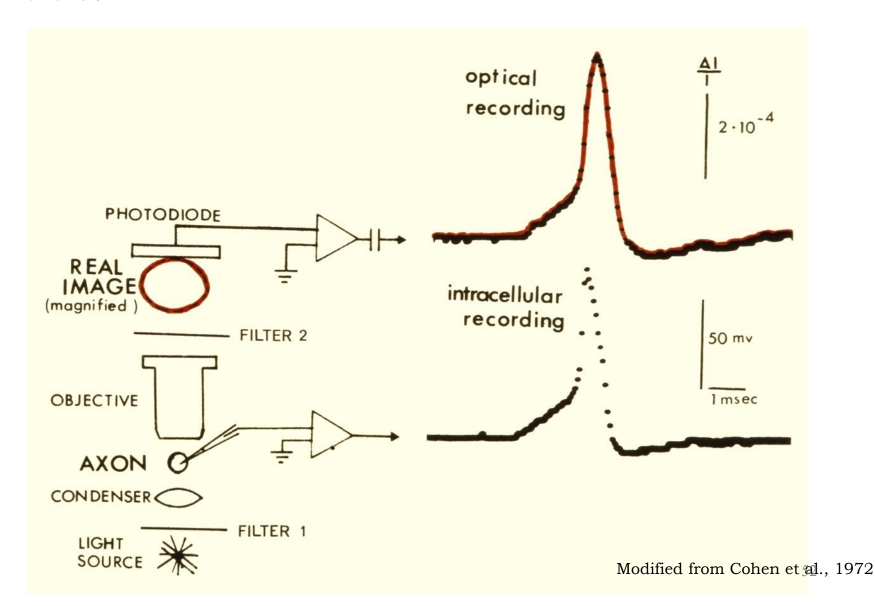
MEASURES:

the weighted sum of membrane-potential changes in neuronal somata, dendritic and axonal arbors, and often glia

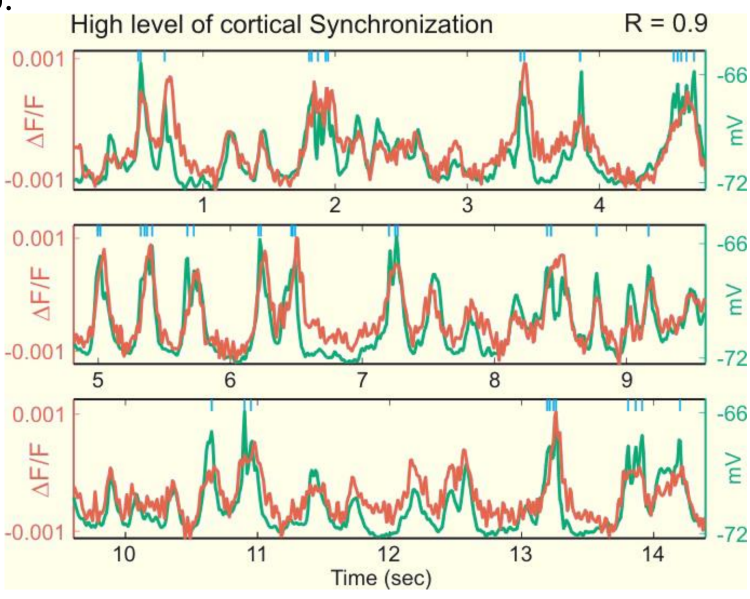
the dye signal is restricted to the site of the electrical activity

This signal mainly reflects the synaptic potentials in the dendrites

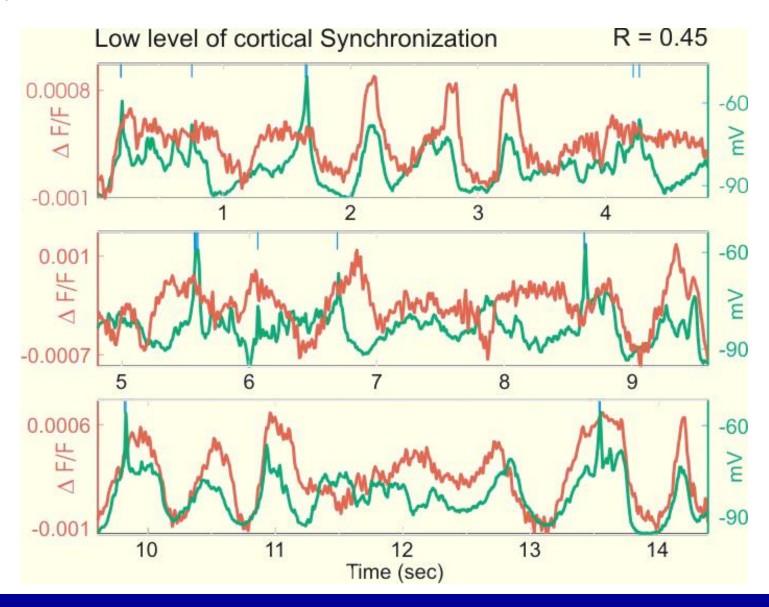
In-vitro:



In-vivo:

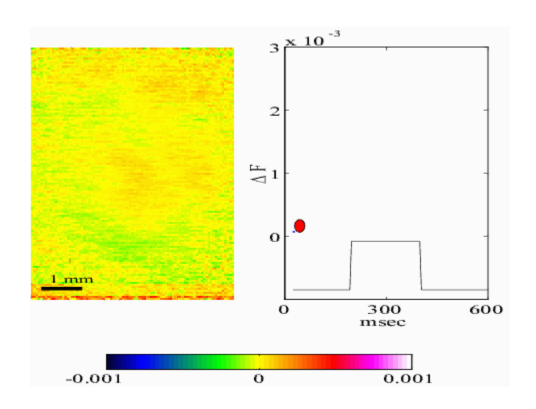


In-vivo:



In-vivo:

Example: Surround Inhibition in the Rat Barrel Cortex

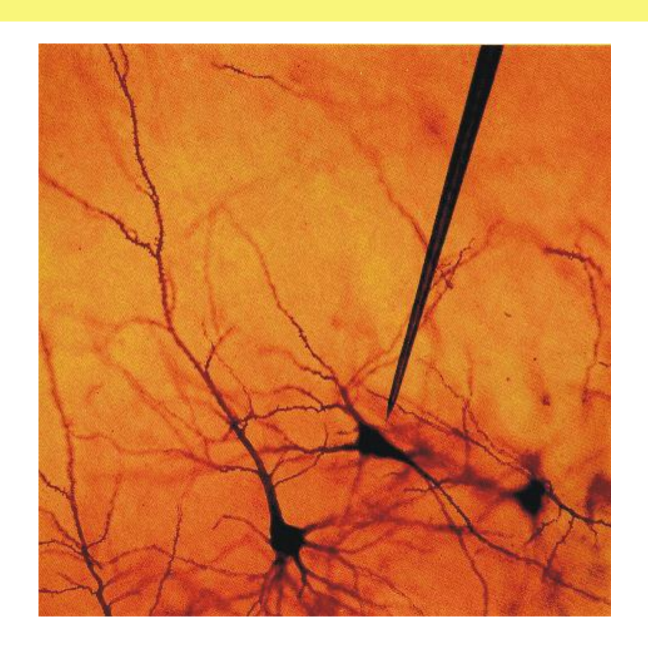


Methods table

Measuring neural activity

	Spatial resolution				Temporal resolution				
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	<u>device</u>	<u>signal</u>	<u>spikes</u>	heuristic	
behavior	brain	brain	> 10 ¹¹	station	1 ms	10 ms	10		
2DG, c-fos	10 μm	10 μm	1		NA	30 min	> 10 ⁶		
fMRI	> 1 mm	> 200 µm	> 10 ⁵		1 s	> 10 s	1000	10 ms	
EEG	10 mm	100 mm	> 10 ⁹	station	< 1 ms	50 ms	50	1 ms	
MEG	10 mm	100 mm	> 10 ⁹		1 ms	50 ms	50		
ECoG	10 mm	1-10 mm	> 10 ⁴⁻⁶		< 1 ms	10 ms	10		
Intr. Signal	10 μm	200 μm	> 1000		< 1 ms	1 s	1000	10 ms	
VSD Imaging	10 μm	30 μm	< 100		< 1 ms	10 ms	< 1		
μElec	50 μm	10 μm	1	10 μm	< 1 ms	1 ms	1		
μDialysis	< 1 mm	100 μm	> 10 ⁴		>1 min	100 ms	> 10 ⁶		
Intracell elec	10 μm	10 μm	< 1		< 1 ms	< 1 ms	< 1		
Ca imaging	1 μm	1 μm	< 1		1 ms	> 100 ms	> 100		

Micro-electrode recordings



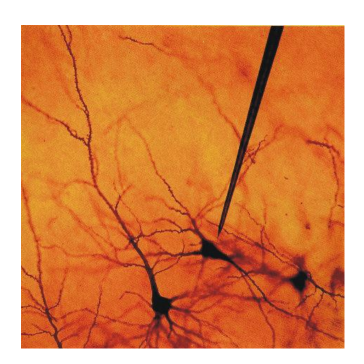
Micro-electrode recordings

From the tip of the microelectrode one can record:

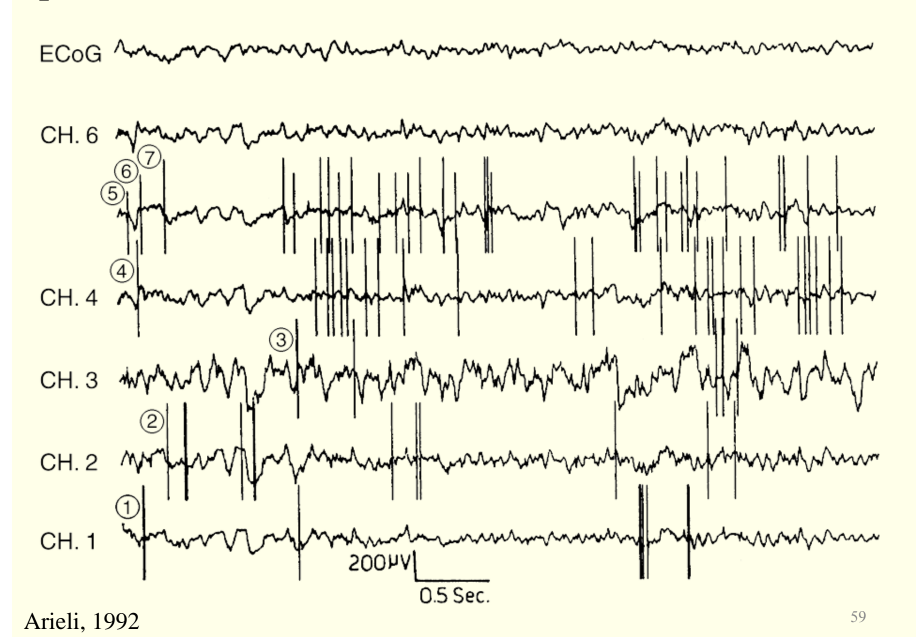
LFP – local field potential

MUA – multi-unit activity

SUA – single-unit activity (using spike sorting)

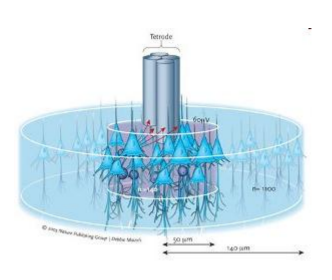


Spikes, LFP and ECoG



Micro-electrode recordings

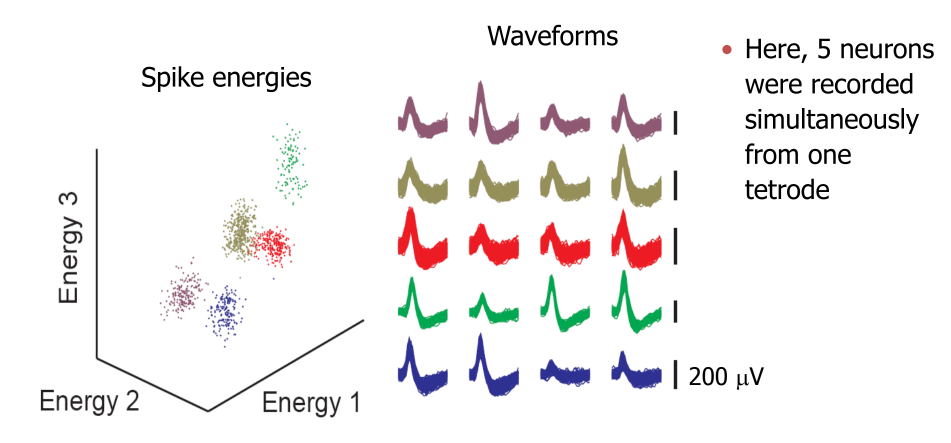
Multiple single-unit recordings using tetrodes



