Logical modelling of *Drosophila* signalling pathways†

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A limited number of signalling pathways are involved in the specification of cell fate during the development of all animals. Several of these pathways were originally identified in *Drosophila*. To clarify their roles, and possible cross-talk, we have built a logical model for the nine key signalling pathways recurrently used in metazoan development. In each case, we considered the associated ligands, receptors, signal transducers, modulators, and transcription factors reported in the literature. Implemented using the logical modelling software *GINsim*, the resulting models qualitatively recapitulate the main characteristics of each pathway, in wild type as well as in various mutant situations (e.g. loss-of-function or gain-of-function). These models constitute pluggable modules that can be used to assemble comprehensive models of complex developmental processes. Moreover, these models of *Drosophila* pathways could serve as scaffolds for more complicated models of orthologous mammalian pathways. Comprehensive model annotations and *GINsim* files are provided for each of the nine considered pathways.

Introduction

Deciphering the regulatory mechanisms enabling the development of sophisticated multicellular animals and plants from a single cell (a fertilised egg or seed) constitutes one of the most intriguing and challenging tasks in biology. In this respect, the use of *Drosophila* as a model system has been particularly fruitful in molecular genetic and more recent genomic studies. This has revealed a blueprint for animal development, leading to the identification of crucial regulatory components, from homeobox transcriptional factors to signalling pathways, which turn out to be highly conserved throughout metazoans.1–3

Such studies have progressively led to the delineation of large intertwined regulatory networks underlying developmental processes, whose properties are difficult to grasp intuitively due to the inherent pleiotropy, cross-talk and redundancy within these systems. To deal with this complexity, diverse mathematical approaches are increasingly used to study developmental networks, ranging from statistical or probabilistic methods to mine large data sets, to dynamical modelling frameworks to integrate relevant components into full fledged mechanistic and predictive models.

Early *Drosophila* development has been the focus of a large number of dynamical modelling studies, often using differential equations (see e.g. ref. 4–10 and references therein). However, these studies have typically considered relatively limited numbers of regulatory components and required a quantitative determination of poorly documented parameters. In this context, formal qualitative modelling approaches constitute an interesting alternative, at least as a first step towards more quantitative modelling. In particular, logical modelling (Boolean or multilevel) has been applied to various regulatory networks of increasing sizes (see e.g. ref. 11–26 and references therein). Logical models have been proposed for several *Drosophila* developmental processes, focusing on early stages of embryo segmentation,27–31 on boundary and compartment formation in imaginal discs,32,33 and also on the specification of the mesoderm during stages 8 to 10 of *Drosophila* development (Mbojd et al., in prep.). Because signalling pathways are known to play important roles in these developmental processes (see e.g. ref. 34–38 and references therein), these models contain several signalling components (e.g. Wg, Hh, Notch, and a limited number of downstream components) belonging to different signalling pathways.

Noticeably, although *Drosophila* has been a prominent model system to decipher cell signalling, in particular in the context of development, *bona fide* dynamical models for these...
pathways are still largely lacking. Furthermore, information about pathway organisation and function is spread across many publications and various databases (for links pointing to Drosophila signalling pathways in public knowledge repositories, see Table 1). In particular, Flybase and Interactive Fly provide rich textual descriptions of each pathway. However, computational biologists must perform substantial analyses to convert this vast array of information into a format that can be used to derive formal graphical and dynamical models. There is thus a clear need for reference models for each of the recurrently used signalling pathways in Drosophila, which would undoubtedly constitute a valuable resource for the modelling of developmental networks.

To address this issue, we have developed reference logical models for the major Drosophila developmental pathways, with extensive references as hyperlinks to relevant articles and database entries. These pathway models can be used as building blocks to assemble more comprehensive models, as we are currently doing for mesoderm specification during Drosophila embryogenesis (Mbodj et al., in prep.). As they represent a manually curated collection of the current knowledge on each signalling cascade, they can also be used for teaching purposes.

Hereafter, we first introduce the logical framework used to define the qualitative dynamical models of Drosophila signalling pathways (see Methods), which are then briefly described in the following section (Results). As comprehensive descriptions of each of the nine presented models would be very lengthy, these are provided in ESI†. The last section (Conclusions and prospects) is devoted to the evocation of further developments and potential exploitations of this model collection.

### Methods

Logical modelling is adapted to the qualitative nature of genetic data. It can offer a global view on the dynamics of large systems. The logical modelling approach has been introduced in genetics by Stuart Kauffman39,40 and René Thomas.41,42 Here, we used an extension of Thomas’ original Boolean approach, which enables the consideration of multi-level components, e.g. to account for differential effects of different signal levels on their targets.11,42–44

In short, regulatory networks and their dynamics can be represented using two kinds of graphs called logical regulatory graphs (model structure) and state transition graphs (dynamical behaviour).

