

Information processing by biochemical networks: a dynamic approach

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Understanding how information is encoded and transferred by biochemical networks is of fundamental importance in cellular and systems biology. This requires analysis of the relationships between the stochastic trajectories of the constituent molecular (or submolecular) species that comprise the network. We describe how to identify conditional independences between the trajectories or time courses of groups of species. These are robust network properties that provide important insight into how information is processed. An entire network can then be decomposed exactly into modules on informational grounds. In the context of signalling networks with multiple inputs, the approach identifies the routes and species involved in sequential information processing between input and output modules. An algorithm is developed which allows automated identification of decompositions for large networks and visualization using a tree that encodes the conditional independences. Only stoichiometric information is used and neither simulations nor knowledge of rate parameters are required. A bespoke version of the algorithm for signalling networks identifies the routes of sequential encoding between inputs and outputs, visualized as paths in the tree. Application to the toll-like receptor signalling network reveals that inputs can be informative in ways unanticipated by steady-state analyses, that the information processing structure is not well described as a bow tie, and that encoding for the interferon response is unusually sparse compared with other outputs of this innate immune system.

Keywords: biochemical reaction networks; stochastic kinetics; information; conditional independence; modularization; signalling

1. INTRODUCTION

Imagine a record over time of the trajectory of the number of molecules of each type of biochemical species present in a reaction network, together with a record of the time-varying, external signals received by the cell. The network may be large, is not assumed to be in a ‘steady’ or stationary state, and is subject to intrinsic stochasticity. Current understanding in this context of how to analyse the relationships between the various trajectories is very limited. We describe how to identify conditional independence relationships between the trajectories of groups of species. These are robust network properties that provide important insight into how information is encoded and transferred by cellular networks—an area whose importance has been widely recognized [1,2].

Both the external environment and internal state of the cell are inherently dynamic. Signalling and regulatory networks must detect and coordinate responses to both external and internal fluctuations using stochastic biochemistry. The multiple signals arriving at the cell surface are expected to exhibit complicated dynamics and are themselves noisy carriers of information.

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Advances in understanding information processing within the cell therefore require both a dynamic (see [3]) and a probabilistic approach. Hence the focus here is on the probabilistic relationships between species and module trajectories. There is clear evidence that key components of the cellular signal transduction machinery exhibit involved dynamic behaviour and that different temporal patterns are associated with differential regulation of downstream targets. Spiking and oscillations are exhibited by, for example, cytosolic calcium, cyclic AMP, nuclear NF- κ B, and nuclear ERK [4]. Frequency/amplitude modulation has been established for the control of both gene expression and mitochondrial redox state by calcium [5], and for the nuclear translocation of cAMP-dependent protein kinase [6]. Recently, Ashall *et al.* [7] report that altering the frequency of stimulation by tumour necrosis factor TNF α also results in different patterns of nuclear translocation of NF- κ B and of NF- κ B-dependent gene expression.

Two related types of question arise from these observations. First, the inferential one concerning what may be learnt from the trajectories of outputs and intermediaries about the dynamics of the input signals and hence about the external state of the cell. This connects with recent work [8,9] that interprets biochemical networks as implementing inference in order to provide a basis for cellular decisions. Second, the question of

how external information is encoded by and transferred within biochemical networks exposed to multiple inputs.

The need for a rigorous, dynamic, information-based framework for relating the stochastic trajectories of biochemical networks and for quantifying their information content is thus clear. There has been little previous work in this area (although note [10]). We describe how to identify conditional independences, written $A_t \leftrightarrow_{D_t} B_t$, between the trajectories (up to some time t) of groups of biochemical species A , B and D (see §2.2). Such an independence means that the information contained in or encoded by the trajectory D_t makes any information in A_t irrelevant for B_t (and vice versa). That is, A_t and B_t contain no mutual information given the trajectory D_t . In this sense, the species in D play the role of informational intermediaries or ‘information carriers’ between A and B within the network. Further discussion of the importance of such conditional independences in the context of signalling networks is given in §2.1. Although probabilistic and information theories have been used to quantify the mutual information content between inputs and outputs for single neurons [11], a small signalling network [12] and simple regulatory transcriptional mechanisms [13,14], the approach adopted here is different. We consider general, potentially large biochemical networks and the sequential information transfer via intermediaries that is central to signal transduction mechanisms. A stochastic process treatment of (bio)-chemical networks that focuses on mathematical proof of dynamic independence properties is outside the scope of this paper and is given in Bowsher [15].

In developing our approach to a quantitative theory of information processing by biochemical networks, it has proved fruitful to adopt a modular perspective. This is unsurprising as such networks are increasingly thought of as having a modular structure [16–19], a separable architecture that can be decomposed into units that perform to some extent independently of one another. Furthermore, it is recognized both that understanding such modularity requires a dynamic approach [20] and that rigorous definition and identification of dynamic modularizations is a difficult problem ([21], ch. 3). We show how to exactly decompose or modularize an entire network using informational properties. We view a modularization here as a collection of groups of biochemical species in which, for each module, all information relevant to the trajectory of the module is encoded by the trajectory of the module’s overlap (interface) with the rest of the network. With this definition, it turns out that valid conclusions may be drawn by studying the kinetics of each module in isolation from the rest of the system.

As would be expected of a functionally important property of a naturally evolved network, our modularizations and analysis of information processing are robust to (unaffected by) changes in rate parameters. The MIDIA algorithm is developed (§2.3) which allows automated identification of modularizations for large networks, control over the degree of coarse graining, and visualization using a tree that encodes the conditional independences. Only stoichiometric

information is used and neither simulations nor knowledge of rate parameters are required. A bespoke version of the algorithm for signalling networks identifies routes of sequential encoding between inputs and outputs, visualized as paths in the tree.

Previous dynamic approaches to module identification adopt various perspectives. Network motifs [22] are an important concept, but the manner in which individual motif function depends on the context within and connections to the rest of the network remains an open question. Graphical, community detection-based methods [18,19] are extended by Saez-Rodriguez *et al.* [23] to kinetic models of signalling networks—edges are interpreted as retroactivities and ‘inter-modular retroactivity’ is minimized in order to partition species between modules. Some bi-directional, local dynamic influences between modules remain after decomposition. Correlated reaction sets (or ‘Co-sets’; [24]) arise where interest is in a region of the solution space of steady-state fluxes (or reaction rates)—e.g. a Monte Carlo sample from that space [25,26] or the collection of all ‘extreme pathways’ [27]. A Co-set is a group of reactions that have non-zero flux at all points in the solution region under consideration and that thus function together when the system is at steady state—i.e. when species concentrations and reaction rates are constant over time.

Using their stoichiometric reconstruction of the toll-like receptor (TLR) signalling network, Li *et al.* [28] employ a steady-state flux balance analysis (FBA) to identify signalling pathways for different input–output (I–O) pairs. The objective flux maximized is the flux of the particular reaction regarded as the output of signalling in each case. Although a virtue of this and the other constraint-based methods is that they require knowledge only of the stoichiometric matrix, the steady-state assumption is a strong one that considerably limits their generality. In many settings, signalling and gene regulatory networks are inherently dynamic and non-stationary, as has already been noted. Furthermore, none of the module identification methods discussed allows for stochasticity, which is both an important characteristic of gene expression and signalling dynamics (e.g. [29–33]) and essential for a consideration of how the cell processes noisy information.

2. RESULTS

2.1. Informational encoding, transfer and decomposition—a preview

The approach of the paper relies on two particular properties of the probabilistic relationships between species trajectories in biochemical networks. By the trajectory of a species we mean its dynamic evolution or time course in continuous time. These properties are as follows.