- Neurons are spike-sorted based on relative amplitudes on the 4 tetrode channels (amplitude differences are caused by physical proximity of neurons to different tetrode wires)
- Up to 25 cell can be well-separated per tetrode
- More typically: 5 10 cells per tetrode
- Can record > 100 neurons overall (in 10-20 tetrodes) in a freely behaving, freely moving animal

Micro-electrode recordings

Multiple single-unit recordings using tetrodes



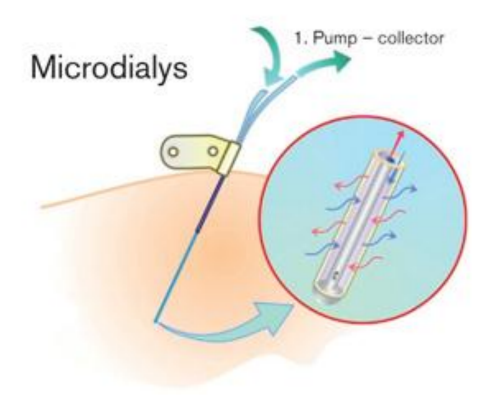
Methods table

Measuring neural activity

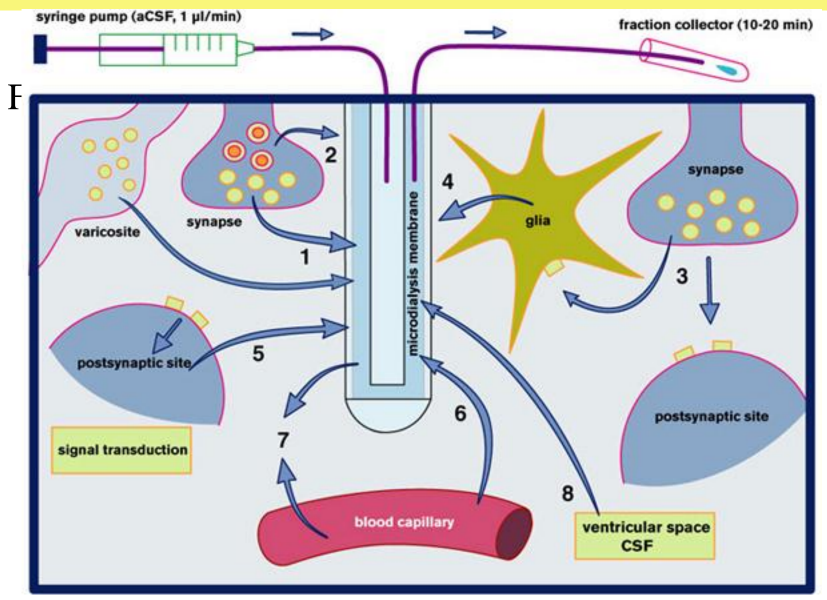
	Spatial resolution				Temporal resolution				
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	<u>device</u>	<u>signal</u>	<u>spikes</u>	heuristic	
behavior	brain	brain	> 10 ¹¹	station	1 ms	10 ms	10		
2DG, c-fos	10 μm	10 μm	1		NA	30 min	> 10 ⁶		
fMRI	> 1 mm	> 200 µm	> 10 ⁵		1 s	> 10 s	1000	10 ms	
EEG	10 mm	100 mm	> 10 ⁹	station	< 1 ms	50 ms	50	1 ms	
MEG	10 mm	100 mm	> 10 ⁹		1 ms	50 ms	50		
ECoG	10 mm	1-10 mm	> 10 ⁴⁻⁶		< 1 ms	10 ms	10		
Intr. Signal	10 μm	200 μm	> 1000		< 1 ms	1 s	1000	10 ms	
VSD Imaging	10 μm	30 μm	< 100		< 1 ms	10 ms	< 1		
μElec	50 μm	10 μm	1	10 μm	< 1 ms	1 ms	1		
μDialysis	< 1 mm	100 μm	> 10 ⁴		>1 min	100 ms	> 10 ⁶		
Intracell elec	10 μm	10 μm	< 1		< 1 ms	< 1 ms	< 1		
Ca imaging	1 μm	1μm	< 1		1 ms	> 100 ms	> 100		

Micro-dialysis recordings

From the tip of the micro-dialysis probe one can record concentrations of chemicals



Micro-dialysis recordings



Methods table

Measuring neural activity

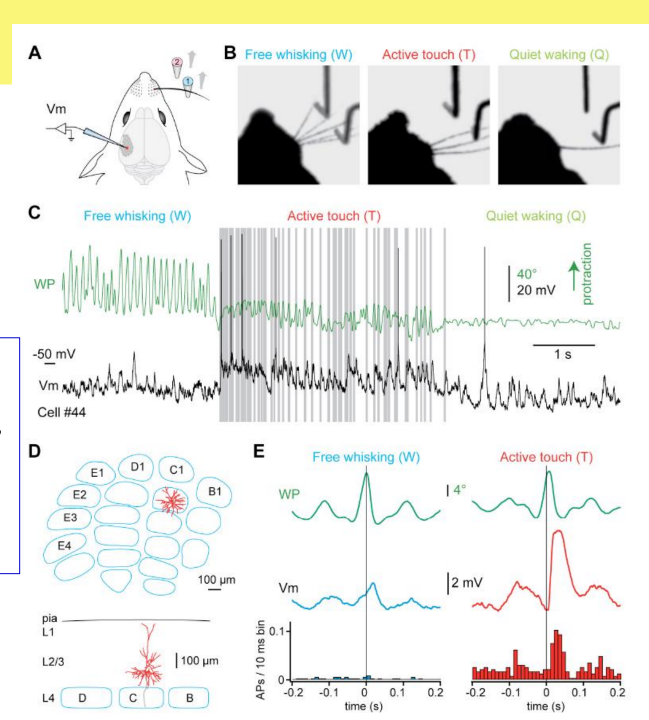
	Spatial resolution				Temporal resolution				
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	<u>device</u>	<u>signal</u>	<u>spikes</u>	heuristic	
behavior	brain	brain	> 10 ¹¹	station	1 ms	10 ms	10		
2DG, c-fos	10 μm	10 μm	1		NA	30 min	> 10 ⁶		
fMRI	> 1 mm	> 200 µm	> 10 ⁵		1 s	> 10 s	1000	10 ms	
EEG	10 mm	100 mm	> 10 ⁹	station	< 1 ms	50 ms	50	1 ms	
MEG	10 mm	100 mm	> 10 ⁹		1 ms	50 ms	50		
ECoG	10 mm	1-10 mm	> 10 ⁴⁻⁶		< 1 ms	10 ms	10		
Intr. Signal	10 μm	200 μm	> 1000		< 1 ms	1 s	1000	10 ms	
VSD Imaging	10 μm	30 μm	< 100		< 1 ms	10 ms	< 1		
μElec	50 μm	10 μm	1	10 μm	< 1 ms	1 ms	1		
μDialysis	< 1 mm	100 μm	> 10 ⁴		>1 min	100 ms	> 10 ⁶		
Intracell elec	10 μm	10 μm	< 1		< 1 ms	< 1 ms	< 1		
Ca imaging	1 μm	1 μm	< 1		1 ms	> 100 ms	> 100		