In a logical regulatory graph, nodes (or vertices) usually denote regulatory components or their targets. They can represent different types of biomolecules (genes, mRNA, proteins, complexes, etc.) or functional processes (e.g. entry in a specific cell cycle phase). A logical function (or a logical rule) is associated with each regulatory component. By default, a node can have two alternative levels: active (1) or inactive (0). However, more levels can be considered when justified by available data. When all entities take their values within the [0, 1] interval, the model is said to be Boolean.

Signed arcs connecting graph components denote regulatory interactions. Positive or negative signs denote activatory or inhibitory effects on their targets, respectively. Multiple arcs between a pair of nodes can be used to account for dual regulatory interactions, or for differential qualitative effects of increasing regulator concentrations. When a source node may take more than two levels, a threshold (1 per default for the Boolean case) must be further associated with each outgoing arc.
Arcs may represent different types of influences, from transcriptional regulation (e.g. the activation or the repression of the transcription of a particular gene), to biochemical reactions (de/phosphorylation or degradation of a protein, etc.), to subtler situations (e.g. modulation of the activity of another component, co-factors participating in a complex, regulation through implicit components, etc.).

The regulatory graph allows a first glance at a logical model, but a logical model fundamentally consists of a set of logical rules associated with the components, thereby enabling qualitative simulations. Logical rules combine positive and negative influences using NOT, AND and OR operators and define the target value of a component depending on the values of its regulators.11,15,16

The dynamical behaviour of a logical model is represented in terms of state transition graphs, where each node corresponds to a state of the system, i.e. a vector giving the levels of all the components of the regulatory graph (integers between 0 and the corresponding maximal levels). Arcs connecting states represent transitions enabled by the logical rules. More precisely, starting from a given initial state (or a set of initial states), the successor states are computed by comparing the values of the different components for the current state with those returned by the logical rules. Whenever the current value of a component differs from that returned by its logical rule, there is an updating call on this component. The computation of the next state(s) then depends on the updating mode. In biology, only unitary changes (+1 or −1) are...
usually considered, while two main updating modes are generally used. According to the \textit{synchronous mode},\textsuperscript{40} all updating calls are considered simultaneously. Network behaviour is then deterministic since each state can have at most one successor state (although several paths may lead to the same state). In the case of the \textit{asynchronous mode},\textsuperscript{11} updating calls are considered individually. A successor state is then created for each updating call.

Various solutions have been proposed to refine logical simulations and avoid the complexity of fully asynchronous schemes, for example by distinguishing between fast and slow subnetworks simulated sequentially,\textsuperscript{30} or by grouping transitions into ranked (a) synchronous classes.\textsuperscript{15}

A simulation can lead to a \textit{stable state}, i.e. a state without successor. From a biological point of view, a stable state may correspond to a specific cell type. Alternatively, a simulation can lead to a (simple or complex) \textit{cyclic attractor}, which may denote periodic behaviour, as in the case of the cell cycle.\textsuperscript{16}

Whatever the updating mode, a logical model keeps the same stable states, but cyclic attractors may differ. Interestingly, powerful algorithms have been recently proposed to compute stable states in large regulatory networks.\textsuperscript{45,46}

To ease the simulation of large networks, we have developed a flexible model reduction method, which allows hiding selected components of the regulatory graph. Reduction is then applied in an iterative way to hide multiple components. When a component is masked, the effects of its regulators are transferred to its targets, for which new logical functions are then computed. By avoiding the masking of auto-regulated components, most

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig1.png}
\caption{Boolean model for the Drosophila Wingless (Wg/Wnt) pathway. (A) Wg pathway is displayed from regulatory (Wf, Dly and Daily) and ligand (Wg) components on the left to the main downstream effectors (Dsh, Arm, Pan) and a generic “Targets” node, along with inhibitory and activatory partners on the right. Red blunt and green normal arrows denote activatory and inhibitory interactions, respectively. The blue dot arrow denotes the fact that Pangolin (Pan) can activate or inhibit different sets of target genes. (B) In the absence of Wg binding to its receptors, Arm is phosphorylated by the Axin–Sgg–APC complex and degraded by the proteasome; Pan then forms a complex with Gro, which inhibits target genes. (C) Binding of Wg to Arrow or Frizzled triggers the accumulation of Arm, which forms a complex with Pan and other cofactors (Lgs, Nej, Hx and Pygo) for transcription activation. Grey nodes and arcs denote the lack of activity or influence depending on Wg pathway activity. Note that both inhibitory and activatory complexes are modelled implicitly by considering interactions from their components onto their targets and using proper logical rules (i.e. using the AND operator).}
\end{figure}
essential dynamical properties, including the stable states, are preserved upon model reduction.\textsuperscript{47}