- For a given network, there are usually many ways to partition the network species into non-overlapping groups A , B and D so that *the information encoded by the trajectory of D* makes any information in the

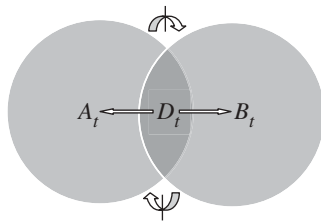


Figure 1. Illustrative example of a network with two modules. Each circle represents a module, i.e. a group of biochemical species. The network species are grouped into three non-overlapping sets A , B and D as shown, in such a way that $A_t \leftrightarrow_{D_t} B_t$ —i.e. the trajectories (up to any time t) of A and B are independent given the trajectory of D . Hence A_t and B_t contain no mutual information given D_t , all information transfer between the two modules being conveyed via D_t .

trajectory of A irrelevant for that of B (and vice versa). The species in D thus play the role of informational intermediaries or ‘carriers’. Such species may be identified by establishing the conditional independence between trajectories, $A_t \leftrightarrow_{D_t} B_t$.

- Furthermore, the entire network can be decomposed exactly into modules on *informational* grounds, and the modules arranged in a tree structure that encodes conditional independences between the trajectories of (groups of) modules. The path in the tree from one module to another then identifies the route and species involved in sequential information processing between the two modules.

We will denote the state of the molecular network by $x(s) = [x_1(s), \dots, x_k(s), \dots, x_n(s)]$, where $x_k(s)$ is the number of molecules of type k present at time s . The trajectory $X_{k,t} = (x_k(s); 0 \leq s \leq t)$ records the dynamic evolution of the k th species. Figure 1 shows a simple illustration of a network with two modules. The network species are grouped into the three sets A , B and D in such a way that the conditional independence relationship $A_t \leftrightarrow_{D_t} B_t$ holds. We stress the informational aspect of the relationship—namely that the trajectories A_t and B_t contain no mutual information given D_t , and the species in D play the role of ‘information carriers’ between A and B within the network. The network is thus said to decompose exactly into the two modules $M_1 = [A, D]$ and $M_2 = [B, D]$ (shown as circles), with all information relevant to each module’s trajectory thus encoded by their region of overlap, D .

Importantly, our approach is well suited to the analysis of information processing by cell signalling networks. It is commonplace to refer heuristically to information flow within such networks. However, a precise, quantitative understanding of how (and which) internal species of the cell encode information about the trajectories of signalling inputs (ligands) is lacking. Denote by I_t^* the stochastic process by which multiple signalling ligands enter and depart the cell’s surroundings, and by O some target output species of the signalling network. We refer to I_t^* as the external input process. By identifying a conditional independence $A_t \leftrightarrow_{D_t} B_t$ as above, where the trajectory A_t (resp. B_t) implies that of I_t^* (resp. O_t), we are able to

identify encoding species D such that the conditional independence $I_t^* \leftrightarrow_{D_t} O_t$ holds. (Furthermore $I_t^* \leftrightarrow_{E_t} O_t$ whenever the species D are contained in E .)

The implication is that, given the trajectory of the species in D , the trajectories I_t^* and O_t contain no mutual information. This may be interpreted in two ways. First, the encoding of information by the trajectory D_t makes the *original information in the external input process I_t^* irrelevant for the outputs O_t* . Furthermore, this irrelevance holds no matter what other trajectories in addition to those of D are also considered. Second, inferences about the multiple signals received, I_t^* , based on D_t are unaffected by taking the outputs O_t into account. These properties of the trajectory D_t thus capture the essence of signal ‘transduction’ by intermediary species.

In general, the signalling network modularizations we identify are represented using a tree structure that allows us to trace *sequential* routes of information transfer through the network. Denote the groups of species on the path in the tree between signalling inputs (I^*) and outputs (O)—i.e. the relevant ‘edges’ in the tree—by S_1, S_2, \dots, S_N . Then an important result (see §2.4) will be that $I_t^* \leftrightarrow_{S_{j,t}} O_t$ for $j = 1, \dots, N$, i.e. conditional independence holds for all of the species groups S_j , and moreover $(I_t^*, S_{1,t}, \dots, S_{j-1,t}) \leftrightarrow_{S_{j,t}} (S_{j+1,t}, \dots, S_{N,t}, O_t)$. The latter property means that (S_1, S_2, \dots, S_N) is a *sequence of encoders*, each one of which makes the information in both I_t^* and all earlier encoders in the sequence irrelevant for both all later encoders in the sequence and the outputs O_t . A comparative analysis varying the groups I^* and O for a given network then reveals important differences of information processing for the various I–O combinations.

2.1.1. Module kinetics may be studied in isolation. The problem of when it is appropriate to study smaller subsystems without reference to the network in which they are embedded arises frequently in systems biology. We therefore mention the following property of our network decompositions (without further development). It turns out that, given the definition of a decomposition or modularization here (§2.3), valid conclusions may be drawn by studying the kinetics of each module in isolation from the rest of the system. Consider again the two-module network in figure 1 (where it is assumed that conditions 1, 2 and 3 of §2.2 apply). Observing the trajectories of all species in the first module $M_1 = [A, D]$ alone, without also observing the trajectories of any species in B , allows many of the rate parameters of the reactions governing the dynamics of M_1 to be estimated using maximum likelihood. (We abstract from practical issues of data collection such as discrete sampling.) More specifically, the estimators may be computed for the rate parameters of all reactions that change some species in A and of reactions that change species in D alone. This is a surprising result and one that arises from the ability to decompose the likelihood function along the lines of the conditional independence $A_t \leftrightarrow_{D_t} B_t$ —see ([15], theorem 4.5). (The only reactions changing M_1 whose rate parameters cannot be

estimated are those that change both B and D but not A —inevitably there is some loss from not observing B .) These comments extend straightforwardly to decompositions with greater than two modules.

2.2. Identifying dynamic conditional independences

The first stage in identifying conditional independences between trajectories for a given reaction network is to construct its *kinetic independence graph* or KIG. The nodes in this directed graph correspond to the different biochemical species. The graph is constructed in order to display, for each species, those nodes whose concentration influences the instantaneous, stochastic kinetics of that species (see the electronic supplementary material, SM1.2; [15]). An arrow (directed edge) is therefore drawn from species j to species k whenever j is a *reactant* in a reaction that either results in the overall production or consumption of molecules of k . In the former case k usually participates in the reaction only as a product whereas in the latter case j and k react together. It is an obvious but important point here that only the concentrations of reactants influence instantaneous reaction rates or ‘intensities’. (Reverse reactions are treated as separate reactions in the formalism.)

For any collection of species A , it is possible to show that the instantaneous kinetics of A depend only on the current concentration (or ‘levels’) of A itself and those species that are parents of A in the KIG—i.e. those species (outside A) that lie at the tail of an arrow that points to some node in A . Construction of the KIG of a reaction network requires only limited stoichiometric information about the network—often all that is needed is a signed binary, $(0, \pm 1)$ -version of the stoichiometric matrix identifying, for each reaction, those species that are overall consumed and produced. Importantly, no knowledge of rate parameters is required. Figure 2 shows the KIG for the NF κ B signalling network example discussed below.

The following means of identifying conditional independences applies to a general class of stochastic biochemical reaction networks termed (standard) stochastic kinetic models. The class includes, but goes considerably beyond, the class corresponding to the stochastic simulation algorithm of Gillespie [34]. (The SKM class is described formally in the electronic supplementary material, SM1.1.) Let G be the (undirected) graph obtained from the KIG of the reaction network by substituting lines for all arrows. A reaction is said to *change* A if it causes the level of some species in A to change (no matter which particular species). For a partition $[A, D, B]$ of the species in the network, the conditional independence of trajectories $A_{t \leftrightarrow D_t} B_t$ holds whenever two conditions are both satisfied (for mathematical proof, we refer to [15], corollary 4.6 and proposition 4.9):

- A and B are separated by D in the graph G —i.e. every path along lines in G that begins at a node in A and (without revisiting any node) ends at a node in B visits some node in D (condition 1); and

- for any two reactions that change D and result in identical changes to all species in D , either both reactions change A or neither changes A , and either both change B or neither changes B (condition 2).

