Crown 85: Visual Perception: A Window to Brain and Behavior



Lecture 3: Techniques for Studying Brain and Behavior

Crown 85 Winter 2016

Visual Perception: A Window to Brain and Behavior

Lecture 3- Techniques for Studying the Brain

Reading:Amherst College CourseStanford College Course

Looking: Optogenetics ECoG

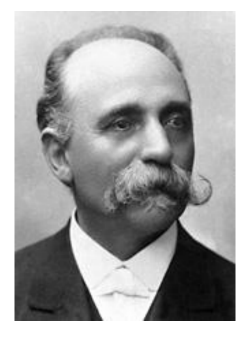
Understand the basic principles upon which the following techniques and the kinds of information about neural processing which their application can provide:



not expecting U to become an



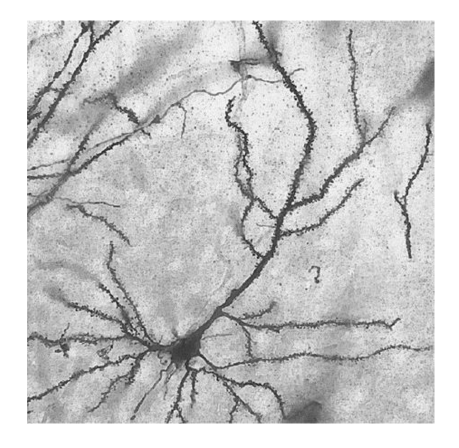
1.Anatomical a.Neuron staining b.Electron microscopy c.Pathway tracing



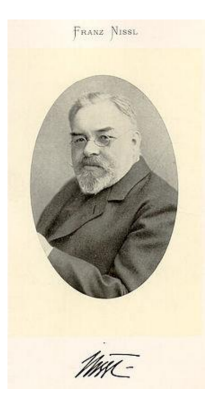
Camillo Golgi 1843-1926

Golgi

staining of **entire neuron** with silver chromate. only stains a subset of cells



http://www.nature.com/npp/journal/v32/n10/covers/index.html



Franz Nissl 1860-1919

Nissl

staining of **cell body** with dyes (e.g. cresyl violet) that interact with RNA (and DNA).

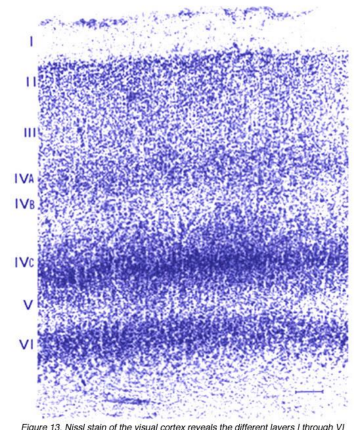


Figure 13. Nissl stain of the visual cortex reveals the different layers I through VI quite clearly.

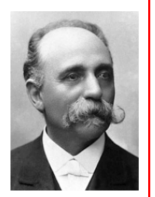
http://webvision.med.utah.edu/imageswv/nissl2.jpg



The Nobel Prize in Physiology or Medicine 1906 Camillo Golgi, Santiago Ramón y Cajal

Share this:

The Nobel Prize in Physiology or Medicine 1906





Camillo Golgi Prize share: 1/2

Santiago Ramón y Cajal Prize share: 1/2

The Nobel Prize in Physiology or Medicine 1906 was awarded jointly to Camillo Golgi and Santiago Ramón y Cajal *"in recognition of their work on the structure of the nervous system"*

Photos: Copyright © The Nobel Foundation

OPINION

The contribution of Santiago Ramón y Cajal to functional neuroscience

Rodolfo R. Llinás

Santiago Ramón y Cajal — arguably the most accomplished anatomist in the history of neuroscience — became recognized as such not only because of his incredible anatomical skills and his indefatigable working habits, but also because of his uncanny sense of the functional implications of his work, a sense that made him a true genius in the field of biology

http://www.utdallas.edu/~tres/memory/intro/llinas.pdf

Anatomy: electron microscopy

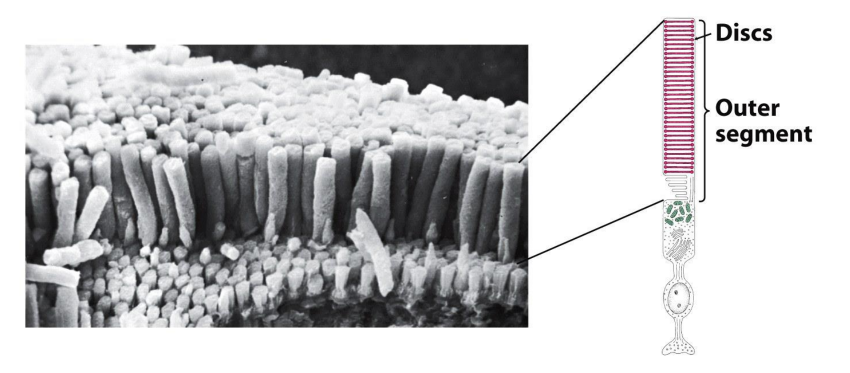


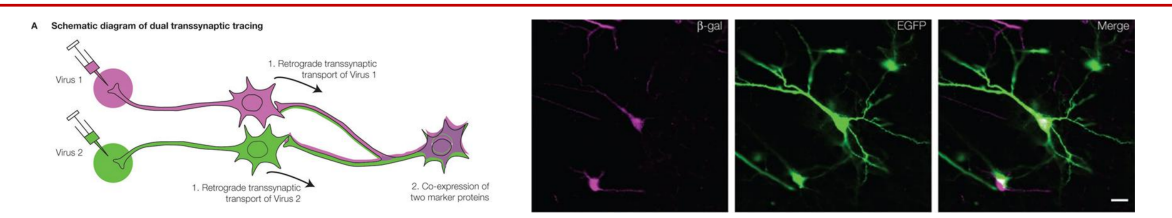
Figure 33.19 Biochemistry, Seventh Edition © 2012 W. H. Freeman and Company