The software \textit{GINsim} (for Gene Interaction Network simulation) implements the logical formalism and various analysis tools, including stable state computation and model reduction.\textsuperscript{48} \textit{GINsim} further supports the annotation of regulatory graphs with free text and URLs, thereby enabling extensive documentation of the considered regulatory elements and interactions. Once a model is defined, the user can launch simulations, which are presented in the form of state transition graphs. \textit{GINsim} also supports the annotation of regulatory graphs with free text and URLs, thereby enabling extensive documentation of the considered regulatory elements and interactions. Once a model is defined, the user can launch simulations, which are presented in the form of state transition graphs. \textit{GINsim} also supports the annotation of regulatory graphs with free text and URLs, thereby enabling extensive documentation of the considered regulatory elements and interactions. Once a model is defined, the user can launch simulations, which are presented in the form of state transition graphs. 

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**Fig. 2** Logical models of \textit{Drosophila} Hh, Notch, JAK/STAT and Dpp signalling pathways. Ellipses and rectangles denote Boolean and multilevel nodes, respectively. Red blunt and green normal arrows denote activatory and inhibitory interactions, respectively, while the blue dot arrows denote dual effects of the source nodes on their targets.
allows blocking the expression levels of specific components to simulate different types of perturbations (e.g. loss-of-function, gain-of-function, and combinations thereof).

All signalling pathway models described hereafter have been implemented and annotated using GINsim. Extensive documentation is provided in ESI† for each of the nine pathway models. Furthermore, we are providing these models in a dedicated repository, in the form of an archive file that can be opened using GINsim (release 2.4 or higher), which can be downloaded from the GINsim web site (http://ginsim.org).

Results

Table 2 lists the nine main Drosophila signalling pathways covered by this study, namely Wg, Hh, Notch, JAK/STAT, Dpp, EGF, FGF, VEGF, and Toll pathways, mentioning the main ligands, receptors, and effectors, along with reference developmental processes. Based on an extensive literature analysis, and predominantly focusing on the roles of signalling pathways during Drosophila embryonic development, we have developed a logical model for each of these pathways. Hereafter, for sake of space, a general presentation of these models is given, complemented by a more detailed presentation provided in ESI†.

The number of components considered in the model is mentioned in the last column of Table 2, along with some key model properties. The regulatory graphs corresponding to the nine considered pathways are displayed in Fig. 1–4, using similar conventions. In all regulatory graphs, ellipses denote Boolean components (i.e. components with two possible values, 0 or 1, standing for absent/inactive or present/active, respectively), while rectangular contours denote multi-level components (here all ternary, i.e. taking three possible values, 0, 1 and 2, corresponding to low/insignificant, middle and high activity levels, respectively); furthermore, components with yellow background denote molecules (e.g. signals) acting from the exterior of the cell considered.

Logical rules have been defined for the components of these models, so that they qualitatively recapitulate the effect of the reported perturbations (effects of loss-of-function or gain-of-function of pathway components).

Note that in all cases, only single signalling components are explicitly modelled, while associations in complexes are implicitly modelled by arcs outgoing from complex partners onto their targets, along with adequate logical rules (using the AND operator).

When one of these models is used to simulate the behaviour of the pathway for various input configurations (potentially in the presence of genetic perturbations such as loss-of-function...
or gain-of-function), the levels/activities of pathway components are encoded into a discrete vector or a logical state, while temporal changes are represented by state transitions (cf. Methods). Such simulations can be reproduced using the GINsim files provided in ESI† Hereafter, we focus on a limited number of representative results.

Fig. 1A displays the logical regulatory graph for the Wingless (Wg) pathway, which acts recurrently during Drosophila development.49,50 In the absence of Wingless, the protein complex composed of Axin (AXN), Shaggy (SGG or ZW3) and APC sequesters Armadillo (ARM), enabling its ubiquitination and a Slimb-dependent degradation by the proteasome. This absence of ARM facilitates the formation of a Pangolin (PAN)–Groucho (GRO) inhibitory complex, leading to the inhibition of Wg target genes. This corresponds to the situation shown in Fig. 1B (an inactive Wg pathway), where inactive (or irrelevant) regulatory components and interactions are shown in grey.

