Comprehensive Logic Based Analyses of Toll-Like Receptor 4 Signal Transduction Pathway

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Abstract

Among the 13 TLRs in the vertebrate systems, only TLR4 utilizes both Myeloid differentiation factor 88 (MyD88) and Toll/Interleukin-1 receptor (TIR)-domain-containing adapter interferon-β-inducing Factor (TRIF) adaptors to transduce signals triggering host-protective immune responses. Earlier studies on the pathway combined various experimental data in the form of one comprehensive map of TLR signaling. But in the absence of adequate kinetic parameters quantitative mathematical models that reveal emerging systems level properties and dynamic inter-regulation among the kinases/phosphatases of the TLR4 network are not yet available. So, here we used reaction stoichiometry-based and parameter independent logical modeling formalism to build the TLR4 signaling network model that captured the feedback regulations, interdependencies between signaling kinases and phosphatases and the outcome of simulated infections. The analyses of the TLR4 signaling network revealed 360 feedback loops, 157 negative and 203 positive; of which, 334 loops had the phosphatase PP1 as an essential component. The network elements’ interdependency (positive or negative dependencies) in perturbation conditions such as the phosphatase knockout conditions revealed interdependencies between the dual-specific phosphatases MKP-1 and MKP-3 and the kinases in MAPK modules and the role of PP2A in the auto-regulation of Calmodulin kinase-II. Our simulations under the specific kinase or phosphatase gene-deficiency or inhibition conditions and phosphorylations of the elements in the model is extremely sparse; nonetheless, as large as of TLR4 knowledge of kinetic parameters/concentrations of the elements in the model is extremely sparse; nonetheless, the interaction details among the elements of the TLR4 network are relatively well characterized. Such knowledge of reaction stoichiometry was used for building qualitative Boolean logic based models independent of kinetic parameters or concentration of the network components [6]. By adopting such logical modeling formalism, which utilizes the available molecular interaction details and reaction stoichiometry of the network to convert the mental information was converted to logical connections [7]. The further updated with recent experimental findings. The experimental reports. The preliminary information was gathered from the comprehensive map of TLR signaling [8] which we utilized information derived from hundreds of independent experimental reports. The preliminary information was gathered from the comprehensive map of TLR signaling [8] which we further updated with recent experimental findings. The experimental information was converted to logical connections [7]. The model identified a total of 360 operational feedback loops of which most of the loops (positive or negative) owe their origin to four phosphatases, MKP-1, MKP-3, PP1 and PP2A. We calculated the relative contribution of each phosphatase in determining the total number of feedback loops in the system by systematically carrying

Introduction

Toll-Like Receptors (TLRs), thirteen in vertebrates, are the members of pattern recognition receptor family (PRRs) and recognize the pathogen-associated molecular patterns (PAMPs) [1,2]. Upon ligand binding TLR signal is processed in a Myd88-dependent and MyD88-independent (TRIF dependent) manner [3], where MyD88 and TRIF are the adaptor molecules differentially recruited to the TLRs. Among all the TLRs, only TLR4 utilizes both Myd88-dependent and TRIF-dependent pathways [3]. This makes TLR4 signal processing a relatively complex process as compared to other TLR family receptors.

In such complex biological systems of signal processing and integration, understanding the inter-regulation between the signaling elements by computational reconstruction of the signaling networks enables one to systematically investigate the regulatory principles associated with the network and thus to build hypothesis that can subsequently be tested through experiments. For capturing the quantitative system level properties of signaling network dynamic modeling approaches [4,5] are best suited. But for signaling networks

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Abstract

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Introduction

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out several in-silico knockout studies and subsequently analyzing the changes in global dependency (positive/negative/neutral) among the network elements. The model was useful in extracting previously unreported/less emphasized paths of signal propagation for providing plausible explanations to paradoxical experimental results. For example, the counteractive roles for ERK-1/2 in the augmentation or inhibition of IL-10 could be explained by extracting novel signaling pathways from the model. We tested the predictive power of the model by successfully simulating effects of various experimentally reported knockout conditions. Finally, we subjected the model to in silico perturbations that mimic the effect of infections by Escherichia coli and Torrisonia pests. The model at its current state is open to experimental tests and thus further fine-tuning; as it was observed that training such logical models with high throughput experimental data remarkably increases its predictive power [9].

**Results**

**Constructing the logical model of TLR4 signaling**

We started the model building by extracting the information on the TLR4-specific interactions from the comprehensive map of TLR signaling [8]; newer information (published later, after [8]) was subsequently incorporated in the model (Table S1). All the reactions were next converted to logical connections (hypergraphs), creating 181 species (nodes) and 263 reactions (edges) based on Boolean logic functions AND, OR, NOT and their combinations [6,10] (Figure 1). The logical functions were implemented in accordance to the nature of biological connections between the network elements using CellNetAnalyzer [10]. Table S2 and Table S3 show the logical reactions and species, respectively, involved in the TLR4 signaling. The nodes in signaling network are kinases, phosphatases, adapter molecules, transcription factors, cytokines etc. The edges are the unidirectional hypergraphs, which represent the direction of signal flow.

In the logical model the state 1 and 0 represents on (active) and off (inactive) state, respectively.

In the simplest kind of interaction where an upstream directly activates its downstream, activation state 1 of the upstream activator is transferred to the downstream, representing the signal flow. Consider for example, a set of molecules X, Y and Z; 1] if activation of Z requires activation of both X and Y simultaneously, then the logic gate AND connects A, B to C as: X AND Y = Z; 2] If Z is activated when any of its upstream X or Y is active, then the Boolean gate OR connects X, Y, Z as: X OR Y = Z; 3] If Z is active only when X is active but Y is NOT active then the NOT gate connects the elements as X+NOT Y = Z. Figure 2 (methods section) shows a set of representative molecules from the constructed TLR4 network whose biological interactions required representation utilizing AND, OR and NOT gates.