http://www.ncbi.nlm.nih.gov/books/NBK22541/

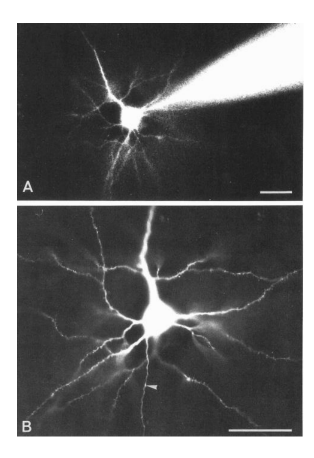
Figure 32.20The Rod Cell

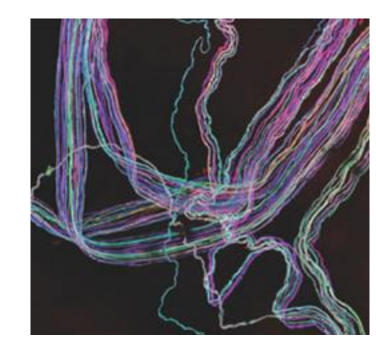
(Left) Scanning electron micrograph of retinal rod cells. (Right) Schematic representation of a rod cell. [Photograph courtesy of Dr. Deric Bownds.] Pathway Tracing:

Injecting, via fine needle, a 'tracer substance' into or near a neuron which is then transported down the axon (anterograde: *soma* \rightarrow *axon terminal*) or up the axon (retrograde : axon terminal \rightarrow soma). The pathway is then visualized by the color or radiographic 'footprint' of the tracer. The color may come from a tracer that is itself a dye or one that is produced by a subsequent 'developing' reaction.

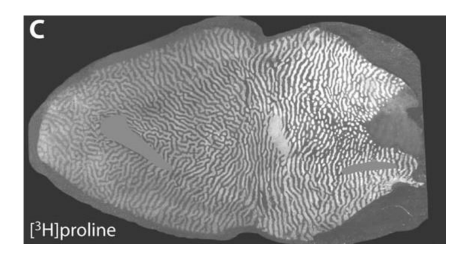


http://journal.frontiersin.org/article/10.3389/neuro.01.032.2009/full





http://www.nature.com/nrn/journal/v9/n6/ full/nrn2391.html



http://vision.ucsf.edu/hortonlab/ResearchProgram.html

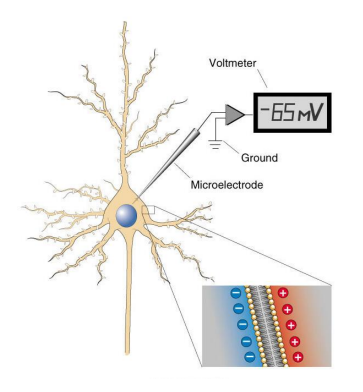
https://www.hccfl.edu/media/153423/handbook%20of %20chemical%20neuroanatomy%20vol8%20ch5.pdf 1. Anatomical

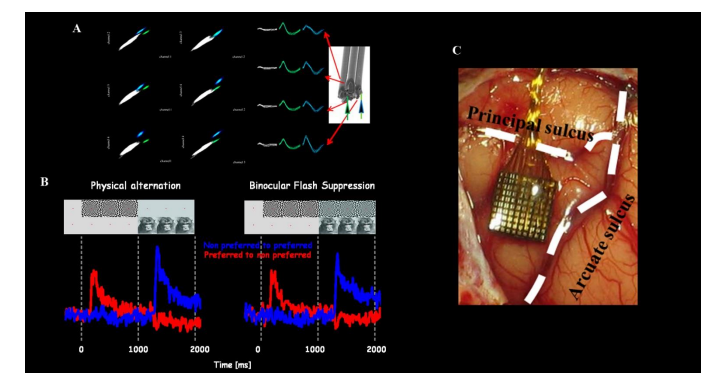
a.Neuron staining
b.Electron microscopy
c.Pathway tracing

- 2. Electrophysiological recording of neural activity
 - a. Single cell recording in neurons
 - b. Electroencephalography (EEG)
 - c. Magnetoencephalography (MEG)
 - d. Electrocorticography (ECoG)

