**BONAFIDE CERTIFICATE**

Certified that this Project report titled “**PATTERN MATCHING IN BIOLOGICAL DATA TO DETECT ABNORMALITIES**” is the bonafide work of “**VINAY KUMAR REDDY K, ARVIND SRINATH K, ELAVARASAN A J”** undermy supervision in partial fulfilment for the award of the degree of Bachelor of Engineering in Computer Science and Engineering. Certified further that to the best of my knowledge the work reported here in does not form part or full of any other thesis or dissertation on the basis of which a degree or award was conferred on an earlier occasion to this or any other candidate.

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**INTRODUCTION**

In the past decade though whole genome sequencing is completed to analyse the genome due its vast requirement of memory for storing the genome while processing the genome to exploit the various properties of DNA. Now, with the introduction of Big Data it became easier to handle the date and many algorithms have been proposed for analysing the genome which led to detection that the tandem repeats play an important role in abnormality. If we can detect the abnormalities earlier and accurately it would aid for getting treatment which could save many lives.

**DEOXYRIBONUCLEIC ACID (DNA)**

Deoxyribonucleic acid is molecule that contains the genetic instructions used in the growth, development, functioning and reproduction of all known living organisms and many viruses. It is made up of two strands called polynucleotides. Since they are made of simple monomers called nucleotides. Nucleotides are mainly made up of four components Adenine (A), Guanine (G), Cytosine (C) and Thymine (T).

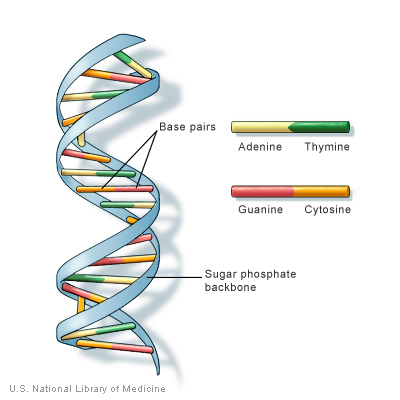


Fig. 1 - Sample DNA Strand

**TANDEM REPEATS:**

Tandem repeats occur in DNA when one or more nucleotides are repeated and the repetitions are adjacent to each other. In most organisms, the bulk of the intergenic DNA is made up of repeated sequences. Repetitive DNA can be divided into two categories: interspersed repeats, where repeat units are distributed in the genome in an apparently random fashion, and tandem repeats, where repeat units are placed next to each other in an array. The functions of DNA repeats are still largely unclear, but, in recent years, it has been found that a number of tandem repeats are related to diseases and play an important role in gene regulation.

Next-generation sequencing (NGS) machines do not read out an entire genomic template. Instead, they produce short random fragments of nucleotide sequences known as sequence reads. A quality score, expressing the confidence that a particular nucleotide has actually been found at a specific location, is associated with each nucleotide in a sequence read. This raw sequencing data generated by NGS machines is commonly stored in FASTQ files. Recently, several tools for FASTQ compression have been developed, which – without performing any kind of internal alignment or assembly – are approaching the Kolmogorov complexity. FASTQ data can be aligned with tools such as BWA, SAMtools or Bowtie2. Subsequently, the data is stored in Sequence Alignment/Map (SAM) format files which, compared to FASTQ files, contain the alignment/mapping information as well as additional metadata. Mapped sequencing data represented in SAM files contains more redundancy, as multiple sequence reads are typically mapped to the same location on the donor genome. The average amount of reads mapping to the same location is referred to as coverage.

**VARIABLE NUMBER TANDEM REPEATS**

Is a location in an genome where a small nucleotide is organised as a tandem repeat. They often show variations in length among each individual.

**QUALITY SCORES**

The FastQ format provides a simple extension to the FastA format, and stores a simple numeric quality score with each base position. Despite being a “standard” format, FastQ has a number of variants, deriving from different ways of calculating the probability that a base has been called in error, to different ways of encoding that probability in ASCII, using one character per base position.

**PHRED scores**

Quality scores were originally derived from the PHRED program which was used to read DNA sequence trace files, and linked various sequencing metrics such as peak resolution and shape to known sequence accuracy. The PHRED program assigned quality scores to each base, according to the following formula:

Q\_PHRED=-10log10(pe)

where Pe is the probability of erroneously calling a base. PHRED put all of these quality scores into another file called QUAL (which has a header line as in a FastA file, followed by whitespace-separated integers. The lower the integer, the higher the probability that the base has been called incorrectly.

While scores of higher than 50 in raw reads are rare, with post-processing (such as read mapping or assembly), scores of as high as 90 are possible. Quality scores for NGS data are generated in a similar way. Parameters relevant to a particular sequencing chemistry are analyzed for a large empirical data set of known accuracy. The resulting quality score lookup tables are then used to calculate a quality score for de novo next-generation sequencing data.

**Solexa scores**

The Solexa quality scores, which were used in the earlier Illumina pipelines, are calculated differently from the PHRED scores:

Q-SOLEXA=-10log10(pe/1-pe)

**CLASSIFICATION**

Instead of processing the genome sequence which is made of millions of base pairs it would be better if it is processed once the repetitive parts are removed which would increase efficiency in both time and cost.

**PROBLEM DEFINITION**

Though human genome is more than 99 percent common in all humans the current available methods of processing biological data reprocess the complete sequence (on average of 3 million base pairs) each time. Human analysis of the genome is more error prone.

**OBJECTIVE**

When processing the data it is enough to consider only the parts of genome that has changes when compared to reference genome. Integration of intelligence in order to analyse the genome in order to find any abnormalities.

**LITERATURE SURVEY**

The authors **“Zhang, Jingsong, Yinglin Wang, Chao Zhang, Yongyong “**  described a paper titled **“Mining contiguous sequential generators in biological sequences”**.In this paper they have designed an algorithm to remove the repeated sequences. The advancement in the big data field has led to design of algorithms to process the sequential data but they were not considering the fact that the genome is made up of repetitive segments called tandem repeats. To overcome this situation they have designed an algorithm called ConSgen [1] an effective algorithm for discovering contiguous sequential generators. It adopts the n-gram model, called shingles, to generate potential frequent subsequence’s and leverages several pruning techniques unpromising parts of search space. The algorithm works on three-step manner. In the first step, each sequence is split into a series of snippets which preserve the original ordering of items. The length of the snippets is fixed within one complete scan, while it incrementally increases by step length 1 according to n-gram model in subsequent scans. In the next step, by exploring the some properties of contiguous sequential generators, three effective pruning techniques, redundant snippet pruning, max-prefix-suffix pruning and support pruning are proposed. In the third step, all contiguous sequential non-generators are distinguished successively from the set of contiguous sequential patterns by performing the lower-cluster checking. The problem is to check whether there exist a super-pattern absorbing smaller patterns. **Redundant snippet pruning** In real-world datasets, some snippets often appear multiple times, not only among different sequences but also inside one sequence. For each newly split snippet, we check the previous snippets to see whether the new one already exists. The repeated snippets can be easily identified and discarded on-the-fly. **Max-prefix-suffix pruning** Unlike classic candidate enumerate-and-test paradigm, max-prefix-suffix pruning, for a new length-k snippet s, does not need to check all its subsequences if there exist an infrequent one. Instead, it checks only the frequency of the max-prefix-suffixes of s. The length-k snippet is pruned immediately if its max-prefix or max-suffix is infrequent. **Support pruning** A snippet is called a promising candidate if it meets: (1) it is distinct from the previous snippets (already split snippets); and (2) both the max-prefix and the max-suffix of the snippet are frequent. Such snippets satisfying the above two conditions can be shifted to count their supports and check whether they are frequent.

