Chapter 9
Cloud-Based Computing Architectures for Solving Hot Issues in Structural Bioinformatics

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ABSTRACT
Bioinformatics as a scientific domain develops tools that enable understanding the wealth of information hidden in huge volumes of biological data. However, there are several problems in bioinformatics that, although already solved or at least equipped with promising algorithms, still require huge computing power in order to be completed in a reasonable time. Cloud computing responds to these demands. This chapter shows several cloud-based computing architectures for solving hot issues in structural bioinformatics, such as protein structure similarity searching or 3D protein structure prediction. Presented architectures have been implemented in Microsoft Azure public cloud and tested in several projects developed by Cloud4Proteins research group.

1. INTRODUCTION
Bioinformatics is an interdisciplinary scientific domain, which develops algorithms, methods and software tools for discovering knowledge that is hidden in the wealth of biological data. For example, bioinformatics tools used for comparison of two genes, genomes or proteins help in understanding of evolutionary relationships between two organisms. Such information is not apparent at first glance when looking at raw data and requires appropriate software tools and algorithms in order to become visible. Bioinformatics helps in understanding genetic basis of unique diseases, like Alzheimer’s, cancer, diabetes mellitus, providing information on mutations in nucleotide sequences of DNA or RNA strands. Based on the information scientists can distinguish types of particular diseases and medical doctors are able to adjust their therapies to the type of disease. Knowledge of protein structures and other biological molecules is a foundation for the
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creation of new drugs, which translates into better quality of life. Therefore, it is not surprising that bioinformatics has become a part of many areas of biology, medicine and modern pharmacy.

Structural bioinformatics is the branch of bioinformatics, which focuses on the analysis, comparison and modelling of the three-dimensional structures of proteins, DNA and RNA acids, and other molecules and molecular complexes. This sub-discipline of bioinformatics has emerged as a result of rapid increase in the number of three-dimensional macromolecular structures available in databases such as the Protein Data Bank (http://www.pdb.org) (Berman et al., 2000). Structural bioinformatics provides general purpose methods for processing information about biological macromolecules with aim to create new knowledge and a high-resolution understanding of biology. It relies on the belief that having this high-resolution structural information about biological systems will allow scientists to precisely reason about the function of these systems and the effects of modifications or perturbations (Gu & Bourne, 2009). However, performing analyses on such detailed level requires increased computational resources that are sometimes available only in big data centers. Bioinformatics is developing very dynamically in recent years and the development is also supported and accelerated by development of new computing architectures and new computing models. The concept of Cloud computing became one of the elements catalyzing the development of the domain.

Following the definition of NIST, Cloud computing is “a model for enabling convenient, on-demand network access to a shared pool of configurable computing resources (e.g., networks, servers, storage, applications, and services) that can be rapidly provisioned and released with minimal management effort or service provider interaction” (Mell & Grance, 2011). In practice, cloud computing allows to run applications and services on a distributed network using virtualized system and its resources, and at the same time, allows to abstract away from the implementation details of the system itself. The use of cloud platforms can be particularly beneficial for companies and institutions that need to quickly gain access to a computer system which has a higher than average computing power. In this case, the use of cloud computing services can be more cost effective and faster in implementation than using the owned resources (servers and computing clusters) or buying new ones. For this reason, cloud computing is widely used in business. However, the concept of cloud computing is also becoming increasingly popular in scientific applications for which theoretically infinite resources of the cloud allow to solve the computationally intensive problems. This chapter shows several, tested on Microsoft Azure platform, cloud architecture patterns for solving hot issues in structural bioinformatics, including protein structure similarity searching or 3D protein structure prediction. Presented architectures have been implemented and tested in several projects developed by Cloud4Proteins research group.

2. BACKGROUND

Today’s bioinformatics is struggling with many problems and use a variety of approaches to solve them. This chapter provides broad definitions of the problems, which Cloud4Proteins group tries to solve, and focuses on technologies that are used to achieve the solutions.

Proteins and Computationally-Intensive Problems

From the perspective of biological processes, functioning of living organisms is tightly related to existence and activity of protein molecules. Proteins are important molecules that play a key role in all biochemical reactions in organisms’ cells. They are involved in many processes, e.g.: reaction catalysis, energy storage, signal transmission,