

Poster presentation

Open Access

## Reduced models of striatal neurons: dopamine modulation and dynamics

Mark D Humphries\*, Nathan Lepora, Ric Wood and Kevin Gurney

Address: Adaptive Behaviour Research Group, University of Sheffield, Sheffield, S10 2TN, UK

Email: Mark D Humphries\* - [m.d.humphries@shef.ac.uk](mailto:m.d.humphries@shef.ac.uk)

\* Corresponding author

from Eighteenth Annual Computational Neuroscience Meeting: CNS\*2009  
Berlin, Germany. 18–23 July 2009

Published: 13 July 2009

BMC Neuroscience 2009, **10**(Suppl 1):P321 doi:10.1186/1471-2202-10-S1-P321

This abstract is available from: <http://www.biomedcentral.com/1471-2202/10/S1/P321>

© 2009 Humphries et al; licensee BioMed Central Ltd.

Loss of dopamine cells in Parkinson's Disease and its animal models leads to profound motor deficits. An intact dopamine system also seems to be critical for many forms of learning [1]. Much work on understanding these roles of dopamine has focused on the striatum, the main input nucleus of the basal ganglia, as the striatum is the main locus of dopamine's action in the vertebrate brain [2]. Damage to the striatum itself also impairs both motor action and learning [3]. Thus, the twin problems of understanding the computational roles of dopamine and the striatum are inseparably intertwined.

Understanding dopamine's effects on the complex striatal microcircuit ideally requires large-scale models that replicate the neuron types, numbers, and connectivity at one-to-one scale. To build at such scales, we require individual neuron models that are simple enough to be computationally tractable, but sufficiently complex to capture key membrane properties that contribute to the characteristic behavior of a neuron species. Our neuron model of choice is the recent canonical spiking model of Izhikevich [4]. However, it has not yet been extended to account for the action of neuromodulators.

We extend the striatal medium spiny (MSN) and fast-spiking (FS) interneuron models of [5] to account for dopaminergic modulation of intrinsic ion channels and synaptic inputs. We use data from a recent 189 compartment model of the MSN [3] to tune our simple model of that neuron under both current injection and spiking

input regimes with varying activation of dopamine D1- and D2-type receptors. The reduced models capture the input-output relationships for both current injection and spiking input with remarkable accuracy. We derive a full set of stability properties for the original and dopamine modulated forms of the MSN model. We use these to establish that the dopamine models do not change the stability properties and hence the models predict that the MSN is not bistable in either baseline or dopamine-saturated conditions. Our extensions to the simple model of the FS interneuron are consistent with the existing data, but tuning the new parameters is made difficult by the lack of quantitative results from experimental work. Our work thus establishes reduced yet accurate dopamine-modulated forms of MSN and FS interneuron models, suitable for use in large-scale models of the striatum. Moreover, these also provide a tractable framework for further study of dopamine's effects on individual neuron computation.

### Acknowledgements

We thank Jason Moyer for generously sending us the input-output data from the multi-compartment medium spiny neuron model. This work was funded by the EU Framework 6 Project IST-027819-IP, EPSRC Research Grant EP/C516303/1, and the EPSRC "CARMEN" e-Science Pilot Project.

### References

1. Schultz W: **Behavioral dopamine signals.** *Trends Neurosci* 2007, **30**:203-210.
2. Dawson TM, Gehlert DR, McCabe RT, Barnett A, Wamsley JK: **D-1 dopamine receptors in the rat brain: a quantitative autoradiographic analysis.** *J Neurosci* 1986, **6**:2352-2365.

3. Yin HH, Knowlton BJ: **The role of the basal ganglia in habit formation.** *Nat Rev Neurosci* 2006, **7**:464-476.
4. Izhikevich EM: *Dynamical Systems in Neuroscience Cambridge, MA: MIT Press; 2007.*
5. Moyer JT, Wolf JA, Finkel LH: **Effects of dopaminergic modulation on the integrative properties of the ventral striatal medium spiny neuron.** *J Neurophysiol* 2007, **98**:3731-3748.

Publish with **BioMed Central** and every scientist can read your work free of charge

*"BioMed Central will be the most significant development for disseminating the results of biomedical research in our lifetime."*

Sir Paul Nurse, Cancer Research UK

Your research papers will be:

- available free of charge to the entire biomedical community
- peer reviewed and published immediately upon acceptance
- cited in PubMed and archived on PubMed Central
- yours — you keep the copyright

Submit your manuscript here:  
[http://www.biomedcentral.com/info/publishing\\_adv.asp](http://www.biomedcentral.com/info/publishing_adv.asp)