The authors **“Zhou, Hongxia, Liping Du, Hong Yan”** described a paper titled **"Detection of tandem repeats in DNA sequences based on parametric spectral estimation".** In this paper they explain an algorithm using a spectral method. Tandem repeats [2], which occur frequently in genomes, are related to the gene regulatory function and various diseases. Since a DNA sequence consists of ordered nucleotides represented as letters A, C, T, and G, string matching methods including those based on the Hamming or edit distance, data compression, and color coding, are useful for tandem repeat analysis. The algorithms that have been proposed used Hamming distance and edit distance. However, the main limitation of these algorithms is excessive running time since a genome sequence can easily consist of millions of neucleotides. In order to reduce the run-time, heuristic algorithms have been proposed however these algorithms are not sufficient by themselves because they require a period of the repeat or the basic pattern of the region to be specified in advance. To overcome this problem a computer program known as tandem repeat finder (TRF) has been proposed which works based on probabilistic model. It consists of a screening phase and a verification phase. Algorithms, other than these measuring the sequence edit distance, may also be used. If the four letters A, C, T, and G, in a DNA sequence are assigned numerical values, then signal processing operations can be used for tandem repeats. In the tetrahedral representation, each of the four letters is assigned to a vertex of a regular tetrahedron in space. An optimal criteria like use of a weight vector to obtain the sequence can also be used. Numerical representation of DNA sequences opens up the possibility of applying signal processing techniques to the analysis of genomic data. Recently, signal processing methods have also been used for DNA repeat identification. These techniques include the discrete Fourier transformation (DFT), the short time periodicity transfer (STPT), and the exact periodic subspace decomposition (ESPD). A computer program called spectral repeat finder is also developed for this purpose. An efficient method for tandem repeat detection using spectrogram of a DNA sequence is analysed based on the autoregressive model. Most signal -processing-based methods use the FT in spectral analysis for tandem repeats detection since they are able to process a large amount of data quickly. However, the FT is well known to produce data truncation articrafts and poor resolution for spectral analysis. To overcome this problem a repeat detection method based on autoregressive (AR) model. It can offer a high resolution since it has the capability of extrapolating the autocorrelation function of the input.

The authors **“Deng, Yue, Zhiquan Ren, Youyong Kong, Feng Bao, Qionghai Dai”** described a paper titled **"A hierarchical fused fuzzy deep neural network for data classification".** In this paper they have designed a DNN for learning deep features of DNA.Over the past few years there is so much growth in the field of bioinformatics with the completion of genome sequencing. As the analysis of genome is unveiling various abnormalities. deep learning method has been widely used in many fields of science researches. Many different deep learning models are applied for different applications, e.g., deep neural networks (DNN)[3] and convolutional neural network (CNN) . DNN has a classical auto encode (AE) structure which can be used to extract deep feature automatically. An AE is composed with several stacked RBM, The RBM is an undirected graphical model that defines the distribution of visible units using binary hidden units. In training process, each RBM can be trained independently and the output of the bottom RBM can be used as the input data of the upper RBM. After getting all the parameters for each layer, they can get a DNN to carry out feature extraction and dimension reduction. Because the input unit is binary, the input data sent to the input unit should be bounded from 0 to 1. This peculiarity just adapts the characteristic of DNA methylation data. The output getting from equation 4 are also bounded, which makes the features finally extracted from the top layer are still in accordance with the characteristics of methylation data. In order to visually assess the effect of feature selection, they adopted the distributed stochastic neighbour embedding (t-SNE)[3] method to visualize the high-dimensional data in two-dimensional space. t-SNE is a nonlinear dimensionality reduction technique which has been used in a wide range of applications, including computer security research, music analysis, cancer research, and bioinformatics. It is particularly well suited for embedding high-dimensional data into a space of two or three dimensions, which can then be visualized by a scatter plot. Especially, it models each high-dimensional object by a two- or three-dimensional point in such a way that similar objects are modelled by nearby points and dissimilar objects are modelled by distant points. In order to make fair comparisons, they also applied the k-means, the Gaussian mixture model (GMM), and the SOM method to cluster the features extracted by PCA, NMF, and DNN methods, respectively.

The authors **“Gosztolya, Gábor, László Tóth”** described a paper titled **"DNN-based Feature Extraction for Conflict Intensity Estimation from Speech"**. In this paper they have designed a DNN for Speech recognition. The standard computational approach, developed over the years on various paralinguistic tasks[4], is to extract several thousand general, utterance-level features from the speech excerpts, and use these to train general machine learning methods such as Support-Vector Machines (SVM) or Deep Neural Networks (DNNs) to perform classification or regression. In the DNN each neuron corresponds to a particular person. They have proposed to use only two classes, corresponding to a zero-or-one speaker, and a two-or-more speaker case. Next, they extracted features from the frame-level DNN outputs, and these features will be used for a specific time window instead of the whole utterance. Although using the amount of speaker overlap may prove to be beneficial for conflict intensity estimation, we should not discard all other features.

The authors **“Ramakrishnan, Nithya, Ranjan Bose”** described a paper titled **"Analysis of healthy and tumour DNA methylation distributions in kidney-renal-clear-cell-carcinoma using Kullback–Leibler and Jensen–Shannon distance measures".** They have praposed an algorithm for detecting kidney related cancer using kullback-leibler and jensen-shannon distance measurement. DNA methylation [5] ia an epigenetic phenomenon in which methyl groups gets bounded to the 5-carbon of the cytosines ring to result in 5-methyl cytosine (5-mC) of the DNA molecule altering the expression of the associated genes. Abnormalities like cancer is linked with hypo or hyper-methylation of specific genes as global changes in DNA methylation. In human somatic cells, 5-mC occurs in CpG sites and islands. A CpG site in a location within a DNA sequence in which a cytosine and guanine appear consecutively. When a CpG island island in the promoter region of a gene is methylated, the gene expression is switched off. DNA methylation patterns are either inherited (maintenance) or developed newly (de-novo methylation). Irregular changes in DNA methylation have been linked with many abnormalities. Based on the kullback-leibler and jensen-shannon distance measurement a popular abnormality kidney-renal-clear-cell-carcinoma is detected.

