

# Biological Models of Security for Virus Propagation in Computer Networks

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## **Abstract**

*This article discusses the similarity between propagation of pathogens (viruses and worms) on computer networks with proliferation of pathogens in cellular organisms (organisms with genetic material contained within a membrane encased nucleus). It introduces several biological mechanisms that are used in these organisms to protect against such pathogens and presents security models for networked computers inspired by biological paradigms. Models for computer security can be based on several different biological paradigms, including, genomics (RNA Interference), proteomics (pathway mapping) and physiology (Immune System). In addition, the study of disease control under the rubric of epidemiological models can inspire models for controlling the spread of pathogens across multiple nodes of a network. The paper discusses these models and presents results based on the authors' research in immune system modeling.*

## **Introduction**

The analogy of computers and communication networks with living organisms is an enticing paradigm that researchers have been exploring for some time. Fred Cohen (1984) first put the word “computer virus” into print, although within the paper he credits Len Adleman with coining the term used to describe the malicious pieces of code that can proliferate on a network and infect disparate computers. Since then, advances in bioinformatics<sup>1</sup> have helped to more precisely define these analogies to the point where results in bioinformatics can often be leveraged for use in computer networking and security. The challenges faced in both bioinformatics and computer network security are quite similar. Several mechanisms have been devised in biological organisms to protect against pathogen invasion. It is important to learn from these biological phenomena and devise innovative solutions to protect the computer systems from software pathogens.

Virus detection systems prevalent today are based on data analysis that looks for the presence of specific patterns in data. The data may be comprised of header information in incoming packets at a firewall, data resident on a node or behavioral patterns of programs resident on a computer. In most cases, the patterns of behavior (signatures) are defined a priori based on knowledge of existing pathogens that have been previously discovered. The signatures are usually gleaned from virus code by teams of virus experts who dissect the virus code and identify characteristic strings that uniquely identify the virus. The signature database in virus detection programs becomes obsolete very rapidly as new virus strains are released continuously and is periodically updated as these new viruses are discovered. However, with the speed of virus propagation growing as is evident from the spread of the *Slammer Worm* that infected more than 90% of vulnerable hosts in 10 minutes, this mechanism is proving inadequate to effectively control the spread of viruses and consequent loss of data and services. It is imperative to develop new virus detection software that does not rely solely on external intervention and can detect new strains of viruses by organically generating “antibodies” within a node. Physiology of cellular organisms contains several paradigms that can be used as inspiration for developing such autonomous security systems in computer networks. Several streams of research on automatic detection of virus (and worm) signatures are in progress (Kim, 2004), however, this research is still preliminary and not mature enough for commercial deployment.

One of the initial areas explored in the realm of biological models of computer security involves the work of Forrest et al. (1994) with regard to virus detection. Here the similarities are strikingly clear, regarding the need

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<sup>1</sup> We consider “bioinformatics” to be modeling of biological processes as well as storage, retrieval, and analysis of biological data through the use of information technology.

to quickly and efficiently identify viruses, generate “antibodies” and remove them from the system before they cause damage and propagate throughout the system. Prior to the advances of Forrest et al. (1994) in this area, Stuart Kaufman et al. (1969) had been focused on understanding and modeling the mechanics of gene transcription and translation within the cell. The concept of a complex network of interactions describing gene regulation had been born in the form of the Boolean Network model. Now that the human genome has been fully sequenced, the task of determining gene function is a significant focus. However, specific genes identified in the sequence can interact with other genes in complex ways. Some portions of the genome can turn off the expression of other genes. These portions are called the structural and regulatory genes. This is thought to be a defense against foreign sequences, perhaps passed on from ancient viruses, from being expressed and potentially harming the organism (Hood, 2004). In fact, in this mechanism one can draw upon concepts that apply directly to network security, namely, the idea of defensive code that can be inherently activated to turn off dangerous code or viruses within the network. One of the problems in virus protection systems is the result of false positives where portions of the code that provide legitimate functionality may be turned off accidentally. The authors propose use of surrogate code that can replicate the functionality of the pieces of code that are shut off maintaining continuity in the operations of the node. Specifically, fault tolerant networks are capable and surviving attacks and dynamically reconstituting services. Bush (2003) explores the ability for a communication network to genetically constitute a service. The network service evolves in real-time using whatever building blocks are available within the network. Thus, a service damaged by a virus attack may be genetically re-constituted in real-time. The general concept was illustrated using a specific example of a genetic jitter control algorithm that evolved a 100-fold decrease in jitter in real-time.