Additionally, we require that a weak regularity condition, condition 3, holds for the set of reactions that change both A and B , namely that these reactions be ‘identified by consumption of reactants’ (see the electronic supplementary material, SM1.1). Condition 1 is equivalent to there being no species in A that is a reactant in a reaction that results either in the net production or consumption of some species in B (i.e. in a reaction that changes B), and vice versa. When this fails to hold, there is a direct influence of A on the instantaneous kinetics of B , or of B on A (or both). Condition 2, together with the first, ensures the conditional independence of the trajectories up to time t (for all $t > 0$). A full understanding of the role of this condition is rather involved but the following point is apparent—given the trajectory of D , the possible occurrence of two reactions that change D identically can be distinguished using the trajectory of B if one of them changes B but the other does not; this information is then often relevant for the trajectory of A (contradicting the conditional independence of A and B).

In the mathematical literature, the dynamic conditional independence relationship $A_{t \leftrightarrow D_t} B_t$ is usually written as $A_t \perp\!\!\!\perp B_t | D_t$ (and, technically, is a relationship between σ -fields). We prefer the more intuitive notation $A_{t \leftrightarrow D_t} B_t$ in this context. Note that the relationship concerns the trajectories up to time t . (There may, owing to a lag effect, be information in A_t relevant for the trajectory of B beyond time t and up to time u (when D_t is given) but this information will be encoded in D_u so that $A_{u \leftrightarrow D_u} B_u$ holds at the later time u .)

2.2.1. Illustrative example—NF κ B signalling. The exposition of the results and methods of the paper includes, as a running example, a kinetic model of the core mechanisms of NF κ B signalling. An important contribution of the work lies in the ability to analyse information processing by large, complex reaction networks (see the TLR example in §2.4.1). However, the smaller NF κ B example is useful both in aiding rapid understanding of the techniques and in revealing the often subtle informational properties of biochemical networks.

The network used is based on a state-of-the-art, stochastic dynamic model recently developed using extensive experimental data by Ashall *et al.* [7]. Essential aspects include the TNF ligand-induced activation of I κ B.NF κ B and degradation of pI κ B, leading to release of NF κ B; and translocation to the nucleus of NF κ B. Three negative feedback loops are set up as NF κ B itself activates transcription of I κ B α , I κ B β , and A20. The latter inhibits conversion from the inactive to the neutral form of IKK. Complete details of the reaction network (together with species names) may be found in the electronic supplementary material, SM2.

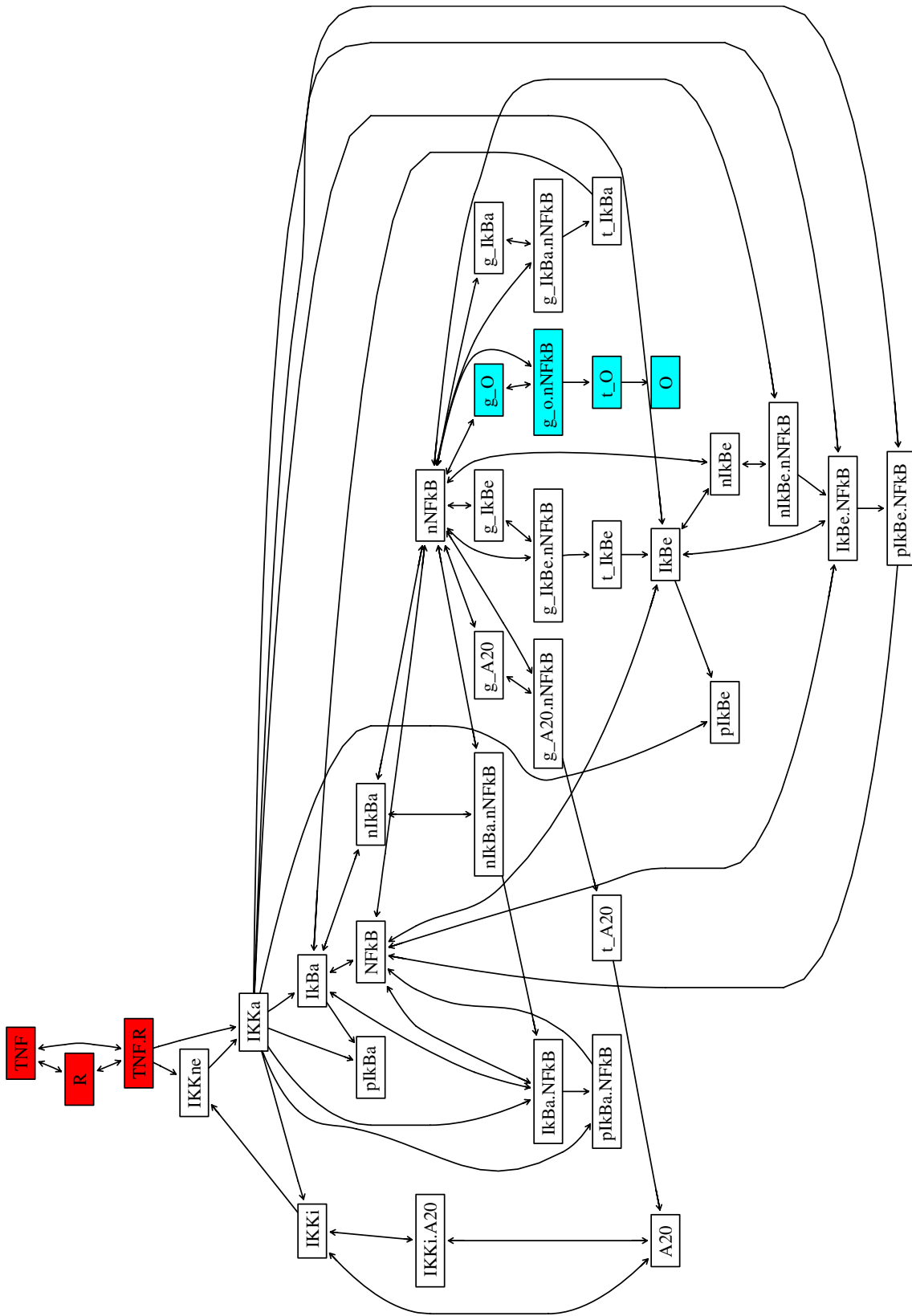


Figure 2. Kinetic independence graph of the NFκB signalling network. TNF is the tumour necrosis factor ligand, R is the ligand's receptor, IKBa is IkBe, and IKBe is IkBe; IKK has three forms (ne neutral, a active, and i inactive); O denotes a generic output protein; the prefix g denotes gene, t transcript, p phosphorylation and n nuclear location. For ease of viewing, input- and output-associated species are coloured red and blue, respectively. Complete details of the reaction network are given in the electronic supplementary material, SM2.

Figure 2 shows the associated KIG. To represent generic outputs of the signalling network, the subgraph $g_{O.nNF\kappa B} \rightarrow t_O \rightarrow O$ is included (the NF κ B-activated expression of protein O). Much more complicated ‘output graphs’ would not alter our conclusions—the important aspect is that the output species are separated from all other species in the KIG by $g_{O.nNF\kappa B}$ (or its equivalent for multiple promoters)—feedback from O is thus ruled out. Throughout we use \mathcal{V} to denote all species in a given network; the notation \setminus should be read as ‘excluding’ (and corresponds to set difference).