The authors **“Voges, Jan, Marco Munderloh, Jörn Ostermann”** described a paper titled **"Predictive coding of aligned next-generation sequencing data".** They have designed an algorithm for high throughput next-generation sequencing. Due to novel high-throughput next-generation sequencing technologies, the sequencing of huge amounts of genetic information has become affordable. On account of this flood of data, IT costs have become a major obstacle compared to sequencing costs. High-performance compression of genomic data is required to reduce the storage size and transmission costs. The high coverage inherent in next-generation sequencing technologies produces highly redundant data. Due the above reason there is a need for the design of a compression algorithm for aligned sequence reads. They have proposed an algorithm TCS[6]. The algorithm is solely requires SAM files which contain regions that are sorted by their mapping positions. The algorithm is based on the unwinding of the sequence reads by using the mapping positions and the additional alignment information provided by the CIGAR strings. The process of expanding the nucleotide sequence yields a modified CIGAR string, which they call stogy, and the expanded nucleotide sequence, called exs. We call this mapped file as tcs file. After encoding the first read, the mapping position pos and the expanded read exe are pushed to a circular buffer which implements the sliding window. The tsc records are a new sequence read representation are ultimately passed to a dictionary-based code in order to exploit the redundancy along the record stream. Specifically, error-free and properly aligned sequence reads produce a tsc record stream which is highly suitable for dictionary-based compression because of repetitions in the data stream and the compression is based on the DEFLATE algorithm which uses Huffman coding and LZ77 algorithm.

The authors **“Deng, Yue, Zhiquan Ren, Youyong Kong, Feng Bao, Qionghai Dai”** described a paper titled **"A hierarchical fused fuzzy deep neural network for data classification".** In this paper they have described a fuzzy logic and deep learning to extract more information and to provide more accurate results. Typical deep learning is a fully deterministic model that sheds no light on data uncertainty reductions. In this paper, they show how to introduce the concepts of fuzzy learning[7] into DL to overcome the shortcomings of fixed representation. The bulk of the proposed fuzzy system is a hierarchical deep neural network that derives information from both fuzzy and neural representations. Then, the knowledge learned from these two respective views are fused altogether forming the final data representation to be classified.

The authors **“Azim, Md Aashikur Rahman, Costas S. Iliopoulos, M. Sohel Rahman, M. Samiruzzaman”** described a paper titled **"A Simple, Fast, Filter-Based Algorithm for Approximate Circular Pattern Matching".** In this paper they have explained an algorithm forn circular pattern matching. They have also explained other algorithms for this purpose and their advantaged and disadvantages. String matching or pattern matching problem is a classical problem in computer science with extensive applications in different branches of science and engineering. This problem is interesting as a fundamental computer science problem and is a basic requirement of many practical applications. In this paper, they present a fast and efficient algorithm for the approximate circular pattern matching problem based on some filtering techniques. They presented SimpLiFiCPM that is a filter-based algorithm to handle the exact version of the circular pattern matching problem. Given a pattern P of length m and a text T of length n, both being sequences of characters drawn from a finite character set , the goal of this problem is to find all the occurrences of P in T. They use SFF (Simple Fast Filter-based Algorithm) [8] to match the sequence.

The authors **“Pérez, Sandino Vargas, Fahad Saeed”** described a paper titled "**A parallel algorithm for compression of big next-generation sequencing datasets".** In this paper they have described a method for compressing the huge amount of DNA sequence. With the advent of high-throughput next-generation sequencing (NGS) [9] techniques, the amount of data being generated represents challenges including storage, analysis and transport of huge datasets. One solution to storage and transmission of data is compression using specialized compression algorithms. They have designed an parallel processing algorithm for compressing the DNA files. However, these specialized algorithms suffer from poor scalability with increasing size of the datasets and best available solutions can take hours to compress Gigabytes of data. In this paper we introduce paraDSRC, a parallel implementation of DSRC using a message passing model that presents reduction of the compression time complexity by a factor of O(1/p). Our experimental results show that paraDSRC achieves compression times that are 43% to 99% faster than DSRC and compression throughputs of up to 8.4GB/s on a moderate size cluster. For many of the datasets used in our experiments super-linear speedups have been registered, making the implementation strongly scalable. They also show that paraDSRC is more than 25.6x faster than comparable parallel compression algorithms.

The authors **“Ku, Chee Seng, En Yun Loy, Agus Salim, Yudi Pawitan, Kee Seng Chia”** described an article titled **"The discovery of human genetic variations and their use as disease markers: past, present and future"**. In this article they have explained the genetic markers that were used in the past, in the present and in future. The field of human genetic variations has progressed rapidly over the past few years. It has added much information and deepened our knowledge and understanding of the diversity of genetic variations in the human genome. This significant progress has been driven mainly by the developments of microarray and next generation sequencing technologies. The array-based methods have been widely used for large-scale copy number variation (CNV) [10] detection in the human genome. The arrival of next generation sequencing technologies, which enabled the completion of several whole genome resequencing studies, has also resulted in a massive discovery of genetic variations. These studies have identified several hundred thousand short indels and a total of thousands of CNVs and other structural variations in the human genome. The discovery of these ‘newer’ types of genetic variations, indels, CNVs and copy neutral variations (inversions and translocations) has also widened the scope of genetic markers in human genetic and disease gene mapping studies. In this article they have summarize the latest developments in the discovery of human genetic variations and address the issue of inadequate coverage of genetic variations in the current genome-wide association studies, which mainly focuses on common SNPs.

**PROPOSED WORK**

1. Design and Implementation of an algorithm to remove the tandem repeats that make the most part of the genome which in turn reduces the memory and time while analysing the genome for abnormalities.
2. To find the count of tandem repeats in a genome by using an open source tool.
3. Design and Implementation of an algorithm to detect whether there are abnormalities.
4. Design and Implementation of a Deep Neural Network(DNN) to find the type of abnormality and the stage of abnormality (if any).

**TOOLS USED**

* **Galaxy**
* **RepeatExplorer**

**Introduction**

Galaxy is a web based analysis and workflow platform designed for biologists to analyse their own data. It comes with most of the popular bioinformatics tools already installed and ready for use. There are many Galaxy servers around the world and some are tailored with specific toolsets and reference data for analysis of human genomics, microbial genomics, proteomics etc.

**Upload Data into Galaxy**

There are two main ways to get your data into Galaxy

1. **Upload a file from your own computer**

* From the Galaxy tool panel, click on **Get Data -> Upload File**
* Click the Choose File button
* Find and select the Contig\_stats.txt.gz file you downloaded and click Open
* Set the “file format” to tabular
* Click the Start button
* Once the progress bar reaches 100%, click the Close button

1. **Upload a file from a URL**

* From the Galaxy tool panel, click on **Get Data -> Upload File**
* Enter the URL into the Text box
* Set the file format to fastqsanger
* Click Start
* Once the progress bar has reached 100%, click Close

1. Convert Fastq file to sangerFastq sequences

* Goto the TOOLS panel
* Select NGS:QC and Manipulation
* Select Fastq groomer, Select input file and click Execute.