Another biological paradigm that lends itself well to conceptualization of a computer security paradigm is protein pathway mapping. Living organisms have complex metabolic pathways consisting of interactions between proteins and enzymes, which may themselves have multiple subunits, alternate forms, and alternate specificities. Molecular biologists have spent decades investigating these biochemical pathways in organisms. These pathways usually relate to a known physiological process or phenotype and together constitute protein networks. These networks are very complex with several alternate pathways through the same start and end point. The partitioning of networks into pathways is, however, often arbitrary, where the start and finish points are chosen based on “important” or easily understood compounds. The models for biochemical pathways that have been developed thus far primarily demonstrate the working of the cellular machinery for specific tasks, such as, metabolic flux and signaling. Several different modeling techniques are used for understanding these models 1) classical biochemical pathways (e.g. glycolysis, TCA cycle) 2) stoichiometric modeling, e.g. flux balance analysis 3) Kinetic modeling, e.g. CyberCell, E-cell. More recently, cell metabolism is being studied using cellular networks that are defined from large-scale protein interaction and gene expression measurements.

Similar to the cellular networks in organisms, computer networks are complex in nature and collectively exhibit complex behavior. In these networks, start and end points can be arbitrarily chosen and multiple paths may exist between the same nodes. Protein networks are predetermined and stay fairly static while computer networks are constantly evolving with the addition of new nodes and network links. In protein networks interactions among proteins, enzymes, and catalysts result in the culmination of specific events. Analogous to protein networks, interactions among nodes of computer networks result in the manifestation of specific events or conditions in the network. The events may include propagation of viruses, denial of service attacks, as well as congestion on the network. Investigation of network pathways along which the events propagate will enable us, in forensic analysis, to determine the root cause of the failures as well as help in developing intelligence for prediction of network events.

One biological paradigm that is not directly related to physiology of living organisms is epidemiology that involves statistical analysis of disease propagation in population. There are three basic models of disease propagation in epidemiology that have been used extensively for the studies of propagation of infectious diseases. Kephart and White (1991) first used these epidemiological models to study the spread of viruses on computer networks. Williamson and Léveillé (2003) have also developed virus spread models in computer

networks using the epidemiological metaphor. Since then, several researchers have used variations of these basic models for studying the spread of computer viruses on computer networks.

The authors' (Goel & Bush, 2003) have used the biological paradigm of the immune system coupled with information theory to create security models for network security. Information theory allows generic metrics and signatures to be created that transcend the specific details of an individual piece of code or system. They compare information theoretic approaches with traditional string matching techniques. They also provide an analytic model using the epidemiology paradigm to study the behavior of the nodes. This article discusses several different biological paradigms that inspire defense against pathogens that invade computer networks, however, it focuses on in depth analysis of the immune system model. Some of the other innovative biological models that are currently being researched would be discussed in depth in a series of articles following this article.

## **2.0 Immune System Models**

The role of the human immune system is to protect our body from pathogens, such as, viruses, bacteria, microbes, etc. The immune system consists of different kinds of cells, which operate autonomously and through interaction with each other to create complex chains of events leading to the destruction of pathogens. At a high level, cells can be categorized into two groups: detectors and effectors. The role of detectors is to identify pathogens and the role of effectors is to neutralize the pathogens. There are two kinds of immune responses evoked by the immune system: innate response and specific response. The innate immune response is the natural resistance of the body to foreign antigens and is non-specific towards invaders in the body. During this response, a specialized class of cells called phagocytes (macrophages and neutrophils) is used. These specialized cells, which have surface receptors that match many common bacteria, have remained unchanged throughout evolution. This system reacts nearly instantaneously to detect pathogens in the body. However, it is incapable of recognizing viruses and bacteria that mutate and evolve.

The innate immune response is complemented by the adaptive immune response, in which antibodies are generated to specific pathogens that are not recognized by the phagocytes. The adaptive response system uses lymphocytes, which have receptors for a specific strain instead of having receptors for multiple strains as phagocytes do. Lymphocytes are produced in the bone marrow, which generates variants of genes that encode the receptor molecules and mature in the thymus. When an antigen is encountered, it is presented to the lymphocytes in the lymphatic system. The lymphocytes that match proliferate by cloning and subsequently differentiate into B-cells that are used for generating antibodies and T-cells that destroy infected cells and activate other cells in the immune system. Most effectors that proliferate to fight pathogens die; only 5-10% are converted into memory cells that retain the signature of the pathogen that was matched. This leads to a rapid response the next time a similar pathogen is encountered which is the principle used in vaccinations and inoculations. The number of memory cells produced is directly related to the number of effector cells in the initial response to a disease. While the total amount of memory cells may be large as an organism is exposed to new pathogens, newer memory cells may take the place of older memory cells (due to competition for space) (Ahmed, 1998). The decrease in memory cells leads to weakened immunity over time. Another reason that weakened immunity results is an immune response rate, which is not sufficiently rapid enough to counteract the spread of a powerful exotoxin, such as that which is produced with tetanus (Harcourt et al., 2004). Lymphocytes have a fixed lifetime and if during this period they do not match a pathogen, they automatically die.