Let $TNF_t^* = I_t^*$ be the external ligand or input process. We will show how to use conditions 1 and 2 above to address the following hypotheses or conjectures concerning information processing by the NF κ B network (the associated conditional independence $A_{t \leftrightarrow D_i} B_i$ is shown in parentheses).

- All information relevant to signalling output is encoded by the trajectory of the promoter occupancy of output genes ($[t_O, O]_{t \leftrightarrow g_{O.nNF\kappa B}, [\mathcal{V} \setminus \{g_{O.nNF\kappa B}, t_O, O\}]_t}$).
- The encoding of information by the trajectory of the active kinase, $IKKa_t$, makes all information about the external input process TNF_t^* and about ligand binding and unbinding by receptors irrelevant for the rest of the network, including the output ($S_{t \leftrightarrow IKKa_t} [\mathcal{V} \setminus \{S, IKKa\}]_t$, where $S = \{TNF, R, TNF.R\}$ and TNF_t^* is implied by S_t).

Both hypotheses perhaps appear plausible given knowledge of the reaction network—however, only the first is correct. In the case of the first, the graphical separation required by condition 1 is satisfied (figure 2), as is condition 2 since no two reactions cause the same change in $g_{O.nNF\kappa B}$ (see the electronic supplementary material, SM2). In particular, all information about TNF_t^* relevant to output is encoded in the trajectory of the promoter occupancy of output genes. The second hypothesis is false. Condition 1 fails (owing to the edge $TNF.R \rightarrow IKKne$ in the KIG)—the trajectory of the neutral form of the kinase, $IKKne_t$, in fact encodes additional information about $TNF.R_t$. The reason is that the reaction list of the network (see the electronic supplementary material, SM2) means that although the trajectory $IKKa_t$ implies the history of the activation reaction $TNF.R + IKKne \rightarrow IKKa + TNF.R$, it does not imply knowledge of the production and hence trajectory of $IKKne$. Since activation of IKK involves both $TNF.R$ and $IKKne$, the trajectory $IKKne_t$ provides additional information about the trajectory of receptor occupancy, $TNF.R_t$.

It is straightforward to verify using conditions 1 and 2 that $S_{t \leftrightarrow IKKa_t, IKKne_t} [\mathcal{V} \setminus \{S, IKKa, IKKne\}]_t$, from which we are able to conclude that $TNF_t^* \leftrightarrow_{IKKa_t, IKKne_t} [t_O, O]_t$. That is, all information about the external stimulus relevant for outputs is encoded in the trajectory of the distribution of IKK across its different isoforms (the level of free plus complexed $IKKi$ being implied by the other two forms). This interesting observation demonstrates that thinking in terms of a signal transduction or activation ‘path’ such as $TNF.R \rightarrow$

$IKKa \rightarrow pIKB.NF\kappa B \rightarrow NF\kappa B \rightarrow nNF\kappa B$ can deliver misleading conclusions about informational properties of the network. Note that $A20$ does not appear—in particular $IKKi.A20_t$ need not be included in the conditioning information because the relevant effect of the negative feedback loop involving $A20$ is ‘impounded’ in the trajectory $IKKne_t$. The example serves to emphasize that a rigorous, mathematical framework is both indispensable for reasoning correctly about information encoding and transfer, and able to deliver biologically relevant conclusions.

2.3. Identifying exact network decompositions

Using our methods, as was emphasized in §2.1, a given biochemical network can usually be decomposed exactly into modules and the modules arranged in a tree structure encoding conditional independences between the trajectories of groups of modules. This then provides a framework for analysing biochemical information processing and is particularly beneficial for larger networks. A modularization here, $\{M_d\}$, is a collection of groups of species with the following two properties: (i) all network species are assigned to at least 1 module; and (ii) the trajectory of each module M_d is conditionally independent of the trajectory of the other modules, given the trajectory of the overlap of M_d with the rest of the network—i.e. $M_{d,t} \leftrightarrow_{S_{d,t}} [\cup_{e \neq d} M_e]_t$, where the overlap (or interface) $S_d = M_d \cap \{\cup_{e \neq d} M_e\}$, and \cap, \cup denote the intersection and union of sets, respectively. Note that this dynamic conditional independence is equivalent to $[M_d \setminus S_d]_{t \leftrightarrow S_{d,t}} [\mathcal{V} \setminus M_d]_t$. All information relevant to a module’s trajectory is encoded by the module’s overlap region with the rest of the network. (It should be noted that this definition of a modularization is considerably stronger than the mathematically abstract one proposed by [15]. Although computationally much more demanding, this definition is the biologically relevant one.)

We have developed a modularization identification by dynamic independence algorithm (MIDIA) based on graphical decomposition methods. It implements automated identification, with the modularization displayed and visualized as a junction tree, \mathcal{T}_M . A junction tree is a (connected, acyclic) undirected graph in which the nodes of the graph are the modules themselves. Furthermore, the overlap or intersection of any two modules of the tree, $M_d \cap M_e$ ($d \neq e$), is contained in every module on the unique path in the tree between M_d and M_e .

Identifying a network modularization by manual inspection of the KIG is far from straightforward even for the small NF κ B signalling network (recall figure 2). With even a modest increase in the size and complexity of the network, a principled and computationally efficient approach to module identification becomes indispensable. Three particular problems must be addressed. First, while using the KIG to find some three-way partitions of the species satisfying condition 1 can be relatively straightforward, it is far from obvious how to assign all network species to modules in a consistent manner so that each module is separated in the KIG from the rest of the network by its interface.

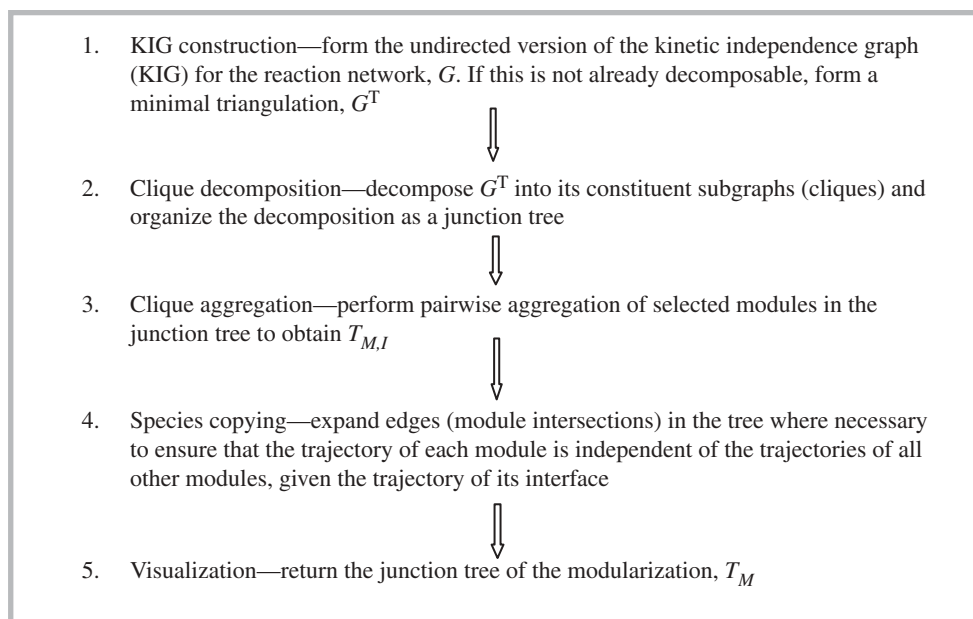


Figure 3. Overview of the stages of the MIDIA algorithm. The algorithm automatically identifies a modularization of the biochemical network on dynamic, informational grounds (see text) and displays it as a junction tree in which the modules are the nodes of the tree. The undirected version of the KIG, G , is formed simply by replacing all arrows with ('undirected') lines. In forming G^T , enough edges are added (but no more) to make the graph triangulated. The cliques of G^T are its maximally complete subgraphs. For these and other concepts in basic graph theory see Cowell *et al.* [37, ch. 4]. The details of the MIDIA algorithm are explained in the electronic supplementary material, SM3.