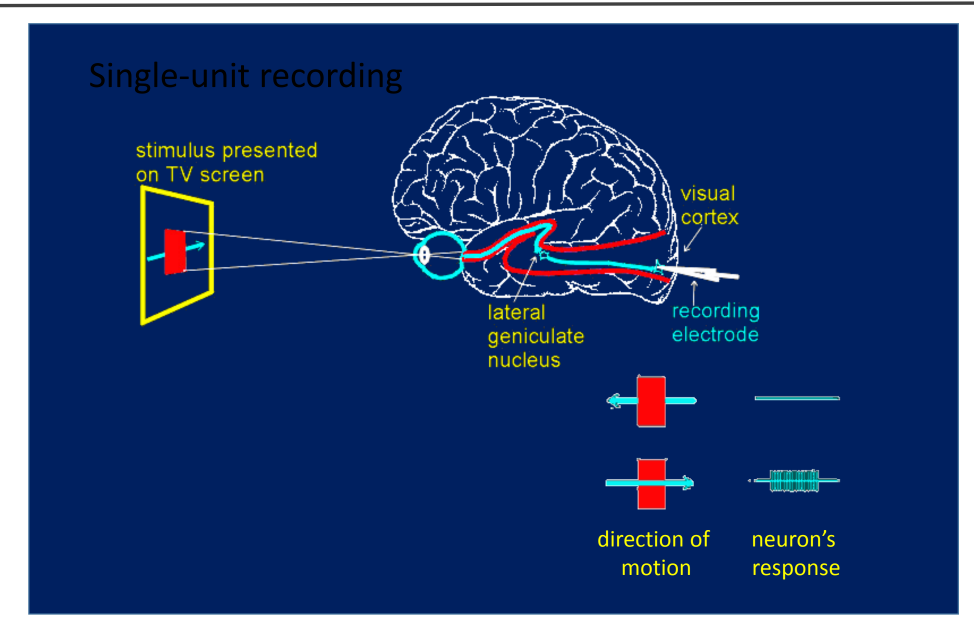
electrophysiology: single cell recordings of neuronal activity

Place an electrode on a single neuron and measure the frequency of firing.





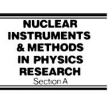
http://www.kyb.tuebingen.mpg.de/nc/de/mitarbeiter/details/theofanis.html



UCSC multielectrode array



Available online at www.sciencedirect.com



Nuclear Instruments and Methods in Physics Research A 501 (2003) 298-307

www.elsevier.com/locate/nima

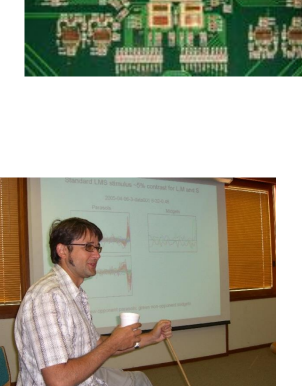
Large-scale imaging of retinal output activity $\stackrel{\scriptscriptstyle \rm tr}{\sim}$

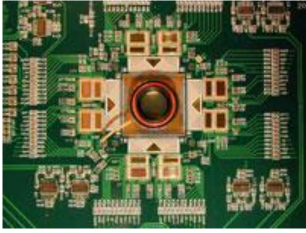
A.M. Litke^{a,*}, E.J. Chichilnisky^b, W. Dabrowski^c, A.A. Grillo^a, P. Grybos^c, S. Kachiguine^a, M. Rahman^d, G. Taylor^a

^a Santa Cruz Institute for Particle Physics, University of California, Santa Cruz, CA 95064, USA ^c Ine Saik Institute, La Jolla, CA 92057, USA ^c Faculty of Physics and Nuclear Techniques, University of Mining and Metallurgy, Krakow, Poland ^d University of Glasgow, Glasgow, UK

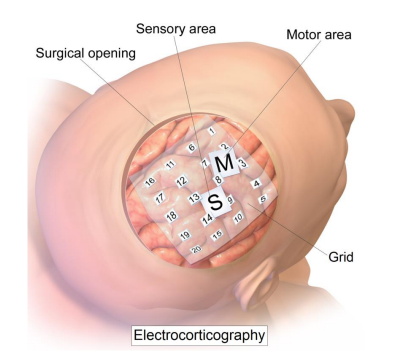


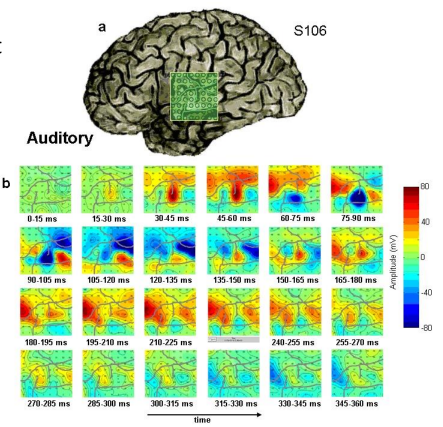
T	Crown 85- Visual Perception: A Window to Brain and Behavior					
IOME	COURSE VISION INFORMATION LECTURES	REPORTS RESEARCH LAB REPORTS INTERVIEWS PRO	DJECTS ECOMMONS			
	One goal of Crown 85 is to introduce its students to researchers and research in the various neuroscience and perception laboratories throughout the UCSC campus. Enrolled students are required to visit one of these researchers and present a class report on the studies being conducted in their laboratory. Laboratories which have agreed to host Crown 85 student interviews include:					
	Enrolled students are required to report on the studies being cond	o visit one of these researchers and ducted in their laboratory. Laborator	present a class			
	Enrolled students are required to report on the studies being cond	o visit one of these researchers and ducted in their laboratory. Laborator	present a class			





Subdural electrodes implanted inside the cranium. For purely clinical reasons, patients with epilepsy are sometimes implanted with such electrodes to localize their seizure onset prior to surgical therapy.



