* Dominic Giannangeli designs a custom MEMS device which allows to investigate neuroplasticity, the spatial and network reorganization of neurons in response of other neurons feedback. The device has three compartments, cells in one chamber is connected to the two others, the two other chambers are only connected to the central one.
* Primary rat cortical neurons are grown in each microfluidic chamber, each side extending their axons through micro-channels with the other side.
* Electrical stimuli are applied to the neurons of one of the extreme chambers and not to neurons in the other one, thus creating neural feedback in only one direction: from one of the peripheral chambers to the central chamber. Neuronal activity and physiological changes in the connections between each well are then studied and compared.
* Organotypic cells compared to cell cultures maintain cytoskeleton and are less limited in term of axon growth or synapse formation. We keep the interconnection of the three chambers used in the initial model, but instead of cortical neurons, rat cortical slices of about 350 m will be cut using a tissue chopper and placed onto petri dishes with three polydimethylsiloxane (PDMDS) wells separated by microchannels. Cultures will be covered with a medium and placed into a tissue incubator. Each slice will be perfused with artificial cerebro-spinal fluid (ASCSF) and drugs. Electrical activity will be stimulated by electrodes in the peripheral dish with carefully calibrated pulses to generate an action potential, the neuronal response will be recorded with microelectrodes placed on the slice of the central well. An electrode will also be attached to the neurons of the dish with no explicit stimuli, to record if neural activation sent by one of the peripheral wells, is transmitted through the central well to the other peripheral compartment, which will allow to have an understanding of the full neural pathway. Signals detected by the electrodes will be amplified and recorded for analysis.
* The previous model is simple but may not fully represent the complexity of interactions between neurons. 2-D cultures may affect the spatial growth of the neural network. In addition, a 3-D cultures like organoids have better cell-cell interactions and simulate cellular functions and neural signaling which resemble more the ones existing in in-vivo tissues. We will also use neural activity optical detection with small florescent molecules like ANEP [6] or ANNINE-6plus dyes, with a careful design protocol to avoid photobleaching or good permeation to the cell membrane.
* It has also been shown that organoids in culture over 9 months exhibit mature astrocytes, showing formation of dendritic spines, and moreover neuronal activity within organoids could be controlled using light stimulation of photosensitive cells [5].
* In addition, the various models previously discussed will show significant limitations to understand the neural network reconstruction which happens for example in stroke or brain injury where some components of the microenvironment are beneficial like blood capillaries, oxygen and neurotrophic factor or damaging like neuroinflammatory. In order to mimic closer, the in-vivo environment in which these cells evolve, the design of a more complex model like organ-on-a-chip may ultimately be required.

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