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1 **A Graphical and Computational Modelling Platform for Biological Pathways**

2

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15 EDITORIAL SUMMARY: This is a biologist-friendly modelling scheme facilitating the capture and
16 visualization of knowledge on biological pathways and how components interact. Moreover, when
17 parameterised, these pathway models can be used directly to run simulations of their activity and test
18 hypotheses.

19 TWEET: A biologist-friendly modelling scheme to visualize and computationally model biological
20 pathways @roslininstitute @mrc_crh

21

22 **[AU: Please highlight to the editor up to 3 key references from your lab that demonstrate the**
23 **development/use of the protocol; we hope to highlight 1 or more of these particular references in the**
24 **'associated links' box on the online article page].**

25 Please use references

26 #1: O'Hara et al PMID: 27509052

27 #14: Raza et al PMID: 20478018

28 #41: Polak et al PMID: 28386100

29

30 **Key words:** Pathway, notation system, model, dynamic modelling, Petri Net, simulation, SBGN, mEPN

31 **Abstract**

32 A major endeavour of systems biology is the construction of graphical and computational models of
33 biological pathways as a means to better understand their structure and function. Here, we present a
34 protocol for a biologist-friendly graphical modelling scheme which facilitates the construction of detailed
35 network diagrams, summarising the components of a biological pathway (such as proteins, biochemicals
36 etc.) and how they interact. These diagrams can then be used to simulate activity flow through a pathway,
37 thereby modelling its dynamic behaviour. The protocol is divided into four sections: 1) Assembly of
38 network diagrams using the modified Edinburgh Pathway Notation (mEPN) scheme and yEd network
39 editing software using pathway information obtained from published literature and databases of molecular
40 interaction data, 2) parameterisation of the pathway model within yEd through the placement of 'tokens'
41 based on the known or imputed amount or activity of a component, 3) model testing through visualization
42 and quantitative analysis of the movement of tokens through the pathway using network analysis tool
43 BioLayout *Express*^{3D}, 4) optimisation of model parameterisation and experimentation. This is the first
44 modelling approach that combines a sophisticated notation scheme for depicting biological events at the
45 molecular level, with a Petri net-based flow simulation algorithm and powerful visualisation engine with
46 which to observe the dynamics of the system being modelled. Unlike many mathematical approaches to
47 modelling pathways, it does not require the construction of a series of equations or rate constants for
48 model parameterisation. Depending on a model's complexity and the availability of information, its
49 construction can take days to months, and, with refinement, possibly years. However, once assembled
50 and parameterised, a simulation run, even on a large model, typically takes only seconds. Models

51 constructed using this approach provide a means of knowledge management, information exchange, and
52 through the computation simulation of their dynamic activity, a means to generate and test hypotheses,
53 and predict a system's behaviour when perturbed.

54 **Introduction**

55 The era of molecular biology has resulted in the generation of vast amounts of data on biological
56 processes, ranging from in-depth studies of one or two molecules and their interactions, to large sets of
57 omics data. These data are currently scattered in the literature and databases and difficult to connect
58 together. Those trying to learn about a particular biological process or pathway often start by studying the
59 primary literature and reviews. However, relying only on the medium of the written word it can often be a
60 struggle to understand the available knowledge as a series of interconnected events. The task is made
61 more difficult as the literature often refers to the same pathway component by different names, which may
62 not or not be their official names (as dictated by nomenclature committees). Representation of biological
63 systems as graphical models, i.e. diagrams, can in principle circumvent these issues by presenting a
64 system in a visually intuitive manner using a standardized notation scheme to represent pathway
65 components and the interactions between them. One of the ultimate goals of a pathway model is the
66 ability to use it for computational simulations, thereby supporting hypothesis generation and experimental
67 design. Use of a system that fulfils these criteria could benefit any scientist working in experimental
68 biology.

69
70 We have developed a modelling platform that combines elements of other approaches¹. The modified
71 Edinburgh Pathway Notation (mEPN) scheme was first published in 2008², refined in 2010³, and is
72 presented here in its current form (BOX 1). Representing the interactions between biological components
73 in the context of a pathway diagram is a challenge, and a number of notation schemes have been
74 proposed³⁻⁹. In an effort to standardise pathway diagrams, the Systems Biology Graphical Notation
75 (SBGN) community proposed standards for pathway depiction, including the process description (PD)
76 language⁷ based on ideas first proposed by Kitano *et al.*⁴. In SBGN-PD diagrams (and in the modelling
77 approach described here), components of a pathway are depicted using a standard set of shapes

78 (glyphs), and both the nature of the interactions between components and the products of those
79 interactions must be shown explicitly. Pathway models are constructed where *entity nodes* represent
80 molecular components, *process nodes* represent the different types of interactions that can occur
81 between the components, and *edges* link entity and process nodes. Since PDs were first described
82 various models have been constructed based on this approach^{2,10-15} and there is a growing number of
83 software tools that support model creation (using SBGN-compliant languages), e.g., CellDesigner^{16,17},
84 NaviCell¹¹, KEGG Mapper¹⁸, ReactomeFiviz¹⁹, iPath²⁰ and SBGN-Ed²¹. There are also a number of
85 centralised databases providing pathway resources of this type²²⁻²⁵. The mEPN scheme used here to
86 model pathway systems is similar to the SBGN 'process description language' but with important
87 differences in how both components and events are represented. In particular, mEPN supports the
88 representation of wider variety of biological components and processes, simplifies the depiction of
89 complexes, promotes the use of standard nomenclature, and importantly, diagrams can be used directly
90 for the computational modelling of system dynamics (for a more complete description of mEPN and
91 comparison to the SBGN-PD language, see O'Hara 2016¹). mEPN pathway models can be drawn using
92 the free graph editing software yEd (yWorks, Tübingen, Germany; www.yworks.com), and since the
93 notation scheme was first described^{2,3} has been formalised so as to support the use of models for
94 pathway activity simulations¹.

95 Numerous mathematical approaches exist to simulate system dynamics including ordinary and partial
96 differential equations, qualitative differential equations and stochastic equations. The Systems Biology
97 Markup Language (SBML) has been developed as an open interchange format for such mathematical
98 models²⁶ and the SBML site also has an extensive²⁶ list of existing tools and resources supporting pathway
99 modelling primarily by equation-based approaches
100 (http://sbml.org/SBML_Software_Guide/SBML_Software_Summary). Most mathematical models require
101 experimentally derived rate constants to feed into equations and significant computational power to solve
102 a series of equations. This generally limits equation-based approaches to modelling relatively small and
103 well characterised systems. Moreover, the level of mathematics skills required to construct and run these
104 models is often a deterrent to adoption by biologists. The platform described here uses Petri nets, as the
105 basis for pathway activity simulations. The primary resource for Petri net modelling is a network diagram

106 consisting of nodes, called 'places', and other nodes representing the interactions between them, called
107 'transitions', to which the user need only add 'tokens', that represent the amount or activity of a place prior
108 to performing a simulation (for more details of Petri nets see BOX 2). There is a long and established
109 precedent for the use of Petri nets in the modelling of biological pathways²⁷⁻³³ and several tools and
110 algorithms are available that allow the user to construct models based on Petri nets³⁴⁻³⁶. The Petri net
111 algorithm employed here was first described by Ruths *et al.*³⁷ who named their approach the signalling
112 Petri net simulator (SPN). It combines elements of a Boolean network simulator³⁸ with a synchronized
113 Petri net model³⁹, and models the stochastic flow of tokens through a network. A great advantage of Petri
114 nets is the relative ease of model parameterisation, the scale to which models can be constructed, the
115 computational speed of simulations, as well as the fact that the user does not need to directly modify the
116 maths when experimenting. The downside to most of the tools that currently support pathway modelling
117 using Petri nets is the inability to represent pathway models in anything but the standard Petri net notation
118 (open circles and black rectangles), and limited options for the visualization of results. Here, we describe
119 how a pathway model drawn according to the rules of mEPN, can then be parameterised for
120 computational modelling by the addition of tokens, whose quantity can be based on experimental results
121 such as quantitative transcriptomics or proteomics data. When a mEPN model is imported into the open-
122 source software BioLayout *Express*^{3D 40} it is visualised in a 3D environment, with nodes now represented
123 as 3D shapes. Simulations can subsequently be performed that calculate the flow of tokens through the
124 pathway over time. Pathway activity can then be visualised as plots or animations, where token
125 accumulation is represented by the size and colour of an entity node (Supplementary Video 1). Altering
126 the simulation parameters can change the flow of tokens, allowing the dynamics of pathways to be
127 modelled under different conditions. The modelling approach described in this protocol can be applied to
128 model any system, large or small, biological or otherwise, that consists of a series of components that
129 interact in a predefined manner. In the case of biological pathways, the mEPN notation scheme allows for
130 the detailed representation of signalling cascades, metabolic pathways, transcriptional networks, as well
131 as feedback/feedforward loops. To date, we have used this approach to model a wide variety of biological
132 pathways, particularly associated with immune signalling, e.g., Toll-like receptors (TLR), NF- κ B,
133 complement activation and antigen presentation, but also biochemical pathways e.g. cholesterol

134 metabolism, TCA cycle, and even pathways spanning multiple organ systems e.g. glucocorticoid,
135 oxytocin/prolactin signalling (see: www.virtuallyimmune.org and O'Hara *et al.*¹ for examples). Many of
136 these models were built as graphical representations of events as described in the literature and as such
137 act as a graphical bibliography, with pathway components or processes hyperlinked to research papers or
138 reports. However, with additional work they can also be used as the basis for performing simulation
139 experiments. These simulations not only test the logic of what is depicted, but also predict the behaviour
140 of the system and its response to perturbation. Using this approach models can be assembled at scale,
141 representing tens or thousands of components and the interactions between them. The overall aim of the
142 modelling approach described here is to, provide a platform for the assembly of information on a
143 particular system into an informative diagram, to allow the use of the diagram explore how the system
144 might operate, and through experimentation, make testable predictions^{41 42}.