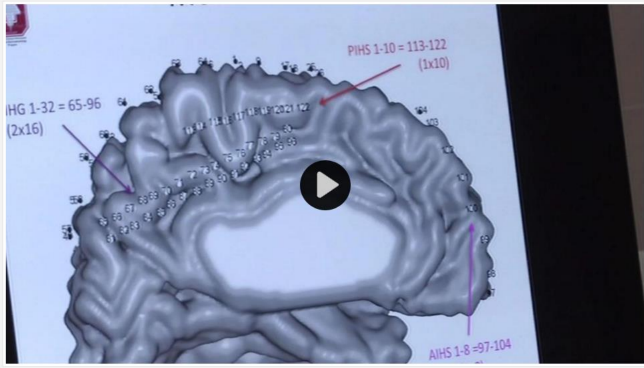


https://www.med.nyu.edu/thesenlab/research-0/intracranial-eeg/

https://en.wikiversity.org/wiki/Blausen_gallery_2014

electrophysiology (neural recordings): electrocorticography (ECoG)



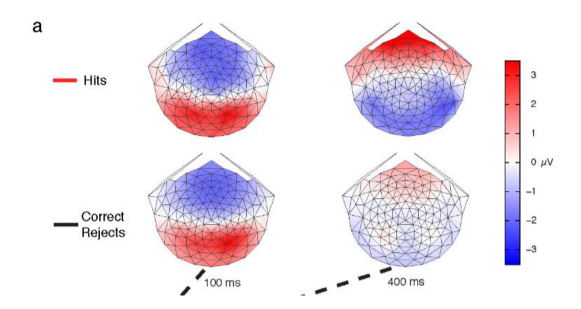


Surgeons at Stanford Hospital helped a Bay Area woman overcome intractable epilepsy and have fewer seizures.

http://abc7news.com/archive/9405444/

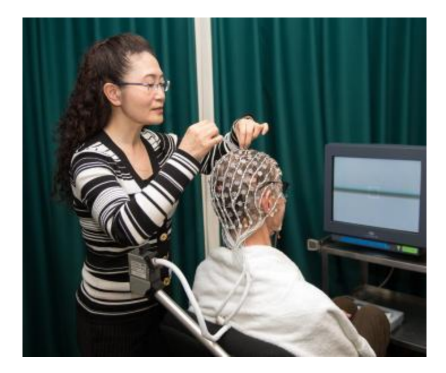
electrophysiological recording: electroencephalography (EEG)

EEG: measures electric fields produced by neural electrical activity



The time course of shape discrimination in the human brain Ales JM, Appelbaum LG, Cottereau BR, Norcia AM

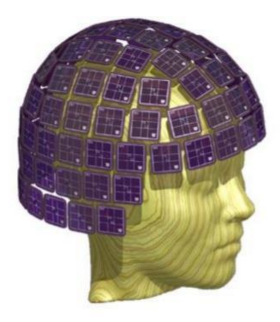
http://www.st-andrews.ac.uk/~jma23/papers/alesInPressNeuroimageShape.pdf



http://www.ski.org/lab/hou-lab

electrophysiological recording: magnetoencephalography (MEG)

MEG: measures the magnetic fields generated by neural activity







http://www.washington.edu/news/author/mollywmc/

http://ilabs.washington.edu/what-magnetoencephalography-meg

- 2. Electrophysiological recording of neural activity
- A. Single cell recording in neurons
 - b. Electroencephalography (EEG)
 - c. Magnetoencephalography (MEG)
- d. Electrocorticography (ECoG)

- 3. Imaging
 - a. Positron emission tomography (PET)
 - b. Functional magnetic resonance imaging (fMRI)
 - c. Calcium dyes
 - d. Voltage sensitive dyes

imaging: tomography (CT, PET, OCT)

collection of absorbed or emitted radiation (or electrons/positrons) from multiple detectors distributed in space; allows one to localize and image the source of the absorption or emission

Gamma ray detectors -

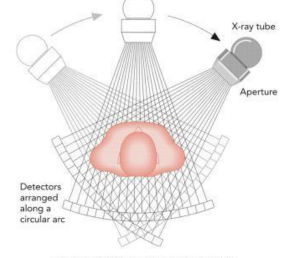


Figure 7-10 Computer tomography

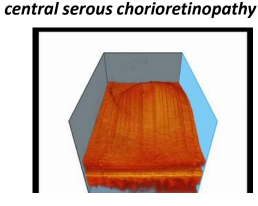
http://130.237.83.53/medicaldevices/album/Ch%207%20Medical%20ima ges/slides/F%207-10%20Computer%20tomography.html

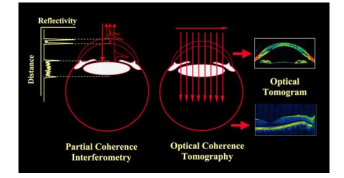
> ComputerTomography: xrays absorbed

Positron Emission Tomography: positrons emitted

http://science.howstuffworks.com/nuclear-medicine1.htm

@2000 How Stuff Works



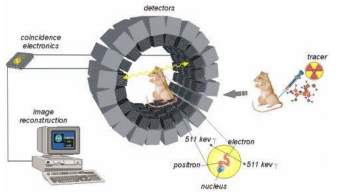


http://vsri.ucdavis.edu/research/retinal/oct

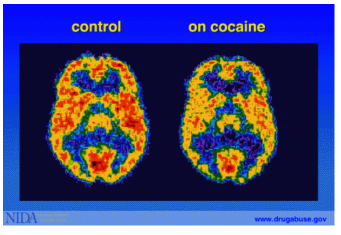
OpticalCoherenceTomography: coherent light

imaging (metabolic): positron emission tomography (PET)