In the presence of its co-receptor Frizzled (FRZ), binding of Wingless (the ligand) to its receptor Arrow (ARR) triggers a set of reactions, starting with the activation of Dishevelled, which in turn inhibits the AXN–SGG–APC complex;51 this leads (with the help of HIPK) to the stabilization and accumulation of ARM. Next, ARM translocates into the nucleus, where it can form an activatory complex with PAN and other co-factors (LGS, NEJ, PYGO and HYX), leading to the activation of WG target genes. This situation (an activated Wg pathway) is illustrated in Fig. 1C.
Fig. 2 shows the logical models for the Hedgehog (Hh), Notch, JAK/STAT, and Decapentaplegic (Dpp) pathways, which are also recurrently used during *Drosophila* development.

The Hh pathway (Fig. 2A) consists of a regulatory cascade leading to the activation of the pathway effectors encoded by the gene *cubitus interruptus* (*ci*). CI can be released in a full-length CI[act] form, which functions as a transcriptional activator, or in a truncated CI[rep] form, which functions as a repressor. The processing of CI is regulated by several kinases (cf. ref. 56).

The Hh and Wg pathways both play key roles in the anteroposterior segmentation of the *Drosophila* embryo (many of their signalling components are encoded by the so called “segment polarity” genes). Consequently, they have been considered in several models for *Drosophila* embryo segmentation, although in very simplified ways.6,29–31

In this context, the more comprehensive pathway models presented here may serve as references to extend already published models, e.g. to recapitulate a broader collection of experimental data, or yet to better ground the simplifications made (e.g. applying model reduction tools to extended pathway modules to obtain consistent simplified models).

The Notch pathway (Fig. 2B) is required during tissue development, cell fate determination, cell proliferation (lateral inhibition), cell differentiation, apoptosis, wing patterning and development.

Here, we particularly refer to its role in the specification of the mesoderm, where it contributes to modulate the level of Twist (TWI).35

The JAK/STAT pathway (Fig. 2C) is involved in various developmental processes, including eye development, cell growth, sex determination, hematopoiesis, spermatogenesis, oogenesis, and signal transduction in larva and adult.34,52–60 Our model considers three different ligands (OS, UPD2 and UPD3), several inhibitors, as well as the differential action of the two forms of STAT92E, the normal form and the truncated form (STAT92E).

Turning to the Dpp pathway (Fig. 2D), our model recapitulates the establishment of the morphogen gradient, by the implicit modelling (using proper logical rules) of different homodimers of DPP itself, Screw (SCW) and Glass-bottom-boat (GBB), which are involved in relatively low signalling, as well as of heterodimers (DPP/SCW, DPP/GBB, SCW/GBB), which are responsible for high signalling.

Dpp and Notch pathways both play important roles in the delineation of boundaries and compartments in *Drosophila* imaginal discs, which prefigure the organisation of adult structures such as the wings, legs or eyes. Consequently, these pathways have been implemented, albeit in very simplified ways, in models of these differentiation processes (see e.g. ref. 32, 61 and 62).

### Table 3

Sample results from simulations of EGF and FGF pathway models. For each initial or final state, only active components are listed (omitted components are thus absent or inactive). In the case of multi-level (ternary) components, high levels (value 2) are specified whenever it is used under initial conditions or reached at final states; when not otherwise specified, the listed components are set to the value 1. In all simulations, all internal nodes were set to zero at the initial states. Significant changes in activity levels between initial conditions or between initial and final states are emphasised in bold.

#### EGF signalling pathway

<table>
<thead>
<tr>
<th>Initial state</th>
<th>Final (stable) state</th>
</tr>
</thead>
<tbody>
<tr>
<td>No ligand</td>
<td>Spi, Cnk, Src42, Ksr, Msk</td>
</tr>
<tr>
<td>Medium Spi</td>
<td>Spi, (2), Cnk, Src42, Ksr, Msk</td>
</tr>
<tr>
<td>High Spi</td>
<td>Spi (2), Cnk, Src42, Ksr, Msk</td>
</tr>
<tr>
<td>High Spi in the presence of Sprouty (Sty)</td>
<td>Spi (2), Cnk, Src42, Ksr, Msk</td>
</tr>
<tr>
<td><em>cnk</em> loss-of-function</td>
<td>Spi (2), Cnk, Src42, Ksr, Msk</td>
</tr>
<tr>
<td><em>aop</em> gain-of-function in the presence of high Spi</td>
<td></td>
</tr>
</tbody>
</table>

