

Rheinisch Westflische Technische Hochschule Aachen  
Informatik 5 (Information Systems)



## **Master Thesis Proposal**

### **Capturing Data Curation Process in Cancer Genomics**

**Jasim Waheed Ansari**

**Aufgabenstellung:** Prof. Dr. Stefan Decker

**Betreuer:** Dr. Oya Deniz Beyan

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# Chapter 1

## Introduction

Cancer is one of the leading causes of death around the globe. It is considered to be the most complex disease that human species has to deal with. It is characterized by undesirable growth in cells which are caused by changes in genomes and exposure to environment[2]. Each cancer type has unique architecture of genetic variation - somatic mutations, copy number alterations, gene expression profiles and different epigenetic aberration. Therefore, there is a strong need for better diagnosis, personalized medicines for patients have arose, which correlates with better understanding the changes at genetic level in tumors. All thanks to technological advancements in high-throughput techniques for genome-wide interrogations such as Microarray and Next-Generation Sequencing(NGS) platforms have provided means to target personalized healthcare solutions for patients[5] [4].

### 1.1 Data Integration Challenges in the era of Cancer Genomics

With the deluge of multiple projects from different organizations catering to Cancer Genomic has grown over years, one such is The Cancer Genome Atlas(TCGA). It comprises of genomic, epigenomic, and proteomic data from over 10 thousand samples from 33 kinds of cancer, with a target to support bioinformaticians and medical researchers in understanding molecular foundation of cancer development. Since the data available in TCGA qualifies the 3Vs i.e. Volume, Velocity and Veracity principle of Big Data, integration of comprehensively characterizing the whole genomes at low cost is a big challenge[1][3]. Performing mining on the big data extracted through TCGA is overwhelmingly difficult by the traditional tools like R and Python. Furthermore, the data collected in TCGA site exists in native raw form, therefore there is a need to perform further preprocessing which needs background in both biological domains as well as technical expertise for programming. TCGA provides ample of intuitive web-based tools, such as TCGA-Assembler, cBioPortal, Firebrowse etc which supports clinicians and researcher in making knowledgeable insights of all the data types in a significant manner. Visual analytical tools has features which offers integrative data mining capabilities in the following modes:

- **Discovery mode:** Finds global correlations across different data types.
- **Confirmation mode:** Analysis on results from few genes.

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## Chapter 2

# Background & Related work

### 2.1 What is Cancer Genome?

### 2.2 Current state-of-art

#### 2.2.1 The Cancer Genome Atlas

cBioPortal

Firebrowse

### 2.3 Big Data Fundamentals

#### 2.3.1 Technologies for Big Data

#### 2.3.2 Shortcomings in traditional Data Warehousing

#### 2.3.3 Need for Data Lakes in Big Data

#### 2.3.4 Big Data Visualization



## Chapter 3

# Solution

### 3.1 Development of Big Data Lake Platform using Kylo

#### 3.1.1 Data Ingestion Pipeline and Design with Apache Nifi

#### 3.1.2 Data Standardization and Validation with Apache Spark

#### 3.1.3 Metadata Tracking

### 3.2 Development of interactive dashboard using Apache Zepelin





## Chapter 4

# Evaluation

### 4.1 Distinguishing Breast Cancer Tumor Subclasses of Gene Expression Patterns with Clinical Implications

#### 4.1.1 Background

#### 4.1.2 Dataset in Detail

#### 4.1.3 Identification of Tumor Subtypes by Hierarchical Clustering

#### 4.1.4 Correlation to Clinical Results

### 4.2 Results & Discussion



## Chapter 5

# Time Plan



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