The TLR4 model comprised of 40 input nodes. These inputs correspond to TLR4 receptor; TLR4 ligand, co-receptor CD14 and lipopolysaccharide binding protein (LBP) and the rest 36 are the intermediates for which direct activators or inhibitors are not know clearly. Table S2 shows the list of all input nodes used in the model. The intermediate nodes carry the signal from the input layer which mostly comprises components of various evolutionarily conserved kinase-signaling pathways. Complexity of the signaling increases as the signal goes down from the membrane to the cytoplasm and then to the nucleus as more and more intermediate signaling components is activated by the incoming signal. Finally, the signal converges to output elements comprising various pro- and anti-inflammatory cytokines, transcription factors and various target genes, trigger of cell division, transcription factors, apoptosis and regulation of the RNA metabolism.

In biological system activation of signaling elements in a pathway is temporal in nature which is determined by the hierarchy of signal flow from membrane to nucleus. To mimic the dynamicity (or pseudo dynamics) of signal flow like a real system we have opted time scale scenario formulation by selectively assigning different time scale to different signaling elements in a hierarchical manner [11]. We implemented time scales as below:

**Time scale 0:** Formation of ligand receptor complex and activation of housekeeping species.

**Time scale 2:** Activation of the all-cytoplasmic reactions taking place after recruitment of ligand to the receptor.

**Time scale 4:** Activation of reactions corresponding to the nuclear translocation of specific cytoplasmic kinases that leads to triggering of transcription factors or activation of other target molecules/genes.

**Time scale 7:** In this time scale signaling events in the nucleus that are triggered by the MyD88 dependent pathways were activated. List of the reactions involved in the MyD88 dependent pathway is given in Table S2.

**Time scale 7.1:** Activation of the TRIF dependent pathway was assumed to occur at later as suggested by experimental studies [12]. List of reaction involved in the TRIF dependent pathway is given in Table S2.

**Time scale 8:** Finally in the time scale 8 the cytoplasmic and nuclear phosphatases and inhibitors of the TLR4 signaling were activated. This leads to complete shutdown of information flow in the system.

In the in-vivo settings, the cytoplasmic/nuclear phosphatases can be activated earlier to the time required for the signal to reach the nucleus or they remain constitutively active causing an attenuation/suppression of the signal strength, thus imparting dynamically achieved finer regulations [4,5,11,13], so activation of phosphatases attenuates the signal flow in most cases but may not always abolish it. However, owing to the nature of our modeling approach (output is either 0 or 1), such finer modulations of the processed signal cannot be captured [7,14]. Nonetheless using the logical models analyses can be performed on various structural aspects of regulations in the signaling network. Here we focused our analysis on understanding role of the phosphatases in regulating the global interdependencies among the elements of the TLR4 signaling network.

**TLR4 signaling network contains a large network of positive and negative feedback loops**

Feedback loops are pivotal for the functioning of almost all the signal transduction networks. Various dynamic analyses of signaling systems show that positive feedback loops lead to signal amplification, multi-stability and hysteresis or tolerance [15,16] whereas negative feedback loops trigger signal attenuation, adaptation, oscillations or delay in activation [17,18]. From the large signaling systems with information on reaction stoichiometry, logical analysis can identify the feedback loops and their nature (positive or negative). A feedback loop- whereby a signal from a signaling intermediate “I” flows through the intermediate “L” and comes back to influence the activity of “I”- is an interaction graph-based property of the model [6]. So, during the calculations, the number of times such conditions are satisfied yielded the number of feedback loops in the system. An important element in the feedback loops are the phosphatase of the network [14]. Our simulations identify 360 feedback loops, of which 157 were negative and 203 were positive feedback loops. Most of these feedback loops emerged from the kinase-phospha-
tase interactions in the network. We explored the role of phosphatases involved in TLR4 signaling and calculated their contribution in the total feedback loops of the network. Figure 3 shows the schematics of relative contribution of different phosphatases, alone and in combinations in determining the total number of feedback loops in the system. To decipher the relative contribution of various phosphatases we systematically knocked out a target phosphatase and calculated the remaining feedback loops for each of the knockout conditions (Figure 3). The phosphatase PP1, as we found from the in-silico knock down, is the most contributory phosphatase in determining the total number of feedback loops (Figure 3) as ~93% (334 out of 360) feedback loops required PP1 as an essential component.

When all the phosphatases of the system were knocked out, 25 feedback loops remained. These 25 feedback loops owed their origin purely to kinase-mediated interactions of which 5 were negative and 20 were positive feedback loops. The interactions that gave rise to the 25 feedbacks were next extracted and shown; 11 reactions were found to comprise the 5 negative feedback loops and 31 reactions comprised the 20 positive feedback loops, respectively (Table S4). Relative contribution of each of the reactions in the 25 feedback loops was also calculated and shown (Table S4). Notably, 4 out of 5 negative feedback loops comprised NF-κB as an essential component and Vav1 was found to be essential element in more than 60% of the 20 positive feedback loops.

Interdependency among the network elements: wild type and phosphatases knockout conditions

As interdependencies among the signaling intermediates characterizes cell signaling [9,19], any local changes imposed in the system can possibly lead to global alteration of interdependencies among the remaining elements in the network in varying degrees[9,11]. Experimental investigations and hypothesis testing commonly involve targeted knockout studies [20–22] although systems level implications of such knockout studies are less understood. As our analyses suggest (Figure 3), knockout of different phosphatases do change the number of feedback loops differentially, indicating that the interdependencies among the network elements plausibly changes with each perturbation.