145 The protocol provided here complements the paper published recently by O'Hara *et al.*¹, which describes
146 the development of and underlying concepts associated with this approach. This modelling platform may
147 not be appropriate in situations where many of the interactions between components are not known due
148 to the requirement to define both components and interactions, or where there is need to use specific rate
149 parameters to regulate the dynamics of a system, as the approach does not allow for the integration of
150 rate constants for specific reactions. Other limitations of Petri net-based approaches are the requirement
151 that all tokens and transitions behave the same way. In other words how a protein binding event is
152 modelled, is the same as how an enzymatically catalysed biochemical reaction would be modelled, where
153 outputs are determined by the number of tokens on the reactants. This is not likely to be an issue for
154 many applications, but could limit the approach's applicability in certain circumstances requiring a more
155 complicated concepts to be built within the model.

156

157 **Experiment Design**

158 First we describe how to construct a graphical model of a biological pathway using the mEPN scheme
159 (steps 1-8). We then explain how to convert this purely graphical representation into a resource that

160 supports computational modelling of the system (steps 9-11). Next, we present how to test the dynamic
161 properties of the model through running simulations and visualizing results (steps 12-20), and in the final
162 section (steps 21-25), describe how to optimise and validate the pathway model. The workflow is shown
163 schematically in Figure 1. To illustrate our approach, we use a model of interferon- β signalling. The model
164 is small and simple, but encompasses many of the basic concepts associated with pathway construction
165 and motifs such as a negative feedback loop, a common feature of many biological systems⁴³. However,
166 we encourage the examination of other larger models we have constructed, covering a range of biological
167 systems, in order to appreciate the scale and complexity models can achieve (examples can be found at
168 www.virtuallyimmune.com). Before embarking on model construction, users should search the literature
169 and pathway databases, such as those listed in Table 1, for existing diagrams of their system of interest.
170 Careful consideration should be given to the initial scope of the model, the level of detail to be
171 represented and what the model is to be used for once constructed. For instance, given the
172 interconnectivity of biological pathways, it is easy to begin with the aim of modelling one thing and end up
173 spending a lot of time modelling something entirely different, because it is one way or another related to
174 the first. Having said this, models will inevitably evolve as information is gathered and assimilated, and
175 the journey taken is part of the reward of modelling.

176

177 **Materials**

178 **Equipment**

179 A computer with Windows, Apple Mac or Linux operating system (preferably 64-bit), internet connection
180 and a web browser with JavaScript enabled. The hardware configuration may limit the size of models that
181 can be displayed within yEd, as well as when running pathway simulations within BioLayout *Express*^{3D},
182 where it will influence the speed of simulations and the frame rate for animations of flow. We
183 recommended >4Gb main RAM, a Dual-core CPU, NVidia GeForce / Quadro series or ATI equivalent

184 graphics card for advanced visualization with GLSL Shaders, preferably two monitors capable of
185 displaying at 1,600 x 1,200 resolution and a three-button mouse to aid navigation.

186

187 **Equipment setup**

188 **Installation of yEd Graph editor**

189 yEd is a free and intuitive software application that can be used to create high-quality network diagrams, it
190 runs on all major platforms: Windows, Unix/Linux and Mac OS X. Download and install the latest release
191 of the yEd Graph editor from the yWorks (Tübingen, Germany) website www.yworks.com. yEd will use up
192 approximately 215 MB of hard disk space. If you encounter any problems with the installation of yEd
193 contact: support@yworks.com

194 **Loading the mEPN palette**

195 Download the GraphML (.graphml) file containing the mEPN glyphs (Supplementary Data 1) and load it
196 into yEd by selecting Edit → Manage Palette → Import Section. This will provide the standard palette of
197 mEPN glyphs that can be selected as required when constructing a pathway model. To display the mEPN
198 symbols palette select from the menu bar Windows → Palette. Alternatively, create each node type
199 afresh by adding a node and changing its visual properties [F6]. As an example, see the list of
200 components present in the interferon- β pathway (Figure 2A).

201 **Installation of BioLayout *Express*^{3D}**

202 BioLayout *Express*^{3D} software allows the visualization and analysis of large network graphs in two and
203 three-dimensional space and supports the computational modelling of networks using the signalling Petri
204 net (SPN) algorithm³⁷. BioLayout *Express*^{3D} runs on Windows, Mac OS X and Linux platforms. Java SE 6
205 or 7 is required and can be downloaded from <http://www.java.com/getjava>. BioLayout will use up
206 approximately 41 MB of hard disk space. To download the BioLayout *Express*^{3D} installer, navigate to
207 <http://www.biobioinformatics.org/download/> and download an installer for Windows (.exe) or Mac OS X (.dmg).
208 For Linux platform use the universal JAR file that may be run without an installer. When BioLayout

209 *Express*^{3D} runs for the first time it creates a preferences file that can be changed and saved at any time
210 from the menu option Tools → Save Preferences. Users can customize many options selecting from the
211 menu bar Tools → General Properties (Shift+P). Further details on the software interface and its
212 customization are available in the BioLayout *Express*^{3D} manual that can be downloaded from the tools
213 support pages. If you encounter any problems with the installation of BioLayout *Express*^{3D} contact:
214 support@biolayout.org

215

216 **Procedure**

217 **Pathway model construction (timing: hours to months depending on complexity** 218 **of model)**

219 1. Source information for pathway construction. A pathway model should aim to provide a
220 comprehensive and reliable view of the current state of knowledge about the system. To
221 achieve this collect and extract the relevant information about the pathway from the literature,
222 databases and existing diagrams. Possible sources to consult are presented in Table 1. A
223 comprehensive list of databases for data mining can be found at www.pathguide.org. For
224 some pathways, data are available from multiple species and/or cellular systems, therefore
225 users must decide whether to piece together information from heterogeneous sources or to
226 restrict their model to reflect a particular species, cell type or developmental stage. To keep
227 track of the data, create a spreadsheet that includes: molecular identifiers, e.g. HUGO,
228 Entrez IDs, details about nature of the molecular interactions, sources of information, e.g.
229 PubMed ID, the quality of evidence (as assessed by number of publications supporting a
230 given interaction and the reliability of the assays used) and any additional information that
231 may be relevant.

232 **!Troubleshooting**

233

234

235 2. Identify the types of pathway information. Divide details of the pathway of interest into the
236 categories defined the mEPN notation scheme (see BOX 1). Find the 'real' name of pathway
237 components. The use of standard gene/protein names is essential in defining the exact
238 identity of components, especially if models are to be used in the interpretation of omics data
239 where the use of standardised nomenclature systems is standard practice (see BOX 3).
240 Record and characterise the type of interaction between components.
241

242 3. Commence drawing - addition of entity nodes. Molecular components are represented using
243 entity nodes. To add an entity node to the diagram, select and drag the appropriate glyph
244 from the mEPN palette (BOX 1). Edit a node's properties by selecting it and pressing [F6].
245 The node Properties dialogue will appear. Add the component's name to the General tab,
246 where necessary changing the size of the node to fit the label, record the reference source or
247 insert a brief description about a given component in the Data tab. Also add a hyperlink to an
248 external site (for example NCBI's Gene database), which can then be activated by selecting
249 the node and pressing [F8]. A description of the component, if available, will be shown in a
250 pop-up window when the mouse is placed over the node in yEd.
251

252 4. Draw the interactions between entity nodes. The nature of an interaction between
253 components may be represented using a combination of process nodes and edges. To add a
254 process node to the diagram, select and drag the appropriate glyph from the mEPN palette
255 (BOX 1). As with components (entity nodes) additional information may be added to the
256 process node by selecting it and opening the Properties dialogue [F6]. Pathway modules are
257 a special type of process node. They represent multi-reaction processes or events and are
258 represented using octagons with a label identifying the name of the process they represent.
259 They might be used to represent such pathway as signalling cascades, endocytosis,
260 compartment fusion, etc. Edges are lines that join entity and process nodes. Edges denote
261 the type of interaction (activation, catalysis, inhibition) and their directionality establishes
262 inputs and outputs from entity/process nodes. To add an edge, click on a node using left

263 mouse button and keep held down, then drag the mouse to move the edge to the target node
264 and release. If you release the mouse button on the way to the target node, a pivot point will
265 be introduced. To change the appearance of an edge (colour, thickness, arrow type or to add
266 text/hyperlink), select the edge by clicking on it and open the edge properties dialogue [F6].
267 The sample *Interferon_components.graphml* file can be used to try out the procedures
268 described in this step (Supplementary Data 2). **Note:** yED supports the import of data in
269 Excel or .CSV files in a variety of formats (see <http://yed.yworks.com/support/> for details).
270 This functionality may help initially in defining a set of pathway components and the
271 interactions between them, prior to manual editing of node/edge properties and layout.