- insert radioactive tracer into blood stream e.g. 2-deoxy-2-(¹⁸F)fluoro-Dglucose (FDG)
- FDG (viz 'glucose' and metabolism) concentrates in areas of high neural activity = high metabolism
- ¹⁸F isotope emits positrons ('electrons' with + charge)
- detectors map areas of positron emission







http://www.drugabuse.gov/publications/teachingpackets/understanding-drug-abuse-addiction/section-ii/2positron-emission-tomography-pet-scan-person-us

http://healthyscientist.blogspot.com/2009/11/wonders-of-pet-scan.html

imaging (metabolic): functional Magnetic Resonance Imaging (fMRI)

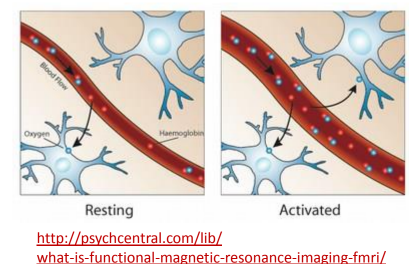
THE **BOLD** SIGNAL : **B**LOOD**O**XYGEN**L**EVEL**D**EPENDENT **fMRI**

- Magnetic Resonance Imaging involves putting the sample (*i.e. your head*!!) in a large external magnet
- Brain imaging is most often down with a spectrometer that measures the magnetic properties of hydrogen (in water) in various parts of the brain
- The hydrogen nucleus behaves like a tiny magnet with different energies when aligned with or against an external magnet. However the exact energies also depend on nearby molecules that provide 'local' magnetic fields. The 'flip' energies are measured using radio wave pulses.
- (via the hemoglobin molecule) Deoxygenated blood is magnetic while oxygenated blood is not magnetic (TAKE CHEM 1B!!).

Also See: <u>What is Functional Magnetic Resonance Imaging (fMRI)?</u>, Hannah Devlin <u>fMRI an Introduction: Michael Firbank</u> imaging (metabolic): functional Magnetic Resonance Imaging (fMRI)

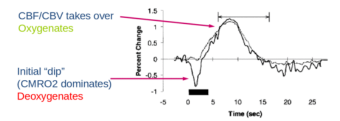
THE **BOLD** SIGNAL : **B**LOOD**O**XYGEN**L**EVEL**D**EPENDENT **fMRI** (cont.)

- Near inactive neurons there is both oxygenated and deoxygenated blood due to normal metabolic activity
- When a neuron fires there is a momentary increase in the nearby deoxygenated (magnetic) blood but then a rush of blood flow brings fresh a great concentration of oxygenated (nonmagnetic) blood in nearby capillaries
- fMRI monitors the BOLD signal for parts of the brain with active vs inactive neurons



fMRI BOLD and Haemodynamic effects

- · Increased energy consumption extracts oxygen
- \cdot $\,$ Vasodilation and CBF increase oxygen supply



http://www.slideshare.net/christina 101/fmri-introduction-newcastleuniversity-newcastle-upon-tyne

• Hemodynamic response time of ~3s

imaging (metabolic): functional Magnetic Resonance Imaging (fMRI)



The neural correlates of maternal and romantic love

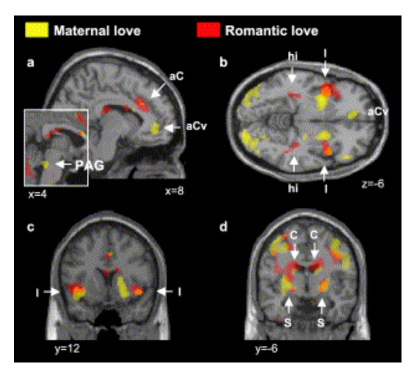
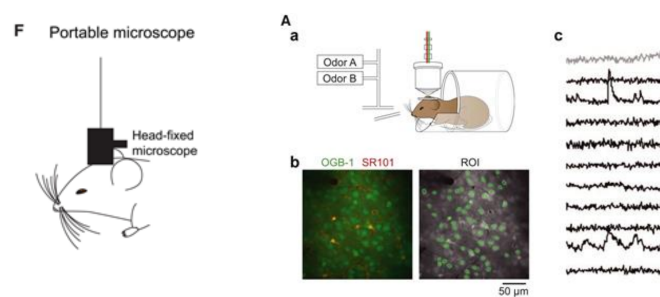


Fig. 3. Overlap between activity of maternal love and romantic love. Activity obtained in this study (contrast: cO vs. cA) was colored in yellow and overlaid on sections through a template brain, along with activity obtained in our previous study on romantic I...

Andreas Bartels, Semir Zeki

NeuroImage, Volume 21, Issue 3, 2004, 1155–1166

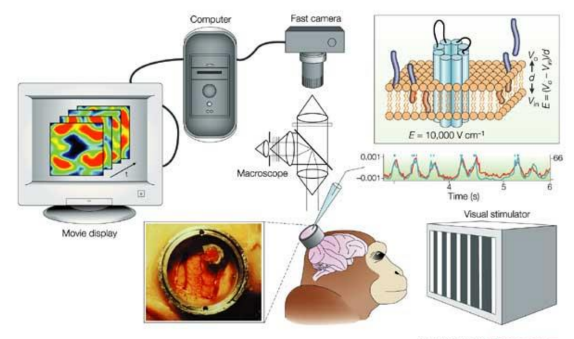
- the flow of Ca²⁺ ions is associated with neuronal activity (e.g. synaptic vesicles, remember !!)
- certain molecules that change their fluorescence ("glow") when attached to Ca²⁺
- these molecules are inserted into neurons
- the brain preparation is excited with UV light and the neurons fluorescence ("glow") with an intensity depending on concentration of Ca²⁺
- cameras used to follow the action potentials fluorescence as Ca²⁺ 'flows'

