A Feasibility Study of Challenges and Opportunities in Computational Biology: A Malaysian Perspective

R.J. A. Richard and N. Sriraam
Centre for Multimedia Computing, Faculty of Information Technology
Multimedia University, Cyberjaya, Malaysia

Abstract: The term computational biology refers to the knowledge derived from computer analysis of biological data that includes identification of genes in DNA sequence of different organisms, prediction of structural and functional mechanism of proteins, feature extraction and classification of genomics and proteomics. Computational biology is a rapidly developing branch of science and is highly interdisciplinary, using techniques and Concepts from informatics, mathematics, chemistry, physics, statistics and biochemistry. This field has arisen in parallel with the developments of automated high throughput methods of biochemistry and biological discovery that yield a variety of forms of experimental data, such as DNA& RNA sequences, gene expressions patterns and chemical structures. The field’s rapid growth is spurred by the vast potential for new understanding that can lead to new technological treatments, new agro-crops cultivation and new pharmaceutical drugs discovery. In the recent years, most bioengineering disciplines are started adopting the information technology oriented curriculum due to its high performance computing, data inter-operability, web-based platform compatibility and secured a suitable job opportunities. This study discusses the challenges to setup an interdisciplinary oriented curriculum by merging life sciences and information technology in a university level. It also provides the career opportunities for different life science disciplines like drug development, microbial genome applications, biotechnology, forensic analysis of microbes.

Key words: Computational biology, genomics, proteomics, interdisciplinary oriented curriculum

INTRODUCTION

The term Computational biology refers to the knowledge derived from computer analysis of biological data that includes identification of genes in DNA sequence of different organisms, prediction of structural and functional mechanism of proteins, feature extraction and classification of genomics and proteomics. Computational biology is a rapidly developing branch of science and is highly interdisciplinary, using techniques and concepts from informatics, mathematics, chemistry, physics, statistics and biochemistry[1-5]. Computational Biology today consists of collection of methods and computational approaches and tools that have been developed in very disparate areas to analyze genomic data. This technology has been evolved to provide functional genomics support in terms of data management, the decision process, database integration and access to many of the large databases being produced with today’s technology innovations[2]. In recent years, Bioinformatics, a branch of computational biology is emerging as a hottest research field among the Biotechnology and information technology researchers[6]. This field has arisen in parallel with the developments of automated high throughput methods of biochemistry and biological discovery that yield a variety of forms of experimental data, such as DNA& RNA sequences, gene expressions patterns and chemical structures. The field’s rapid growth is spurred by the vast potential for new understanding that can lead to new technological treatments, new agro-crops cultivation and new pharmaceutical drugs discovery. In the recent years, most bioengineering disciplines have started to adopt the information technology oriented curriculum due to its high performance computing, data inter-operability, web-based platform compatibility and secured a suitable job opportunities[2,3]. This study discusses the functional research areas of life science that requires information technology platforms and highlights the scope, challenges and opportunities for computational biology in Malaysian region.

Primary areas and functions: The following key areas of computational biology enumerate its major function:

Genomics

Description: Genomics is one of the leading research areas in life science, which includes the range of research from the molecular basis of disease to biochemistry and to developmental biology.
Proteomics

Description: High-throughput genomics technologies such as genome sequencing and whole-genome expression analysis are retransforming the biological sciences now. The genome analysis and sequencing helps the scientists to identify similar RNA and DNA sequences and turn into cloning of new genes.

Scope and growth: High-throughput genomics technologies will help to merge them in sequences. The sequence analyzing tools and development of user interfaces that bring together thousands of fragments that must be correctly "stitched" together. The sequence analyzing tools and algorithms will help to merge them in sequences.

Scope and growth: Standardization of biological sequences into XML formats, simplified the process of sequence gathering and comparison by the cross references of biological databanks. The growth of BioXML, BSML, etc., assures the interoperability among the biological data banks.