272 **CRITICAL STEP.** In general (and absolutely so when constructing a diagram to be used for
273 computational modelling), nodes comprising a pathway should be arranged as a bipartite
274 graph i.e. entity nodes should be connected exclusively to a process nodes and vice versa.
275 This structure is the same structure used by Petri nets (places must be connected to
276 transitions) and it is essential if the model is to be used for simulation experiments.

277

278 5. Add compartments. Components should be represented as existing within a given cellular
279 compartment. Drag the desired compartment node from the mEPN palette and enlarge it to
280 cover the section of the pathway diagram which represents a given cellular compartment
281 such as the plasma membrane, cytoplasm or nucleus. Move the selected compartment
282 behind the diagram by choosing Edit → Lower selection. Name the compartment, placing the
283 name between asterisks (*compartment name*). The asterisks inform the BioLayout
284 *Express*^{3D} parser to treat these nodes differently: they are displayed as a translucent
285 background to the pathway and cannot be selected within this tool. If they are labelled as
286 follows *compartment*N* where *N* is a numerical value, e.g. 100, when viewed in BioLayout
287 the compartment becomes a 3D container where the N value determines its depth (Z-value).
288 The decision on a compartment's 'depth' is based on purely on final aesthetics as
289 compartments do not effect model dynamics. Generally the cell membrane would be the

290 largest component and its internal organelles are smaller compartments that sit within it, but
291 other than this the values given are subjective. Suggested cellular compartment colours are
292 defined in the mEPN palette. Generally, when compiling a model it is best to add
293 compartment nodes at the end or at least put them to one side when editing, as they tend to
294 get in the way. The completed pathway should look similar to the interferon- β pathway
295 example in Figure 2B.

296
297 6. **(Optional)** Add negative feedback loops. Feedback loops are network motifs common to
298 many systems and involve the activation of a pathway component that goes on to inhibit an
299 earlier step in the process. The inhibition of the interferon- β receptor by SOCS1 whose
300 transcription is activated by the interferon signalling pathway represents such a negative
301 feedback loop (Figure 2B). To represent an inhibitory activity such as this using the mEPN
302 scheme, place an inhibitor edge from the inhibitor molecule to a process node representing
303 the step that is inhibited.

304
305 7. Optimise model layout. It is essential that a pathway model is compact and easy to follow i.e.
306 be readable by a human. How this is best achieved will be influenced by a model's size and
307 complexity. First organise the pathway components based on where they reside within the
308 cell, i.e. their cellular compartment. Then attempt to separate out 'modules' based on
309 connectivity amongst a group of nodes, e.g. a particular signalling cascade or other series of
310 events. This helps with a model's readability and facilitates model expansion as new data
311 becomes available. Further information on layout optimisation can be found in BOX 4. In
312 practice, model optimisation is normally an iterative and time consuming process often
313 requiring a degree of trial and error in how best to layout the diagram. When in the dynamic
314 modelling phase it may for example be necessary to add in motifs that in effect delay the
315 passage of tokens from place to another in order, to model processes that are not explicitly
316 shown but may influence the order or timing of an event.

317

318 8. Save and export the pathway models. The pathway model should now represent a
319 diagrammatic version of known events and should be saved in the GraphML file format
320 choosing File → Save. Within yEd a model can also be exported as an image in PNG or JPG
321 format, as PDF or as HTML by selecting File → Export and selecting the preferred format.

322

323 **Model testing - conversion of a graphical model into a computation model**

324 **(timing: minutes to hours)**

325 9. Set the initial parameters. To convert a graphical representation of a pathway into a
326 computational model, you must define the initial state of the system through a process of
327 'parameterisation'. To parametrise the assembled model, add defined numbers of tokens to
328 entity nodes at the beginning of network i.e. nodes that have no 'parents' (upstream
329 connections). To define the initial state of a component, place an input node (depicted as a
330 black rectangle which functions as a transition node) upstream of the component to be
331 parameterised and connect it with a standard edge. Define the number of tokens to be added
332 to the node by selecting the edge between an input node and component. Open the edge
333 properties dialogue [F6] and type the desired number of tokens into the edge name (text)
334 (Figure 2C). Token values can in theory range from 0 to millions but in essence represent the
335 relative amount or activity of a given component under initial conditions. Ideally, the initial
336 parameters should be set with reference to some experimental data providing information on
337 the relative initial concentrations of the pathway components where known. However, in the
338 initial stages of pathway parameterisation and model testing, it is often sufficient to place an
339 arbitrary number of tokens on components, e.g. 1000, just to check the connectivity between
340 inputs and outputs is not compromised in any way.

341

342 10. Defining inhibitory reactions. An inhibitor edge originates from an inhibitory molecule and
343 terminates at the process to be inhibited, tokens present on the inhibitor node preventing
344 token flow through the process node. During a simulation tokens will not be lost through an

345 inhibitor edge and therefore tokens accumulate on an inhibitor node and irrevocably block the
346 process to which it is connected. However, if a sink node is connected to the inhibitor it
347 serves to give the inhibitory molecule a 'half-life' (in practice any process node will serve the
348 same purpose, but use of the sink node helps visually define the process involved). In the
349 absence of further input into the inhibitor node, such as during the 'off phase' of negative
350 feedback system, tokens will now be lost from the inhibitor. The result is that its inhibitory
351 effect will lessen and the blocked transition will eventually open and tokens may flow again
352 through it. In presence of a constant input this can cause token flow in negative feedback
353 systems to oscillate. There are two types of inhibitor edge included in the notation scheme
354 that perform differently in the modelling environment; the non-competitive inhibitor edge (red
355 with perpendicular bar at end) and a competitive inhibition edge (red with open diamond end).
356 The non-competitive inhibitor edge completely blocks token flow through the target
357 transmission if any tokens are present on the inhibitor node. In contrast the competitive
358 inhibitor edge works by deducting the number of tokens residing on the inhibitor away from
359 the number of tokens flowing through the target transition. The behaviour of negative
360 feedback systems is not only dependent on the type of inhibitor edge used but also the
361 distance between the input of tokens and the inhibitory step. The greater the distance the
362 more tokens are able to accumulate in the system and the greater the time taken between
363 the opening and closing of the inhibited transition, i.e., the longer the wavelength and the
364 higher the amplitude of the oscillating signal. Other factors that can affect the oscillatory
365 behaviour of the feedback loop are the number of inhibitors acting on the pathway and
366 assumptions about the stochasticity of token flow.

367
368 11. Save the parameterised model in the GraphML file format choosing File → Save. GraphML
369 files can be loaded directly into BioLayout *Express*^{3D}. A parser within the tool translates the
370 mEPN nodes into their 3D equivalent shapes such that they can now be visualised within the
371 tool's 3D environment. It also differentiates between nodes in the diagram that act as Petri
372 net 'places' and that are 'transitions', and reads the parameterisation markings that define

373 initial token inputs. BioLayout *Express*^{3D} is also able to perform stochastic flow simulations
374 using a modified version of the signalling Petri net algorithm³⁷. A file containing simple Petri
375 net models of all primary motifs found in pathways is provided as a means to better
376 understand the flow characteristics of this algorithm (Supplementary Data 3).

377

378 **Running simulations using BioLayout *Express*^{3D} (timing: 5-15 min)**

379 12. Load the saved GraphML file into BioLayout *Express*^{3D}. Supplementary file 4 is the
380 interferon- β pathway model shown in Figure 3C and as such is 'simulation ready'.
381 Following opening of the file answer yes to the dialogue window "This looks like a
382 Signalling Petri Net (SPN) pathway. Would you like to run a SPN simulation now?" (Figure
383 3A). This opens the SPN simulation dialogue (Figure 3B). The dialogue can also be
384 selected from the main menu under the Simulation menu or by pressing the "RUN SPN"
385 button on the sidebar.

386 **!Troubleshooting**

387

388 13. Set the SPN simulation options. Choose the number of time blocks and the number of runs.
389 A 'time block' is when all transitions are fired exactly once in a random order and tokens
390 moved as a result. A series of time blocks is referred to as a 'run', the more time blocks the
391 longer the run (Figure 3B1). The bigger the model or the more conditions you want to test
392 within a simulation, the more time blocks you will require for a simulation. It is good practice
393 to check the nodes furthest away from token input points to ensure that token accumulation
394 has plateaued or in the case of negative feedback circuits, that enough time blocks have
395 been run to evaluate the oscillatory behaviour of the system. The Petri net algorithm
396 employed here is stochastic in nature. That is to say that the number of tokens passed on,
397 when a transition is 'fired' is variable depending on the algorithms stochastic setting (see
398 below), and furthermore the order in which transitions are fired is random. Therefore, the
399 result of individual runs can be highly variable. For this reason a simulation is generally

400 comprised of multiple runs, where the outcomes from individual runs are averaged to
401 calculate the mean number of tokens present on a given node at each time block (Figure
402 3B2). The more runs used the less variable the results between simulations, but the more
403 time it will take to perform a simulation. To visualise the variation associated with a given
404 simulation check the 'Calculate Variance' and pick either standard deviation or standard
405 error (Figure 3B3).