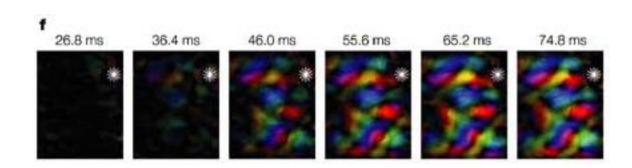


Imaging Calcium in Neurons Christine Grienberger Arthur Konnerth Neuron 73, pp 862-885 (2012)

functional imaging: voltage sensitive dyes

- certain dye molecules change their optical properties [for example the color that they 'glow' fluoresce)] when in the strong electric field of an action potential
- these dyes are painted on the surface of the cortex
- cortical areas with neurons responding to a specific stimulus fluoresce with different wavelengths (colors) than do inactive areas





http://www.nature.com/nrn/journal/v5/n11/full/nrn1536.html

3.Imaging

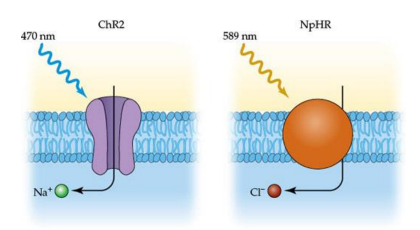
- a. Positron emission tomography (PET)
- ✓ b. Functional magnetic resonance imaging (fMRI)
- ✓ c. Calcium dyes
- ✓ d. Voltage sensitive dyes

- 4. Neural activation by external stimulus
 - a. Optogenetics
 - b. Intracranial electrical stimulation
 - c. Transcranial magnetic stimulation (TMS)

- sensory input (light, sound, rat whiskers !!)
- pharmacology (drugs, etc.)
- stimulating electrode (as in neuron lectures)
- optogenetics
- intracranial electrical stimulation
- transcranial magnetic stimulation

- genetically modify neurons to express LIGHT SENSITIVE ion channels e.g. Na⁺ or Cl⁻ channels
- activate neuron (depolarize or hyperpolarize) by light source (e.g. laser)
- image subsequent neural activity (e.g. Ca2+ imaging) or behavior in awake animal

genetically modify neurons to express LIGHT SENSITIVE ion channels
e.g. Na⁺ or Cl⁻ channels



Channelrhodopsin-2 (ChR2) and halorhodopsin (NpHR) and light-sensitive proteins used in optogenetic studies. ChR2 is a cation channel that opens in response to blue light, whereas NpHR is a chloride pump that is activated by yellow light.

http://sites.sinauer.com/psychopharm2e/webbox 09.03.html

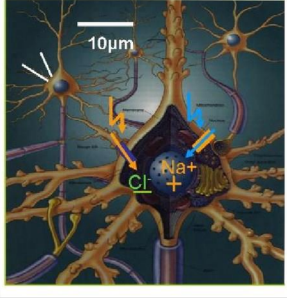
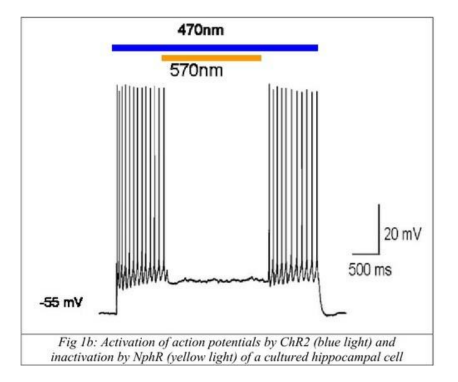
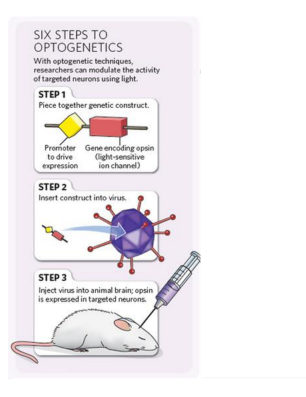


Fig 1a: Schematic representation of the action of Channelrhodopsin2 and Halorhodopsin on neural cells.

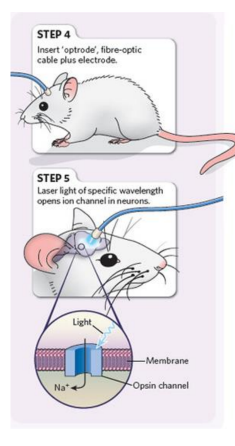


http://optogenetics.weebly.com/why--how.html

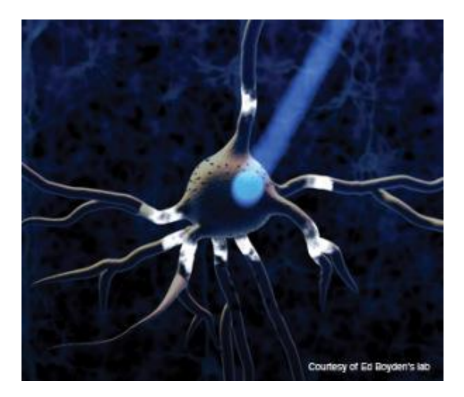
genetically modify neurons to express LIGHT SENSITIVE ion channels
e.g. Na⁺ or Cl⁻ channels