Typically in large-scale networks a remote downstream kinase could be an activator/inhibitor of an upstream molecule and the process of activation/inhibition comprises multiple intermediates [17,18]. Here positive or negative influence of a network element on the rest of the network elements is calculated using the dependency matrices [6,10]. For example, a species X can influence any other species Y of a network in various ways: X could be a total activator (TA), total inhibitor (TI), one of the inhibitors (I) or one of the activators (A). Also species X can have no influence (N), or have both positive and negative influence (U, called as ambivalent factor) on species Y, all of which could be calculated using the dependency matrix [6].

The model of the TLR4 signaling network contained 181 species, so the size of the dependency matrix is 181×181; making its visual comprehension and representation inconvenient. We
Figure 2. Conversion of reactions from the comprehensive map to the logical reactions. Set of representative interactions extracted from comprehensive map of TLR signaling and represented as interaction graph and interaction hypergraph.
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Figure 3. Relative participation of phosphatases in feedback loops. Figures show number of remaining feedback loops in different phosphatases knockout conditions.
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thus selected a subset of the total elements of the complete dependency matrix comprising molecules such as the MAP kinases (ERK, p38MAPK, JNK), Akt, NIK, IKK and CAMKII which are representative of distinct signaling pathways activated during TLR4 signaling, together with the phosphatases MKP-1, MKP-3, PP1, PP2A. The chosen kinases are also pivotal biologically, as they carry out growth, differentiation, proliferation and are involved in several diseased conditions [23–28].

Effects of various in-silico phosphatases’ knockout and resultant alterations of the positive/negative dependencies among species are detailed (Table S5). The complete dependency matrices for wild type and various knockout conditions are also shown (Supporting information). A total of 16 conditions were simulated: wild type, MKP-3+, PP2A−/−, PP1−/−, MKP-1−/−, (MKP-3+ MKP-1)−/−, (MKP-3+PP2A)−/−, (MKP-3+PP1)−/−, (MKP-1+ PP2A)−/−, (MKP-1+PP1)−/−, (PP1+PP2A)−/−, (MKP-3+MKP-1+PP2A)−/−, (MKP-3+PP2A+PP1)−/−, (MKP-1+PP2A+PP1)−/− and (MKP-3+MKP-1+PP2A+PP1)−/−. Several interesting behaviors were observed in the knockout conditions all of which cannot be discussed due to the volume of the information. So, we state some interesting regulatory behaviors exhibited by the system in different knockout conditions that can be easily tested experimentally.

> ERK-1/2 acted as an ambivalent factor of MEKK3 and IKK in the wild type condition but in the MKP-3 knock out condition, ERK-1/2 become an activator of MEKK3 and IKK; this suggests that the negative effect from ERK-1/2 to MEKK3 and IKK is mediated in an MKP-3 dependent manner. In addition, in the MKP-3 knockout condition ERK-1/2 becomes an inhibitor of PP2A and p38MAPK whereas in the wild type condition ERK-1/2 is ambivalent to both. ERK-1/2 becomes an activator of CAMKII and inhibitor of Akt in MKP-3 knock out condition whereas in wild type condition ERK-1/2 is ambivalent to both (Figure 4A–E).

> When PP1 is knockout MEKK3 becomes an activator of PP2A and Akt and inhibitor of itself, IKK and CAMKII respectively, whereas in the wild type condition its influence is ambivalent for all of these kinases (Figure 5C). An experimentally testable hypothesis from this the analysis would be: MEKK3 inhibits PP2A and Akt but activates itself, IKK and CAMKII (Figure 5A–E) in the wild type setting which could be tested through perturbation studies like overexpression and/or siRNA mediated inhibition of MEKK3 and compare the effect of such perturbation on these kinases with their respective wild type counterparts.

> In PP2A knockout conditions, CAMKII becomes its own inhibitor and its influence on Akt is completely abolished (N) whereas in the wild type condition CAMKII ambivalently affects itself and Akt. Thus the positive self-regulatory loop of CAMKII must have PP2A as an essential element. Also PP2A has a dual role (positive and negative) in CAMKII mediated regulation of Akt in the wild type condition. Further, knockout studies in the model show that the positive loop from CAMKII to Akt is abolished when MKP-1+PP1 is knocked out (Figure 6A–E).

> All the signaling paths from MKP-1 to MEKK3, CAMKII and Akt have PP2A as an essential element as in the PP2A knockout condition MKP-1 is no longer connected to these kinases (Figure 7A–C). Such coherent regulation between the phosphatases can also be tested experimentally by systematic knock down studies.

Coexisting signaling paths show plausible positive and negative pathways for ERK-1/2 mediated IL-10 production

Independent experiments show that in TLR4 signaling ERK-1/2 regulates IL-10 production both positively [29,30] and negatively [31]. As analysis of logical models show that complex connectivity among the elements of large networks lead to ambivalent effect [14], we explored whether both positive and negative signaling paths between ERK-1/2 and IL-10 potentially exists in the wild type condition. Due to the overwhelming complexity of connections in the network the total number of signaling paths from ERK-1/2 leading to IL-10 were found to be 2500, out of which exactly 50% (1254) were positive and the rest 50% were negative. Notably, all of these signaling paths contained PP2A and MKP-3 as essential components, because knockout of either of these phosphatases resulted in 0 signaling paths between ERK-1/2 and IL-10. Figure 8 shows the shortest paths that connect ERK-1/2 to IL-10 both positively and negatively. Similarly, we extracted a pathway that positively connects ERK-1/2 to IL-12, which is mediated through NF-κB (Figure 9) [32,33].

