The NeuroML ecosystem for standardised multi-scale modelling in neuroscience

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28/02/2026

What we've been up to on the NeuroML front

- Google summer of code (GSoC)
- Paper and general improvements

Google Summer of Code

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 - Conversions of models to NeuroML (to allow them to be re-used and featured on OSB)
 - Get folks looking into and using NeuroML in different use cases
 - · Get folks to improve NeuroML where possible

GSoC: Anuja: convert Allen Institute models to NeuroML

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- Steps:
 - Automate download of models using the Allen SDK
 - Automate conversion to NeuroML
 - Plot comparison graphs to validate conversions (LIF models, Detailed)
 - Document comparison, usage, update OMV tests for CI

GSoC: Anuja: example figure I

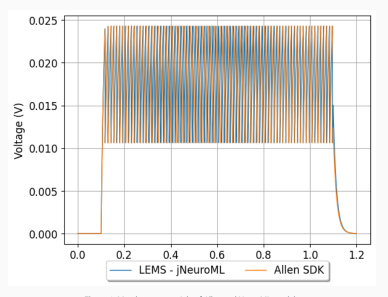


Figure 1: Membrane potentials of Allen and NeuroML models

GSoC: Anuja: example figure II

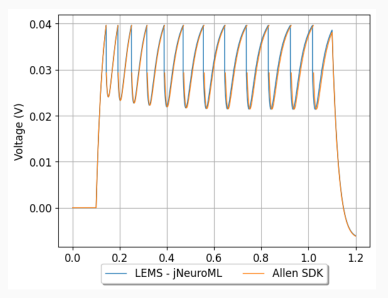


Figure 2: Membrane potentials of Allen and NeuroML models

GSoC: Anuja: example figure III

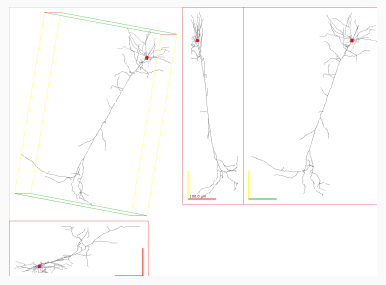


Figure 3: Morphology of example cell

GSoC: Anuja: example figure IV

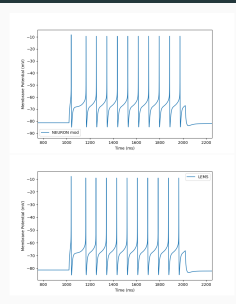


Figure 4: NEURON vs NeuroML output

GSoC: Shayan: convert BahlEtAl2012 (Reduced L5 Pyr Cell) to NeuroML

MSc/BSc from IIT Kharagpur (Maths and Computing)

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- MSc/BSc from IIT Kharagpur (Maths and Computing)
- Source models: OSB/BahlEtAl2012
- Steps:
 - Convert ion channels to NeuroML, validate, compare with NEURON mod files
 - Implement single compartment cell model with passive channels
 - · incrementally add ion channels
 - · compare with NEURON model
 - · Implement multi-compartmental cell, repeat
 - · Document comparison, usage, update OMV tests for CI
 - Interactive notebook to reproduce figures from paper using NeuroML models

GSoC: Shayan: example figures

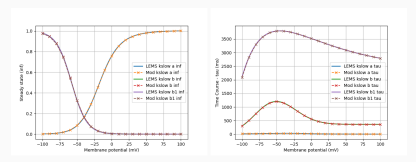


Figure 5: Comparing ion channels: steady state, time constant

GSoC: Shayan: example figures II

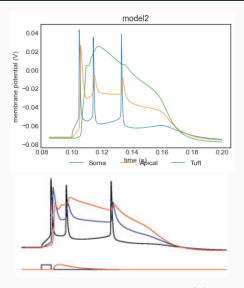


Figure 6: Replicating figures from the paper (8d)

GSoC: Rahul: convert HH tutorial to Jupyter notebook

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- Aerospace engineer (MTech, IIT Kharagpur), working in South Korea
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- Steps:
 - · Investigate Jupyter widgets
 - Convert pure Python HH tutorial to use Jupyter widgets
 - · Investigate NeuroML based tutorial
 - Investigate conversion of NeuroML to Jupyter widgets
 - Update sphinx documentation for ReadTheDocs site
 - Document usage.