The key to the functioning of the immune system is the detection mechanism that is employed within the immune system. Recognition is based on pattern matching between complimentary protein structures of the antigen and the detector. The primary purpose of the genetic mechanism in the thymus and bone marrow is to generate proteins with different physical structures. The immune system recognizes pathogens by matching the protein structure of the pathogen with that of the receptor. If the receptor of the antigen and the detector fit together like a three-dimensional jigsaw puzzle, a match is found. A fundamental problem with the

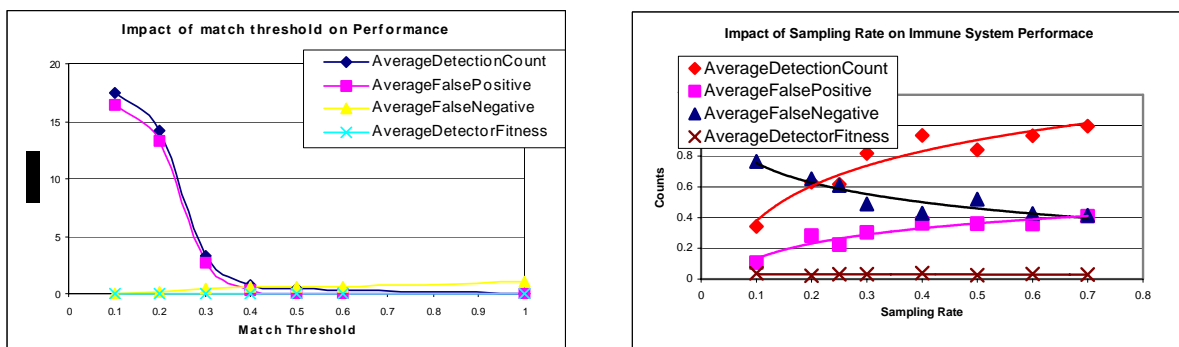
detection mechanism of the immune system is the computational complexity. For example, if there are 50 different attributes with four different values, over six million different detectors are required to cover the entire search space. The virus structures that can arise due to different protein configurations are virtually infinite. In spite of high efficiency in creating detectors and pattern matching at the molecular level, the problem of maintaining a detector for each possible pathogen protein structure is computationally infeasible. The human immune mechanism solves this problem by using generalizations in matching where some features of the structure are ignored during the match. This is called specificity of match; the more the features are ignored during the match, the lower the specificity. The lower the specificity, the fewer the number of detectors required for matching a population of pathogens and the more non-specific is the response. An explanation of specificity is elegantly described in Holland et al.'s description of classifier systems (2000). To cover the space of all possible non-self proteins the immune system uses detectors with low specificity. This enables the immune system to detect most pathogens with only a few detectors; however, it results in poor discrimination ability and a weak response to pathogen intrusion. The immune system counters this problem by employing a process called affinity maturation (Bradley & Tyrell, 2000). Several methods have been proposed for analytic representation of matching pathogen signatures in the immune system, such as bit-strings (Farmer, 1986); De Boer, Segel, & Perelson, 1992), Euclidean parameter spaces (Segel & Perelson, 1988), and polyhedron models (Weinand, 1991) and more recently, Kolmogorov Complexity (Bush, 2002; Goel & Bush, 2003).