Intra-cellular recordings

In awake behaving mouse

- Identifying cell type
- Recording the inputs (reflecting individual inputs, network state, network activity)
- revealing mechanisms (exinh, ...)

C. Petersen and colleagues, 2010



Methods table

Measuring neural activity

	Spatial resolution				Temporal resolution				
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	<u>device</u>	<u>signal</u>	<u>spikes</u>	heuristic	
behavior	brain	brain	> 10 ¹¹	station	1 ms	10 ms	10		
2DG, c-fos	10 μm	10 μm	1		NA	30 min	> 10 ⁶		
fMRI	> 1 mm	> 200 µm	> 10 ⁵		1 s	> 10 s	1000	10 ms	
EEG	10 mm	100 mm	> 10 ⁹	station	< 1 ms	50 ms	50	1 ms	
MEG	10 mm	100 mm	> 10 ⁹		1 ms	50 ms	50		
ECoG	10 mm	1-10 mm	> 10 ⁴⁻⁶		< 1 ms	10 ms	10		
Intr. Signal	10 μm	200 μm	> 1000		< 1 ms	1 s	1000	10 ms	
VSD Imaging	10 μm	30 μm	< 100		< 1 ms	10 ms	< 1		
μElec	50 μm	10 μm	1	10 μm	< 1 ms	1 ms	1		
μDialysis	< 1 mm	100 μm	> 10 ⁴		> 1 min	100 ms	> 10 ⁶		
Intracell elec	10 μm	10 μm	< 1		< 1 ms	< 1 ms	< 1		
Ca imaging	1 μm	1 μm	< 1		1 ms	> 100 ms	> 100		

Ca imaging using 2-photon microscopy

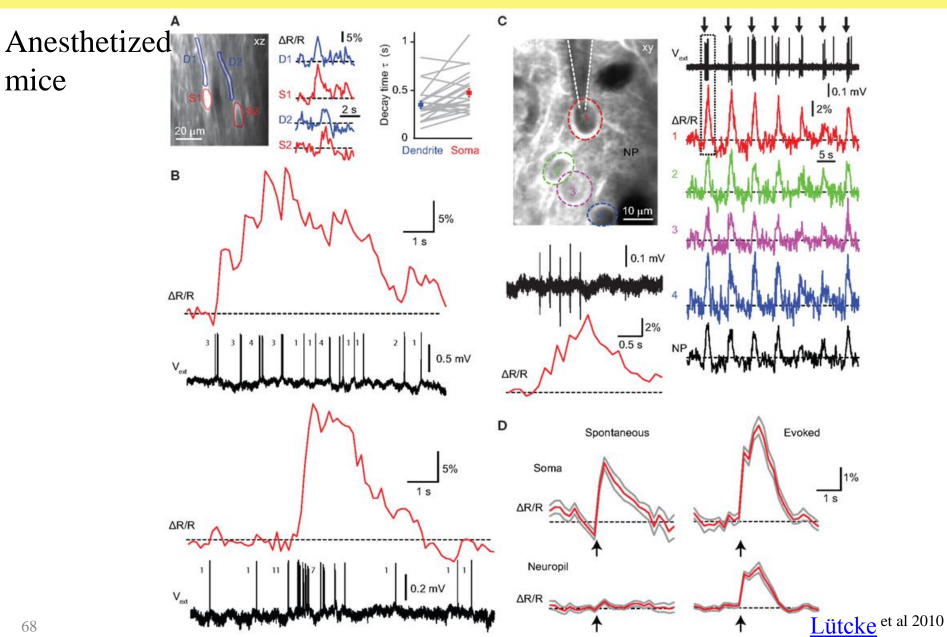


Figure 5. Single-cell and population YC3.60 Ca²⁺ signals in L2/3 of barrel cortex. (A) Simultaneous two-photon Ca²⁺ imaging in soma and dendrites of L2/3 neurons using vertical (xz-)imaging. Examples of spontaneous somatic (S, red) and apical dendritic (D, blue) YC3.60 Ca²⁺ transients for the cells depicted in the left image. Right: Mean decay times in dendrites compared to somata for 23 measurements (gray lines; mean \pm SEM). (B) Simultaneous juxtacellular voltage recording and two photon Ca²⁺ imaging from a neuron showing rare events of sustained and high-frequency AP firing that are accompanied by large YC3.60 Ca²⁺ transients with peak amplitudes of up to 30% ΔR/R. Top: Sustained AP firing leads to prolonged elevation of the fluorescence ratio. Bottom: A short burst of 11 APs is accompanied by a fast Ca²⁺ transient, which returns to baseline following a stereotypical exponential decay. (C) Two-photon Ca²⁺ imaging of a small population of neurons during sensory stimulation (seven times five air-puffs to contra-lateral whiskers at 5 Hz). Large Ca²⁺ transients in cell 1 (red trace) correlated with the spiking activity observed in the simultaneous juxtacellular voltage recording. Concomitant Ca2+ transients were also evoked in neighboring neuronal somata and in the nearby neuropil (NP). The response to the first stimulation episode (dashed box) is shown on expanded scale in the lower left, indicating that YC3.60 resolves the individual steps in the accumulated Ca²⁺ response. (**D**) Event-triggered average Ca²⁺ traces from somata and adjacent neuropil for spontaneous (n = 37 events of 1–3 APs) and evoked (n = 32 events of 1-5 APs) action potentials. Multi-whisker air puff evoked Ca2+ transients in somata were significantly larger than those in the neuropil while spontaneous spikes were accompanied by somatic but no neuropil transients. Errors are shown as SEM.