1. **Quality Check**

* Goto the TOOLS panel
* Select NGS:QC and Manipulation
* Select FastQC, Select input file and click Execute.

1. **Quality Filter**

* Go to NGS:QC and Manipulation
* Select Filter FastQ, Select input file and click Execute

1. Mapping/Short Read Alignment

* Go to NGS:Mapping
* Select Bowtie2
* Select weather the data is single or paired end
* Select the filtered fastq file
* Click yes on write unaligned reads (in fastq format) to separate file(s) to separate unaligned reads to separate file
* Select Build-in genome index option and select corresponding reference genome name.
* Specify the read group for the file by clicking on yes and specify sample name, library, readgroup id, and platform .
* Select analysis mode as default settings.
* Click on Execute

1. SAM/BAM File Manipulation

* Filter **SAM/BAM** file using **SAMTools**
* Go to **NGS:SAMTools**
* Select **Filter SAM/BAM**
* Select input file and click **Execute.**
* Mark duplicates using picard
* Go to **NGS:picard**
* Select **Mark Duplicates**
* Select file from history, and click **yes** in **do not write duplicates to the output file** option
* Click on **Execute.**
* Remove PCR duplicates from SAM/BAM file
* Go to **NGS:SAMTools**
* Select **RmDup**
* Select BAM dataset and specify whether data is single or paired end
* Click on **Execute.**

1. VCF File Creation

* Go to **SAMTools**
* Select **MPileup**
* Select Reference genome
* Select BAM file.
* If Perform **Genotype Likelihood Computation** is selected, output will be in BCF/VCF. If do not perform **Genotype Likelihood Computation** is selected, output will be in pileup format.
* If we select **Perform INDEL calling** or **do not Perform INDEL calling** , we can perform INDEL calling and SNP calling together or SNP calling alone respectively.

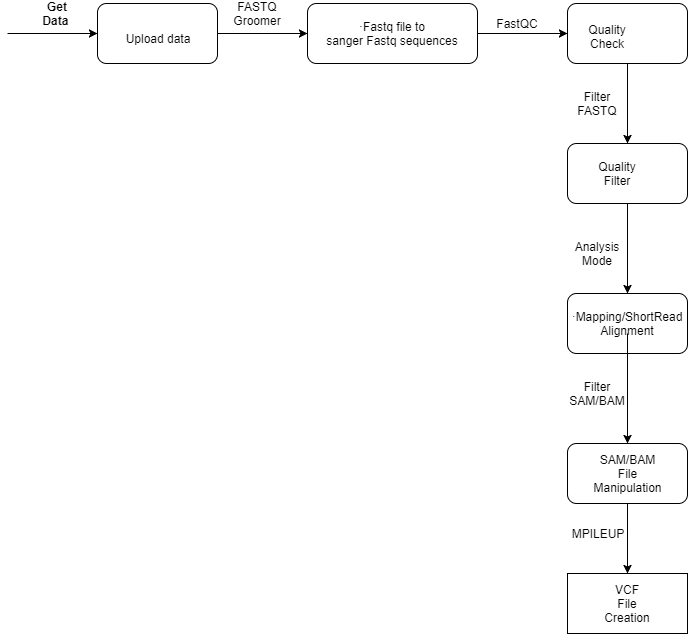


Fig. 2 – Flow diagram of removing repeating patterns

**RepeatExplorer**

RepeatExplorer is a computational pipeline for discovery and characterization of repetitive sequences in eukaryotic genomes. The pipeline uses high-throughput genome sequencing data as an input and performs graph-based clustering analysis of sequence read similarities to identify repetitive elements within analyzed samples. It should be noted that although the repeat identification algorithm generally works for any genome, some parts of the pipeline (e.g. protein domain-based classification of mobile elements) were primarily developed for application to plant genomics. However, there is a possibility to supply a custom repeat database to improve sensitivity in classification of non-plant repeats.

**Steps**

* **Getting your data to server**
* Datasets can be downloaded directly from the EBI Short Read Archive using **Get Data 🡪 EBI SRA** tool. Enter the ENA accession number in the search window, locate the corresponding dataset and select download link in the "Galaxy" column
* **Pre-processing of sequence reads**
* The clustering analysis requires a single file containing read sequences in FASTA format as an input.
* If such a file can be uploaded by the user, no pre-processing is required
* However, data obtained from sequencing facilities or downloaded from public archives are usually in FASTQ format combining nucleotide sequence information with sequencing quality scores.
* There is a number of programs for analysing and pre-processing raw sequence reads in **Tools 🡪 NGS: QC and manipulation 🡪 FastQC:Read QC**
* **Grooming the data**
* **The FASTQ illumine format must be converted to FASTQ sanger format using  Tools 🡪 NGS: QC and manipulation 🡪 FASTQ Groomer**
* **Trim the data**
* **If there is any discrepancy in the sequence then the sequence should be trimmed using Tools 🡪 NGS: QC and manipulation 🡪 FASTQ Trimmer**
* **FASTQ to FASTA**
* **The FASTQ format is converted into FASTA using Tools 🡪 NGS: QC and manipulation 🡪 FastQC:Read QC 🡪 FASTQ to FASTA converter**
* **TAREAN**
* Tandem repeat analyser (TAREAN), a computational pipeline which was built on the principles of graph-based repeat clustering, enhanced and supplemented with additional tools facilitating unsupervised identification and characterization of satellite repeats from unassembled sequence reads
* The pre-processed FASTA file is passed into **Tools 🡪 TAREAN 🡪 Tandem Repeat Analyser**

