

Order in disorder: A key feature of dendritic organization in the brain

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Local Spatial-Temporal Correlation Spatial Preferences Trials Synaptic Event Stimulus Visual Space b 0.3 Trial-to-Trial Correlation $\lambda = 4.1 \mu m$ 0.2 0.1 5 10 15 0 Dendritic Distance (µm)

(a) An example dendritic branch is shown with responsive dendritic spines, colored by spatial preference. Co-active synaptic events occur across repeated



presentations of a bar at a single spatial location. (b) The trial-to-trial correlation between spines is dependent on dendritic distance. Credit: Max Planck Florida Institute for Neuroscience

Neurons are the basic information processing structures in the brain and are composed of three main parts: dendrites, axons, and the soma. Dendrites receive and integrate synaptic inputs that are subsequently processed in the soma and relayed to other neurons via axons. A longstanding question has been to understand how synaptic inputs are arranged on the elaborate dendritic structures of individual neurons in order to better comprehend how dendrites process and transform inputs, ultimately influencing how we perceive the world around us. Recent discoveries in the laboratory of David Fitzpatrick, Ph.D., Scientific Director and CEO at the Max Planck Florida Institute for Neuroscience (MPFI), have shed more light on this issue and the universal properties of cortical neurons.

Benjamin Scholl, Ph.D. and Dan Wilson, researchers in the Fitzpatrick lab, have been studying the spatial arrangement of synaptic inputs onto dendrites and how this arrangement impacts computations in visual cortex. Previous work by Wilson et al. (2015) revealed that orientation selectivity and functional clustering of synaptic inputs in visual cortical neurons correlates with localized dendritic events, sharpening somatic tuning. In their recent paper, published in *Neuron* on November 2nd, Scholl and Wilson examined whether this clustering exists at a finer scale and across the diversity of functional inputs cortical cells receive.

To investigate fine-scale synaptic clustering, the researchers presented simple visual stimuli to activate individual excitatory inputs onto individual dendritic spines, small mushroom-shaped protrusions on the dendritic arbors. These visual stimuli allowed the authors to map spatial



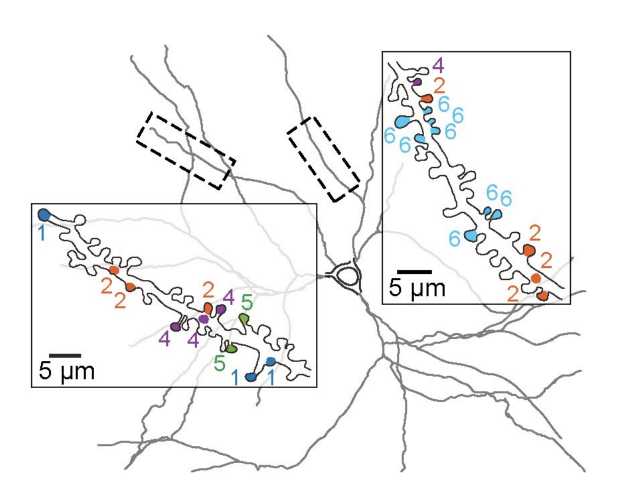
receptive fields - regions in visual space where stimuli elicit responses from particular neurons or synaptic inputs - and polarity - the preference for luminance increments (ON) or decrements (OFF). Large numbers of spines on individual layer 2/3 neurons were mapped and visualized using in vivo two-photon imaging of a genetically encoded calcium indicator, providing a measure of subcellular and cellular activity.

Within individual dendritic branches and across the dendritic field, they found no evidence for a global organization. Upon closer inspection, however, they discovered robust functional synaptic clustering locally. Neighboring spines separated by less than 10 µm share similar functional properties across the diversity of sensory features they encode. Further, neighboring spines exhibit spontaneous and sensory-driven co-activity that can occur independent of global dendritic calcium events, driven largely by action potentials initiated at the soma. Distance-dependent relationships for these features exist at a spatial scale of 5-10 µm, and this omnipresent spatial length constant may indicate a fundamental biological limit to short-range plasticity mechanisms operating within the dendrite. "The clustering of inputs appears more universal than we might have previously thought," said Scholl. "We are just beginning to appreciate the complexities of the dendritic field."

They also revealed, via machine learning, that fine-scale functional synaptic clusters exist within the broader population of spines, and these clusters appear to be selective for features both resembling and distinct from the somatic output. Clustered inputs could amplify such functional features, but the role of synaptic clustering in shaping somatic response is more complex. These distributed functional clusters could enhance the computational power of neurons by allowing greater flexibility of responses to a wide range of sensory inputs, increasing dendritic sensitivity without sacrificing dynamic range, and facilitating nonlinear interactions within individual dendrites.



Appreciating the integration and processing of distributed networks at the subcellular level is key to understanding organization at the cellular level and beyond. "It is critical that we understand how the inputs to a given neuron are arranged within its dendritic field so that we can understand how <u>individual neurons</u> integrate the information arising from thousands of synaptic inputs," said Fitzpatrick. What are the consequences of such an arrangement for the function of a neuron and for the network in which it's embedded? Currently, there is a large gap in our scientific knowledge when it comes to perceiving the role that dendrites play in neuronal circuits. "What makes this paper really exciting," said Fitzpatrick, "is that it bridges that gap, putting cellular principles in the context of how a network operates."





The dendritic locations of a subset of spines. Each spine is colored to indicate the group label from k-means clustering; the group label from k-means clustering is also indicated next to each spine. Credit: Max Planck Florida Institute for Neuroscience

Moving forward, the lab is interested in exploring the development and maturation of spines and synaptic clusters. Ongoing work involves chronic imaging of the structure and function of individual spines during development and changes to visual experience to determine whether emerging spines adopt the same functional properties of their more adjacent, stable predecessors. Such research could provide greater insight into the role of synaptic clustering during development and plasticity mechanisms throughout an individual's lifespan.

More information: *Neuron* (2017). <u>DOI:</u> 10.1016/j.neuron.2017.10.017

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