Second, it is desirable to have control over the degree of coarse graining of the modularization identified. Third, ensuring that condition 2 is satisfied for every module interface S_d is particularly challenging. The solutions to these problems are, respectively, clique decomposition, clique aggregation and species copying within the junction tree.

An overview of the various stages of the MIDIA approach is given as a flowchart in figure 3 (see the electronic supplementary material, SM3, for further details). Stages 1 and 2 involve decomposition of the (undirected, minimally triangulated) KIG into a junction tree in which the nodes of the tree are the cliques. Use is then made of results concerning properties of the clique junction tree for a triangulated graph. These properties turn out to be stable when neighbouring modules in the tree are 'merged' or aggregated in stage 3. The selection of module pairs for aggregation is automated and allows the degree of coarse graining of the network to be controlled by setting a minimum size for the module residuals, $M_d \setminus S_d$. The resultant junction tree, $\mathcal{T}_{M,I}$, constitutes a modularization in which the instantaneous kinetics of each module residual are independent of all the network species outside the module. It is an important output in its own right, giving insight about the kinetic structure of the network. Stages 4 and 5 are explained in the following subsection.

The junction tree, $\mathcal{T}_{M,I}$, for the NF κ B network (§2.2.1) is shown in figure 4. The benefits of a principled modularization in revealing the architecture of the network implicit in the KIG are obvious even for this small network. Working from the top to the bottom of $\mathcal{T}_{M,I}$,

the NF κ B network is decomposed into modules associated with: the TNF receptor and IKK; the A20 negative feedback mechanism; cytoplasmic NF κ B, I κ B and their complexes; nuclear I κ B α and its gene expression; nuclear I κ B ϵ and its gene expression; and finally the signalling outputs. Each edge in the tree contains species determining the instantaneous kinetics of the two adjacent module residuals. The central roles played by IKK α , nuclear NF κ B, and nuclear import and export processes are thus made clear.

We emphasize that for moderately sized or large networks an automated tool for investigating the modular architecture of the network based on its dynamic properties is essential—manual inspection of reaction mechanisms and their raw graphical representations becomes infeasible (consider, for example, the TLR signalling network in §2.4.1). The junction tree in figure 4, obtained from the fully automated algorithm, illustrates MIDIA's efficacy in this regard using the example of a relatively small network for transparency.

2.3.1. Encoding dynamic conditional independences in junction trees. This section explains how to read conditional independences between species trajectories from the junction tree returned by the MIDIA algorithm. As is usual, each edge between adjacent modules in a junction tree is associated with the modules' overlap which we denote for modules d and e by $S_{de} = M_d \cap M_e$. Consider first the intermediate tree, $\mathcal{T}_{M,I}$, returned by stage 3. Imagine cutting any one of its edges, say the one between the modules M_d and M_e . Denote the species present in each of the two

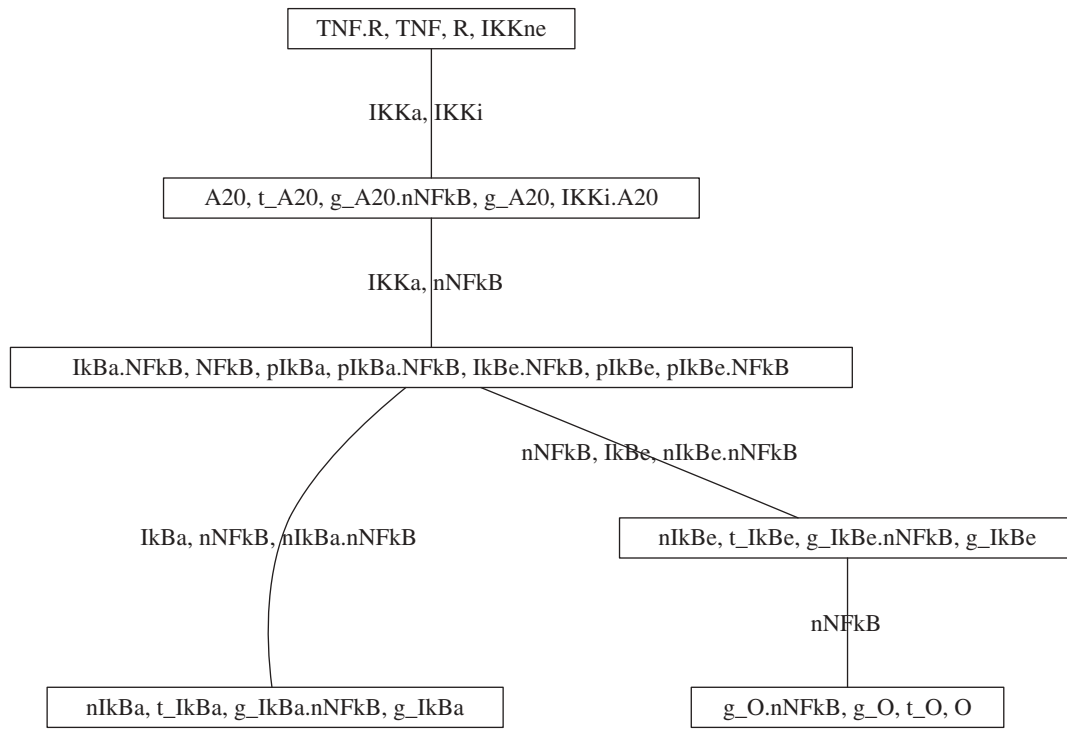


Figure 4. Modularization $\mathcal{T}_{M,I}$ for the NF κ B signalling network. The modularization is the one returned after stage 3 of the MIDIA algorithm (using a minimum residual size of 4 (except for the ‘root’ module)), and is based on instantaneous kinetics alone (see text). Each module M_d is labelled with the corresponding residual ($M_d \setminus S_d$); each edge is labelled with the intersection of the two modules it connects.

subtrees obtained from the cutting procedure by V_{de} and V_{ed} , respectively. Then an important property is that V_{de} and V_{ed} are separated in the undirected KIG by the modules’ intersection—there is no direct influence of either ‘subtree’ on the instantaneous kinetics of the other. The species copying procedure (stage 4) is designed (i) to preserve this property in the final tree, \mathcal{T}_M , thus allowing application of our condition 1 (§2.2) and (ii) to ensure via condition 2 that for the two subtrees obtained by cutting any one of its edges, the conditional independence of trajectories $V_{de,t} \leftrightarrow S_{de,t}$ holds (see theorem 1 of the electronic supplementary material, SM1.3). It is then possible to show for each module M_d in \mathcal{T}_M that $M_{d,t} \leftrightarrow S_{d,t} [\cup_{e \neq d} M_e]_t$ (see theorem 2 of the electronic supplementary material, SM1.3). These dynamic conditional independences, of course, constitute the defining property of our modularizations.