406
407 14. Select the token stochasticity setting. The possible modes for token flow can be selected as
408 shown in (Figure 3B4) and are described below:

409
410 **Uniform Distribution:** Each time a transition is fired, an entirely random number of
411 tokens between zero and the maximum number of tokens are moved from an input place
412 to the output place (assuming there no other inputs on the transition which may influence
413 flow). This mode is as originally described by Ruths et al.³⁷ in their description of the
414 SPN algorithm.

415 **Standard Normal:** Each time a transition is fired, the number of tokens moved between
416 input and end places will be randomly chosen from a standard normal distribution around
417 50% of the number of tokens on the input place.

418 **Deterministic:** This moves exactly half of the tokens from input place to the output place
419 each time a transition is fired.

420
421 Normally we would use the standard normal distribution setting, as a halfway house between
422 the other two modes of token flow. In some settings, the average result of simulations is
423 similar with all these settings although variation between runs is greater with the more
424 stochastic token flow settings, especially the uniform mode. However, when simulating
425 feedback loops the mode of token flow can have a marked effect on the behaviour of such
426 systems. The mode you select may be based on which best models your system of interest.

427

428 15. Selection of SPN simulation transition type. Token movement between places is via transition
429 nodes which all operate using the same set of rules governing token flow. We have introduced
430 two options, consumptive transitions and original transitions, which differ in how they operate
431 with respect to token accumulation (Figure 3B5). Generally, we use the consumptive transition
432 mode as this prevents the accumulation of tokens on entities where there is a constant input of
433 tokens throughout the simulation but where flow through the target transition may be
434 intermittent.

435
436 **Consumptive Transitions:** Tokens are consumed from place nodes irrespective of
437 whether the transition is 'open' or not i.e. if there are two inputs into a transition and one
438 has tokens and the other does not, tokens will still be lost from the input place with tokens
439 as if flow were unrestricted.

440 **Original Transitions:** Tokens accumulate on input nodes where flow from them is
441 blocked i.e. if there are two inputs into a transition and one has tokens and the other does
442 not, tokens will not be lost from the input with tokens. This mode was as originally
443 described by Ruths et al.³⁷.

444
445 15. Run the simulation. Press the 'Run the Simulation' button to initiate the computation of the SPN
446 algorithm (Figure 3B6).

447 **Critical Step.** The time it takes to run a computation depends on the number of time
448 blocks/runs, the size of the pathway model and hardware on which the simulation is run.
449 However, for most small to medium size pathways (10's-100's of entity/process nodes) and
450 hardware configurations, the time taken is usually a few seconds or less for a typical modern
451 laptop.

452 **!Troubleshooting**

453

454 16. Save the results. Once the SPN simulation algorithm has finished, a Simulation Results
455 dialogue appears (Figure 3C). Token level per node per time block results can be saved as a
456 *.txt* or *.spn* file by ticking the “Save SPN Results” or pressing ALT+S (Figure 3C7). Saved
457 simulation results files can be loaded pressing ALT+L in the main window. *.spn* result files can
458 also be opened in programs such as Excel as a spreadsheet, or viewed and edited in a text
459 editor. An mEPN model can also be exported as a Systems Biology Graphical Notation
460 (SBGN)⁴⁴ diagram via File -> Export -> SBGN file. This can be opened in any SBGN compliant
461 software, e.g. VANTED with SBGN-ED add-on²¹.

462 17. Visualize token flow as node output graphs. To visualize the simulation results for a selected
463 entity/place node, close the SPN simulation results dialogue (Figure 3C8), position the cursor
464 over the node of interest and a pop-up window will appear showing the token flow associated
465 with that node (Figure 3D). To view and compare token flow in multiple nodes, select the
466 nodes of interest by pressing Shift+left mouse button and dragging the select window over the
467 nodes of interest or by pressing the Shift+ALT+left mouse button to select multiple nodes. The
468 corresponding flow graphs can be viewed using the Class Viewer by pressing CTRL+C
469 (Figure 3E) or the button with cog icon on left menu bar of the main window (Figure 3G).

470 A range of options are available within BioLayout to adjust graph appearance. Shift+> or
471 shift+< will increase or decrease node size, respectively; under the General tab you may turn
472 on or off the visualisation of the compartments (yEd Graphml Container Rendering); and by
473 pressing the ‘Render Plot to File button’ on the top of the Class viewer window the graph can
474 be saved as *.jpg* or *.png* image file (Figure 3E9).

475 18. Visualize token flow as an animation. Open the Simulation Animation Control window (ALT
476 + A) when the simulation has finished (Figure 3F). Use options provided within BioLayout
477 *Express*^{3D} tool to control simulation visualization:

478 **Node Animation** (Figure 3F10). Choose which nodes are animated (all, selected or
479 pathway components only), and the type of animated transition that takes place between
480 the node value associated with one time block and the next (discrete, linear, polynomial).

481 **Timing** (Figure 3F11). Define how many time blocks per second are displayed and
482 therefore the speed of the animation. If necessary also define which time block the
483 animation begins from.

484 **Size Transition** (Figure 3F12). Set maximum size of nodes during the visualization of token
485 flow, i.e. when the number of tokens is at its maximum. The 'Set (fixed) node value' is the
486 number of tokens on a node at which the maximum node size/colour is reached. The default
487 value for the 'Max value' is determined by the maximum number of tokens that accumulates
488 on any node during a simulation. It is often the case that some nodes accumulate tokens
489 much in excess of others, e.g. when their output is blocked. This can result in the majority of
490 nodes seemingly to change little in size or colour during a simulation. Click on this value and
491 add a value of your choice, and click on the associated check box, to maintain this value for
492 subsequent runs.

493 **Colour Palette Spectrum Transition** (Figure 3F13). A number of colour palettes are
494 available, or one can be loaded by user, to colour nodes so as to reflect their token value.
495 Select colour spectra from dropdown menu, load your own, or more normally, use default.

496

497 19. Visualize token flow as an animation. Select 'Start Animation' (Figure 3F14) to watch tokens
498 flow through your model (Figure 3G and Supplementary Video 1).

499 **!Troubleshooting**

500

501 **Model optimisation, parameterisation and validation (timing: days to months)**

502 20. Check for errors. Errors in a diagram's structure (predominately a failure to adhere to the strict
503 requirement for a bipartite graph or improper logic), can lead to bottlenecks in token flow.
504 When the bipartite graph structure is not maintained (e.g. an entity node is directly linked to
505 another), tokens will accumulate on the node upstream of the issue and tokens are not passed
506 downstream of the error. Check the reactions preceding any entity node whose token output is
507 zero (Figure 4A) and correct mistakes. Place and transition spacer nodes are available to
508 position between two nodes of the same type where the graphical description of events leads
509 to this situation. Another common issue encountered is where the presence a pathway
510 component is under the control of the system in which it operates. This can lead to a situation
511 whereby for component to be synthesised it needs the pathway to be active, but the pathway
512 is not active because it requires the activity of that component. In these circumstances it may
513 be necessary to 'prime' the system adding tokens to the component in question prior to
514 beginning the simulation. Mistakes and errors in logic are easy to make, but equally easy to
515 spot and rectify with this approach. It is normal practice to run a simulation, find out where the
516 issues are, edit the model in yEd and rerun the simulation. There will likely be a need to repeat
517 this process a number of times.

518

519 21. End of the line. Without a downstream transition, a component at the end of a line of flow will
520 simply accumulate tokens (Figure 4A). A final transition node is required to dissipate tokens
521 from such entities. One option is to place a 'pathway node' at these points by dragging and
522 dropping from the palette to allow indication of what happens next without showing it in detail.
523 Alternatively, a 'sink' node can be placed as described above to signify that a component is
524 removed from the system, e.g., the destruction of a protein by proteosomal degradation.

525

526 23. Amplify or reduce token flow at specific sites. To simulate the amplification or reduction of a
527 signal at specific sites in the network, add a numeric value to a transition-to-place edge. Select

528 the edge, press [F6] and write a number in the Text field of the edge Properties dialogue.
529 When a transition fires, the number of tokens produced on the downstream entity will
530 correspond to the number of input tokens multiplied by the weight of the output edge, e.g., an
531 edge weight of 2 will result in a doubling in the number of input tokens, where as an edge
532 weight of 0.5 will halve the number of tokens going forward. For example, one can amplify
533 tokens as means to model the production of numerous protein molecules from a single
534 transcript during protein translation (Figure 4C).

535

536 24. Varying token input during a simulation. To simulate variation in the level of an input signal at
537 different time blocks of a simulation, assign tokens to an input edge (as described in Step 9)
538 using the following notation : a-b,c;d-e,f where 'a-b' defines the first and last time blocks that
539 the number of tokens 'c' will be added to the model and 'd-e' are the first and last time blocks
540 that you would like the number of tokens 'f' to be added to the model. For example: 0-5,0;6-
541 15,100,16-20,0 translates into, add no tokens between time blocks 1-5, 100 tokens between
542 time blocks 6-15, and then remove token input until the end of the run, time block 20. Any
543 number of these statements may be added to an input. This allows modelling of a system
544 before and after a stimulus, or when a stimulus is transient or delayed.