Ca imaging using 2-photon microscopy

Freely moving mice

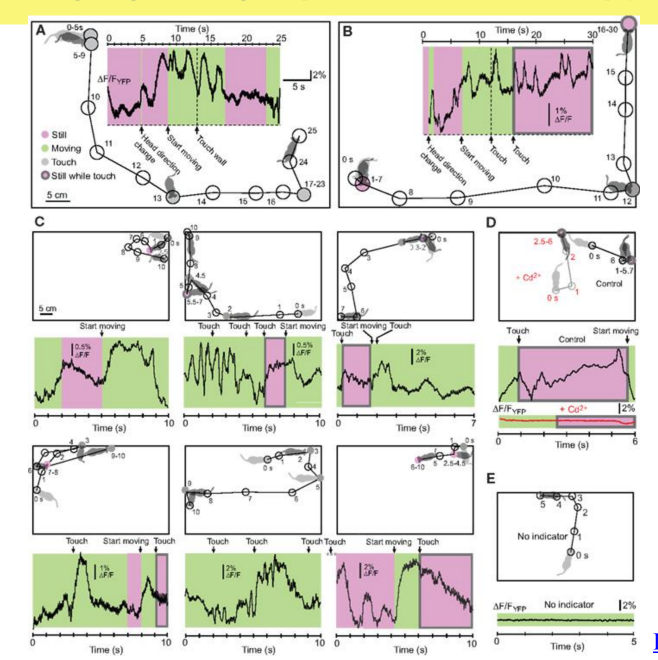
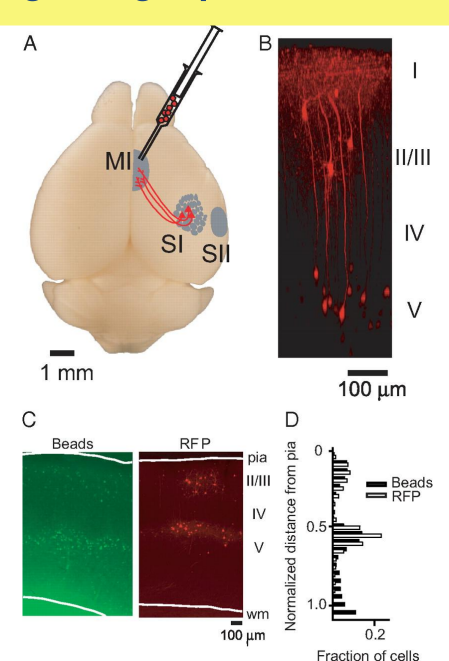


Figure 7. Fiber-optic recording of brain area activity in freely moving mice using YC3.60. (A,B) Two examples of fiber-optic recording of YC3.60 signals in awake, freely moving mice. Bulk Ca²⁺ signals indicating neuronal activity were recorded in somatosensory cortex through a singlecore optical fiber as shown in Figure 6C. Fluorescence changes in the YFPchannel are shown during 25–30 s periods together with the position of the mouse in an open field box. Animal behavior (sitting still, moving, touches, or having contact to the wall) is indicated by background colors. The trajectory of the animals' movement is indicated with selected time stamps. (C) Six more examples of Ca²⁺ imaging from three mice, together with corresponding behavioral observations. Changes of the animal's behavioral state (e.g., start of movement) were frequently associated with marked discontinuities in the fluorescence trace, indicating complex underlying Ca^{2+} dynamics. (**D**) Control experiment showing that Ca^{2+} signals are blocked by local perfusion of the cortical region with Cd²⁺. (**E**) Control experiment demonstrating that a flat fluorescence trace is observed in the absence of YC3.60 expression.

Ca imaging using 2-photon microscopy

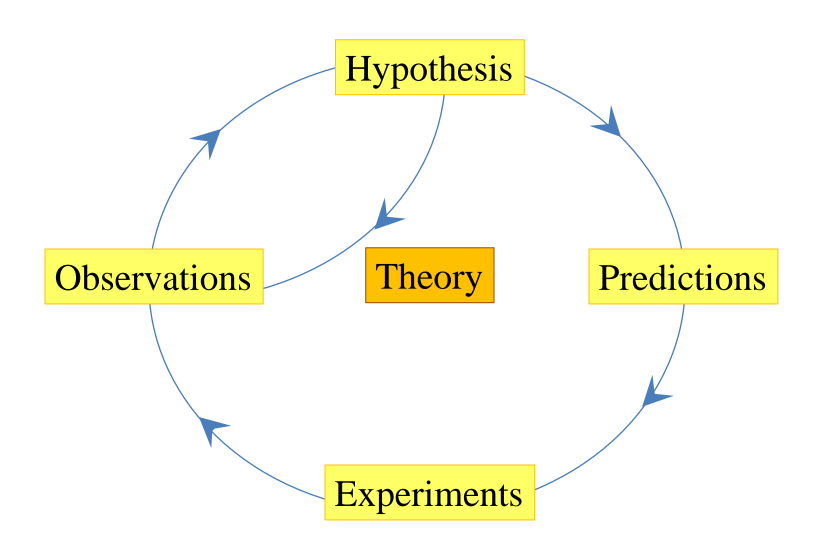
Selecting only projecting neurons



• **Figure 1.** Retrograde labeling with a virus expressing a red fluorescent protein. A, Schematic showing retrograde labeling of neurons in SI by injection of the retrograde virus HSV1 into MI. B, In vivo image of RFP+ neurons (maximum-intensity side projection of an image stack of RFP+ neurons; 512 x 128 x 96; section spacing, 8 µm). C, Distribution of labeled neurons in SI barrel cortex after bead (left) or virus (right) injection into MI. The white lines indicate the pia and the border between the cortex and the white matter. D, Normalized distribution of labeled neurons after bead (black, 1293 neurons) and HSV (white, 808 neurons) injection into MI.