Logical steady state analysis and model validation using experimental results

Logical steady state analysis (LSSA) was performed to understand the characteristics of signal flow through the TLR4 network. In LSSA the state 1 means an on state and a state 0 means off state. In LSSA, the state of each model element is consistent with the value of its associated Boolean function, and therefore, once a Boolean network has moved into a logical steady state, it will stop to switch and then retain this state [6]. The signal starting at input node (TLR4 receptor/ligand) traverse the entire span of the network through the series of logical gates and the cumulative result of such logical interactions decide the 0 or 1 state of each element of the network. Logical steady state was calculated at different time scales (biological logic behind the time scales 0, 2, 4, 7, 7.1, 8 is explained in the model building sections and also given in supporting information). For example, we showed the LSS corresponding to time scales 0, 2, 4, 7, 7.1 in the wild type condition and subsequently effect of different knockout/knockin conditions at the time scale 7.1. At time point 7.1 [as explained in methods section] all the elements of the network are activated, hence global effect of perturbations was captured in time scale 7.1 (Table S6). Simulation of the model with the next higher time scale (time scale 8) corresponds to complete shutdown of the signal in the system.

As demonstrated earlier [14], results of LSS can be qualitatively compared to experimental results. Here we imposed various experimentally observed scenarios for TLR4 signaling and compared the simulated outcomes with the experimental reports. Table S7 lists several simulation results that corroborated with the experimental results. When the simulation results matched the experimental results, we extracted the signaling path that links the perturbed species to the affected species. This was done because in experimental studies only the cause (perturbed species) and effect (changes in target species) are observed and the signaling path/information processing route that connects both the species are often considered as a black box. For example, it was recently reported that in the TRADD−/− mice phosphorylation of the MAPKs’ (p38MAPK, ERK-1/2, Jnk) and I-kb is reduced [34]; this is qualitatively observed in our model as 0 activation states of the MAPK’s and I-kb when TRADD “0. We found 6 positive signaling paths that connect TRADD to MAPK’s and 110 signaling path that connects TRADD to I-kb. The signaling paths unravel the plausible intermediates involved in transmitting the signal from TRADD to the affected molecules, which is open to experimental validation.
Therapeutic utility of the model: Pathogen specific perturbations in the logical network and proposition of drug targets

Several pathogens are known to perturb TLR4 signaling during their host invasion. We tested two cases of infection; infection by uropathogenic E.coli and Yersinia pestis and investigated whether the model predictions fall in the same lines as the experimental reports on the same.

**Yersinia pestis infection.** *Yersinia pestis* is a Gram-negative bacterium that causes bubonic plague. YopJ protein found in *Yersinia* inhibits MKK6 and IKK by acetylation [35]. YopJ is also shown to inhibit the TRAF6 and TRAF3 by deubiquitination [36]. Yersinia YopJ protein also inhibits MKK6 and IKK [37]. We computed the LSS for such pathogenic conditions by assigning zero value to the YopJ affected molecules (MKK6, IKK, TRAF6 and TRAF3). LSS values of different species in the condition of infection are shown (File S1). LSS shows that pro-inflammatory cytokines production is inhibited while anti-inflammatory cytokine IL-10 remained uninhibited which was also observed experimentally [35,37]. *Yersinia* plausibly uses this strategy to overcome the defense mechanism of the immune system by selective inhibition of the pro-inflammatory cytokines’ production and allowing only the production of anti-inflammatory cytokine such as IL-10. IL-10 production in the infection condition is ERK1.2 dependent, which is activated through a path comprising Rac.GTPase. To alter the infection condition one strategy would be to suppress the ERK-1/2 activation which in our model would mean ERK-1/2 = 0. We thus calculated the MIS (Minimum Intervention Set [6], see methods section) for IL-10 production by assigning the MIS goal to be ERK1.2 = 0, while keeping MKK6, IKK, TRAF6 and TRAF3 = 0 such that the effect of *Yersinia* infection is considered as starting condition in the model. Table 1 shows a set of molecules that are calculated as MIS suitable for inhibiting IL-10 production under *Yersinia* infection.

**Uropathogenic E.coli infection.** Strains of *Uropathogenic E.coli* are responsible for the urinary tract infection [38]. They secrete Tcpc protein containing the TIR like domain that binds to Myd88 making it functionally ineffective [39]. In the model we have implemented the condition by assigning zero value to Myd88 and then simulated the model to calculate the LSS (File S1). Experiments show that during *Uropathogenic E.coli* infection TNF-α production is abrogated [37,38]. Simulation of the model with Myd88^−/− condition shows that TNF-α production is zero, even when the TRIF dependent pathways were considered activated. We performed MIS calculation by setting a MIS goal of reactivating TNF-α in a Myd88 independent manner. We found 363 MIS sets for this goal. Set of molecules whose coupled activation/inhibition would result in reactivation of TNF-α is given in Table S8.

Figure 4. Influence of ERK1/2 on various kinases in wild type and MKP3 knockout condition. A. ERK1/2 regulates MEKK3 both positively as well as negatively in wild type condition. In the MKP3 knockout condition the regulation of MEKK3 by ERK1/2 is only positive; similarly the effect of the ERK1/2 on IKK (B), p38MAPK (C), Akt (D) and CAMKII (E), in wild type and MKP3 knockout condition is shown. Black arrow represents positive and red arrow represents negative influence respectively.