#### FGF signalling pathway

<table>
<thead>
<tr>
<th>Initial state</th>
<th>Final (stable) state</th>
</tr>
</thead>
<tbody>
<tr>
<td>No ligand</td>
<td>Tbs, Pyr, Cnk, Src42, Ksr, Msk</td>
</tr>
<tr>
<td>Htl activation</td>
<td>Bnl, Cnk, Src42, Ksr, Msk</td>
</tr>
<tr>
<td>Bnl activation</td>
<td>Tbs, Pyr, Bnl, Cnk, Src42, Ksr, Msk</td>
</tr>
<tr>
<td>Inhibition by Sprouty (Sty)</td>
<td></td>
</tr>
<tr>
<td>in the presence of all ligands</td>
<td>Tbs, Pyr, Cnk, Src42, Ksr, Msk</td>
</tr>
<tr>
<td><em>Htl</em> loss-of-function in the presence of Pyr and <em>Tbs</em></td>
<td><em>Ras</em> gain-of-function combined with <em>stumps</em> loss-of-function in the presence of <em>Tbs</em> and <em>Pyr</em></td>
</tr>
</tbody>
</table>
Fig. 3 shows the logical models derived for the EGF, FGF and VEGF signalling pathways. For sake of simplicity, the processing of the EGF ligand Spitz (SPI) is not modelled. Noticeably, only two ligands have been selected when several redundant ligands have been characterised (e.g. Gurken, Keren). Various inhibitors known to function in the establishment of a signalling gradient between expressing cells and receiving cells are explicitly considered (Sprouty, Argos, Kekkon, etc.).

From these graphical representations, one can easily see that the three pathways share an extensive core cascade (MAPK cascade) encompassing eight components, including homologs of the notorious vertebrate oncogenic factors RAS and RAF (components coloured with light green). This emphasises the fact that the same signalling cascade can respond to different signals, with contextual regulatory variations, to finally affect target genes through common effectors (i.e. the transcriptional factors AOP and PNT). Examples of simulation results for EGF and FGF pathway models are given in Table 3.

Finally, Fig. 4A shows our current logical model for the Drosophila Toll signalling pathway, which is involved in early dorso-ventral embryo differentiation. The Toll pathway also plays a key role in innate immunity. Indeed, it can be activated upon infection by various kinds of pathogens, a process involving complex signals converging onto the production of Spaetzle (SPZ), the main ligand of Toll in Drosophila (Fig. 4B). At this stage, however, our logical model does not yet account for potential differential effects of SPZ depending on immunological challenges.

All nine pathway models are provided in ESI† (ZGINML files, which can be opened using the public software GINsim), along with extensive documentation (included in the model files, but also provided in html and pdf formats). To ease simulations, each model file includes sample parametrisations for typical signalling situations.

Conclusions and prospects

We have presented a collection of logical models for the prominent signalling pathways involved in Drosophila development. Each of these models can be considered as a building block to assemble more complex models for specific developmental processes. In their present form, these logical modules can be used to extend recent models for pattern formation in Drosophila embryo or in imaginal discs.

Overall, although the nine pathway models reported here will require refinements as new data are gathered, or depending on specific applications, they already constitute a comprehensive knowledge repository that should facilitate the delineation of sophisticated models by combining inter-cellular signalling and other regulatory modules involved in Drosophila development. In this respect, simplified versions of seven of these signalling modules have been integrated in a novel model for mesoderm specification (Fig. 5), which will be reported in detail elsewhere (Mbondj et al., in prep.).

Several studies indicate that the proteins implicated in these pathways are shared between species. Our collection of pathway modules could thus be used to perform comparative analyses or to build models of orthologous pathways in other model organisms.

By and large, our modelling work mainly relied on genetic data (gene expression patterns and mutant phenotypes). As can be seen in Table 2 and Fig. 1–4, we modelled most signalling
components by Boolean variables. This is certainly an oversimplification, which could be refined as novel data become available, thanks to the recent technological advances (transcriptomics, proteomics, interactomics, or high-throughput functional assays such as RNAi screens).

Furthermore, the mutant developmental phenotypes used to identify pathway components may only reflect extreme situations (loss-of-function or gain-of-function). This presumably led to overlook many potential unknown pathway actors and regulators, which can now be identified using high-throughput methods.76

The pace of acquisition, the quantity and the nature of the resulting data pose novel challenges to biologists. Indeed, efficient exploitation of the resulting datasets and their integration with relevant lower-throughput results is far from trivial. To do so, several groups set to progressively complete and extend canonical pathways with novel interactions and components identified by large-scale interaction screens.77,78 Although still in its infancy, there is no doubt that integration of large-scale datasets along with more detailed molecular studies will open novel avenues for signalling pathway mapping and modelling. In this respect, the collection of signalling pathways proposed here may be considered as seeds to build more comprehensive maps and ultimately delineate more quantitative and predictive models.

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