Immune systems in computer science

Imunski sistemi v računalništvu

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Abstract: A simplified description of the immune system is as follows: this is an organic system intended for protecting the host organism from the threats posed to it from pathogens and toxic substances. The architecture of the immune system is such that a series of defensive layers protect the host. Once the pathogen makes it inside the host, it must contend with the innate and adaptive immune system. These two immunological sub-systems are comprised of several types of cells and molecules produced by specialized organs and processes to address the self non-self problem. Artificial Immune Systems (AIS) are computational models, based on natural immune systems. They tend to solve specific problems in computer science by resembling the natural mechanisms. These systems are more widely applied within problem domains like clustering, pattern recognition, classification, optimization, and machine learning. Modern AIS are inspired by one of three sub-fields: clonal selection, negative selection, and immune network algorithms. In computer science, the clonal selection pattern can be used for pattern matching and optimization. The negative selection algorithm was designed for change detection, novelty detection, and intrusion detection. Immune network algorithms are inspired by the immune network theory of the acquired immune system and it is an upgrade of the clonal selection theory. The objective of the immune network process is to prepare a repertoire of discrete pattern detectors for a given problem domain, where better performing solutions suppress low affinity solutions within the same network. This is an interactive process of exposing the pattern to external information to which it responds. This article explains the biological background, the mechanisms of AIS, and presents their real-world applications. It presents an overview of those important applications of AIS for solving problems from problem domains like data analysis, anomaly detection, intrusion detection, and others.

Key words: natural immune system, artificial immune system, antigen, antibody, algorithm, somatic hyper-mutation.

1. Introduction

The following article presents the usages of immune system concepts in computer science. Immunology is a relatively new science. Its origins have been attributed to Edward Jenner, who in 1796 discovered that by introducing a small amount of vaccine in an animal would induce protection against the often lethal disease smallpox. When Jenner discovered the process of vaccination, very little was known about the functioning of the immune system. In the 19th century the discoveries of Koch and other researchers contributed to the development of the science of immunology, and this development is still continuing.

Artificial Immune Systems (AIS) is a sub-field of Computational Intelligence (CI) motivated by Immunology which emerged during the early 1990s \cite{1,2}. It is based on a proposal for applying theoretical immunological models to automated problem solving and machine learning \cite{3}. The first attempts were inspired by immune network theoretical models and were applied to machine learning, control, and optimization problems. The more successful attempts were those that proposed the immune systems as analogies for information protection systems within the field of computer security. The two prominent examples include Forrest et al.’s Computer Immunity \cite{4} and Kephart’s Immune Anti-Virus \cite{5}. Immune systems are inspired by one of three sub-fields: clonal selection, negative selection, and immune network algorithms. The techniques are commonly used for clustering, pattern recognition, classification, optimization, and other similar machine learning problem domains.

This article analyses the concepts of modern artificial immune systems. Chapter 2 explains their biological background and the basics of immune systems. The chapter is divided into innate and adaptive immune systems for providing the basic biological explanations of those mechanisms which are used within the field of artificial immune systems. Chapter 3 focuses on artificial immune systems and their differentiation and principles. The correlations between the natural and artificial immune systems are explained as to how and where they work. Chapter 4 upgrades the previous chapter by presenting applications of immune-based algorithms in practice. It also includes some new fields by considering immune system investigations that might be important in the future.

2. Biological basics of immune systems

The immune system can be divided into two basic components, the innate and adaptive immune systems. \cite{6} The innate immune system lacks the specificity. Therefore it is also called non-specific response. The adaptive immune system adapts its response during an infection and improves its recognition of a pathogen after the first encounter. This improved response is then retained in the form of an immunological memory. The immunological memory allows the adaptive immune system to act much quicker and stronger when there is a second encounter with the same pathogen \cite{7}. Table 1 presents the basic differences between the innate and adaptive immune system.