In stage 4, the MIDIA algorithm works ‘backwards’ through the junction tree $\mathcal{T}_{M,I}$, imposing condition 2 on each edge of the tree, S_{de} , so as to preserve the junction tree property and to leave edges treated in previous steps unchanged. Each edge or intersection of neighbouring modules, S_{de} , is enlarged by ‘copying’ the appropriate species from a module containing them, M_g say, both to M_d (which lies on the unique path between g and e) and to M_e . The species are also copied to all other modules on that path. For example, in the case of the NF κ B network, the edge {IKKa, nNF κ B} in $\mathcal{T}_{M,I}$ is enlarged by including g_A20 also in the third module. (The whole tree \mathcal{T}_M returned after stage 5 for the NF κ B network is shown in the

electronic supplementary material, SM4.) Cutting this edge in \mathcal{T}_M and applying theorem 2 then reveals the non-obvious conditional independence $S_t \leftrightarrow [\text{IKKa, nNF}\kappa\text{B, } g_A20]_t [\mathcal{V} \setminus \{S, M_1, M_2\}]_t$, where $S = \{\text{TNF, R, TNF.R}\}$ as before, and M_1, M_2 are the first and second modules (from the top) of \mathcal{T}_M . As is the case for the trajectory of the distribution of IKK across its isoforms, the trajectory $[\text{IKKa, nNF}\kappa\text{B, } g_A20]_t$ is thus another encoder with only a few species that makes the information in S_t irrelevant for the trajectories of all species containing cytoplasmic and nuclear NF κ B, for the trajectories of the IKB’s (i.e. their protein complexes, and genes and transcripts) and for the trajectories of the output module. This encoder is equivalent to the trajectory of active IKK and nuclear NF κ B, both free and bound to the A20 promoter. Space precludes further discussion of the NF κ B example here. Sequential information processing is considered in general and in the context of the TLR signalling network example below.

2.4. Identifying paths of sequential information processing in signalling networks

As was discussed earlier in §2.1, our modular approach and the MIDIA algorithm are well suited to analysing information processing by signalling networks, especially large ones. Let M_{I^*} be a module identified whose trajectory contains that of an external input process, I_t^* , and let M_o be a module containing the target output O . Consider the (unique) path in the junction tree \mathcal{T}_M from M_{I^*} to M_o , which we write as

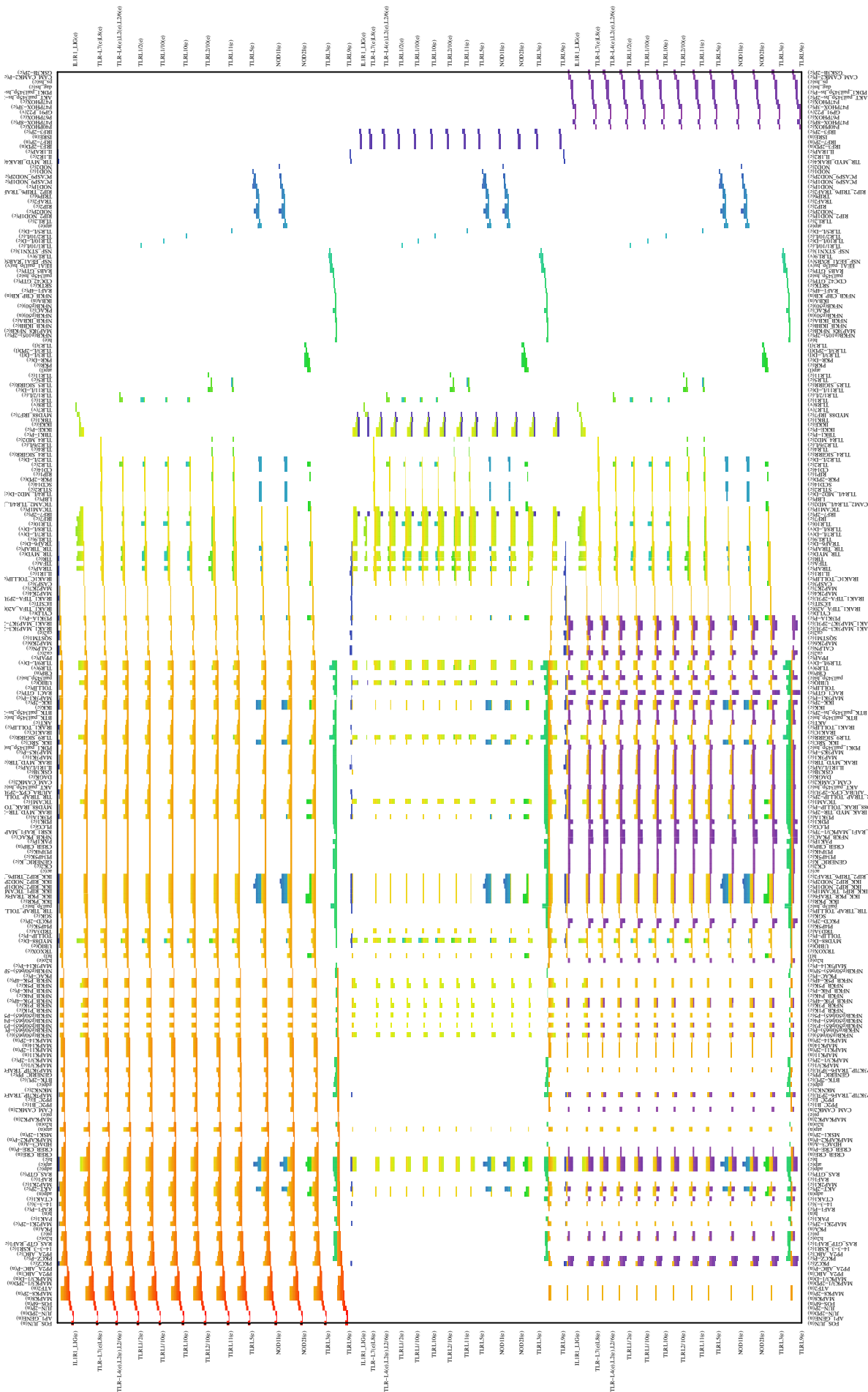


Figure 6. Sequential information processing by the TLR network: the I–O path matrix. Each column corresponds to a species (named as in [28], table S1) and input processes are labelled on the vertical axis. The upper, central and lower regions of the plot correspond to the outputs API-JUN, ISRE-IRF3 and NADPH-OX-3P, respectively. Each row in the matrix indicates the species composition of a given edge on an I–O path, with edges colour-coded (as in figure 5). Rows of the matrix are grouped according to the individual I–O combinations and ordered within a group from top to bottom according to the sequence of the edges in the corresponding I–O path in T_M in figure 5.

$M_{I^*} - S_1 M_1 - \dots - S_{N-1} M_{N-1} - S_N M_o$, N being the number of edges on the path. The sequence of edges (S_1, S_2, \dots, S_N) reveals the sequential process by which the network encodes and transfers information. Imagine cutting the edge S_j —then applying the result of §2.3.1 on the subtree trajectories implies that $I_t^* \leftrightarrow_{S_{j,t}} (S_{j+1,t}, \dots, S_{N,t}, O_t)$ for $j=1, \dots, (N-1)$, and furthermore implies the main result that $(I_t^*, S_{1,t}, \dots, S_{j-1,t}) \leftrightarrow_{S_{j,t}} (S_{j+1,t}, \dots, S_{N,t}, O_t)$ for $j=2, \dots, N$. To see this, recall that (using this notation), for example, $S_{j+1,t} = M_j \cap M_{j+1}$, and therefore the species (S_{j+1}, \dots, S_N, O) are in the ‘subtree’ $V_{j,j-1}$ while $(I^*, S_1, \dots, S_{j-1})$ is ‘contained in’ the subtree $V_{j-1,j}$.

The main result here then, $(I_t^*, S_{1,t}, \dots, S_{j-1,t}) \leftrightarrow_{S_{j,t}} (S_{j+1,t}, \dots, S_{N,t}, O_t)$, means that (S_1, S_2, \dots, S_N) is a sequence of encoders, each one of which encodes information that makes both I_t^* and all earlier encoders in the sequence irrelevant for all later encoders in the sequence and for the outputs O_t . Using the junction tree of the modularization identified by MIDIA we are thus able to identify truly sequential routes of information processing by the network. Since identifying where the information resides is a prerequisite for understanding how it is encoded and determining the extent of any information loss involved, this is an important step in the development of a quantitative theory of cellular information processing.