545

546 25. Validate the model by comparing it to experimental data. Once a model has been constructed,
547 checked for structural errors and parameterised according to known variables, the first
548 question is whether the model recapitulates the known activity of the system. For example,
549 check if genes are expressed as transcriptomics data suggests, or does the flow of metabolic
550 pathways under different conditions reflect what is known? The simulation of pathway
551 dynamics should recapitulate the known activity of the system. If not, the obvious conclusion is
552 the model is wrong. This could be because it is poorly constructed or parameterised, in which
553 case the model needs improving. More interestingly, it could reflect the fact that there is part of

554 the system that is as yet undiscovered. Once a model is working, i.e., it verifies the known
555 characteristics of the pathway, it can be used to test known perturbations of the system e.g.
556 the effect of knocking down/out a gene or inhibiting an enzyme. With confidence in a model's
557 characteristics it is then reasonable to use it to predict the effect of perturbing it, proving
558 results that can be tested experimentally: one of the ultimate aims of dynamic modelling.

559

560

561 **Troubleshooting**

562 Troubleshooting information can be found in Table 2

563

564 **Table 2: Troubleshooting table.**

Step	Problem	Possible reason	Solution
1	Information about a part of the pathway is unavailable.	The experiments to elucidate the process have not been done.	Use a 'pathway module' node to indicate that a process occurs but the details of which are undefined.
12	BioLayout <i>Express</i> ^{3D} fails to run.	Incompatible hardware or software configuration.	Contact support@biolayout.org . Improve hardware specification.
15	'Error with Vertex weight!' error popup is shown during simulation.	Token input to node(s) is not readable by software.	Check that token input is numerical (token input can be any positive number, including decimals) at all token input nodes.
19	Flow stops and downstream nodes do not accumulate tokens.	Bipartite graph structure has not been adhered to.	Identify bottleneck node by watching BioLayout flow animation. Return to yEd graph to edit model and rerun.
	Node accumulates tokens at linear rate.	Node has no output.	Return to yEd graph to add sink node or other transition to node accumulating tokens.
	Token flow occurs, but the size of nodes changes little.	The maximum token value is set too high.	Open Animation Control window and lower value in 'Set Max Value' dialogue box.

565

566

567 **Timing**

568 Steps 1 to 8, Information mining and pathway construction: Hours to months

569 Steps 9 to 11, Conversion of the graphical model into a computable format: Minutes to hours

570 Steps 12 to 19, Visualization of pathway and running simulations using BioLayout *Express*^{3D}: 5 to 15
571 minutes

572 Steps 20 to 25, Model optimisation and parameterisation: Days to months

573

574 **Anticipated Results**

575 This protocol describes the generation of pathway models using the mEPN language for graphically
576 representing biological systems. Graphical models can be used both as a resource to store and display
577 what is known about a pathway and can be considered an end point in their own right, that can be
578 updated or extended as new information becomes available. In addition, they can be converted to a
579 computational model by setting parameters to define the initial state of the pathway. Using the sample
580 interferon- β components GraphML file (Supplementary Data 2) a pathway model can be produced that
581 represents events from the binding of interferon- β to its receptor and the signalling pathway leading the
582 activation of target genes. The resulting model can be parameterised by adding tokens to obtain a
583 simulation-ready model (Supplementary Video 1). This is a relatively small diagram; we also provide a
584 model of the hedgehog signalling pathway as an example of a larger model (Supplementary Data 6).
585 This was produced as part of a 10 week elective course by an undergraduate student with no previous
586 modelling experience, indicating our modelling scheme can easily be performed by biologists with no
587 prior modelling knowledge. Other pathway models are available at: www.virtuallyimmune.com.

588 The BioLayout *Express*^{3D} software is fully compatible with the mEPN notation and can be used for
589 pathway visualization and to perform stochastic flow simulations using a modified version of SPN

590 algorithm³⁷. Running simulations provides insights into the dynamic behaviour of the system by
591 enabling users to visualize the signal flow within the network. The signal flow is simulated by the
592 accumulation of tokens at entity nodes (places) and can be visualized in 2D graphs (as seen in Figure
593 4) or by 3D animation (as seen in Supplementary Video 1). The inflation and contraction of the entity
594 nodes represents the accumulation and degradation of reactants in the pathway. In this way, complex
595 biological processes with multiple components can be modelled.

596

597 **The interferon- β signalling network - a feedback control system**

598 Biological systems display a variety of dynamic behaviours ranging from stable steady states to
599 oscillations. Oscillations in protein concentrations or gene expression levels are commonly associated
600 with the presence of negative feedback loop(s) in the regulatory network⁴⁵. Based on the analyses
601 performed using this system many factors can affect the amplitude, frequency and stability of oscillations.
602 For instance, the time-delay (path length) between token input and inhibitor, the type of inhibition edge
603 used (competitive or non-competitive), the number of inhibitors present and their half-life, may all affect
604 how a model containing a negative feedback loop will operate in practice.

605 As an example, the simple model of the interferon- β signalling pathway is presented. Interferon- β is a
606 cytokine released by immune cells in response to pathogens. It acts as an autocrine and paracrine
607 signalling system that triggers the activation of host defence systems. In the provided model it operates
608 as a damped oscillator⁴⁶, where low-dose IFN stimulation yields oscillations of lesser amplitude that are
609 damped faster than those induced at a high-dose (Figure 5). To reproduce these observations users can
610 look at the different versions of IFN signalling feedback model provided in Supplementary Data 5 or
611 modify the token input or topology of the Petri net examples provided in Supplementary Data 3.

612

613 **Contributions of the authors:** A.L., L.O'H., M.E.P developed and wrote the protocol based on their
614 experience in using the approach for modelling their own pathway systems of interest, T.A. developed a
615 number of the features within BioLayout *Express*^{3D} including SBGN export and refinement of the Petri net

616 algorithm, D.W. developed and has helped maintain the VirtuallyImmune.org website that is associated
617 with this work, and L.B.S. helped with writing and editing the manuscript. T.C.F. has lead the
618 development of the mEPN notation scheme and its use in modelling a variety of pathway systems;
619 oversaw the implementation of model import into BioLayout *Express*^{3D}; model visualisation within this tool,
620 refactoring and refinement of the Petri net algorithm, and conceived of and assisted in writing the paper.
621

622 **Figure legends**

623 **Figure 1. Workflow with steps described in the Procedure.**

624 **Figure 2. Construction of a simple pathway model describing type-1 interferon signalling. (A)**

625 Components of the interferon- β signalling pathway drawn using mEPN notation (BOX 1). The information
626 necessary to construct the interferon- β pathway has been highlighted in the pathway description: entity
627 and transition nodes, edges and cellular compartments. **(B)** interferon- β pathway diagram assembled in
628 yEd software using the pathway parts shown in A. **(C)** Parameterisation of interferon- β pathway by the
629 addition of token inputs and a sink node output on SOCS1 inhibitor node (highlighted by red rectangles).
630 Also shown is the edge properties dialogue in yEd where tokens can be added to an input edge. Model
631 adapted from O'Hara *et al.*, 2016¹.

632 **Figure 3. Visualization of token flow. (A)** When a model is loaded into BioLayout *Express*^{3D} it is

633 displayed in a 3D environment using 3D equivalents to the 2D node glyphs as rendered in yEd. The
634 software automatically recognizes a diagram as having been parameterised for computational modelling
635 (based on the presence of process nodes as defined in the notation system) and prompts users to run a
636 SPN simulation. **(B)** In the SPN Simulation dialogue users can set constraints on how to run the SPN
637 simulation algorithm. (1) Defines the number of time blocks in a simulation; (2) Defines the number of
638 runs in a simulation; (3) Calculates the variance in token flow between runs; (4) Defines the nature of the
639 stochastic flow of tokens; (5) Defines the rules governing token flow through transitions; (6) Run the
640 simulation. **(C)** SPN Simulation Results dialogue box summarises the simulation. (7) SPN results may be
641 saved before the user chooses to (8) run the simulation again, close the dialogue box or proceed to

642 animate the simulation. **(D)** After a simulation has been run token accumulation at specific nodes can be
643 visualised by placing the cursor over the node. **(E)** The flow of tokens across one or a number of selected
644 nodes can be plotted using the Class Viewer showing plots of token flow over the time course of the
645 experiment for selected nodes. The name and class of selected nodes is also displayed, and below are a
646 range of options available for node selection and data export. (9) The token plot can be saved as a .png
647 or .jpg image file. **(F)** Animation Control dialogue. (10) Options for the type of nodes to be animated and
648 interpolation of flow between time blocks; (11) Speed of animation (time blocks per second); (12) Node
649 size at maximum token number; (13) Colour palette selection to highlight change in token number; (14)
650 Animation control: start, pause, step, stop. **(G)** BioLayout *Express*^{3D} can produce animations of token flow
651 through the model across time blocks. Node size will increase/decrease depending on the number of
652 tokens passing through them and the colour of the node will also change according to a predefined
653 spectrum of colours.

