

Poster presentation

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## Influence of external input on waxing and waning of neuronal network oscillations

Oscar J Avella Gonzalez\*<sup>1</sup>, Ronald van Elburg<sup>2</sup>, Huibert Mansvelder<sup>1</sup>, Jaap van Pelt<sup>1</sup> and Arjen van Ooyen<sup>1</sup>

Address: <sup>1</sup>Department of Integrative Neurophysiology, VU University Amsterdam, 1081 HV The Netherlands and <sup>2</sup>Department of Artificial Intelligence, University of Groningen, 9700 AB, The Netherlands

Email: Oscar J Avella Gonzalez\* - oscar.avella@cncr.vu.nl

\* Corresponding author

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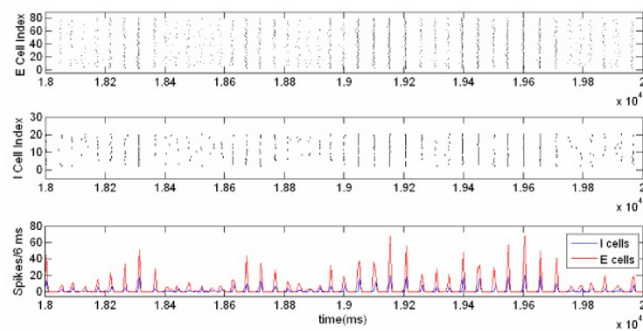
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### Introduction

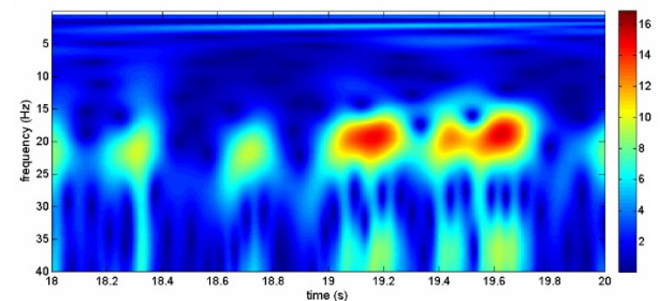
Experimental observations have reported modulation of cortical oscillations as phases of high synchronization (waxing) followed by periods of reduced synchronization (waning) [1-3]. Although the phenomenon is present in almost all frequency bands, it is still not understood how this is driven. Here we study whether this phenomenon can occur in a network of inhibitory (I) and excitatory (E) cells and what effect external inputs have.

### Methods

Using NEURON, we model a network of  $N_e$  excitatory and  $N_i$  inhibitory cells such that  $N_e/N_i = 4$ . The cells have a single compartment, and include passive channels and voltage dependent  $\text{Na}^+$ ,  $\text{K}^+$  channels. Synaptic connections are random, projecting GABA synapses from I to I and I to E cells and AMPA synapses from E to E and E to I cells. To stimulate the network, each cell receives a baseline of current and a stream of spikes delivered at random intervals across the simulated period.



**Figure 1**  
Raster plots of E (top) and I (middle) populations, during waxing and waning of a beta oscillation, and firing rate histograms (bottom).



**Figure 2**  
Wavelet transform of the activity in the E population for the same time period as shown in figure 1.

## Results

We show that in a stable oscillatory network, waxing and waning occurs without the need for other synaptic mechanisms than the spike generating  $K^+$  and  $Na^+$  channels. The phenomenon can be modulated by changing the characteristics of the external input, such as number of spikes, mean inter-spike interval, randomness and whether E or I cells receive the external input. See figures 1 and 2.

## Acknowledgements

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## References

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