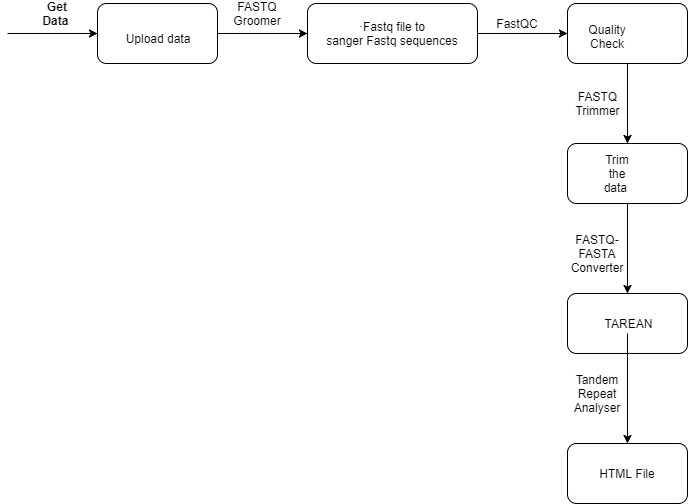


Fig. 3 – Flow diagram of tandem repeat analysing

**PROGRAMING LANGUAGE USED**

**Java**

**JAVA**

**Java** is a general-purpose [computer programming language](https://en.wikipedia.org/wiki/Programming_language) that is [concurrent](https://en.wikipedia.org/wiki/Concurrent_computing), [class-based](https://en.wikipedia.org/wiki/Class-based_programming), [object-oriented](https://en.wikipedia.org/wiki/Object-oriented_programming), and specifically designed to have as few implementation dependencies as possible. It is intended to let application developers "[write once, run anywhere](https://en.wikipedia.org/wiki/Write_once,_run_anywhere)" (WORA), meaning that [compiled](https://en.wikipedia.org/wiki/Compiler) Java code can run on all platforms that support Java without the need for recompilation. Java applications are typically compiled to [bytecode](https://en.wikipedia.org/wiki/Java_bytecode) that can run on any [Java virtual machine](https://en.wikipedia.org/wiki/Java_virtual_machine) (JVM) regardless of [computer architecture](https://en.wikipedia.org/wiki/Computer_architecture). As of 2016, Java is one of the most [popular programming languages in use](https://en.wikipedia.org/wiki/Measuring_programming_language_popularity), particularly for client-server web applications, with a reported 9 million developers. Java was originally developed by [James Gosling](https://en.wikipedia.org/wiki/James_Gosling) at [Sun Microsystems](https://en.wikipedia.org/wiki/Sun_Microsystems)(which has since been [acquired by Oracle Corporation](https://en.wikipedia.org/wiki/Sun_acquisition_by_Oracle)) and released in 1995 as a core component of Sun Microsystems' [Java platform](https://en.wikipedia.org/wiki/Java_(software_platform)). The language derives much of its [syntax](https://en.wikipedia.org/wiki/Syntax_(programming_languages)) from [C](https://en.wikipedia.org/wiki/C_(programming_language)) and [C++](https://en.wikipedia.org/wiki/C%2B%2B), but it has fewer [low-level](https://en.wikipedia.org/wiki/Low-level_programming_language) facilities than either of them. Excel is the very popular file format created by Microsoft. Although it is not an opened file format, Java applications can still read and write Excel files using the **Apache POI - the Java API for Microsoft Documents**, because the development team uses reverse-engineering to understand the Excel file format. Hence the name POI stands for Poor Obfuscation Implementation.

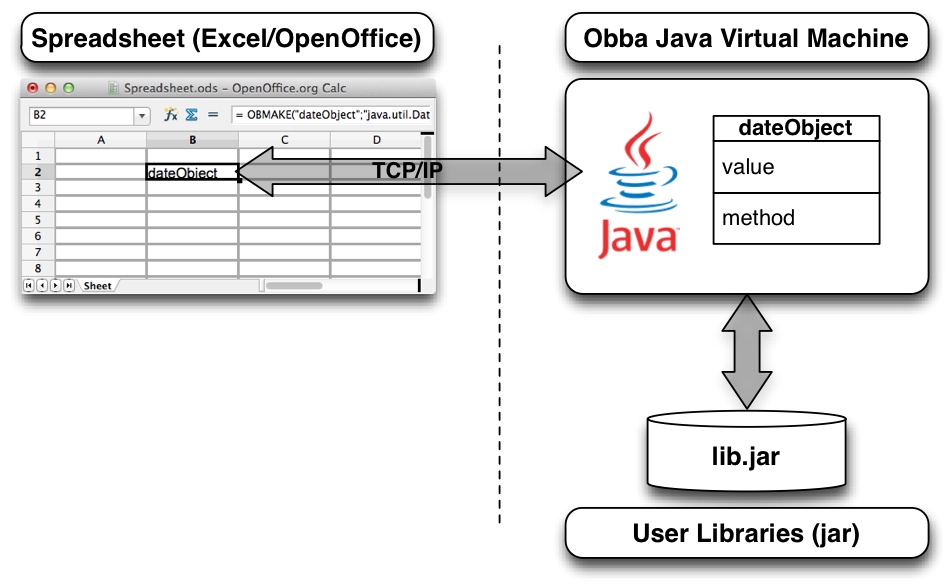
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Fig. 4 – Architecture of Java integration with Excel Sheet

**ARCHITECTURE DIAGRAM**

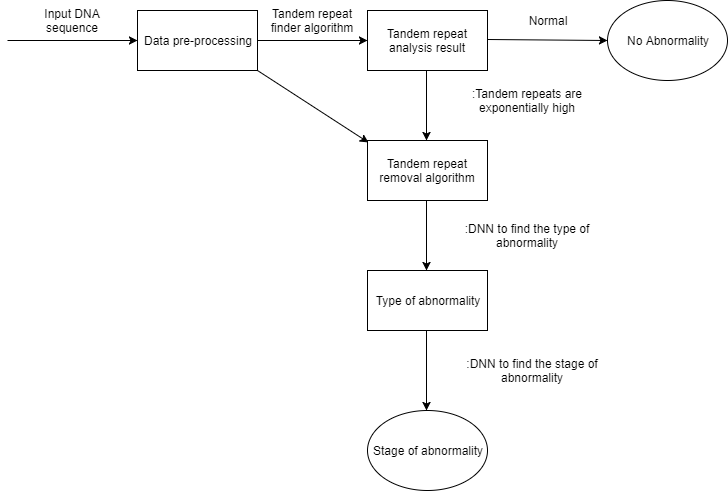
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Fig. 5 – Architecture diagram of the proposed project

**ALGORITHM**

**Abnormality Type Detection algorithm**

Step 1: Extract the features from the vcf file.

Step 2: For each record in the vcf file read the different parameters.

Step 3: Once the parameters are extracted based on the values of the different parameters the records are compared with the reference records of the human genome (Since we are trying to detect the abnormalities in human genome).

Step 4: For each record assign the reference gene.

Step 5: Write the record into an flat file.

Step 6: Open the flat file.

Step 7: For each record in the flat file read the gene.

Step 8: Compare it with the reference Genes of abnormalities which is stored in another flat file.

Step 9: If any matching is found then display the abnormality.

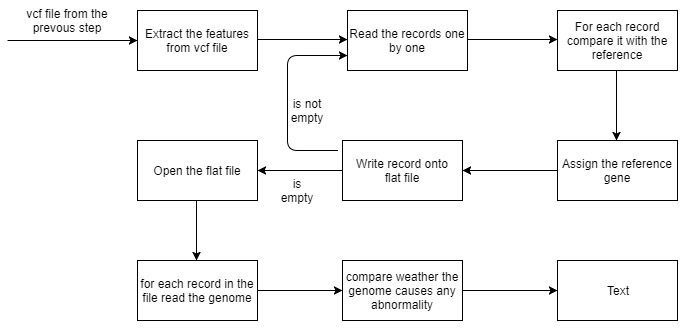
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Fig 6 – Flow diagram of abnormality type detection algorithm

**IMPLEMENTATION**

The input is a DNA sequence which is generated based on one of the standard method of DNA sequencing. This file contains a lot of unnecessary data that need to be removed for this purpose the DNA sequence need to be filtered before doing any processing. By using the galaxy tool this can be done. Once the useful sequence is checked for quality and the base pairs that have a low quality are discarded which leads to better accuracy.