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Discussion

Here, we present the logical model of TLR4 signaling, comprising 181 nodes and 263 edges, which is to the best of our knowledge the largest logical model of signaling network built till date. Despite having incredibly complex interactions among the molecules of the network it was possible to extract several novel regulatory properties that owe their origin to stoichiometry of interactions and their deduced Boolean relations based on such stoichiometry matrix. Implementation of logical modeling methodology for large scale signaling networks is a relatively recent advancement in systems biology [6]. Till date, the prominent logic based models of large scale signal transduction are- T cell signaling network with 94 nodes and 123 interactions [7]. Apoptosis signaling network with 86 nodes and 125 interactions [14], EGFR/ErbB signaling network with 104 nodes and 204 interactions [9]. All these models showed that logic based modeling can be of immense help in extracting a systems regulatory design stemming from the topology of interaction among its components.

TLR4 signaling network comprises at least 181 nodes and 263 interactions, making it one of the most complex signal processing systems to be subjected to quantitative modeling. In an earlier attempt, semi-quantitative model of TLR4 signaling was constructed where effect of controlled perturbation of input on the

Figure 5. Auto regulation of MEKK3 and its influence on different kinases/phosphatase in wild type and knockout conditions.
Influence of MEKK3 on PP2A (A), Akt (B), MEKK3 (C), CAMKII (D) and IKK (E), in wild type and PP1 knockout condition is shown. Black arrow shows positive and red arrow represents the inhibitory influences respectively. doi:10.1371/journal.pone.0092481.g005

Figure 6. Self-regulation of CAMKII.
(A) CAMKII and Akt (B) in the wild type and PP2A knockout out condition are shown. Black arrow shows positive influence; red arrow shows the inhibitory effect and cross mark shows no effect. doi:10.1371/journal.pone.0092481.g006
system’s output was modeled [40], capturing properties of the network like redistribution of signal flow between the MyD88 and TRAM-dependent pathways upon inhibition of either of them. However, some of the crucial regulatory aspects of the TLR4 signaling network such as identifying the feedback loops within the network, multi-species interdependencies in coherently determining the output of the system was not addressed till now.

We built the logic based model of TLR4 signaling and largely focused our analysis on understanding the role of phosphatases in determining the global emergence of feedback loops and interdependencies among the network elements. It was previously shown that phosphatases play crucial roles in the emergence of feedback loops in signaling pathways [41–45]. The in-silico studies performed here using knockout of various phosphatases indicate that phosphatases orchestrate a complex network of feedback loops, implying interdependencies among the phosphatases themselves; indeed, in a smaller network of MAPK signaling in B cells, the phosphatases’ interdependency in regulating kinases’ phosphorylation has been already implicated [45]. Because the phosphatases work in unison in the wild type system, it is hard to decipher the contribution of an individual phosphatase in influencing the interdependency among kinases in the network. We found that systematic knock out of one or multiple phosphatases in combinations differentially altered the interdependency between phosphatases, between kinases and between kinases and phosphatases. We systematically compared the effect of various phosphatase knockout conditions and studied the change in regulations among the pair of network elements whose positive or negative interaction is known for the wild type conditions. We show that the nature of regulations among various elements of the network differentially changes as a result of a phosphatase knockout. Although dynamic models and model-based experimental investigations [5,15,16,45,46] revealed the regulatory mechanisms underlying various experimentally observed phenomena in signaling pathways, global reorganization of the interdependency between network elements in a specific knock out condition remains poorly understood in general. Our analysis of the in silico knockout conditions presented several experimentally testable hypotheses that aim to elucidate the structure based design principles in the TLR4 signal in a finer way.

We also showed the advantage of logical model formalism in extracting the hitherto unknown signaling pathways between the network elements. For example, the experimentally observed counter-regulation of TLR4-induced ERK-1/2-mediated IL-10 production [29–31] is now shown to be due to the positive and negative signaling paths from ERK-1/2 and IL-10, dominance of either type of path could thus result from cell type specific or context specific expression/activation of intermediates connecting ERK-1/2 and IL-10. However the nature of information flow here restricts one from inferring the relative dominance of either of
positive or negative connections between ERK-1/2 and IL-10. This is again because of the definition of on state (\(= 1\)) and off state (\(= 0\)) of a species, as once ERK-1/2 attains an on state it will transmit the information to IL-10 through intermediates irrespective of wiring of signaling intermediates upstream to ERK-1/2. For example when we simulate a system similar (but not identical, as molecular connections may vary in individual pathways) to TLR9 network which signals through MyD88 but not through TRIF or a condition similar to TLR3 which signals through TRIF but not through MyD88 respectively, then in both cases we found that ERK = 1 and also both positive and negative paths exist. This doesn’t allow us to distinguish TLR receptor specific nature of wiring between ERK-1/2 and IL-10. Under such conditions one can consider building TLR receptor specific models, conduct high throughput experiments, train the models with the high throughput data [9] and then compute the signaling paths.

Finally, we tested the model’s ability to reproduce experimental results and demonstrated its potential therapeutic utility. Several knockout conditions were implemented in the model and simulations considering such perturbations corroborated with the experiments. Similarly, we inhibited (\(= 0\)) the network elements that correspond to parasitic infections where our simulations reproduced the experimentally reported inhibition of cytokines. The therapeutic utility of the model was examined by predicting the potential drug targets through MIS analysis. In addition, SHP1-mediated regulation of TLR4 signaling was simulated by adding three additional reactions in our model as SHP1 inhibits TLR4-mediated IRAK1 activation [47,48] regulating the production of various cytokines. But we did not find any changes in the feedback loop mediated regulation of TLR4 signaling (data not shown) as SHP-1 acts in the receptor level [47] and it fulfills the definition of an inhibitor but not of a feedback element.

Taken together, this study helps in understanding the structural organization based working principles governing the whole cell level TLR4 signaling and poses a number of conjectures that can be experimentally verified. As proposed earlier [6,9], the model can further be trained with high throughput experimental data; that will contribute towards enhancing the predictive power of the model.