GSoC: Rahul: example I

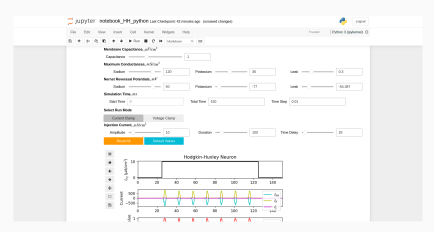


Figure 7: Pure python tutorial converted to use Jupyter Widgets

GSoC: Rahul: example II

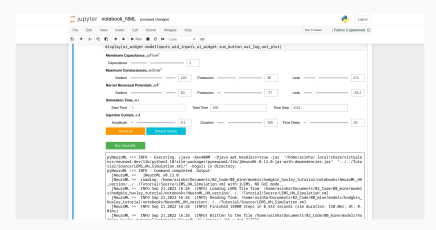


Figure 8: NeuroML tutorial converted to use Jupyter Widgets

Paper and general updates

Convince readers (research community) to use NeuroML for their modelling work*.

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Convince readers (research community) to use NeuroML for their modelling work*.

- *instead/ahead of other tools
- *on a daily basis
 - not as an afterthought for standardisation—once the paper has been published no one has time to re-write model (or re-process data!) to standardise
 - the carrot, "standardisation is good for science", isn't enough in a mostly resource limited academic/research system
 - requires stick: "for this grant, you must ..."; "for this journal, you must ..."

You should use NeuroML because

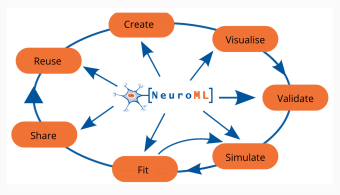


Figure 9: NeuroML overview figure from paper

You should use NeuroML because

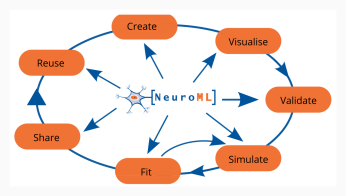


Figure 9: NeuroML overview figure from paper

These claims are all true, and have been for quite a while.

- information on NeuroML needs to be easy to find:
 - website
 - · point of entry for completely new users: first impression
 - replaced with modern looking static page that redirects to individual pages in docs
 - all information migrated to docs

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 - · website
 - · point of entry for completely new users: first impression
 - replaced with modern looking static page that redirects to individual pages in docs
 - · all information migrated to docs
 - docs
 - · re-organised, modernised
 - include tutorials, interactive tutorials via Jupyter note books, how-tos
 - · complete searchable schema docs
 - · still not yet fully complete

- usability: so much can be done, but can it be done easily?
 - · has not been clearly easier to use than other tools
 - Python API exists, but we haven't taken advantage of it enough to make life easier for users yet

Example: create: single neuron Izhikevich network (from docs)

```
nml_doc = NeuroMLDocument(id="IzhSingleNeuron")
 2
 3
     izh0 = Izhikevich2007Cell(
 4
          id="izh2007RS0", v0="-60mV", C="100pF", k="0.7nS_per_mV", vr="-60mV",
 5
          vt="-40mV", vpeak="35mV", a="0.03per ms", b="-2nS", c="-50.0mV", d="100pA")
 6
      nml doc.izhikevich2007 cells.append(izh0)
 8
      net = Network(id="IzhNet")
9
      nml doc.networks.append(net)
10
11
      size0 = 1
12
      pop0 = Population(id="IzhPop0", component=izh0.id, size=size0)
13
      net.populations.append(pop0)
14
15
      pg = PulseGenerator(
16
          id="pulseGen %i" % 0, delay="0ms", duration="1000ms",
17
          amplitude="0.07 nA"
18
19
      nml doc.pulse generators.append(pg)
20
      exp_input = ExplicitInput(target="%s[%i]" % (pop0.id, 0), input=pg.id)
      net.explicit_inputs.append(exp_input)
21
22
23
      nml_file = 'izhikevich2007_single_cell_network.nml'
24
      writers.NeuroMLWriter.write(nml_doc, nml_file)
25
      print("Written network file to: " + nml_file)
26
27
      validate_neurom12(nml_file)
```

Inspect/visualise network



Figure 10: Generated network graph

```
>>> nml doc.summarv()
* NeuroMLDocument: IzhSingleNeuron
   Izhikevich2007Cell: ['izh2007RS0']
  PulseGenerator: ['pulseGen_0']
  Network: IzNet
    1 cells in 1 populations
     Population: IzhPopO with 1 components of type izh2007RSO
    O connections in O projections
    O inputs in O input lists
    1 explicit inputs (outside of input lists)
     Explicit Input of type pulseGen_0 to IzhPopO(cell 0), destination: unspecified
```