654 **Figure 4. Influencing token flow along a linear pathway.** Figure shows the flow of tokens through a
655 small linear pathway motif depicting a gene being transcribed into mRNA, which is then translated into the
656 encoded protein. Addition of a sink node represents the protein's degradation. **(A)** Blocked flow. Top of
657 the three illustrations all is as described above, and over the 20 time blocks of the simulation there is an
658 initial rise in the level of the protein followed by a steady-state, as the number of tokens entering the entity
659 node matches those leaving through the sink i.e. the rate of production of the protein matches the rate of
660 its degradation (blue line). Failure to maintain bipartite graph structure (shown here by connecting two
661 entity nodes for mRNA and protein without a process node between them) causes tokens to accumulate
662 on the first entity node (mRNA) and not pass to the second (protein), which stays at zero tokens
663 throughout the simulation (pink line). In the absence of a sink node, tokens to accumulate on the protein
664 node (green line). **(B)** Modelling time. The diagrams show the effect of increasing the length of linear
665 networks on the rate of token accumulation. Input tokens are introduced into each of the networks at the
666 same time but the protein accumulates at different rates. The delay is proportional to the number of
667 transition steps introduced in the network and such delay motifs may be added where a transition nodes
668 represents a multistep process such as transcription or translation. **(C)** Amplifying or depleting signal.
669 The addition of a value to the edge leaving a transition can be used increase (pink line) or decrease

670 (green line) downstream token flow. In this way one might model one mRNA molecule leading to
671 production of multiple protein molecules, or the inefficient translation of mRNA where only a single protein
672 molecule is produced from multiple mRNAs. Simulation options in A, B and C: 100 tokens, 100 runs, 20
673 time blocks, Normal Standard distribution and with standard deviation of token flow between runs shown.

674 **Figure 5: Effect of parameterisation on activity of feedback loop. (A)** A simple pathway model
675 representing the type-1 interferon signalling pathway constructed and parameterized in yED using mEPN.
676 Graphs show token accumulation on the activated transcription complex ISGF3 (circled in red) following
677 simulations with **(B)** 1000 input tokens (blue line) or 100 input tokens (green line) added to interferon- β
678 with SOCS1 acting as a non-competitive inhibitor or **(C)** as a competitive inhibitor of the activated receptor
679 (dashed red edges). **(D)** Oscillatory activity of the SOCS1 feedback loop with (magenta line) or without
680 (blue line) a delay introduced between the transcription and translation of SOCS1 (dashed black edges),
681 again with SOCS1 acting as a non-competitive or **(E)** competitive inhibitor of the activated receptor.
682 Simulation options: 100 runs, 500 time blocks, Normal Standard distribution and with standard deviation
683 of token flow between runs shown.

684

685

686 **Supplementary Files**

687 **Supplementary Data 1.** mEPN palette for loading within yEd software. This is a graphml file containing
688 all the different types of entity nodes to represent the different classes of molecules that might play a part
689 in a pathway, as well as the process nodes that represent different types of interactions. This file can be
690 loaded into yEd to provide a palette of mEPN nodes for pathway construction (see step 4 of the
691 procedure).

692 **Supplementary Data 2.** Interferon- β signalling pathway components. This is a graphml file containing all
693 the different parts of the simple model shown in figure 2B, to allow practicing model construction. Try
694 assembling the model using only the text below: '*Interferon B (IFNB1) is a cytokine released from many
695 cell types in response to immune stimulation. It homodimerises and binds to its cell surface receptor
696 complex composed of the receptor proteins IFNAR1 and IFNAR2 and the intracellular kinases TYK2 and
697 JAK1. The complex is composed of 2 of each of these proteins. Binding causes a conformation change
698 in the complex resulting in the autophosphorylation of JAK1. Once activated, the complex catalyses the
699 phosphorylation of STAT2 which forms a heterodimer with STAT1. This complex then binds interferon
700 regulatory factor 9 (IRF9), forming the complex often referred to as ISGF3, and translocates to the
701 nucleus. Here it binds to the IRF sequence in the promotor of a number of genes including MX1, MX2,
702 IFIT1, IFITM3, TAP1, OAS1, GBP1, PSMB9, SOCS1. In turn SOCS1 inhibits the autophosphorylation of
703 the receptor thereby inhibiting further activation.*'

704 **Supplementary Data 3.** Primary network motifs drawn in Petri net style for testing SPN algorithm. This is
705 a graphml file containing a series of different network motifs and parameterisations potentially found in
706 pathway diagrams. This includes linear networks, multiple inputs/outputs to transitions and nodes, and a
707 series of models representing a range of feedback loops with varying path lengths between token input
708 and inhibition, inhibition type (competitive vs. non-competitive), and one or multiple feedback inhibitors.
709 The file is designed to allow you to explore the different interaction types and algorithm settings when
710 setting up a simulation run. Certain nodes are coloured such that when a simulation has been run within
711 BioLayout, these nodes may be selected and you can compare results across motifs.

712 **Supplementary Data 4.** Interferon- β signalling pathway. This is a graphml file of the simple model
713 shown in figure 2B.

714 **Supplementary Data 5.** Changing parameters - Interferon- β signalling pathway. This is a graphml file
715 containing six versions of the model shown figure 2B (Supplementary Data 4), each version is
716 parameterised slightly differently, with variation in: type of inhibition (competitive vs. non-competitive);
717 introduction of a delay between SOCs1 expression and protein; and an amplification of signal between
718 gene and mRNA.

719 **Supplementary Data 6.** Example of a more complex model - Hedgehog signalling pathway. This model
720 (given here as graphml file) is a representation of Hedgehog signalling from the binding at its receptor,
721 activation of the GLI protein on the tip of the primary cilium through the activation of various downstream
722 pathways (not shown in detail). It contains 550 nodes and 601 edges and was assembled using pathway
723 resources such as Reactome²² and the primary literature. Its parameterisation, in terms of token
724 placement is arbitrary.

725

726 **Supplementary Video 1.** Movie of the interferon- β signalling pathway (Supplementary Data 4) simulation
727 run within BioLayout. The movie shows the process of model loading, running the simulation, inspecting
728 token accumulation on specific components and watching the flow of tokens run through the model as an
729 animation.

730

731

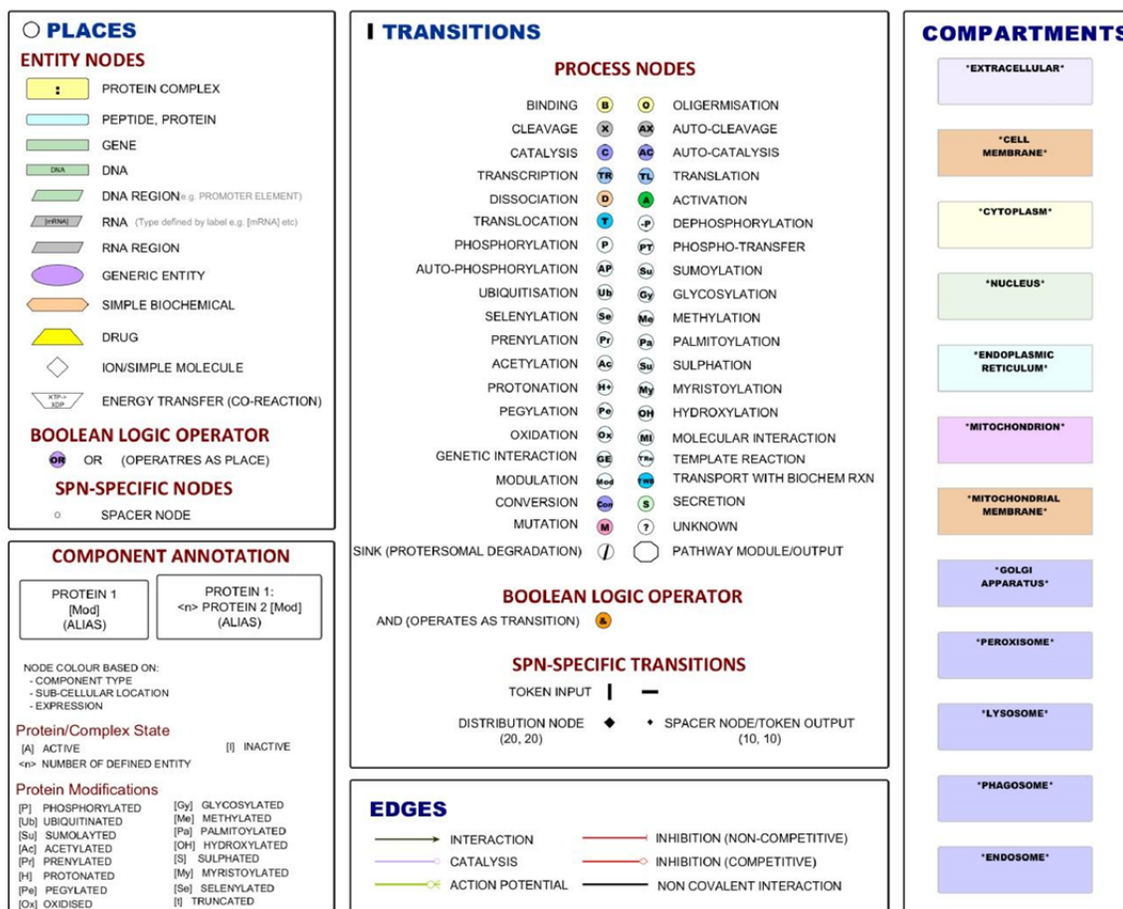
732

733

734

735 **BOX 1: mEPN Notation 2017**

736 The modified Edinburgh Pathway Notation (mEPN) scheme³ is a graphical notation system based on the
 737 concepts of the process diagram⁴, below the glyph library is shown in its current form (reproduced from
 738 O'Hara *et al.* ¹).