Introduction

The scientific method

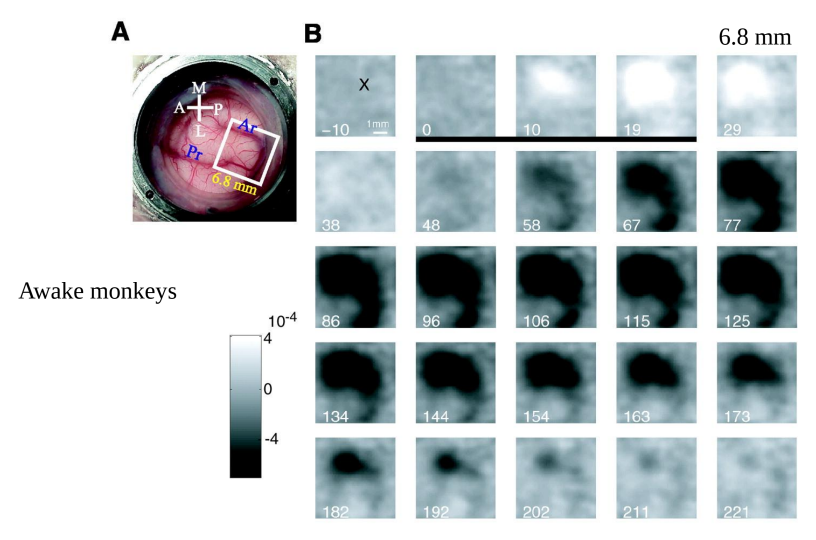


		Spatial re	solution	ı	Temporal resolution				
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	<u>device</u>	<u>signal</u>	<u>spikes</u>	heuristic	
sensory	modality	modality	> 10 ⁹	station	< 1 ms	< 1 ms	< 1		
TMS	100 mm	10 mm	> 10 ⁹	station	< 1 ms	100 ms	100		
μStim	< 10 µm	> 100 µm	> 50		< 1 ms	10 ms	10		
μPharmac	< 10 µm	> 100 µm	> 50		1 ms	> 10 s	10		
single cell	< 10 µm	< 10 µm	1		< 1 ms	< 1 ms	< 1		
sub-cell	>100 μm	< 1 µm	> 50		< 1 ms	< 1 ms	< 1		

		Spatial re	solution		Temporal resolution				
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	device	signal	<u>spikes</u>	heuristic	
sensory	modality	modality	> 10 ⁹	station	< 1 ms	< 1 ms	< 1		
TMS	100 mm	10 mm	> 10 ⁹	station	< 1 ms	100 ms	100		
μStim	< 10 µm	> 100 µm	> 50		< 1 ms	10 ms	10		
μPharmac	< 10 µm	> 100 µm	> 50		1 ms	> 10 s	10		
single cell	< 10 µm	< 10 µm	1		< 1 ms	< 1 ms	< 1		
sub-cell	>100 μm	< 1 µm	> 50		< 1 ms	< 1 ms	< 1		

		Spatial re	solution		Temporal resolution				
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	<u>device</u>	<u>signal</u>	<u>spikes</u>	heuristic	
sensory	modality	modality	> 10 ⁹	station	< 1 ms	< 1 ms	< 1		
TMS	100 mm	10 mm	> 10 ⁹	station	< 1 ms	100 ms	100		
μStim	< 10 μm	> 100 μm	> 50		< 1 ms	10 ms	10		
μPharmac	< 10 μm	> 100 µm	> 50		1 ms	> 10 s	10		
single cell	< 10 µm	< 10 µm	1		< 1 ms	< 1 ms	< 1		
sub-cell	>100 μm	< 1 µm	> 50		< 1 ms	< 1 ms	< 1		

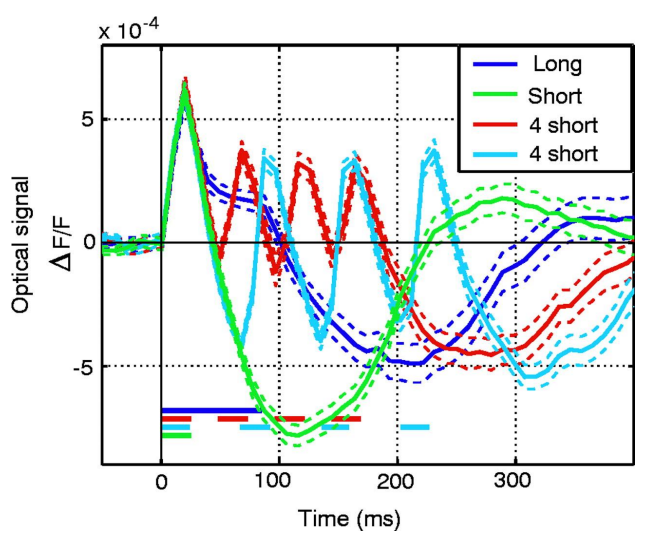
Micro-stimulation



E Seidemann et al. Science 2002;295:862-865

Figure 1 Spatiotemporal dynamics of microstimulation-evoked activity.

Micro-stimulation

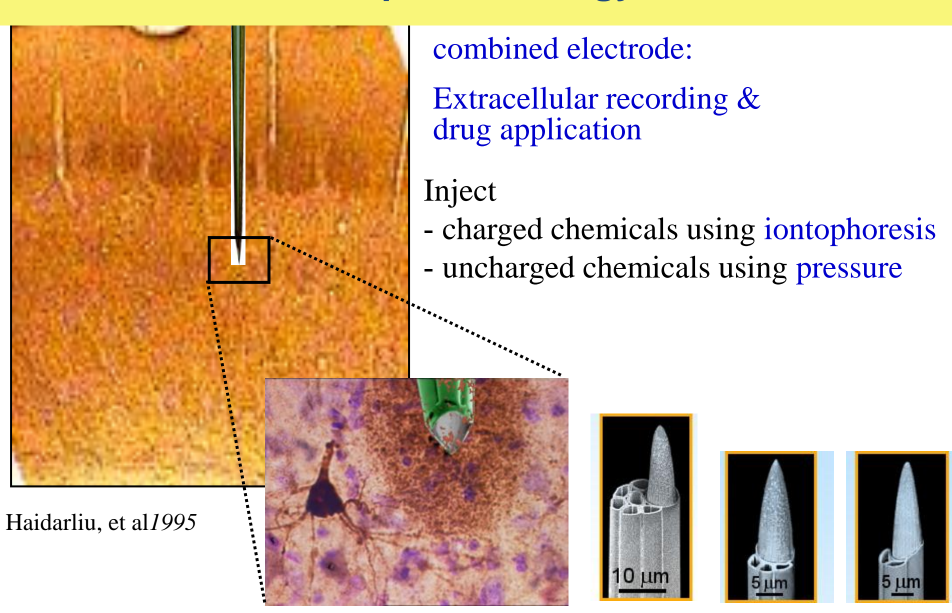


E Seidemann et al. Science 2002;295:862-865

Figure 2 Time course of the response to microstimulation.