- extremely declarative
 - components (Izhikevich2007Cell, Network, Population, PulseGenerator, ExplicitInput) clearly visible
 - components and dynamics fully, formally documented in schema docs
 - component parameters clearly visible
 - units/dimensions explicitly mentioned

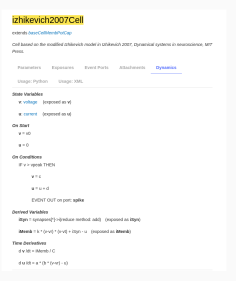


Figure 11: Schema docs on Izhikevich2007Cell

- validation without simulation (no simulator has this)
 - Level 1 validation: units/dimensions, structure of model checked against schema
 - Level 2 validation: extra "logical" checks

- validation without simulation (no simulator has this)
 - Level 1 validation: units/dimensions, structure of model checked against schema
 - Level 2 validation: extra "logical" checks
- easy simulation with different simulators
- easy visualisation and inspection of model: network structure, connectivity matrices, LEMS simulation graph, morphology figures, model summary

- information on components: must switch back and forth between docs and code
 - Not too bad: required for most simulators/programming languages

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 - how do I know that Network \rightarrow Populations \rightarrow ?

- information on components: must switch back and forth between docs and code
 - Not too bad: required for most simulators/programming languages
- unclear how components fit together
 - how do I know that Network → Populations →?
- Too many "under the hood" bits that users are expected to know:
 - nml_doc.izhikevich2007_cells.append(izh0)
 - · how do users know this?
 - · read the schema docs?
 - · read the NeuroML Python API documentation?
 - read the NeuroML Python API source? (currently 64,000 lines of code)

- validation upon model completion
 - using jnml, so requires Java
 - better than errors on run, but still quite late

- · validation upon model completion
 - using jnml, so requires Java
 - better than errors on run, but still quite late
- NeuroML generated NEURON code is less performant than native NEURON code
 - because hand-written mod files can be optimised, while we rely on a template to generate them
 - · TODO: optimisation of template

Example: create: single neuron Izhikevich network (devel)

```
nml_doc = component_factory("NeuroMLDocument", id="IzhSingleNeuron")
      nml doc.info(show contents=True)
 3
 4
     izh0 = nml_doc.add(
 5
          "Izhikevich2007Cell".
          id="izh2007RS0", v0="-60mV", C="100pF", k="0.7nS_per_mV", vr="-60mV",
 6
          vt="-40mV", vpeak="35mV", a="0.03per_ms", b="-2nS", c="-50.0mV", d="100pA")
 8
      izh0.info(show_contents=True)
9
10
      net = nml_doc.add("Network", id="IzNet", validate=False)
11
12
      size0 = 1
13
      pop0 = net.add("Population", id="IzhPop0", component=izh0.id, size=size0)
14
15
      pg = nml doc.add(
16
          "PulseGenerator".
17
          id="pulseGen_%i" % 0, delay="0ms", duration="1000ms",
18
          amplitude="0.07 nA"
19
20
      exp_input = net.add("ExplicitInput", target="%s[%i]" % (pop0.id, 0), input=pg.id)
21
22
      nml doc.validate(recursive=True)
23
24
      nml_file = 'izhikevich2007_single_cell_network.nml'
25
      writers.NeuroMLWriter.write(nml doc, nml file)
26
      print("Written network file to: " + nml_file)
```

The component factory

- single factory function to create new components
- runs extra checks
 - are all arguments (parameters) correct?
 - is the component valid (level 1 validation for each component at build-time)?

```
>>> izh0 = component_factory("Izhikevich2007Cell")
ValueError: Validation failed:
- Izhikevich2007Cell (None): Required value v0 is missing
- Izhikevich2007Cell (None): Required value v1 is missing
- Izhikevich2007Cell (None): Required value v1 is missing
- Izhikevich2007Cell (None): Required value v1 is missing
- Izhikevich2007Cell (None): Required value vpeak is missing
- Izhikevich2007Cell (None): Required value a is missing
- Izhikevich2007Cell (None): Required value b is missing
- Izhikevich2007Cell (None): Required value c is missing
- Izhikevich2007Cell (None): Required value d is missing
- Izhikevich2007Cell (None): Required value C is missing
- Izhikevich2007Cell (None): Required value C is missing
- Izhikevich2007Cell (None): Required value id is missing
```