Materials and Methods

For qualitative analysis of the TLR4 signal transduction pathway, we have used a Boolean logic based modeling approaches for modeling signaling networks, implemented in the MATLAB toolbox CellNetAnalyzer [6]. The first step in the construction of the model was extraction of the information from the comprehensive map of the TLR signaling [8], followed by incorporation of newer information from literature [Table S1]. The information extracted from the comprehensive map of the TLR was converted into the “Logical Interaction Hypergraph” (LIH) [6]. In the logical interaction hypergraph the interaction among the species are the combination of the three simple Boolean functions AND, OR and NOT gates or their case specific linear combinations. Figure 2 shows the conversion of stoichiometric knowledge to 4 different types of Boolean relation [AND, OR, NOT gates], with a representative set of reactions from the total of 263 reactions of the network.
For example, TLR4 receptor and ligand binding steps are represented by two interaction graphs (Figure 2; 2nd row, 2nd column). Biologically it is possible to form a ligand receptor complex [TLR4.L] only when both ligand and receptor come together. Using the Boolean gate AND (Figure 2; 2nd row, 3rd column) we could implement such biological relations where the function AND allows the TLR4.L to be active (state = 1) only when TLR4 = 1 and LIGAND = 1.

On the contrary, there are situations where any one activator (of several activators) is sufficient for triggering activation of a substrate. For example Mnk1 can be phosphorylated either by ERK-1/2 or p38MAPK. So (Figure 2; 2nd row, 3rd column) we could implement such biological relations where the function OR gate captures the biological relation between these two activators. When any one of ERK-1/2 or p38MAPK = 1, the activation state of Mnk1 = 1.

In another scenario, elements like Elk1 requires p38MAPK = 1 and PP2B = 0 for its activation state to become 1 (Figure 2; 5th row, 1st column). Boolean representation of such situations is shown (Figure 2; 4th row, 3rd column) where AND NOT gate captured the biological relation between these three species.

After the conversion of such molecular interactions into the logical hypergraphs, the network was constructed for its pictorial representation using the software CellDesigner [49]. The picture was exported from CellDesigner to CellNetAnalyzer for analysis.

### Table 1. MISs for inhibiting production of the IL-10 in *Yersinia* infection.

| S. No. | Species |
|-------|---------|
| 1     | IKK_a,b,g |
| 2     | c-Fos |
| 3     | !TPL2 |
| 4     | NF-k_b_p50 |
| 5     | TTP |
| 6     | !ERK1.2 |
| 7     | PP-2A,B |
| 8     | PP2A |
| 9     | !IL-10 |
| 10    | MKP3 |
| 11    | !MKK1 |
| 12    | !ERK1.2_Nu |
| 13    | c-Fos_P |
| 14    | NF-kb.p50 IkBa IkBb |
| 15    | ESCIT NF-kb.p50 IkBb |
| 16    | NF-kb.p50 IkBb TRAF3 |
| 17    | NF-kb.p50 IkBb TBK1 |
| 18    | MEKK1 NF-kb.p50 IkBb |
| 19    | !LIGAND Ubc13.Uev1A.TIFA.TRAF6.IRAK1 NF-kb.p50 IkBb |
| 20    | !LIGAND Ubc13.Uev1A.TIFA.TRAF6.IRAK1* NF-kb.p50 IkBb |
| 21    | Ubc13.Uev1A.TIFA.TRAF6.IRAK1 !MKK3 NF-kb.p50 IkBb |
| 22    | Ubc13.Uev1A.TIFA.TRAF6.IRAK1* !MKK3 NF-kb.p50 IkBb |
| 23    | Ubc13.Uev1A.TIFA.TRAF6.IRAK1 !MAPKAPK2 NF-kb.p50 IkBb |
| 24    | Ubc13.Uev1A.TIFA.TRAF6.IRAK1* !MAPKAPK2 NF-kb.p50 IkBb |

! Denote the permanent inactivation of the species and the species without ! sign are species which should be permanently activated for fulfilling target goal.

* Shows activated form of the IRAK1 complex.

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**Computation of the feedback loops in phosphatases knockout condition**

The signaling pathways activated through the TLR4 involved the activation of the various phosphatases of the system. Role of these phosphatases in regulation of feedback loops is calculated in the 16 different knockout situations. To knockout a phosphatase from the system, it was excluded during the feedback calculations by assigning it a fixed value = 0.

**Computation of the dependency matrix**

The dependency matrix of the full-scale network is extremely large (181x181) making the analysis process complicated. However the information of influence of one species on rest of the model elements could be extracted from the dependency matrix using CNA [6]. Thus, we selected a set of kinases; representatives of various pathways activated during the TLR4 signaling and analyzed their interdependencies in the knockout conditions of the phosphatases.

**Computation of minimal intervention set (MIS)**

Minimal intervention set (MIS) in an interaction networks is the (minimal) sets of elements that are to be removed or to be added in order to achieve a certain goal such as activation or inhibition of a target molecule [6]. MISs were calculated in infection condition to abrogate IL-10 production by assigning 0 value to IL-10 node and
declaring the fixed activation state for TLR4, Ligand and other nodes that are required for receptor-ligand complex formation. To minimize the computation time, flag was set for the nodes that were not to be included during MIS calculation, like receptor and ligand.