Bioinformatics

Description: Bioinformatics is the new field of study that seeks to combine Biotechnology with Information Technology. Its basic tools (especially methods for finding relationships between genes) have already proved tremendously useful. Most of the Biological data banks, accessed tremendously by the researchers are dependent on these bioinformatics tools.

Scope and Growth: It is a very young and incomplete field, whose deficiencies are a major anticlimax in the genomics revolution. Functional aspect of bioinformatics is the representation, storage and distribution of data. Intelligent design of data formats and databases, creation of tools to query those database and development of user interfaces that bring together different tools to allow the user to interact with the databanks through web.

Proteomics

Description: A relatively new area, proteomics studies, which do not rely on the entire genome, but rather on the portion of the genome that is, expressed in particular cells. This often involves cutting-edge technology, such as the usage of micro arrays, which allows the expression level of thousands of genes in a cell sample to be quickly determined.

Scope and growth: It targets the location of drug/gene therapy. Bioinformatics specialists work closely with bench scientists to accomplish the "data mining" that lies behind this next wave of the pharmaceutical industry.

Sequence Analysis

Description: Sequence analysis is the most primitive operation in computational biology. This operation consists of finding which part of the biological sequences are alike and which part differs while medical analysis and genome mapping processes.

Scope and growth: Knowledge based single sequence analysis for sequence characteristics, pair wise sequence comparison and sequence-based searching, multiple sequence alignment and phylogenetic inference.

Database Maintenance

Description: Many pharmaceutical companies and research institutes maintain private data banks of gene sequences and other biological and chemical information. These databanks have to achieve common formats.

Scope and growth: These repositories must be continually updated with data generated internally and from outside sources. This is a challenging task and the design and maintenance of these complex databases has become an important part of these complex databases have become an important part of Bioinformatics.

Pharmacogenomics

Description: Pharmacogenomics is the application of genomic approaches and technologies to identification of drug targets.

Scope and growth: New drug discovery using bioinformatics tools and disease control. There are other emerging fields using bioinformatics tools such as Chemeoinformatics, Agro-informatics, Medical-informatics, etc.

Challenges in Computational Biology: The foremost challenge lies in computational biology is the understanding of the structure of a cell. A cell which contains hundreds of thousands of proteins that depends not only on the linear sequence of amino acids, presence of fats, sugar and water in its micro environment, but also on other molecules in their immediate proximity. Complete understanding of the proteome will require new supercomputer architectures; immensely large databases, new data mining methods, new modeling and simulation techniques and the networking expertise to integrate data from desperate sources such in such a manner that it should be effective and affordable.

In this present trend, the rate of change in computer-enabled technological innovation is accelerating and hence practical applications of computing to unravel the proteome and other bioinformatics challenges are growing exponentially. In this regard, bioinformatics should be given a high priority by programmers, systems architects and other computer technology professionals to make an opportunity to take a proactive role in defining and shaping not only their future, but the future of humanity as well.
Challenges in the Sequence Analysis: The primary challenge for the sequence analysis of genomics is the existence of large volume of data\cite{10}. It must provide analysis that can keep up and decipher the inherent structure of information within the data. This is due to the nature of the data that are detailed, complex and voluminous, representing the complete genetic blueprint for a living organism.

The Challenges of Genomics: Genomics is based on automated, high-throughput methods for generating experimental data, ranging from gene sequence, to gene expression levels, to protein-protein interactions\cite{6-7,10}. Genomics technologies are making it possible to study biology as a system rather than as a microcosm. The completeness of the dataset and the systematic nature of the experiments, is designed to enable total analysis of the system and explicit comparison of any pair or any number of different experiments. These characteristics are the fundamental foundation for the next era in biology reassembling the understanding of molecular pieces into predictive models of the biology as a composite system.

Developing a quantitative analysis and understanding of genome is very complex. At the very basic level, it is very difficult to identify the unknown genes through computer analysis of genomic sequences. The biological data and sequences are also very complex and interlinked with each other. A spot on a DNA Array, for instance, is connected not only to immediate information about its intensity, but to layers of information about genomic location, DNA sequence, structure, functions, etc. To find a metallic and biochemical pathway of genes representation in genomes will need collaborative work of computational biologists, computer scientists, mathematicians and statisticians\cite{10}.