739

740

741 **Pathway components**

742 **Types of pathway information.** The information depicted in a pathway diagram drawn using the mEPN
 743 scheme may be divided into the following categories:

- 744
- *Entity*: any component involved in a pathway, e.g. protein, protein complex, nucleic acid
745 sequence (promoter, gene, RNA), simple biochemical, drug etc., and depicted as an 'entity node'.
746 Different shaped nodes are used to represent different types of components. The mEPN scheme
747 consists of twelve different entity nodes (detailed in the top left panel). Also included are a
748 Boolean logic 'OR' operator node and spacer node (represented as a white circle with a black
749 border) that may be required to maintain bipartite pathway arrangement. Entity nodes function as
750 the equivalent of Petri net 'places' and all entity nodes are equivalent in Petri net simulations
 - *Process*: an interaction that occurs between pathway components, where one component
751 interacts with or influences the state of another through its binding, inhibition, catalytic conversion,
752 etc., is depicted as a process node. Processes are generally depicted as a circular node with a 1-
753 3 letter code to indicate the type of process, e.g., P = phosphorylation, B = binds, X =cleavage
754 etc. Included in the scheme are 38 different process nodes (top central panel). Additional to
755 these, but also acting as transitions, are a sink node which is placed at the end of pathway and
756 represents removal of a component from the system; a pathway module node that summarises
757 not one process but a series of events; a Boolean logic 'AND' operator node; token input nodes
758 that are placed at the start of a pathway and oriented either horizontally or vertically to fit in with
759 the pathway layout; a spacer node represented as a black diamond. A larger version of this node
760 can also be used as a distribution node when multiple edges exit from an entity node. Process
761 nodes function as the equivalent of Petri net 'transitions' and all process nodes are equivalent in
762 Petri net simulations.
 - *Interaction*: a directional edge that links an entity node to a process node or vice versa that
764 indicates direction and the nature of the interaction. There are six possible connecting edges
765 (bottom central panel) that represent the nature of the interaction between nodes. The first three
766 (interaction, catalysis and action potential) operate identically within a Petri net and serve to carry
767 tokens between entity and process nodes. The two inhibitor edges act to inhibit the flow of tokens
768 through target process nodes albeit based upon different rules. The non-competitive inhibitor
769 edge completely blocks token flow through the target transmission if any tokens are present on
770 the inhibitor node. In contrast the competitive inhibitor edge works by deducting the number of
771

772 tokens residing on the inhibitor away from the number of tokens flowing through the target
773 transition. Finally, the non-covalent interaction edge can be used to depict two separate entities
774 within a complex. This may be a useful when describing large complexes, but these edges do not
775 operate within the context of a Petri net simulation.

776 • *Cellular compartment*: define where pathway components reside and interactions take place,
777 such as an organelle (e.g. mitochondrion, nucleus) or a transient cellular compartment (e.g.
778 vesicle). In the diagrams they are shown as large coloured nodes that sit behind the interaction
779 model (right panel).

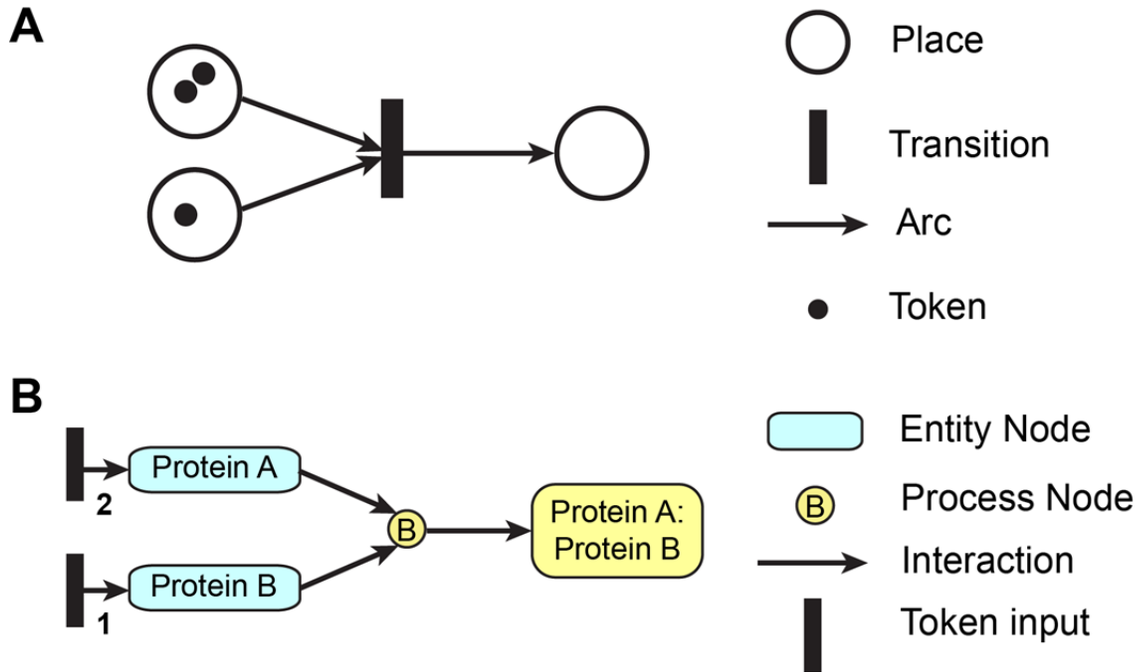
780 End of Box 1

781

782

783 **BOX 2: Petri nets to model biological systems**

784 Petri nets (panel A) are a mathematical approach for describing distributed systems and have been used
785 extensively in the modelling of many different kinds of systems including biological pathways. There are
786 numerous types of Petri net algorithms and software that support modelling using them. The Petri net
787 algorithm employed here was first developed and described by Ruths *et al.*³⁷, a modification of a
788 synchronized Petri net model and firing policy, they called the signalling Petri net (SPN). Petri nets share
789 a number of common features. Models are constructed as directional bipartite networks where nodes are
790 considered to be either 'places' or 'transitions' connected by 'arcs'. Places usually represent an entity or
791 state and by convention are represented as a white circle. Transitions represent interactions between
792 entities or the transition of an entity from one state to another and are usually represented as a black
793 rectangle. Arcs are directional arrows that connect places to transitions and *vice versa*. The availability of
794 an entity and its abundance can be represented by the initial placement of tokens. The flow of information
795 through the network is represented using tokens that move between places through transitions, following
796 in the direction of the arcs. In the context of biological pathways places represent pathway components,
797 transitions correspond to processes that modify the components in some way, such as phosphorylation,
798 binding etc., and are referred to as 'process nodes'. The interactions between molecules are depicted by
799 edges, equivalent to arcs in Petri net parlance. Panel B shows how the notation for representing
800 pathways using mEPN map on to Petri nets.



801

802 **Rules determining token flow through transitions.** Activity flow is represented by the movement of

803 tokens between places. The state of each place (component) is determined by the number of tokens held

804 by it. When a transition is fired, tokens are moved from each input place and redistributed downstream,

805 the transition acting as rule-based controller of flow. A transition will pass on tokens only if all the input

806 places contain tokens and where one input has less tokens than others, the passage of tokens will be

807 governed by the input place holding the least number of tokens. In the case of the Petri net algorithm

808 employed here, the movement of tokens is also stochastic. This is because during a time block all

809 transitions in a model are fired once but in a random order, and the number of tokens taken forward when

810 a transition is fired will be a random number between zero and the maximum available (although we have

811 implemented other versions of this rule, see step 13). Due to the stochastic nature of token flow, a

812 simulation usually comprises of a number of runs, the answer being based on the average token flow

813 across runs. Furthermore, when a transition is fired the number of tokens moved forward through the

814 transition will be subtracted from the amount available on the input places. One innovation not found in

815 most other Petri net simulators, is our implementation of a consumptive transition mode. When running in

816 this mode (for us this is standard), a constant input of tokens is applied to an entity node throughout a

817 simulation, but token levels on the node remain constant (unless others are fed in from another source),

818 as tokens are lost from it at the same rate they are added even when there is no flow through the target
819 transition. In this way places representing entities such as enzymes (or indeed any other molecule) can
820 receive a constant input of tokens throughout a simulation without accumulating or losing tokens.

821 **Modelling time.** Time in Petri nets is measured in abstract units called time blocks. When constructing a
822 model, it is useful to consider how many experimental seconds, minutes, or hours correspond to a time
823 block. Timing depends on the network topology and the further away a node is from the start of flow the
824 longer it will take tokens to reach it. To simulate such time delays users can create a linear network that
825 alternates transitions with spacer nodes multiple times. When tokens are passed through such a linear
826 network the number of output tokens corresponds to the input but the time taken for tokens to reach the
827 end is proportional to the number of spacer nodes (Figure 4B).

828 End of Box 2

829

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832

833 **BOX 3: Component Annotation**

834 Multiple names are frequently employed to describe molecular species. This is particularly the case for
835 one of the main components of biological pathways: proteins. Any given protein may be referred to in the
836 literature by a number of different names concurrently. The use of non-standard nomenclature frequently
837 leads to confusion in written texts and diagrams. If the naming of pathway components is not clear, then
838 uncertainty arises as to what exactly is being depicted in a diagram and it ends up representing little more
839 than a series of abstract concepts.