<table>
<thead>
<tr>
<th>Innate immune system</th>
<th>Adaptive immune system</th>
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<tbody>
<tr>
<td>Non-specific response</td>
<td>Antigen specific response</td>
</tr>
<tr>
<td>Exposure leads to immediate maximal response</td>
<td>Lag time between exposure and maximal response</td>
</tr>
<tr>
<td>No immunological memory</td>
<td>Immunological memory after exposure</td>
</tr>
<tr>
<td>In nearly all forms of life</td>
<td>Only in jawed vertebrates</td>
</tr>
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</table>

2.1. Innate immune system

The innate immune system provides a non-specific response, which is maximal even at the first encounter with an intruder. It is not able to adapt its response. This means that at any encounter with the same pathogen, the response will be the same. The innate immune system relies on four important characteristics: surface barriers, inflammation factors, complement system, and some specially designed cells.
The surface barriers provide chemical, mechanical, and biological protection against intruders. Human skin, the flushing actions of tears and urine, and the surface of a respiratory tract secreting mucus are all good examples of surface barriers. Inflammation factors include eicosanoids and cytokines, which are released by injured or infected cells. They are important for the production of fever and dilatation of the blood vessel associated with inflammation. Common cytokines include chemocines that promote chemotaxis (directed movement of cells) [8].

The third important part of the innate immune reaction is the complement system. This is an important cascade of molecules that attack the surfaces of foreign cells. It contains over 20 different proteins and is a major component of the innate immune response [9]. This catalytic cascade that works in a positive feedback mechanism opsonizes (coats) in the surface of a pathogen and marks it for destruction.

Lastly, specially designed cells also take part in innate response. These cells are macrophages, dendritic cells, mast cells, neutrophils, eosinophils, basophils, and natural killer cells. These cells of innate immunity play an important role in activating and shaping the adaptive immune response upon activation of their innate immunity receptors. Before the discovery of special receptors on these cells, called Toll-like receptors (TLRs), innate immunity was seen as a crude and unsophisticated part of the immune system. TLR receptors are a class of single membrane-spanning receptors and they recognize structurally conserved molecules derived from microbes. After the activation of TLRs, immune cells produce cytokines. Cytokines are involved in cellular activation and therefore we can say that the innate immunity is responsible for triggering the adaptive immunity [10]. Dendritic cells are antigen-presenting cells with a unique ability to induce and control the development of adaptive immune response. They capture and transfer information from the outside world to the cells of the adaptive immune system. Their main function is to process antigens and present it to the specific clones of lymphocytes, thus inducing their proliferation and expansion. They are antigen-presenting cells and act as messengers between the innate and adaptive immune system [11]. They have many spine-like projections and are mainly located in the skin, nose, lungs, stomach, and intestines. Macrophages are also phagocytes, and basophils secrete chemical mediators that are involved in defending against pathogens (Figure 1) [8].

![Figure 1. Neutrophils, eosinophils, and basophils. Collectively they are referred to as granulocytes because of the small granules inside which contain chemical mediators.](image)

2.2. Adaptive immune system

The adaptive immune system remembers a pathogen by its signature. This signature is called the antigen and so this is an antigen-specific response. Antigen is a part of pathogen or self-molecules that evokes the adaptive, antigen-specific immune response, be it the production of antibodies or T-cell mediated immunity. Each antibody or T-cell receptor binds to a specific antigen by way of an interaction similar to the fit between a lock and a key. There is a lag-time between exposure and the production of antibodies, and during this lag-time the immunological memory is formed and the response is upgraded. The two main parts of the adaptive immune system are lymphocytes B and T. We will describe some important mechanisms of an adaptive immune response which have inspired computer science.

Central tolerance is the mechanism by which lymphocytes B and T mature. B-cell maturation occurs in bone marrow and T-cell maturation occurs in thymus. These developing B and T cells are first exposed to positive selection to self-antigens. The positive selection selects for survival of those cells which can respond through their receptors. Those that do not respond, fail to survive [12]. In the next stage, developing B and T lymphocytes are exposed to self-antigens in the process called negative selection. The negative selection deletes the clones which respond to self antigens strongly. The affinity part seems to play an important role, since those clones that still respond, but with a lower affinity, will be regulatory clones which spread the tolerance to self signals (Figure 2) [8][13][14].
Figure 2. The clonal selection and the negative selection principle. Some lymphocytes carry the receptors that bind to self-antigens. They are eliminated during the early phase of development before they are able to perform and attract an immune response. This process leads to self-tolerance and is called ‘negative selection’. When the foreign antigen interacts with the receptor within a mature lymphocyte, the cell is activated and commences proliferating. This is the ‘clonal selection principle’. The clone will arise and differentiate between effector and memory cells. During the differentiation these cell undergo changes called ‘somatic hyper-mutation’.