This challenge will define bioinformatics as a science. It is the analysis component for making sense of genomes data and putting it together to make predictive models of biological systems. Fundamentally, this is the real opportunity of bioinformatics; its interesting theory driven by the benefit of one of the biggest, richest waves of systematic complete data ever produced. Theory suffers when data is too sparse or weak to falsify or “confirm” predictions, the data do not challenge new theoretical frameworks, or fundamental knowledge or theoretical tools are missing\cite{15}.

Career opportunities: Recently, demand for trained and skilled personnel has opened up a new career for bioinformaticians, especially in research & development sectors. This demand will clearly become exacerbated with advances in genomics and post genomic research.

The career in bioinformatics is divided into two parts such as: developing software and utilizing it. Bioinformatics graduates are in high demand. Career opportunities in bioinformatics will continue to expand on both the computer and biological science fronts. For computer science, there are opportunities in programming, database development, systems analysis and software engineering. In the biological sciences career opportunities exist for bioinformaticians who assist biologists and medical researchers to interpret biological or medical data, design user interfaces and run laboratory information management and analysis systems. Opportunities also exist in research laboratories in universities, hospitals and the biotechnological and pharmaceutical industries\cite{16,17}.

Feasibility study: Curriculum on computational biology- A Malaysian perspective: Recently computational biology curriculum has been started in universities at postgraduate level to produce bioinformaticians and computational biologists. Malaysian government’s initiatives on bioinformatics will give more opportunities computational biologists, bioinformatics researchers and bioinformaticians. Our study on computational biology explains the importance of this emerging technology. We would like to list down the findings of this study.

* Computational biology, though seems to be the hottest research area, has to reach student community yet.
* It has rapid growth, which results in high job opportunities.
* Merging of information technology with life sciences will bring more job opportunities.
* Information Technology concepts and subjects should be included in Life sciences curriculum to fulfill the future requirements.
* Mathematics and Statistics knowledge are important to analyze the biological sequences. Both areas to be covered while developing computational biology curriculum.
* Governments and industries have initiated research and development programs in Life science area and this emerging field will be the most wanted technology in future.

* Currently most universities in Malaysia have Life Sciences (Biology, Biotechnology, Molecular Biology, Agriculture, etc) and Information Technology courses at under graduate and postgraduate level. Thus they have the infrastructure to start computational biology oriented curriculum at under graduate level.
* With this we could say, Malaysia has consistent development in biotechnology and Government of Malaysia has allotted funding for Biotechnology field. Since Asia-Pacific region has produced rapid growth in this field, Malaysia can take a lead in Bioinformatics in Asia-Pacific region, by collaborating research organizations, academic sectors and industries\cite{18}. According to its vision
2020, the country needs more trained computational biologist and bioinformaticists to fulfill the job requirements. The country is encouraging researchers to work on Agro-informatics, which is focusing on seed technology, new crop cultivation, tissue culture and other researches on agricultural sector. Malaysian Government has initiated BioValley project to work on Biotechnology and Bioinformatics.

**BioValley Malaysia: Targets and goals:** Proposed under the Eighth Malaysia Plan, BioValley Malaysia will be the nucleus of the country’s biotechnology industry. It adopts a cluster approach that has been successfully applied to develop biotechnology industries around the world. The cluster will comprise biotechnology research institutions, universities and specialized companies. This is to foster inter-relationships and the convergence of intellectual expertise and entrepreneurship as well as the sharing of resources, which will enable the industry to establish a centre of excellence. BioValley will integrate existing research resources with new facilities, equipment and human resources that will seed growth and innovation. BioValley will stimulate the growth and application of biotechnology in Malaysia by catalyzing research through the formation of new Institutes and linking Malaysian academic centers to industry.

**NBBnet:** National Biotechnology and Bioinformatics Network (NBBnet) is an initiative by Malaysian government. It gives R&D support to the researchers for conducting Biotechnology and Bioinformatics researches. It allows access to biological databanks and other resources to conduct better researches in this hottest research area. Research institutes and the center of excellences can use these available resources\(^{(19)}\).