Logical Steady State Analysis
Logical steady state shows the activation state of a set of molecules on assigned time scale for given input signal. For computation of the logical steady state in a particular situation, for example, in a Myd88 knockout situation, values = 1 or 0 were assigned to the respective species and then the LSS were computed at time scale 7.1. Time scale 7.1 was selected for analyzing the effect of a knockout or addition of inhibitors in the system because at time scale 7.1 the signal was allowed to flow through the entire span of the network. During the calculation of the LSS few interactions were not considered. For example, p38MAPK and TAK.TAB1 which are activated at the same time scale are connected though a NOT gate, blocking the downstream flow of signal. As the p38aMAPK activation and the TAK.TAB1 activation occur in the time scale 2, and as p38aMAPK inhibits the TAK.TAB1, calculation of LSS at any further time scale above 2 calculates the downstream signaling effect considering TAK.-TAB1 = 0. Hence the LSS calculations were carried out without including the feedback loop from p38MAPK to TAK.TAB1. A total of five interactions (out of total 263 interactions) were removed during the calculation of logical steady state(shown in Table S9).

Requirement of two models for interaction graph and hypergraphs based analysis
As discussed above in the results sections, activation of inhibitory loops on the same time scale would result in the blockage of the signal through the network, thus the inhibitory loops must be removed to allow the signal flow through the network. However, removal of the interactions also results in the change in the topological properties of the network such as the number of feedback loops. To overcome this problem two models were built, one model for the logical steady state analysis where five reactions were not considered (mentioned above). The other model was used for analysis of topological properties such as feedback loops or the interdependencies among the species which contains all the interactions. Nevertheless, simulated logical steady states devoid of the five reactions corroborated with the experimental reports suggesting that the signal flow in the model is preserved in general and it can be used for making biologically testable predictions.

Model validation using experimentally reported scenarios
We implemented results of various experiments in the logical model aiming to validate its predictive power. For example, experimental data shows that in Myd88 knockout mice production of inflammatory cytokine is down regulated. We knocked out Myd88 in the model by assigning it a 0 value and computed the LSS at time scale 7.1 which qualitatively reproduced the experimental observations. Several such examples are shown.

Supporting Information
Figure S1 (A–Q) Dependency matrix in different conditions of phosphatases knockout. In addition, screenshot of the model LSS for E.coli and Yersinia infection is shown. (PDF)

Table S1 Supplementary table S1 shows the list of all reaction involved in the TLR4 signal transduction. (PDF)

Table S2 List of all species of the TLR4 logical model. (XLS)

Table S3 Relative participation of different reactions in positive and negative feedback loops in (MKP3+MKP1+PP2A+PP1) knockout condition. (XLS)

Table S4 Dependency matrix showing influence of a particular kinase (in the vertical axis of the matrix) on other kinases (elements of different rows in the matrix), in different conditions of phosphatases knockout alone or in combinations. (XLS)

Table S5 Logical steady state of all output species on different time scale and in knock in condition of individual inhibitor in a full activated system (at time scale 7.1). (XLS)

Table S6 Logical steady state of the output species in infection condition of Yersinia and Uropathogenic E.coli. (XLS)

Table S7 Validation of Model through comparison of Logical Steady State Analysis (LSSA) results to various experimentally reported results. (DOC)

Table S8 MISs for the production of the TNF-α in the E.coli infection condition. (XLS)

Table S9 List of the reactions, which are eliminated during the calculation of the logical steady state analysis of model. (XLSX)

Model S1 Logical models of the TLR4 signal transduction pathway is presented. The file contains two folders. The model TLR4_LSSA Complete was used for the feedback loop analysis. TLR4_LSSA was used for the logical steady state analysis of the TLR4 signal transduction pathway. The MATLAB toolbox CellNetAnalyzer was used for both the analysis. (RAR)

Author Contributions
Conceived and designed the experiments: MP US. Performed the experiments: MP US. Analyzed the data: MP US. Contributed reagents/materials/analysis tools: MP US. Wrote the paper: MP US BS.

References
1. Akira S, Uematsu S, Takeuchi O (2006) Pathogen recognition and innate immunity. Cell 124: 783–801.
2. Takeuchi K, Akira S (2005) Toll-like receptors in innate immunity. Int Immunol 17:1–14.
3. Kawai T, Akira S (2010) The role of pattern-recognition receptors in innate immunity: update on Toll-like receptors. Nat Immunol 11:373–384.
4. Bhalla U, Iyengar R (1999) Emergent Properties of Networks of Biological Signaling Pathways. Science 283: 381–387.
5. Huang C-YF (1996) Ultrasensitivity in the mitogen-activated protein kinase cascade. Proc Natl Acad Sci 93: 10078–10083.

6. Klamt S, Saez-Rodriguez J, Lindquist JA, Simeoni L, Gilles ED (2006) A methodology for the structural and functional analysis of signaling and regulatory networks. BMC Bioinformatics 7: 56.

7. Saez-Rodriguez J, Simeoni L, Lindquist JA, Hemmercy R, Bommhardt U, et al. (2007) A logical model provides insights into T cell receptor signaling. PLoS Comput Biol 3: e163.

8. Oda K, Kitano H (2006) A comprehensive map of the toll-like receptor signaling network. Mol Syst Biol 2: 2006.0015.

9. Samaga R, Saez-Rodriguez J, Alexopoulos LG, Sorger PK, Klapproth S (2009) The logic of EGFR/Erbb signaling: theoretical properties and analysis of high-throughput data. PLoS Comput Biol 5: e1000436.

10. Klammt S, Saez-Rodriguez J, Gilles ED (2007) Structural and functional analysis of cellular networks with CellNetAnalyzer. BMC Syst Biol 1: 2.

11. Werlen G, Hausmann B, Naeher D, Palmer E (2003) Signaling life and death in the thymus: timing is everything. Science 299:1859–1863.

12. Kagan JC, Sub T, Horng T, Chow A, Akira S, et al. (2008) TRAM couples endocytosis of Toll-like receptor 4 to the induction of interferon-beta. Nat Immunol 9:361–366.

