

Analysis of Short-Read Data from NCBI SRA and Cghub/TCGA Using HIVE-XLD: Evaluation of the Effects of Non-Synonymous Variation in the Human Proteome

Raja Mazumder^{*}

Department of Proteomics, George Washington University, Missouri, USA

DESCRIPTION

Current sequencing technologies are generating petabytes of data that are inaccessible to majority of the research community because of the costs and expertise required to analyze big data. Another roadblock to analyzing such data is the lack of curated information in NGS data repositories such as NCBI SRA. To address the above challenges, we have implemented a low-cost Highperformance Integrated Virtual Environment of eXtra-Large Data (HIVE-XLD) private cloud at GWU and US FDA. Effects of variation on active sites and glycosylation sites will be presented to illustrate the power of integration of big-data with functional objects such as active sites, binding sites and pathways. DNA sequencing technology advances have enabled genetic investigation of more samples in a shorter time than has previously been possible. Furthermore, the ability to analyze and understand large sequencing datasets has improved due to concurrent advances in sequence data analysis methods and software tools. Constant improvements to both technology and analytic approaches in this fast moving field are evidenced by many recent publications of computational methods, as well as biological results linking genetic events to human disease. Cancer in particular has been the subject of intense investigation, owing to the genetic underpinnings of this complex collection of diseases. New massively-parallel sequencing (MPS) technologies have enabled the investigation of thousands of samples, divided across tens of different tumor types, resulting in new driver gene identification, mutagenic pattern characterization, and other newly uncovered features of tumor biology. This review will focus both on methods and recent results: current analytical approaches to DNA and RNA sequencing will be presented followed by a review of recent pancancer sequencing studies.

This overview of methods and results will not only highlight the recent advances in cancer genomics, but also the methods and tools used to accomplish these advancements in a constantly and rapidly improving field.

Cancer is not one disease, but a collection of different diseases that share common features: origination from patients' own cells and a disrupted regulatory program resulting in uncontrolled growth. Cancers also generally have a genetic origin; changes in the biological blueprint initiate the signaling and regulatory alterations that lead to tumorigenesis. Early work at the beginning of the 20th century suggested that certain chromosomal aberrations could cause unregulated growth in sea urchin eggs, leading to a variety of hypotheses regarding the role of chromosomes in cancer. The role of genetic alterations in human cancer was confirmed with the discovery and subsequent classification of the Philadelphia chromosome. This and other genetic discoveries led to theories of the requirement of multiple genetic events driving clonal selection of tumor cells. Advances in molecular manipulation played a role in the cloning and identification of the first oncogenes, followed by the discovery of the exact nucleotide change responsible for the oncogenic phenotype (in this case, the genetic changes resulting in the RAS G12V amino acid substitution). This body of work has cemented the role of genetic alterations, large and small, in the biology of cancer. Cancer genes had been discovered prior to the public release of the human genome using techniques such as positional cloning, biological screening assays, and candidate gene studies. However, the Human Genome Project has greatly increased our understanding of the structure and contents of our chromosomes, allowing for the design and execution of experiments that were not previously realistic. In the continued study of cancer genetics, this led to numerous large-scale investigations of many genes across different cancer types, resulting in the identification of a number of new cancer genes. Initial studies focused on smaller groups of genes, including tyrosine kinases, a more comprehensive set of kinases, and other genes. Using this approach, BRAF was discovered to contain a common mutation in several cancer types, with a high prevalence in melanoma.

Correspondence to: Raja Mazumder, Department of Proteomics, George Washington University, Missouri, USA; E-mail: Mazumde020@wus.edu

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