• activate neuron (depolarize or hyperpolarize) by light source (e.g. laser)



http://optogenetics.weebly.com/ why--how.html

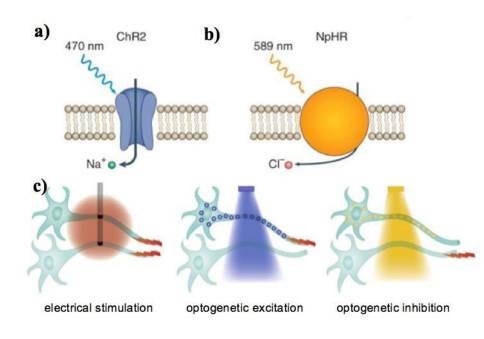


http://sciencecalling.com/2011/08/12/ light-science-optogenetics/

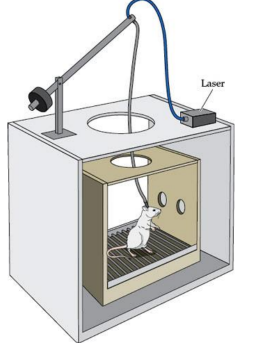


http://web.stanford.edu/group/dlab/optogenetics/

 image subsequent neural activity (e.g. Ca²⁺ imaging) or behavior in awake animal

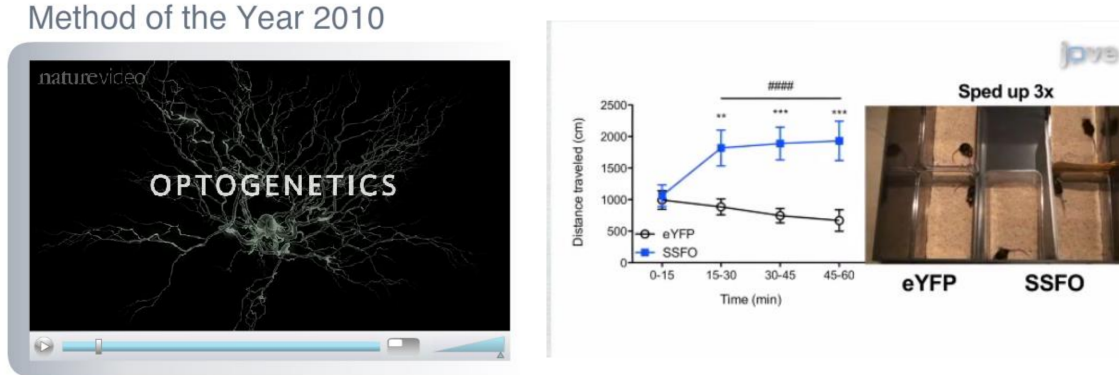


http://nmbl.stanford.edu/research/experiments/optogenetics.htm



http://sites.sinauer.com/ psychopharm2e/ webbox09.03.html

Neurons expressing ChR2 or NpHR can be exposed to light of the appropriate wavelength while the animal is behaving freely in an apparatus such as an operant chamber. Performing an operant response such as a nose-poke turns a laser light generator on for a short period of time to activate light-sensitive proteins in the vicinity of a previously implanted optic fiber 36



http://www.nature.com/nmeth/video/moy2010/index.html

http://www.jove.com/video/51483/in-vivo-optogeneticstimulation-of-the-rodent-central-nervous-system

SANTA CRUZ NEWSCENTER

Ion channel mechanics yield insights into optogenetics experiments

July 06, 2015 By Tim Stephens

SHARE THIS STORY: Y f 8" in 🗇

Optogenetics techniques, which allow scientists to map and control nerve cells using light stimulation, are being used to study neural circuits in the brain with unprecedented precision. This revolutionary technology relies on light-sensitive proteins such as channelrhodopsins, and researchers at UC Santa Cruz have now determined the molecular mechanism involved in the light-induced activation of one of these proteins.

The new findings, published July 3 in two papers in the *Journal of Biological Chemistry*, can help scientists create tailor-made proteins optimized for use in optogenetics, said David Kliger, senior author of both papers and a professor of chemistry and biochemistry at UC Santa Cruz.

"Little was known about the functional mechanism of these proteins even though they are widely used in optogenetics," Kliger said.

The researchers used fast laser spectroscopy to study the function of Channelrhodopsin-2, which is found in a type of marine algae and is widely used in optogenetics experiments. Channelrhodopsins are ion channels that control the flow of ions across cell membranes. There are many kinds of ion channels that serve different purposes in different types of cells. Nerve signals involve ion flow across the





David Kliger's lab uses fast laser spectroscopy to study biologically important molecules such as channelrhodopsin, an ion channel widely used in optogenetics experiments.



Crown 85- Visual Perception: A Window to Brain and Behavior



REPORTS INTERVIEWS

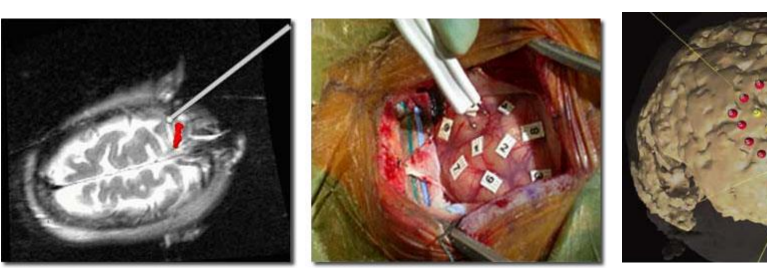
ECTS ECOMMO

One goal of Crown 85 is to introduce its students to researchers and research in the various neuroscience and perception laboratories throughout the UCSC campus. Enrolled students are required to visit one of these researchers and present a class report on the studies being conducted in their laboratory. Laboratories which have agreed to host Crown 85 student interviews include:

Prof. David Kliger Department of Chemistry and Biochemistry Structure and Spectroscopy of **read more** Visual Pigments Early events in Visual Processing • neural excitation by external stimulus: intracranial electrical stimulation

intracranial electrical stimulation: direct electrical stimulation of brain in awake subjects either with temporary or implanted electrodes (in consenting patients often those with epilepsy) in order to:

- map brain areas to guide surgical procedures
- to monitor brain function in patients
- to explore cognitive responses

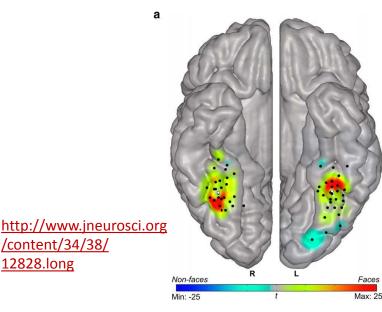


http://golbylab.bwh.harvard.edu/intracranialEEG/EEG.html

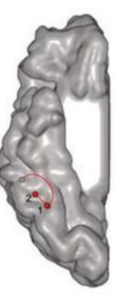
neural excitation by external stimulus: intracranial electrical stimulation

Electrical Stimulation of the Left and Right Human Fusiform Gyrus Causes Different Effects in **Conscious Face Perception**

Vinitha Rangarajan^{1,2}, Dora Hermes², Brett L. Foster^{1,2}, Kevin S. Weiner^{2,3}, Corentin Jacques^{2,3,4}, Kalanit Grill-Spector^{2,3,5}, and Josef Parvizi^{1,2,5}



12828.long



Subject 1 (R)

"Like you weren't vou. You were a different person. I noticed the eyes. I was able to see almost your whole body on your right side."



Subject 2 (R)

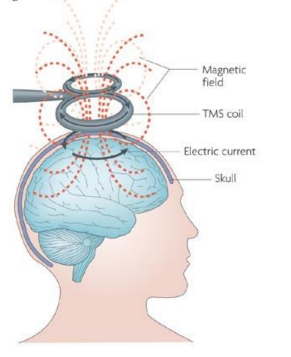
"You turned into someone else. Your face metamorphosed ... your nose got saggy and went to the left."

Bilateral face-selective ECoG responses in the fusiform gyrus. *a*, Face-selective HFB responses in the FG were measured bilaterally

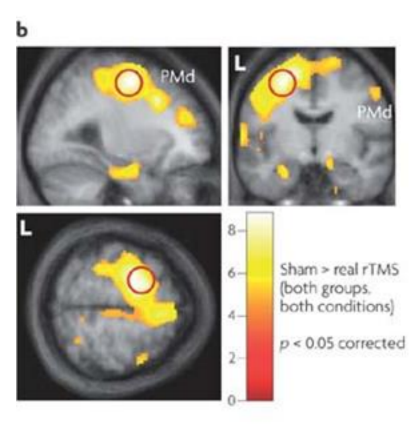


transcranial magnetic stimulation (TMS)

external magnetic coil put on head, pulses direct magnetic field to stimulate brain areas



First developed in 1985, rTMS has been studied as a possible treatment for **depression**, **psychosis** and other disorders since the mid-1990's. Clinical trials studying the effectiveness of rTMS reveal **mixed results**. When compared to a placebo or inactive (sham) treatment, some studies have found that rTMS is more effective in treating patients with major depression but other studies have found no difference in response compared to inactive treatment.



Nature Reviews | Neuroscience

b | Brain images from a study that used positron emission tomography (PET) to measure metabolic activity. The colour coding shows the areas in which activity after a 25 min session of real 1-Hz is less than that seen after a sham rTMS session. There are significant decreases in activity after real rTMS at the site of stimulation (outlined in red) as well as at many distant sites. L, left side of the brain.

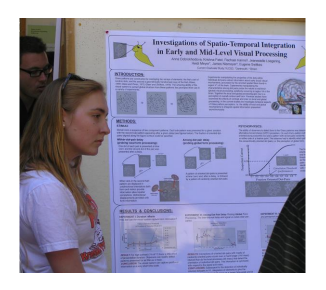
- 4. Neural activation by external stimulus
- a. Optogenetics
- b. Intracranial electrical stimulation
- c. Transcranial magnetic stimulation (TMS)

Psychophysics is the scientific **study** of the relationship between stimuli (specified in physical terms) and the sensations and perceptions evoked by these stimuli. The term **psychophysics** is used to denote both the substantive **study** of stimulus-response relationships and the methodologies used for this **study**.

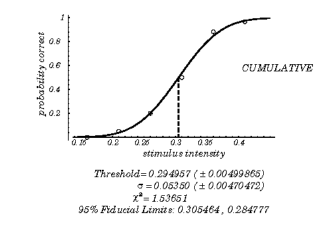
http://www.cis.rit.edu/people/faculty/montag/vandplite/page s/chap_1/ch1p2.html

psychophysics in the switkes group













	fast or slow	resolution	local or global	Invasive nonInvasive
Single cell recording	fast	high	local	invasive
Electroencephalography (EEG)	fast	low	global	noninvasive
Magnetoencephalography (MEG)	fast	moderate	global	noninvasive
Positron emission tomography (PET)	slow	low	global	noninvasive (but involves radioactive material)
fMRI	slow	low	global	noninvasive
Ca ²⁺ dyes	fast	high	intermediate	invasive
optogenetics	fast	high	intermediate	invasive

what types of neural processes would each of these be suited to measure ?