The MIDIA algorithm is tailored to identify the routes of sequential encoding as follows. After performing stages 1 and 2, the modules containing the external input (processes) and outputs are determined, and all edges in the tree that lie on a path between such an input and an output module identified. We call these the I–O paths. In stage 3, clique aggregation is performed according to the following steps: (i) to remove any edges containing either an input or output and (ii) by aggregating as many modules as possible without altering any of the modules on the I–O paths, in order to focus attention exclusively on those paths. Stages 4 and 5 are as usual.

2.4.1. The TLR signalling network. As a proof of principle, the tailored MIDIA algorithm was applied to the stoichiometric reconstruction of the TLR signalling network presented by Li *et al.* [28]. The TLR network is large, having 752 species and 1034 reactions (after accounting for reaction reversibility; for species names see [28], table S1). There are 16 external input processes. These correspond to the extracellular trajectories of the individual receptor-specific ligands (with the exception of the input process for NOD1 which also includes the cytoplasmic trajectory of the NOD1 ligand since some ligand arrivals are owing to an export reaction from the cytoplasm). We focus on three groups of output species of the network—(AP1-JUN, AP1-JUN-FOS), (ISRE-IRF3, ISRE-IRF7) and vacuolar NADPH-oxidase (in two different phosphoforms). The first two consist of active transcription factor (TF) complexes bound to their DNA sites, while the latter is responsible for reactive oxygen species production.

Figure 5 shows the junction tree \mathcal{T}_M and I–O paths for the TLR network. In order to investigate further the structure of the I–O paths while explicitly taking the species content of the edges into account, we introduce a visualization technique called the I–O path matrix. The construction of the I–O path matrix is explained in figure 6, which depicts the matrix for all of the inputs in the TLR network and three outputs (one from each output pair). On-screen viewing of the matrix is recommended. Together, the tree \mathcal{T}_M and the I–O path matrix are powerful, complementary visual tools for investigating information processing by the network.

This is not the place to undertake a lengthy and detailed analysis of information processing by the TLR network. Rather, the utility of our approach is illustrated with a discussion of the insight provided into the following aspects. First, the grouping or clustering of signalling inputs (TLR) is found to be similar on the basis of how information is encoded by the network. Second, as indicated by Kanae & Kitano [35], there is a need to go beyond a ‘bow tie’ conception of the architecture of the network when considering signal transduction. The tree \mathcal{T}_M provides such a ‘structural’ view of how the network processes information between various inputs and outputs. We are able to identify the species involved in information encoding for the various outputs (figure 6), the sequential I–O paths and their associated shapes, and those ‘core’ encoding species in common between the I–O paths. The analysis reveals the breadth of encoding species in less peripheral parts of the I–O paths rather than a bottleneck—the information processing structure is not well described as a bow tie. Third, we find that the extent to which information encoding is distributed broadly across many species differs significantly between the ISRE-IRF and the AP1/NADPH-oxidase outputs (the TLR9 input excluded). Encoding in the case of the ISRE output is much sparser, while the other two outputs share more similar I–O paths (figure 6). Fourth, for a large cluster of 12 TLRs, we identify a surprisingly small group of species responsible for mediating information transfer from the ‘core’ encoding species to the ISRE-IRF outputs. Fifth, we show that in the context of multiple stimuli, inputs can be informative for an output in ways not revealed by steady-state FBA. Each of these aspects is considered in turn below.

The structure of the tree \mathcal{T}_M immediately implies a hierarchical clustering of the input processes on the basis of their associated I–O paths—the dendrogram or tree for the clustering being derived directly from the structure of \mathcal{T}_M in the obvious way. Since the species contents of edges in \mathcal{T}_M can overlap to differing degrees, we note that the I–O path matrix in figure 6 strongly supports this hierarchical clustering of inputs using \mathcal{T}_M . In particular, the following groups of inputs are seen to possess (within each group) extremely similar sequential paths of information processing to all three output types: (TLR-L1/2, L1/10, L10, L2/10), (TLR-L4, L2, L2/6), (TLR-L5, L11), (NOD1I, NOD2I) and (TLR-L7, L8). At a higher level within the clustering hierarchy, there is a large group of 12

inputs contained in the subtree given by cutting edge 29 in \mathcal{T}_M —this group, denoted $\text{TLRI}_{\setminus \text{IL1},7,9}$, contains all but four of the external inputs (the IL1R input and TLR-L7,8,9). TLRL9 stands out as the most exceptional individual input, followed by ILR1L. As can be seen from figure 6, the 12 external inputs comprising $\text{TLRI}_{\setminus \text{IL1},7,9}$ share substantively similar I–O paths.

The breadth of encoding species in less peripheral parts of the I–O paths is apparent, as opposed to any bottleneck or narrow ‘bow-tie centre’. The number of species in the constituent edges of each I–O path in \mathcal{T}_M (and also in $\mathcal{T}_{M,I}$) is roughly ‘diamond-shaped’ going from input to output—i.e. early and late edges have few species while intermediate edges have many. Almost always, the earliest edges in the sequence contain exclusively receptors, ligands and receptor–ligand complexes, while the latest edges contain species closely related to the outputs (the output TFs and binding sites, or the constituents of the NADPH-oxidase complex). The diamond shape might appear to be largely the result of the ‘confluence of multiple paths’ within the tree that could nevertheless retain ‘functionality’ alone—however, this is not the case. The structure of the tree \mathcal{T}_M reflects the edge content and hence path content of the undirected KIG (owing to stages 1 and 2 of MIDIA); hence, even when inputs and outputs are considered only pairwise, an edge on an I–O path in $\mathcal{T}_{M,I}$ is a minimal separator of the particular input and output in the undirected KIG in 92 per cent of all cases. In connection with the breadth of encoding species in less peripheral parts of the I–O paths, we note the existence of edges on the path to each output that contain a relatively large number of species and are shared by all or nearly all of the inputs—e.g. edges 7, 8, 9 and 11 for AP1 outputs, edges 11 and 15 for NADPH-OX outputs, and edges 22 and 27 for ISRE-IRF.

Perhaps the most immediately obvious feature of figure 6 is the similarity between I–O paths corresponding to a given output (the TLRL9 input excluded), compared with the clear differences across outputs. In order to focus on these differences, the following discussion is for the group of 12 external inputs $\text{TLRI}_{\setminus \text{IL1},7,9}$ which share very similar I–O paths. The structure of \mathcal{T}_M and consideration of the species involvement (see figure 6) strongly indicate that the I–O paths are closely related for the NADPH-OX and AP1 outputs while encoding for the ISRE output is much ‘sparser’ (involves fewer species) and is rather different. For the group $\text{TLRI}_{\setminus \text{IL1},7,9}$, consider for each output the edges on the I–O paths in \mathcal{T}_M that are in common between all of the inputs—e.g. the sequence of edges 29, 22, . . . , 501, 504 in the case of ISRE. (The group of species found on all of the I–O paths for a particular output is identical to the species content of these common edges, except for the inclusion of one additional, identical species in each case.) These common edges contain 57 different species for ISRE-IRF—compared with 137 for NADPH-OX and 165 for AP1—of which 48 are found on the I–O paths for all three outputs and only nine are specific to the ISRE output alone. A further 77 species are found on the I–O paths for both

NADPH-OX and AP1 but not for ISRE. There are thus 12 species specific to the NADPH-OX output alone and 40 species specific to AP1 alone.