Several applications based on immune systems outside the area of biology have recently emerged; the most notable of these being computer security, which is a primary focus in this paper. Kephart (1995) was perhaps the first to introduce the idea of using biologically inspired defenses against computer viruses and immune systems for computer security. Forrest et al. (1994) also proposed the use of immune system concepts for design of computer security systems. Forrest et al. (1994) provided an elaborate description of some immune systems principles applicable to security and present three alternate matching schemes: Hamming distance, edit distance, and r-contiguous bits. They argued that the primary premise behind a computer immune system should be the ability to distinguish between self and non-self. They presented (Forrest, 1994) a signature scheme where a data tuple consisting of source IP-address, destination IP-address, and a destination port number were used to identify self-packets from non-self packets. Hofmeyer (1999) presented a detailed architecture of a computer immune system. He analytically compared different schemes for detection of pathogens, such as Hamming distance and specificity. There are several other works in the literature on the use of immune systems for network security including Murray (1988), Kim and Bentley (1999), and Skormin (2001). Kephart and White (1991, 1993) present an architecture for an immune system and the issues involved in its commercialization. They incorporate a virus analysis center to which viruses are presented for analysis through an active network that The Kolmogorov Complexity approach (Goel & Bush, 2003) demonstrated a 32% decrease in the time required to detect a signature over two common Hamming Distance based matching techniques, i.e. a sliding window or number of contiguous bit matches. The Kolmogorov Complexity based technique estimates the information distance of entire code sequences, not just specific segments or bits. Using the entire code sequence makes it more difficult to modify the virus so that it hides in another portion of a legitimate code segment.

Artificial Immune Systems consist of detectors and effectors that are able to recognize specific pathogen signatures and neutralize pathogens. To detect pathogens, the signature of incoming traffic packets is matched against signatures of potential viruses stored in an immune system database. An immune system that is capable of recognizing most pathogens requires a large number of detectors. Low specificity detectors that identify and match several viruses are often used to reduce the number of detectors at the cost of increased false positives. The computational complexity of a computer immune system remains fairly high and individual nodes are incapable of garnering enough resources to match against a large signature set. The computational complexity gets worse as network traffic grows due to use of broadband networks and is straining the capacities of conventional security tools such as packet filtering firewalls. Massive parallelism and molecular-level pattern matching allows the biological immune system to maintain a large number of detectors and efficiently match pathogens. However, artificial immune systems have not achieved these levels

of efficiency. To reduce the computational burden on any individual node in the network, all nodes need to pool their resources, share information, and collectively defend the network. In addition, such inspection should be done within the network itself in order to improve efficiency and reduce the time to react to events in the network. This concept of collective defense, enabled by a unified framework, is the primary premise of this research. To enable this concept of collective network defense we have proposed an approach based on information theory principles using Kolmogorov complexity measures.

To study the parameters, and different schemes of detection and sampling in the immune system, Goel et al. (2004) have developed a simulation model, using RePast (Schaeffer, et al., 2004) which is a simulation tool typically used for modeling self-organizing systems. The simulation models a classical immune system where new signatures are created by mutation of existing signatures and go through a maturation phase. The simulation also models a cooperative immune system where multiple nodes on the network share virus detection information prevalent in the network to improve the efficiency of each immune system. The research will investigate the tradeoff between the additional burden of sharing information across nodes and benefit of improving the scanning efficiency by obtaining intelligence information on active or new pathogens. Figures 1 & 2 show the impact of the match threshold and sampling rate on the performance of the Immune System respectively. Figure 1a shows a high gradient between a threshold match of .2 and .4, which is the practical operating region for the immune system. Figure 1b shows an improved performance with the sampling rate which asymptotes around 70% sampling rate.



**Figure 1:** Plots showing impact of match threshold & Sampling Rate on Immune System metrics

Goel and Bush (2003) have also compared different signature metrics and have demonstrated that Kolmogorov Complexity is a feasible metric for the signature of pathogens. Bush (2003) explores the ability for a communication network to genetically constitute a service. The network service evolves in real-time using whatever building blocks are available within the network. Thus, a service damaged by a virus attack may be genetically re-constituted in real-time. The general concept was illustrated using a specific example of a genetic jitter control algorithm that evolved a 100-fold decrease in jitter in real-time.

**Conclusions:**

The security models for detection and elimination of pathogens that invade computer networks have been based on perimeter defense. Such defenses are proving inept against fast spreading viruses and worms. The current tools are unable to guarantee adequate protection of data and unfettered access to services. It is imperative to complement these existing security models with reactive systems that are able to detect new strains of pathogens reliably and are able to destroy them before they can cause damage and propagate further. Several biological paradigms provide a rich substrate to conceptualize and build computer security models that are reactive in nature. Three specific mechanisms in mammalian organisms present the most potential, these are: 1) the RNAi mechanism, 2) protein pathway mapping, and 3) the immune mechanism. In addition, the models of disease control that study the spread and control of viruses can be used to find ways to throttle the spread of viruses. Current work has mainly focused on the use of immune and epidemiological

models. It is time to beyond these existing models to other innovative models such as the ones based on genomics and proteomics. Such reactive models provide a scalable, resilient, and cost effective mechanism to sustain the constantly evolving security needs.

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