		Spatial re	solution	ı	Temporal resolution				
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	device	signal	<u>spikes</u>	heuristic	
sensory	modality	modality	> 10 ⁹	station	< 1 ms	< 1 ms	< 1		
TMS	100 mm	10 mm	> 10 ⁹	station	< 1 ms	100 ms	100		
μStim	< 10 μm	> 100 μm	> 50		< 1 ms	10 ms	10		
μPharmac	< 10 μm	> 100 µm	> 50		1 ms	> 10 s	10		
single cell	< 10 µm	< 10 µm	1		< 1 ms	< 1 ms	< 1		
sub-cell	>100 μm	< 1 µm	> 50		< 1 ms	< 1 ms	< 1		

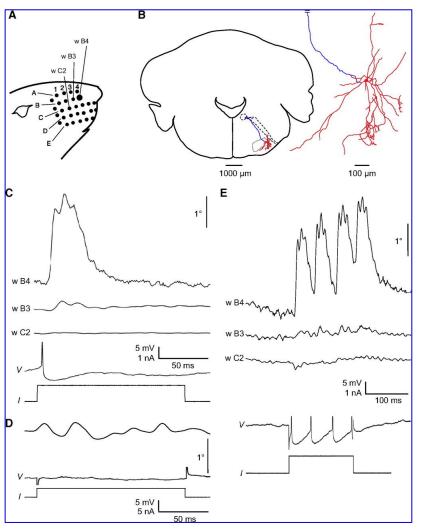
Micro-pharmacology



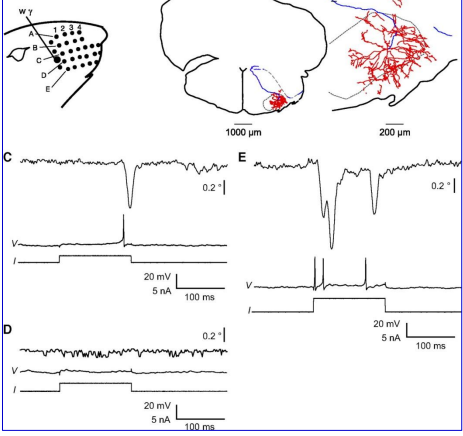
		Spatial re	solution	1	Temporal resolution				
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	device	<u>signal</u>	<u>spikes</u>	heuristic	
sensory	modality	modality	> 10 ⁹	station	< 1 ms	< 1 ms	< 1		
TMS	100 mm	10 mm	> 10 ⁹	station	< 1 ms	100 ms	100		
μStim	< 10 µm	$>$ 100 μm	> 50		< 1 ms	10 ms	10		
μPharmac	< 10 µm	> 100 µm	> 50		1 ms	> 10 s	10		
single cell	< 10 μm	< 10 μm	1		< 1 ms	< 1 ms	< 1		
sub-cell	>100 μm	< 1 µm	> 50		< 1 ms	< 1 ms	< 1		

Nano-stimulation

Examples of whisker protraction and retraction caused by single and multiple spikes

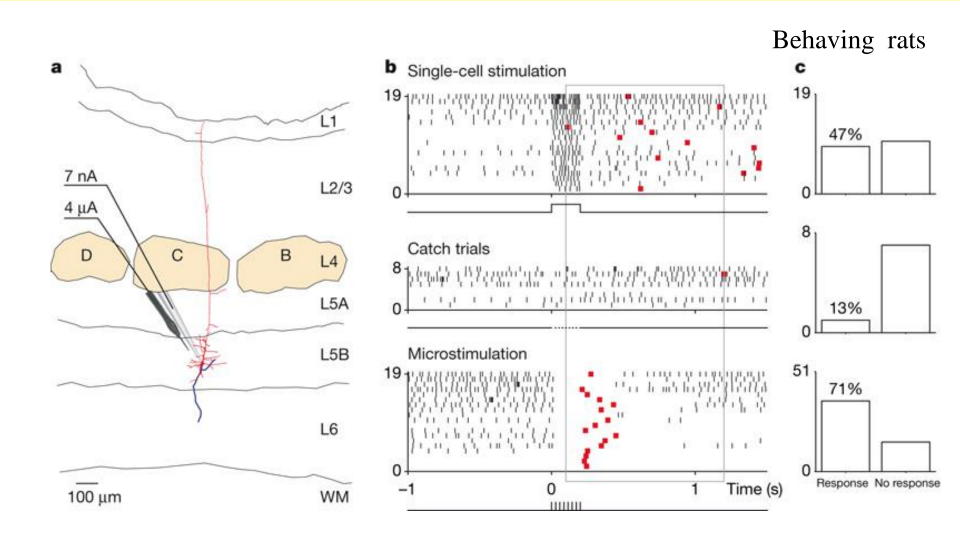


Anesthetized rats



Herfst, L. J. et al. J Neurophysiol 99: 2821-2832 2008; doi:10.1152/jn.01014.2007

Nano-stimulation



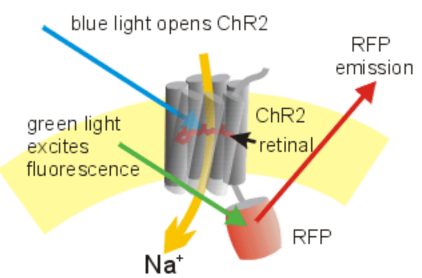
Houweling & Michael Brecht, Nature 2008

		Spatial re	solution		To	Temporal resolution			
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	<u>device</u>	<u>signal</u>	<u>spikes</u>	heuristic	
sensory	modality	modality	> 10 ⁹	station	< 1 ms	< 1 ms	< 1		
TMS	100 mm	10 mm	> 10 ⁹	station	< 1 ms	100 ms	100		
μStim	< 10 µm	$>$ 100 μm	> 50		< 1 ms	10 ms	10		
μPharmac	< 10 µm	> 100 µm	> 50		1 ms	> 10 s	10		
single cell	< 10 µm	< 10 µm	1		< 1 ms	< 1 ms	< 1		
sub-cell	>100 µm	< 1 µm	> 50		< 1 ms	< 1 ms	< 1		

Optogenetic-stimulation

Channelrhodopsins (ChR1,2)

- Channelrhodopsins function as light-gated ion channels.
- They serve as sensory photoreceptors in unicellular green algae, controlling phototaxis, i.e. movement in response to light.
- Expressed in cells of other organisms, they enable the use of light to control electrical excitability, and other cellular processes.
- All known Channelrhodopsins are nonspecific cation channels, conducting H⁺, Na⁺, K⁺, and Ca²⁺ ions.