Lymphocytes B produce antibodies. Antibodies are their way of detecting antigens. Each antibody produced by lymphocyte B has its own unique variable region which allows an antibody to recognize its matching antigen. When an antibody binds to an antigen, this antigen/antibody complex is taken up by B lymphocytes and processed by proteolysis into small particles, peptides.

3. Artificial immune systems

Artificial immune systems simulate the basic concepts of natural immune systems in order to solve real-world problems. Immunology is the researching of immune systems and the body’s protection against viruses, bacteria, cancer and other intruders. Nowadays, artificial immune systems are based on three important types, as follows:

- clonal selection algorithm,
- negative selection algorithm,
- immune network algorithm.

The clonal selection algorithm is inspired by the clonal selection theory of acquired immunity \[15\] \[16\]. This theory proposes that when a lymphocyte binds to an antigen, it starts to proliferate, making many more thousands of copies of itself and differentiates into different cell types (Figure 2).

The important feature of the theory is that when the cell proliferates, it goes through the process of small copying errors \[17\]. This causes the changes in the genome and these changes are called ‘somatic hyper-mutation’ \[18\]. The result of this process is the change of the shape of the expressed receptors and subsequent antigen recognition capabilities. In other words, starting with the initial repertoire of general immune cells, the system is able to change itself in composition and density. The general model of the clone selection algorithm involves the selection of antibodies (possible solutions) and these selected antibodies are subjected to cloning (proliferation) and hyper-mutation. The goal is to produce antibodies which have the highest affinity to a certain antigen pattern. The resultant clonal set competes with the existent antibody population for membership in the next generation. In addition, all low affinity population members are replaced by randomly generated antibodies. The result is antibodies which express the highest affinity for a certain pattern. As seen from the description, the clonal selection pattern in computer science can be used for pattern matching and optimization \[19\].

A negative selection algorithm is inspired by the process of negative selection described earlier in this article. This self-nonself discriminatory behavior is responsible for the fact that only the cells selected for potentially harmful and foreign material in the body are included in the ongoing proliferation (Figure 2). In this way, self-reactive immune cells are destroyed. The negative selection algorithm was designed for change detection, novelty detection, and intrusion detection \[15\]. This is achieved by building a model of changes, anomalies or unknown data by generating patterns which do not match with an existing corpus of available (self or normal) patterns. After that this prepared abnormal model is used to monitor the existing normal data or streams of new data by seeking matches to the abnormal patterns \[20\].
Immune network algorithms are inspired by the immune network theory of the acquired immune system. This is an upgrade of the clonal selection theory. A concern of this theory is that it presumes that immune cells remain idle when there are no pathogens to which to respond [20]. The immune network theory proposes that immune cells are not at rest in the absence of pathogens, instead the antibody and immune cells recognize and respond to each other [21]. Antibodies, both free-floating and surface-bound, possess surface features to which the receptors of other antibodies can bind. As a result of this interaction, antibodies both excite and inhibit each other within complex regulatory networks [chains of receptors] [22]. Regarding this principle, in addition to a cell which reacts directly with pathogen, there are cells that interact with those reactive cells. The objective of the immune network process is to prepare a repertoire of discrete pattern detectors for a given problem domain, where better performing cells suppress low affinity cells within the same network [23]. This is an interactive process of exposing the population to external information to which it responds.