13. Shankaran H, Ippolito DL, Chirdler WB, Rent H, Bollinger N, et al. (2009) Rapid and sustained nuclear-cytoplasmic ERK oscillations induced by epidermal growth factor. Mol Syst Biol 5: 332.

14. Schlatter R, Schmid K, Avolas Vincarza I, Scheurich P, Sauter T, et al. (2009) GNAP controls cAMP production through inhibition of Gs and G0. Proc Natl Acad Sci U S A 107: 21909–21914.

15. Samaga R, Saez-Rodriguez J, Alexopoulos LG, Sorger PK, Klapproth S (2009) The logic of EGFR/Erbb signaling: theoretical properties and analysis of high-throughput data. PLoS Comput Biol 5: e1000436.

16. Kearns JD, Basak S, Werner SL, Huang CS, Hoffmann A (2006) IkappaBepsi-binding protein regulates IkappaBalpha degradation. J Biol Chem 281: 18414–18421.

17. Spiel CE, Buhler A, Schalek R, Simpson J, Schelling M, et al. (2005) Cross talk between ERK and JNK mediates TCR and Src-dependent NF-kappaB activation. Cell 120: 97–108.

18. Kawasaki T, Adachi O, Ogawa T, Takeda K, Akira S (1999) Unresponsiveness of MyD88-deficient mice to endotoxin. Immunity 11:115–122.

19. Shim S-Y, Rath O, Choo S-M, Fee F, McCarron B, et al. (2009) Positive- and negative-feedback regulations coordinate the dynamic behavior of the Ras-Raf-MEK-ERK signal transduction pathway. Journal of cell science 122: 425–435.

20. Saez-Rodriguez J, Alexopoulos LG, Epperlein J, Samaga R, Lauffenburger DA, et al. (2009) Discrete logic modeling as a means to link protein signaling networks with functional analysis of mammalian signal transduction. Mol Syst Biol 5: 331.

21. Miyagi T, Kitano H (2010) CellNetAnalyzer: a process diagram editor for gene-regulatory and biochemical networks. BIOSILICO 1: 159–162.

22. Sarma U, Sareen A, Maiti M, Kamat V, Sudan R, et al. (2012) Modeling and simulation of LPS stimulated JNK signaling network. Mol Syst Biol 2: 2006.0015.

23. Schwartz J, Stobart R, Jr, Wernig M, Mankus K, Schaffner F, et al. (2009) Learning the dynamical behavior of the immunological basal process. J Comput Biol 16: 1199–1221.

24. Chandel NS, Basha C, Lin WJ, Duncan G, Chau H, et al. (2008) Beyond tumor necrosis factor receptor: TRADD signaling in toll-like receptor. PLoS Acad Sci U S A 10: 12429–12434.

25. Mukherjee S, Keitany G, Li Y, Wang Y, Ball HL, et al. (2006) Yersinia YopJ acetylates and inhibits kinase activation by blocking phosphorylation. Science 312: 1211–1214.

26. Keating S (2009) Role of non-degradative ubiquitination in interleukin-1 and toll-like receptor signaling. J Biol Chem 284: 8211–8215.

27. Depaolo RV, Tang F, Kim I, Han M, Levin N, et al. (2008) Toll-like receptor 6 drives differentiation of tolerogenic dendritic cells and contributes to LcRv-mediated plaque pathogenesis. J Cell Sci Microbe 4: 350–361.

28. Ron A (2002) The etiology of urinary tract infection: traditional and emerging pathogens. Dis Mon 48: 149–199.

29. Carić L, Wieser A, Yadav M, Duerst S, Schubert S, et al. (2008) Subversion of Toll-like receptor signaling by a unique family of bacterial Toll/interleukin-1 receptor domain-containing proteins. Nature med 14: 399–406.

30. Sutarina J, Takeda Y, Godha J, Helm Y, Akira S, et al. (2008) Signaling flux redistribution at toll-like receptor pathway junctions. PLoS One 3: e3430.

31. Skonieczny P, Tocchetti CG, Skonieczny P, Tocchetti CG, Flato G, et al. (2008) TLR7 activates a gain-in-strength mechanism of MyD88-dependent IFN-beta production by recruiting SHP-1 to a SYK-TLR4-CEACAM1 complex. PLoS Pathog 4:e1002597.

32. Yoshida K, Takeda Y, Yamada S, Iwamoto Y, Akira S, et al. (2008) Phosphatase SHP-1 promotes toll-like receptor-dependent NF-kappaB activation. Nat Immunol 9:361–368.

33. Ron A (2002) The etiology of urinary tract infection: traditional and emerging pathogens. Dis Mon 48: 149–199.

34. Chen N-J, Chio IIC, Lin W-J, Duncan G, Chau H, et al. (2008) Beyond tumor necrosis factor receptor: TRADD signaling in toll-like receptor. PLoS Acad Sci U S A 10: 12429–12434.

35. Mukherjee S, Keitany G, Li Y, Wang Y, Ball HL, et al. (2006) Yersinia YopJ acetylates and inhibits kinase activation by blocking phosphorylation. Science 312: 1211–1214.

36. Keating S (2009) Role of non-degradative ubiquitination in interleukin-1 and toll-like receptor signaling. J Biol Chem 284: 8211–8215.

37. Depaolo RV, Tang F, Kim I, Han M, Levin N, et al. (2008) Toll-like receptor 6 drives differentiation of tolerogenic dendritic cells and contributes to LcRv-mediated plaque pathogenesis. J Cell Sci Microbe 4: 350–361.

38. Ron A (2002) The etiology of urinary tract infection: traditional and emerging pathogens. Dis Mon 48: 149–199.