Optogenetic-stimulation

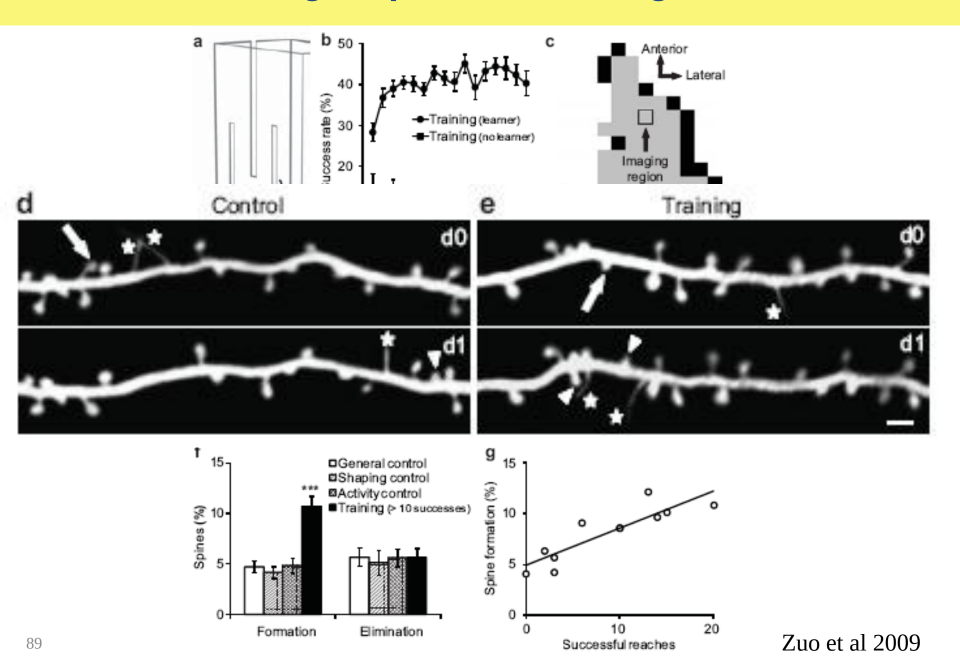
- Variety of excitatory channels: ON short duration, ON long duration, ON/OFF, ON subliminal, ...
- Inhibitory channels
- Linking to identified promoters to be functional in indentified neurons
- Localized infection using viruses
- Combined with dye reporters



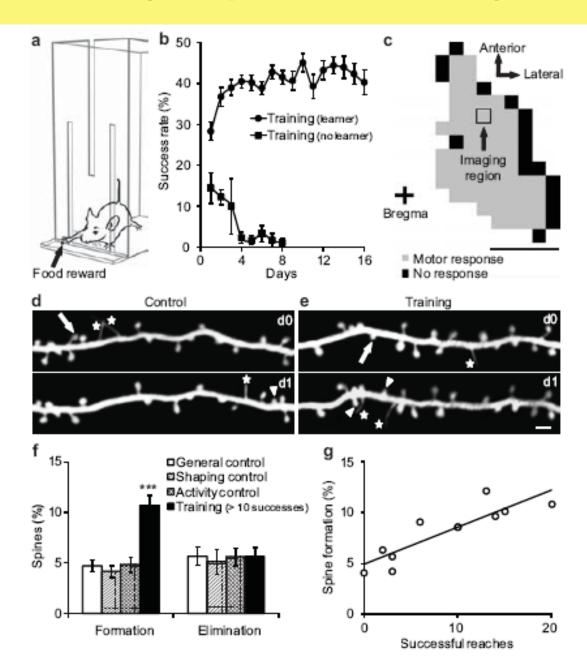
Measuring structure

		Spatial resolution signal neurons heurist			Temporal resolution				
	<u>device</u>	<u>signal</u>	neurons	<u>heuristic</u>	device	signal	<u>spikes</u>	heuristic	
cell density									
receptor dens	sity								
transmitter de	ensity								
tract tracing									
single-cell									
single-spine	< 1 µm	< 1 µm	< 1		<1s	hours			

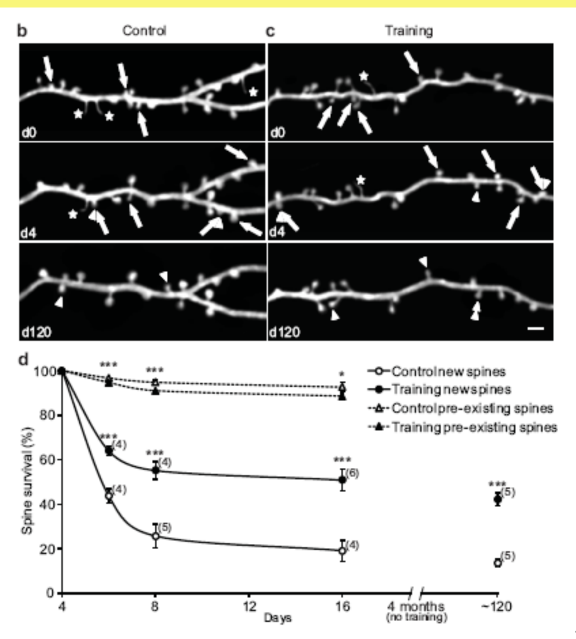
Single-spine monitoring



Single-spine monitoring



Single-spine monitoring



Manipulating structure

		Spatial re	solution	1	Te	emporal i	resoluti	on
	<u>device</u>	<u>signal</u>	neurons	heuristic	<u>device</u>	signal	<u>spikes</u>	heuristic
Neuropsychol	ogy	> 10 mm	> 10 ⁷			months		
lesions	> 100 µm	> 100 µm	> 50		>1s	>1 min		