For the large group of 12 inputs, $\text{TLRI}_{\setminus \text{IL1},7,9}$, we thus identify a ‘core’ of 48 species involved in the three sequences of encoders ending at the respective output modules (and beginning in each case with edge 29). These species, listed in the electronic supplementary material, SM5, are also located in a central region of the tree, in edges 19, 22 and 29 (the latter consisting only of core species). For the ISRE output, the path of sequential information processing involves only these 48 species and (excluding the four that are direct components of the output species themselves) the following: TBK1-P, TBK1, IKK ϵ -P, IKK ϵ and MYD88_TRAF6_IRF7. (Together with cytoplasmic IRF7-2P and ubiquitin the latter group comprises edge 154.) The small group (TBK1-P, TBK1, IKK ϵ -P, IKK ϵ , MYD88_TRAF6_IRF7) can thus be seen as responsible for mediating information transfer from the ‘core’ encoding species to the ISRE output. We view encoding for the other two output types as distributed across many more species trajectories (as less sparse), with the species involved for NADPH-OX typically involved also for AP1. Edge 11, for example, contains 75 species and is common to the sequence of encoders for both outputs.

We conclude this section by drawing a number of contrasts with the steady-state FBA of the same network in Li *et al.* [28]. The FBA analysis there takes a collection of steady-state ‘snapshots’ of the network—in each one, a single input flux is held at a constant non-zero level and the network is assumed to optimize a single output flux. These snapshots are then combined in the manner described in Li *et al.* [28] to obtain discrete signalling (‘DIOS’) pathways. Groupings of inputs are provided by ([28], fig. 4), but the same individual input may appear in groups associated with different DIOS pathways, depending on the output flux chosen as the objective to be maximized. As has already been emphasized, our analysis makes no steady state or stationarity assumption. Nor is it restricted to the separate analysis of particular I–O combinations. Rather, the signalling system is exposed to any number of the possible time-varying input processes and all output trajectories are determined simultaneously by the network biochemistry, as in the living cell. The individual I–O paths in figures 5 and 6 are system properties that hold simultaneously—indeed, \mathcal{T}_M may be used to analyse paths of sequential information processing between a group of inputs (e.g. the group $\text{TLRI}_{\setminus \text{IL1},7,9}$) and a group of outputs, as has just been described.

In steady state, Li *et al.* [28] report that certain individual inputs are unable to affect certain outputs—e.g. no input affects the ISRE-IRF7 output, NOD1 and NOD2 do not affect any of our outputs, and TLRL3 does not affect AP1 outputs. Computational analysis of the KIG for the network reveals by contrast that there is a directed path in the KIG (i.e. one following arrows), not involving ‘currency’ species, from all individual inputs to all outputs. Such paths correspond to chains of direct kinetic influence from input to output (each ‘link’ or edge of which

will be operative at any point in time provided no reactant level is zero for the reaction giving rise to it). Thus, for example, the NOD1 and NOD2 input processes can be informative for all output trajectories, including the ISRE-IRF3 and -IRF7 outputs. One way in which this can occur (revealed by interrogation of the KIG) is that activation of both NOD inputs raises $\text{IKK}_{\alpha,\beta,\gamma}\text{-P}$ levels, which consumes substrates of $\text{IKK}\epsilon\text{-P}$ (such as $\text{NF}\kappa\text{B}(\text{p}50/\text{p}65)$), thus reducing sequestration of $\text{IKK}\epsilon\text{-P}$ and consequently increasing phosphorylation of IRF3 and (in the presence of MYD88-TRAF6-IRF7) of IRF7. The encoders in the rows of figure 6 corresponding to NOD1 and NOD2 render these inputs uninformative (e.g. $\text{IKK}_{\alpha,\beta,\gamma}$ is seen to be an encoding species in the case of the IRF3 output). Similarly, TLR3 can be informative for the AP1 outputs. Thus in the context of multiple stimuli, inputs can be informative for an output in ways not revealed by steady-state FBA.

3. DISCUSSION

Understanding how information is encoded and transferred by the biochemical reaction networks inside cells requires analysis of the relationships between the stochastic trajectories of their constituent molecular (or submolecular) species. We have emphasized the importance in this context of conditional independences between the trajectories or time courses of groups of species, represented by $A_{t \leftrightarrow D_i} B_t$. Such an independence means that the information encoded by the trajectory D_i makes any information in A_t irrelevant for the trajectory B_t (and vice versa). That is, A_t and B_t contain no mutual information given the trajectory D_i , and the species in D play the role of informational intermediaries or ‘information carriers’ between A and B within the network.

We describe how to identify such conditional independences between trajectories using a pair of conditions that are straightforward to check in practice. We have demonstrated, it seems for the first time, that entire networks (including very large ones) can be decomposed systematically into modules so that all information relevant to a particular module’s trajectory is encoded by the trajectory of those species in the region where the module overlaps with the rest of the network. In the context of signalling networks with multiple inputs, the approach identifies the routes and species involved in sequential information processing between input and output modules. The dynamic conditional independences, modularizations and I–O paths we identify are themselves robust features of a given network in three senses emphasized recently by Gunawardena [36]—they are invariant to changes in all rate parameter values ($c_m > 0$), in the initial condition (x_0), and in the ‘dynamical’ or reaction intensity functions ($g_m\{\cdot\}$, eqn (1) of the electronic supplementary material, SM1.1). Since the cellular networks under study are naturally evolved, the absence of any need to fine-tune rate parameters to obtain these properties is consistent with their functional importance.

Furthermore, the methods of analysis presented apply to a rather general form of stochastic dynamics for biochemical networks that includes, but goes considerably beyond, the form employed in a Gillespie [34] simulation algorithm or a variant thereof (for a more technical discussion, see the electronic supplementary material, SM1.1). In particular, each conditional reaction intensity may depend on the entire past trajectory of its reactants (e.g. on the time elapsed since the last jump in some reactant levels) and also on the time evolution of variables taken to be deterministic such as temperature, light level and time itself. The class of stochastic dynamics used can thus accommodate, for example, general memory effects (owing perhaps to inhomogeneity arising from imperfect mixing), non-exponential waiting times, and deterministic circadian or environmental effects.

The MIDIA algorithm is developed which allows automated identification of exact decompositions for large networks, control over the degree of coarse graining, and visualization using a junction tree that encodes the conditional independences. The algorithm relies on construction of the KIG for the network and uses only stoichiometric information. No knowledge of rate parameters or use of simulations are required. A bespoke version designed for signalling networks identifies the process of sequential encoding and visualizes its structure using the collection of I–O paths in the junction tree. Together with the I–O path matrix, this provides a powerful, visual tool for investigating information processing by signalling networks.

The stochastic, dynamic signals received by cells are impounded in the stochastic trajectories of the intermediate species of signalling networks and ultimately in the trajectories of their targets. The use of dynamic conditional independences thus allows us to identify those species trajectories which fully encode the relevant information and to trace the sequential process of information transfer through the network. We have demonstrated using a small $\text{NF}\kappa\text{B}$ signalling network that a rigorous framework is indispensable for reasoning correctly about informational properties.

Application of the techniques to the TLR network establishes both their utility for studying the properties of cell signalling and their computational feasibility for large reaction networks. Joint inspection of the junction tree and the I–O path matrix enables a comparative analysis of different I–O combinations. We find that encoding in the case of the interferon response outputs is much sparser, while the AP1 and NADPH-oxidase outputs share more similar I–O paths. For a large cluster of 12 TLRs, we identify a surprisingly small group of species responsible for mediating information transfer from the ‘core’ encoding species to the interferon response outputs. Overall, the information processing structure is not well described as a bow tie owing to the breadth of encoding species away from the periphery. Our analysis reveals that, in the context of multiple stimuli, inputs can be informative in ways unanticipated by steady-state analyses.

Stoichiometric approaches to network analysis have had a large impact on systems biology, owing in part to the modest knowledge required about reaction

parameters. The dynamic conditional independence-based techniques introduced here provide a modular approach for analysing information encoding and transfer by biochemical networks, while also requiring only stoichiometric information. Importantly, the approach fully incorporates both stochastic and non-steady-state dynamics. We believe it constitutes a significant step in the development of a quantitative theory of cellular information processing.

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