Quality Score is assigned to each base pair using any one of the Scoring methods as follows.

Quality Score Q\_PHRED=-10log10(pe)

Or

Q-SOLEXA=-10log10(pe/1-pe)

Then the Sequence is aligned according to Bowtie2 algorithm. Galaxy an online tool is used to perform the quality check and bowtie2. The steps for processing is shown in tools used.

The sequence is then sorted, marking and removing of duplicates are performed subsequently. In order to remove the tandem repeats by which unnecessary processing of data is eliminated.

Samtools is used to perform the sorting, removing duplicated and piling up the input sequence with the reference genome by using the following commands.

samtools sort filename.bam

samtools rmdup filename.bam

Once the data has got rid of repeats it is compared with the reference genome by using the following command.

samtools mpileup input.bam output.bcf

This file needs to be converted to vcf format because the bcf format is compressed for this purpose another tool called bcftools is used. The command for this is

bcftools view -c -v -g input.bcf ouput.vcf

On the other hand we need to check for tandem repeats for the basic confirmation of any abnormality. For this purpose we use the Galaxy Repeat Explorer. By analysing the result we know whether there is an exponential growth in tandem repeats.

The vcf file is then compared with the reference human data of chromosomes inorder to know the gene expression based on the features of vcf file. Now this reference Gene is compared with that of abnormal genes so that we can find what kind of abnormality it is.

**MODULES**

**Phase - I**

1. Study on DNA properties and their use in bioinformatics
2. Data Pre-processing and deletion of tandem repeats.
3. Detection of abnormality.

**Phase - II**

1. Study on the design and training of DNN’s.
2. Design of Deap Neural Network (DNNs) to find the type and stage of abnormality.

**RESULTS AND ANALYSIS**

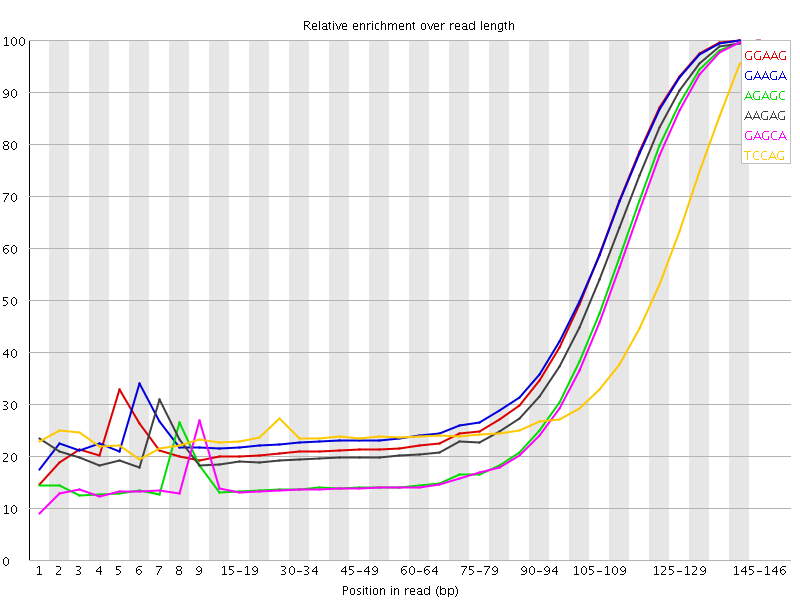


Fig. 7 – K-Mer of the input data

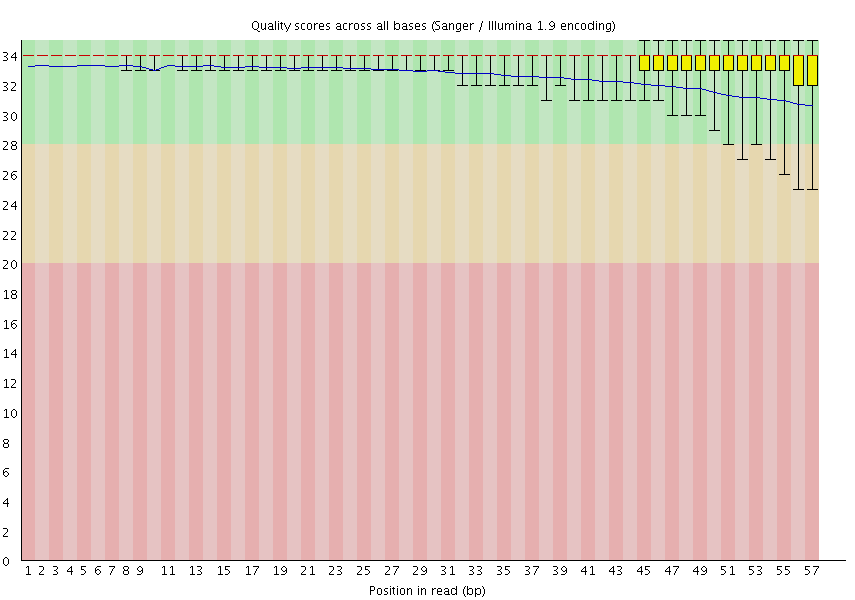
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Fig. 8 – Per base quality of input

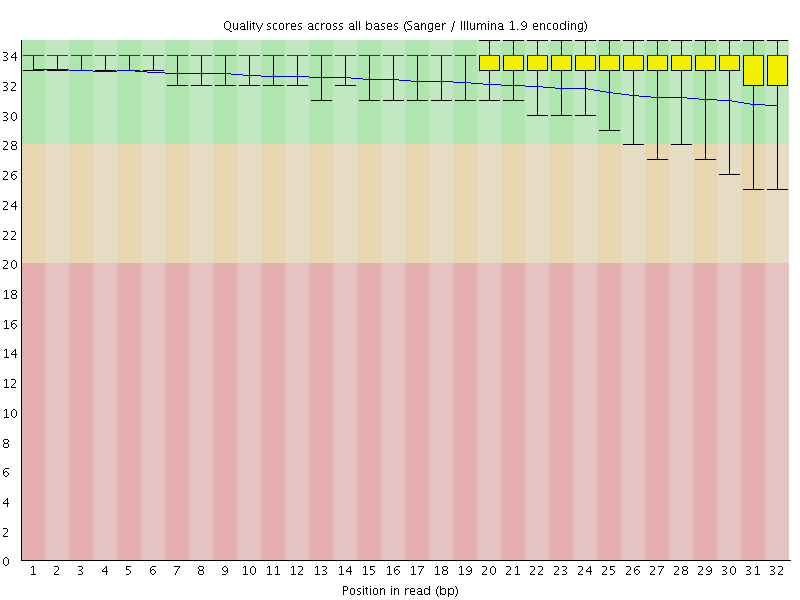
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Fig. 9 – per base quality after pre-processing