840 Our models have generally been focused on human pathways and we have used standard Human Gene
841 Nomenclature Committee (HGNC) names to label nodes representing genes and proteins
842 (www.genenames.org). This and related nomenclature systems such as the Mouse Genome Database
843 (MGD) standard (<http://www.informatics.jax.org/mgihome/nomen/>), now provide a near complete
844 annotation of all human and mouse genes, and their use in the naming of proteins provides a direct link
845 between the identity of the gene and the corresponding protein. Of course, not everyone has adopted
846 these naming systems so where other names (aliases) are in common use, these names are often
847 included as part of the node's label after the official gene symbol in rounded brackets, but generally only
848 on its first appearance in the pathway. Use of standard nomenclature also assists in the comparison and
849 overlay of experimental data (which is usually annotated using standard gene nomenclature) onto
850 pathway models. At the present time there are no standard and universally recognized nomenclature
851 systems available for naming certain types of pathway components. For instance, protein isoforms tend to
852 be named in an *ad hoc* manner by those who study them, and biochemical compounds are known by
853 both their common names or by names that reflect their chemical composition. The IUPAC
854 (www.iupac.org/) provides a standard nomenclature system for organic chemicals, but most names would
855 have little relevance to a biologist. In cases such as these, the important thing is to be consistent and,
856 where possible, to cross-reference the component's ID to other sources such that the identity of the
857 component depicted, where at all possible, is unambiguous. We have used the excellent ChemSpider
858 resource (www.chemspider.com/) as a reference for the naming of biochemical entities, although other
859 resources, e.g., ChEMBL (www.ebi.ac.uk/chembl/) are potentially equally good. Using the node

860 properties dialogue [F6], nodes may be hyperlinked to external web resources and additional notes to
861 nodes can be added using in the Data tab within yEd.

862

863 **Protein state:** The particular 'state' of an individual protein may determine its functional activity. With
864 mEPN, a component's state is indicated as a text addition to the node label using square brackets
865 following the component's name; each modification being placed in separate brackets. The system can
866 be used to describe a wide range of protein modifications like phosphorylation [P], acetylation [Ac],
867 ubiquitination [Ub] etc., and where details of the site of modification are known this may be represented,
868 e.g., [P@L232] = phosphorylation at leucine 232.

869

870 **Protein complexes:** Names of the components are given as a concatenation of the proteins belonging to
871 the complex, separated by a colon. If a complex is commonly referred to by a generic name this may be
872 shown below the constituent parts in rounded brackets. Where a specific protein is present multiple times
873 within a complex, this may be represented by placing the number of times the protein is present within the
874 complex in angle brackets i.e., <n>. A node representing a component may be coloured to impart visual
875 information on the component's type, e.g. to differentiate between a protein and a complex. Similarly,
876 other types of pathway components may be represented using a range of shapes and colours – see
877 palette (BOX 1, downloadable as Supplementary Data 1) for list of glyphs used to represent different
878 entity types. A component may only be shown once in any given cellular compartment (in a given state).
879 A component may however alter from one state to another, e.g., inactive to active, unbound to bound, in
880 which case both forms are represented as separate entities. A different state may be indicated by
881 including the name in square brackets, as described above.

882

883 End of Box 3

884

885

886 **BOX 4: Layout Optimisation**

887 There is a part of pathway modelling that could be considered art, or at least creative cartography. When
888 starting a diagram the number of components is small, and visual comprehension of the system of nodes
889 and edges is relatively easy. However, this situation soon changes as a diagram grows, and one of the
890 greatest challenges is to render the inherently complex connections between components of a network
891 model in a human readable form. This necessitates the careful placement of nodes and edges in the
892 network layout. There are large number of layout algorithms available for network visualization but
893 unfortunately none come close to the results achievable by a skilled human curator. Certain rules can be
894 applied to this process to aid readability of the model:

- 895 • Models should be constructed, where possible, along a horizontal or vertical axis, arranged top down
896 or left to right in the direction of information flow.
- 897 • Nodes should be evenly spaced and aligned along the chosen axis of layout but within the cellular
898 compartment in which they reside.
- 899 • Crossing over of edges should be kept to a minimum and changes in edges direction should be
900 avoided when possible. When multiple edges run parallel to each other it is important to keep the
901 lines straight to maintain an easy-to-follow diagram.
- 902 • Space can be organised effectively by structuring sub-pathways into modules.
- 903 • Modules of the pathway should be arranged so that the connected glyphs are in close proximity, to
904 minimise overlapping of connective edges.
- 905 • Hierarchical relationships between components should be shown in the layout of interactions. To do
906 this, an orientation of pathway flow is chosen (e.g. left to right or top to bottom) and should be
907 maintained throughout the diagram where possible.

908 However, each diagram is essentially unique and each comes with its own challenges. There is no one
909 solution that fits all models so the layout of the diagram must be able to be easily adapted to take in new
910 components and concepts whilst maintaining its readability.

911

912 End of Box 4

914 **BOX 5: Glossary**

- 915 • **Arc:** By convention in Petri net parlance, arcs are the directional edges that connect a place to a
916 transition or vice versa, (but never between places or between transitions). Arcs are referred to as
917 **edges** in mEPN.
- 918 • **Edge:** In mEPN notation, edge is used to refer to any line used to connect entity and process
919 nodes to indicate an interaction (activating or inhibitory), and also the direction of that interaction.
920 The Petri net equivalent is an **arc**.
- 921 • **Entity node:** Entity nodes are glyphs that represent pathway components such as molecules or
922 genes. The Petri net equivalent is a **place**.
- 923 • **Glyph:** a visual representation of an entity, process or transition node.
- 924 • **GraphML:** a XML-based file format used to describe the structural and visual properties of a
925 model.
- 926 • **Layout:** the way in which nodes and edges are set out in 2D or 3D space (physical topology).
- 927 • **Model:** the visual representation of a network.
- 928 • **mEPN:** modified Edinburgh Pathway Notation scheme is a graphical notation system based on
929 the concepts of the process diagram.
- 930 • **Network:** a number of nodes connected by edges.
- 931 • **Notation scheme:** a system of symbols used to represent biological entities and interactions
932 between them in a semantically and visually unambiguous manner.
- 933 • **Parameterization:** defining the initial state of the system through the placement of tokens.
- 934 • **Petri net:** a directed bipartite graph that alternates places (entities) and transitions (events that
935 occur).
- 936 • **Place:** In Petri nets, places represent possible states of the system. They are referred to as
937 **entity nodes** in mEPN notation.
- 938 • **Process diagram:** a diagram used to formally describe the components of a system, their
939 activation state and the interactions between them.

- 940
- **Process node:** In mEPN notation, process nodes are glyphs that represent and define interactions between entities or the transition of an entity from one state to another. The Petri net equivalent is a **transition**.
- 941
- 942
- 943
- **SPN:** Signalling Petri Net as defined by Ruths *et al*³⁷. Please note, in the context of Petri nets SPN is also often used to refer to Stochastic Petri Nets, a class of Petri net algorithms used to model the stochastic flow of tokens, as in the case here.
- 944
- 945
- 946
- **Tokens:** Tokens represent quantitative information that is introduced and distributed through a Petri net. Here they can be thought of representing the amount and/or activity of a pathway component.
- 947
- 948
- 949
- **Topology:** arrangement of various elements of the pathway (nodes, edges, etc.) that illustrates how information flows within the network.
- 950
- 951
- **Transition:** In Petri nets, transitions are events or actions. They are represented as **process nodes** in mEPN notation.
- 952

953

954 End of Box 5

955

956

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964

965 **Conflict of Interest**

966 **The authors declare competing financial interests: details are available in the online version of the**
967 **paper.** There is now a commercial and supported version of BioLayout *Express*^{3D} called Miru, produced
968 by Kajeka Ltd, (Edinburgh, UK) that possesses all the functionality described here for pathway modelling.
969 T.C.F. is a founder and director of Kajeka.

970

971

972

973 **Table 1: A list of resources useful for the compilation of pathway diagrams**

Literature Databases	
NCBI Pubmed	http://www.ncbi.nlm.nih.gov/pubmed/
Web of Science	http://wok.mimas.ac.uk/
Google Scholar	http://scholar.google.co.uk/
Scopus	http://www.scopus.com/
iHop	http://www.ihop-net.org/
Component Annotation	
NCBI Entrez Gene	http://www.ncbi.nlm.nih.gov/sites/entrez
Gene Cards	http://www.genecards.org/
Gene Ontology	http://www.geneontology.org/GO_downloads_annotations.shtml
PubChem	http://pubchem.ncbi.nlm.nih.gov/
Chemspider	http://www.chemspider.com
Interaction Databases	
ConsensusPathDB	http://cpdb.molgen.mpg.de/
BioGRID	http://thebiogrid.org/
IntAct	http://www.ebi.ac.uk/intact/
GeneMANIA	http://www.genemania.org/
Human Protein Reference Database	http://www.hprd.org/index.html
MINT	http://mint.bio.uniroma2.it/mint/Welcome.do
STRING	http://string-db.org/
DIP	http://dip.doe-mbi.ucla.edu/dip/Main.cgi
MIPS CORUM	http://mips.helmholtz-muenchen.de/genre/proj/corum

Pathway Repositories	
KEGG	http://www.genome.jp/kegg/
Reactome	http://www.reactome.org/
Biocarta	http://www.biocarta.com/genes/index.asp
WikiPathways	http://wikipathways.org/index.php/WikiPathways

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