Malaysia’s initiatives in biotechnology will bring the nation to be a top leader in Biotechnology and Bioinformatics in Asia Pacific region. This emerging field will require trained Bioinformaticians and Computer Biologists to fulfill the requirements of the industry. Since computational biology has rapid growth in Asia Pacific region, we strongly feel universities can take initiatives to start up Bioinformatics curriculum at Under Graduate level to give strong foundation in Genomics, Proteomics, Bioinformatics tools and Computational tools for Molecular Biology. Also universities can focus on certain areas according to their specialization, but common courses can be introduced to give basic awareness.

**Proposed computational biology curriculum:** Recent developments in computational and related fields have led to an explosive growth in biological information. Biological data is being generated faster than it can be analyzed and utilized. The importance of this field to drug discovery, biotechnology, agro-informatics, tissue culture, crop cultivation has resulted in a rapidly emerging commercial bioinformatics sector. There is a growing trend for professionals with strong foundation in both information technology and life sciences. The current demand is very high; we strongly feel curriculum on computational biology at under graduation level will educate students with expertise in both computing and life sciences and who have learned to integrate these two disciplines\(^{(20,21)}\). With the above discussion, we would recommend the courses below for the three years computational biology curriculum.

**Year-1 Semester-1**
- Introduction to Molecular Biology
- Basic Mathematics for Computational Biology
- Introduction to Computer Programming Languages (C & BioPerl)
- Database systems

**Year-1 Semester-2**
- Bioinformatics Computing
- Statistics and Probability theory

**Year-1 Semester-3**
- Biophysical Chemistry (I)
- Introduction to sequence analysis
- Representations and Algorithms in Computational Biology

**Year-2 Semester-1**
- Programming in Object Oriented Languages
- Genetics, Genetic Information and Flow Processing
- Protein and Nucleic Acid Structure, Dynamics and Engineering
- Biophysical Chemistry (II)

**Year-2 Semester-2**
- Computer Graphics and Visualization in Computational biology
- Data Mining and machine learning techniques

**Year-2 Semester-3**
- Introduction to system and computational neuroscience
- Biomaterials
- Genetics, Evolution and Behavior
- Study of Human Genome

**Year-3 Semester-1**
- Industrial Training

**Year-3 Semester-1**
- Elective- 1
- Elective -2
- FYP-1

**Year-3 Semester-3**
- Artificial Intelligence
- Advanced Computational Biology
- Elective 3
- FYP-2
- (FYP- Final Year Project)
Optional Subjects: Optional Subjects are listed down in Table 1.

Table 1: Optional Subjects

| Course                        | Targeted Students       |
|-------------------------------|-------------------------|
| Basic Biology (Non Biology Students) | Non Biology students |
| Human-Computer Interaction    | All                     |
| Molecular Biology of Proteins | All                     |
| Software Engineering          | All                     |
| Agroinformatics               | All                     |
| Computer Communication Networks | All                   |
| Neural Networks               | All                     |
| Biochemistry and Metabolic Pathways | All       |
| Study of Human Genome         | All                     |
| Structural Biology            | Non Biology Students    |

CONCLUSION

Computational Biology is a field of science at the cutting edge of two of the fastest growing areas of technology information technology (computer science) and biotechnology (genetics, molecular biology and biochemistry). It is an interface between computing, mathematics, statistics and the Life Sciences and it is responsible for producing new computer systems (databases, software, networks and even hardware) and for using computer systems to solve problems in a wide variety of areas ranging from biology to medicine. The idea of developing/introducing the new curriculum on computational biology is:

* To provide the necessary awareness on computational biology.
* To provide infrastructure and resources for bioinformatics and biotechnology information.
* To produce skilled computational biologists and bioinformaticians for the future requirements and to promote the development of research program.
* To provide bioinformatics tools and resources such as databases, software, network communication, training and consultancy.

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