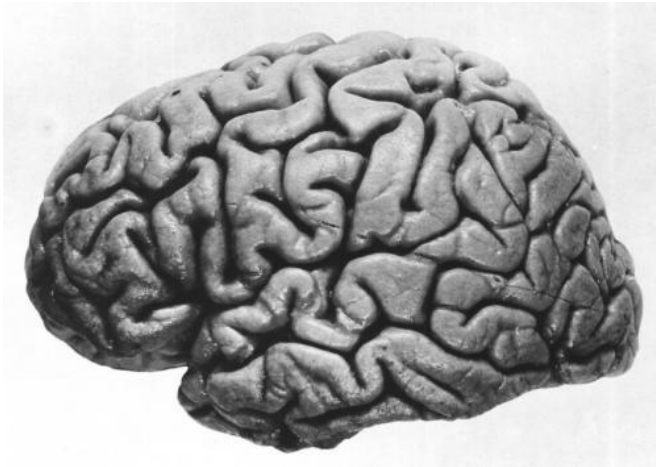


Neuroscientists map brain cell activity that occurs during the delay between sensation and action

8 September 2016, by Julie Cohen



Left hemisphere of J. Piłsudski's brain, lateral view.
Credit: public domain

A UC Santa Barbara researcher studying how the brain uses perception of the environment to guide action has a new understanding of the neural circuits responsible for transforming sensation into movement.

"Mapping perception to a future action seems simple," UCSB neuroscientist Michael Goard. "We do it all the time when we see a traffic light and use that information to guide our later motor action. However, how these associations are mapped across time in the brain is not well understood."

In a new paper, published in the journal *eLife*, Goard and colleagues at the Massachusetts Institute of Technology make progress in mapping brain activity in mice during simple but fundamental cognitive tasks. Although a mouse's brain is much smaller than a human's, remarkable structural similarities exist. The mouse brain is composed of about 75 million nerve cells or [neurons](#), which are

wired together in complex networks that underlie sophisticated behaviors.

The researchers used large-scale calcium imaging to measure the responses of [individual neurons](#) in multiple areas of the brain while mice performed a delayed response task. First, they trained mice to respond to visual stimuli—drifting bars—by either licking or withholding licking, depending on whether the bars moved vertically or horizontally. While the mice performed the task, the investigators recorded neural activity from multiple brain regions thought to be involved—including visual, parietal and frontal motor cortices.

Using a powerful laser-scanning microscope, the team was able to detect the signals from calcium indicators expressed in the neurons well below the brain's surface. Neurons normally have very low concentrations of intracellular calcium, but when they become active, calcium levels rise, increasing the fluorescence of the indicator and enabling measurement of neuron activity. In this way, the scientists were able to see which neurons were active while the mice performed the delayed response task.

"As expected, we found many neurons that responded only during the visual stimulus or the licking action, but we also found a lot of neurons that responded during other parts of the task," said Goard, an assistant professor in UCSB's Department of Psychological & Brain Sciences and Department of Molecular, Cellular and Developmental Biology. "In the frontal motor cortex, we found quite a few neurons that were active during the delay period between the visual stimulus and motor response. This led us to several new interpretations of the role that different [brain regions](#) were playing during performance of the task."

Based on the neural activity in the different brain areas, Goard and his team then used optogenetics—a method of manipulating the nerve cells with light—to inactivate neurons in a temporally precise manner to identify those that function during different parts of the task. This allowed them to figure out which areas were necessary for performing the task. For example, the team determined that the visual and parietal areas are involved in perceiving the stimulus and transforming that into a motor plan, but only the frontal motor cortex is necessary for maintaining the motor plan over the delay period.

"Using this general approach, we hope to map the essential regions for different types of cognitive tasks," Goard explained. "We are particularly interested in how [mice](#) maintain specific types of memories across distributed [brain](#) regions."

More information: Michael J Goard et al, Distinct roles of visual, parietal, and frontal motor cortices in memory-guided sensorimotor decisions, *eLife* (2016). [DOI: 10.7554/eLife.13764](https://doi.org/10.7554/eLife.13764)

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