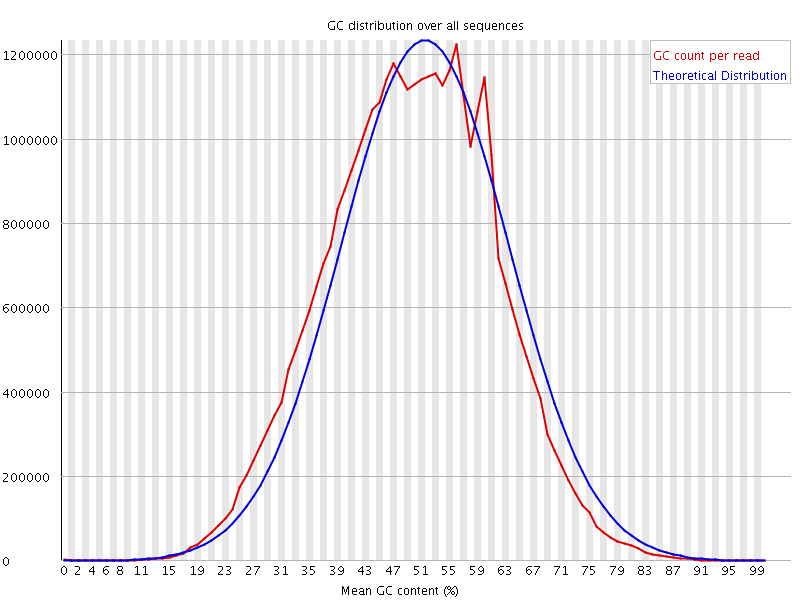
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Fig. 10 – per sequence gc count of input

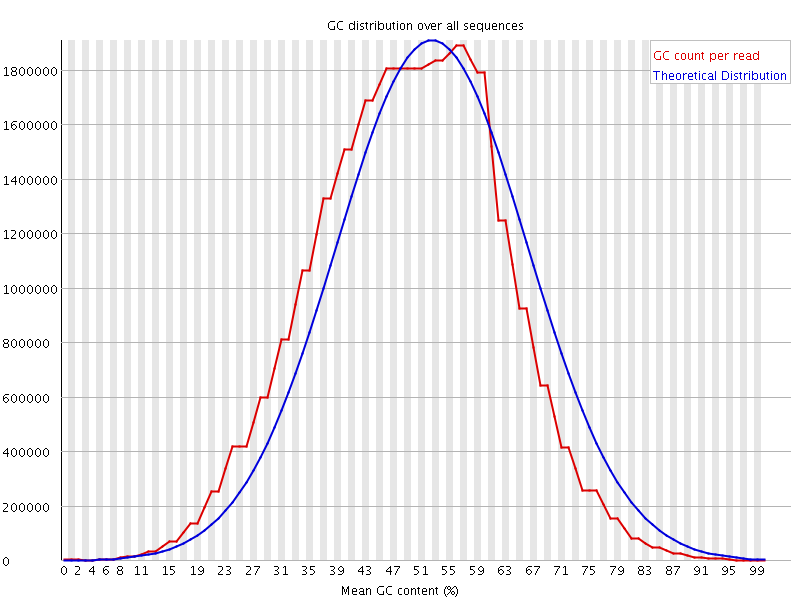
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Fig. 11 – Per sequence gc count after pre processing

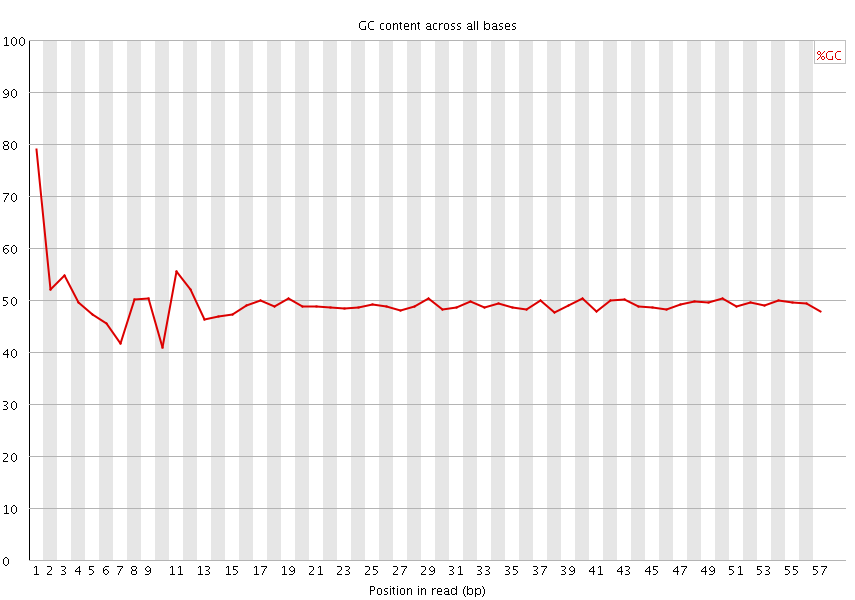
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Fig. 12 – per base gc count of input data

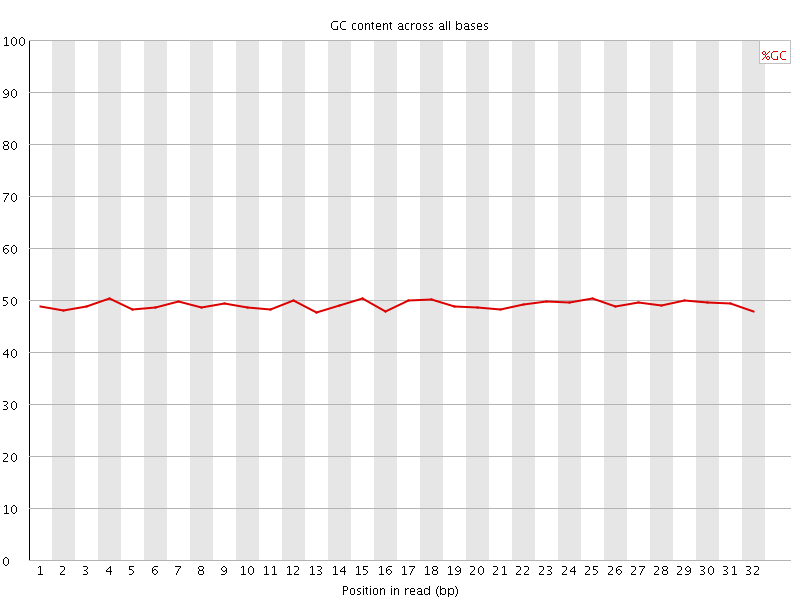
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Fig. 13 – per base gc count after pre processing