The component factory

```
>>> izh0 = component_factory(
    "Izhikevich2007Cell",
    id="izh2007Rs0", v0="-60mV", C="100pF", k="0.7nS_per_mV", vr="-60mV",
    vt="-40ms", vpeak="35mV", a="0.03per_ms", b="-2nS", c="-50.0mV", d="100pA")

ValueError: Validation failed:
    Izhikevich2007Cell (izh2007Rs0): Value "-40mS" does not match xsd pattern restrictions:
    \[['^(-?([0-9]*(\\.[0-9]+)?)([eE]-?[0-9]+)?[\\s]*(V|mV))$']]]
```

New add method

- uses component factory to create a new component and run checks
- smart enough to know where the new element needs to go in the parent

```
1    izh0 = Izhikevich2007Cell(
2    id="izh2007RS0", v0="-60mV", C="100pF", k="0.7nS_per_mV", vr="-60mV",
3    vt="-40mV", vpeak="35mV", a="0.03per_ms", b="-2nS", c="-50.0mV", d="100pA")
4    nml_doc.izhikevich2007_cells.append(izh0)
```

VS

```
1 izh0 = nml_doc.add(
2 "Izhikevich2007Cell",
3 id="izh2007RS0", v0="-60mV", C="100pF", k="0.7nS_per_mV", vr="-60mV",
4 vt="-40mV", vpeak="35mV", a="0.03per_ms", b="-2nS", c="-50.0mV", d="100pA")
```

Inspect each component individually

```
izh0.info(show_contents=True)
Izhikevich2007Cell -- Cell based on the modified Izhikevich model in Izhikevich 2007, Dynamical

→ systems in neuroscience, MIT Press

Please see the NeuroML standard schema documentation at
Valid members for Izhikevich2007Cell are:
* b (class: Nml2Quantity conductance, Required)
       * Contents ('ids'/<objects>): -2nS
* C (class: Nml2Quantity_capacitance, Required)
       * Contents ('ids'/<objects>): 100pF
* c (class: Nml2Quantity_voltage, Required)
       * Contents ('ids'/<objects>): -50.0mV
* d (class: Nml2Quantity_current, Required)
       * Contents ('ids'/<objects>): 100pA
* neuro_lex_id (class: NeuroLexId, Optional)
* metaid (class: MetaId, Optional)
* v0 (class: Nml2Quantity voltage, Required)
       * Contents ('ids'/<objects>): -60mV
* id (class: NmlId, Required)
       * Contents ('ids'/<objects>): izh2007RS0
```

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Component Type info without creating a new component: ctinfo

```
>>> ctinfo("ExpOneSynapse")
ExpOneSynapse -- Ohmic synapse model whose conductance rises instantaneously by ( **gbase** *

→ **weight** ) on receiving an event, and which decays exponentially to zero with time course

→ **tauDecay**
Please see the NeuroMI standard schema documentation at

→ https://docs.neuroml.org/Userdocs/NeuroMLv2.html for more information.

Valid members for ExpOneSynapse are:
* neuro lex id (class: NeuroLexId, Optional)
* gbase (class: Nml2Quantity_conductance, Required)
* metaid (class: MetaId, Optional)
* erev (class: Nml2Quantity_voltage, Required)
* notes (class: xs:string, Optional)
* id (class: NmlId, Required)
* properties (class: Property, Optional)
* annotation (class: Annotation, Optional)
* tau_decay (class: Nml2Quantity_time, Required)
```

Where does this component type fit?

```
>>> ctparentinfo("HHRate")
Please see the NeuroML standard schema documentation at
Valid parents for HHRate are:
* GateHHRates
       * forward rate (class: HHRate, Required)
       * reverse_rate (class: HHRate, Required)
* GateHHRatesInf
       * forward_rate (class: HHRate, Required)
       * reverse_rate (class: HHRate, Required)
* GateHHRatesTau
       * forward rate (class: HHRate, Required)
       * reverse_rate (class: HHRate, Required)
* GateHHRatesTauInf
       * forward rate (class: HHRate, Required)
       * reverse_rate (class: HHRate, Required)
* GateHHUndetermined
       * forward rate (class: HHRate, Optional)
       * reverse rate (class: HHRate, Optional)
```

Additions to make multi-compartment cell building easier

- set_init_memb_potential() instead of cell.biophysical_properties.membrane_properties.init_memb_potential
- add_channel_density()...
- add_segment, add_unbranched_segment_list...
 - also takes care of NeuroLex (now InterLex) Ids, used by NEURON to create "sections".
 - another hidden feature of NeuroML

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- update docs, add more tutorials/recipes

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 - · import works, but cannot modify model
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- Migrate all code to Python (so no more Java required)
 - · long term, multi-year project, requires another grant