4. Applications of AIS

Currently, the negative selection algorithm is the most popular in computer applications. The purpose of negative selection is to provide tolerance for self-cells. The pseudo-code of this AIS algorithm is presented in Pseudo-code 1.

\begin{center}
\textbf{Pseudo-code 1: Algorithm AIS with negative selection}
\end{center}

\begin{verbatim}
01: Input: \( S = \text{set of known self samples} \)
02: Output: \( D = \text{set of collected detectors} \)
03: repeat
04: \quad Randomly generate potential set of detectors \( p \in P \); 
05: \quad Determine the affinity of each member \( p \in P \) with each member of the set \( s \in S \); 
06: \quad if At least one element \( s \in S \) match a detector \( p \in P \) then 
07: \quad \quad Discard \( p \in P \); 
08: \quad else 
09: \quad \quad Add \( p \in P \) to detector set \( D \); 
10: \quad endif
11: until Termination condition has been met;
\end{verbatim}

As can be seen from Pseudo-code 1, let us assume an input set of seen known self-samples, while the set of generated detectors is obtained. In main loop (lines 03-13), the algorithm randomly generates a potential set of detectors \( p \in P \). Then the determining of the affinities of each number in \( p \in P \) against each member of \( s \in S \), needs to be performed (line 04). Finally, detection stage (lines 07-12) is conducted in order to verify whether at least one element in \( s \in S \) matches the detector \( p \in P \). In that case, the detector is rejected while, in contrast, it is added to the set of available detectors \( D \). The principle of this detection stage is illustrated in Figure 3.

As many sources have suggested, AIS is very useful for intrusion [24] or anomaly detection [25]. In line with this, some AIS applications are used for computer virus detection too. On the other hand, very interesting applications also focus on the optimization. For example, Gong et al. [30] applied AIS to numerical optimization. Moreover, Freschi and Repetto [29] solved multi-objective optimization problems with AIS. On the other hand Coello [26] applied AIS to job shop scheduling which is a well-known problem of combinatorial optimization. Some tests have also showed that AIS provides promising results by data analysis [27] and also by clustering [31]. From these facts,
it can be concluded that AIS is a suitable method for
data mining. Actually, data mining is currently one of
the hottest areas of research in computer science.

Nowadays, AIS is also a proven tool in industrial \(^{[28]}\)
as well as control system \(^{[32]}\) domains.

### Table 2. Applications of algorithms based on AIS

<table>
<thead>
<tr>
<th>Application</th>
<th>Author</th>
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<tbody>
<tr>
<td>Intrusion detection</td>
<td>Aickelin et. al. (^{[24]})</td>
</tr>
<tr>
<td>Anomaly detection</td>
<td>Greensmith et. al. (^{[25]})</td>
</tr>
<tr>
<td>Numerical optimization</td>
<td>Gong et. al. (^{[30]})</td>
</tr>
<tr>
<td>Multi-objective optimization</td>
<td>Freschi and Repetto (^{[29]})</td>
</tr>
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<td>Job shop scheduling</td>
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<tr>
<td>Data analysis</td>
<td>Timmis (^{[27]})</td>
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<tr>
<td>Dociment clustering</td>
<td>Tang and Vemuri (^{[31]})</td>
</tr>
<tr>
<td>Industrial applications</td>
<td>Dasgupta and Forrest (^{[28]})</td>
</tr>
<tr>
<td>Control systems</td>
<td>Wei and Zhang (^{[32]})</td>
</tr>
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</table>

AIS are still under development and further
improvement. In the future, we can expect more real
world applications using AIS for solving hard
problems. Therefore, many novel variants and hybrids
of AIS will come out in the following years.

### 5. Conclusion

This paper presented the principles of the natural
immune system which in turn inspire artificial
algorithms for solving complex problems in computer
science. The immune system is a remarkable natural
defense mechanism that exhibits capabilities like
learning, memory, and adaptation. In line with this, it
is hoped that this article has served to convince the
reader that the immune system is worthy of study from
a computational point of view. This article exhibited
three main types of AIS: clonal selection, negative
selection, and immune networks.

The goal of AIS in the future should be to further
extract what can be considered as inspiration from the
components and processes from the immune system
for creating effective and powerful computational
systems. A significant number of concepts could be (or
currently are being) used in the development of AIS.
Some of these concepts are clonal expansion and
affinity maturation, cross-reactivity, epitope, idiotope,
paratope, B-cell and BCR, T-cell and TCR, network
structure, dynamics and meta-dynamics, etc. It can be
seen that the possibilities of extracting useful
metaphors and constructing new AIS theories are still
wide-open.

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