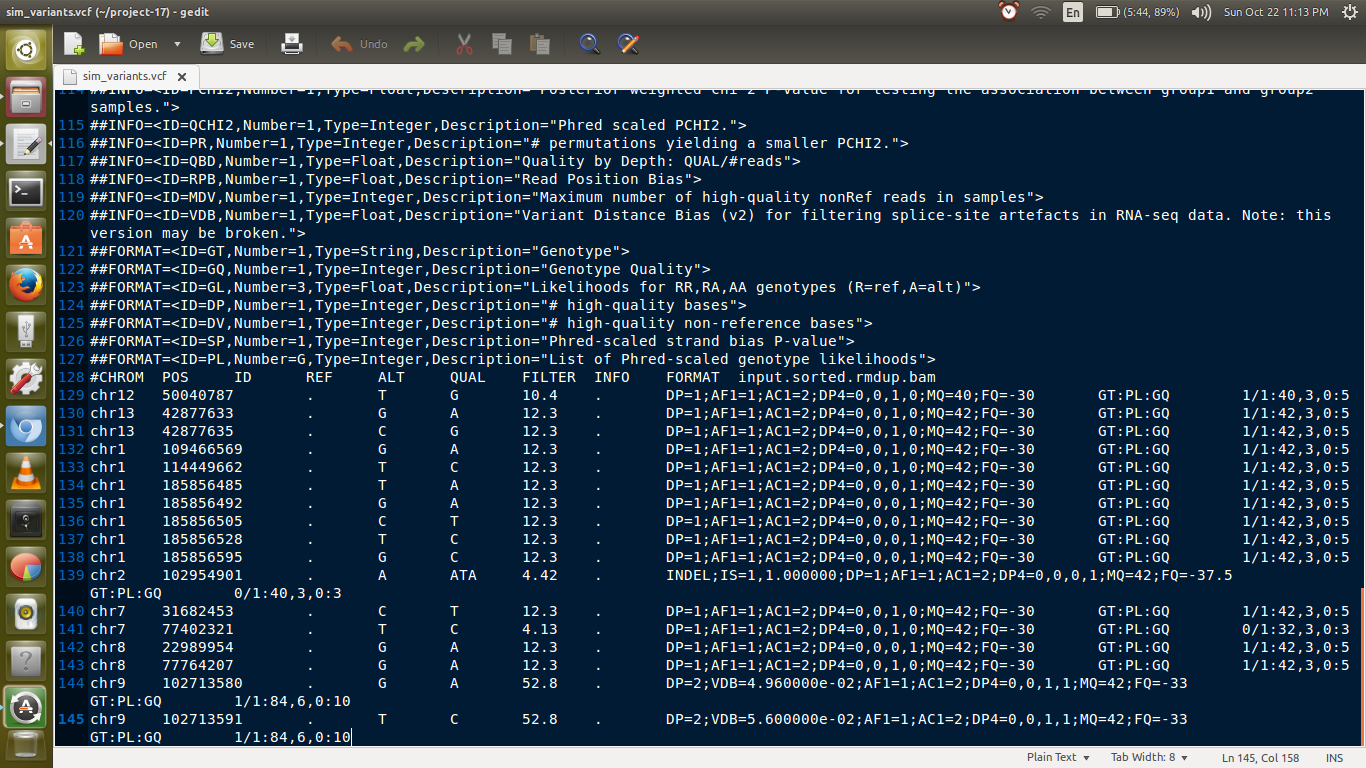
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Fig. 14 – VCF file generated after removal and mpileup

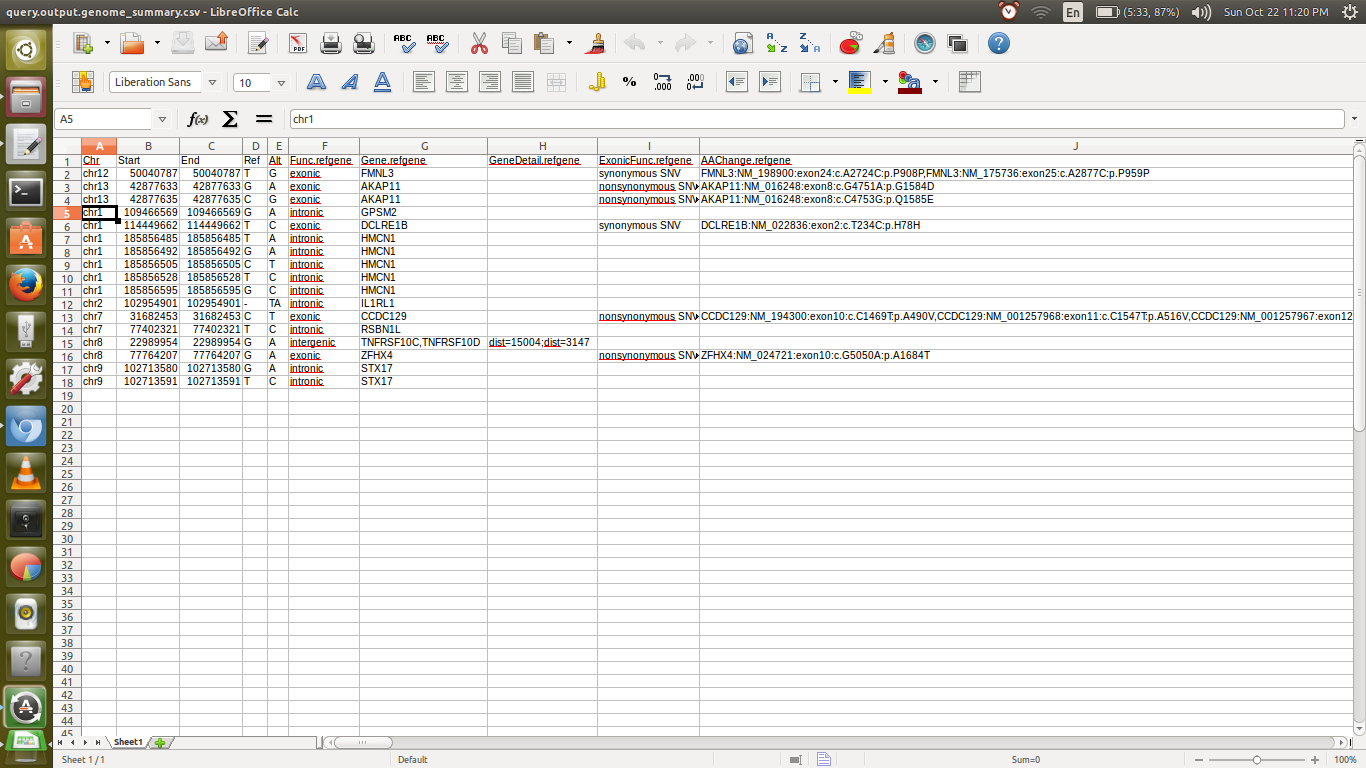
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Fig. 15 – Output of the type of gene for each record in the VCF file

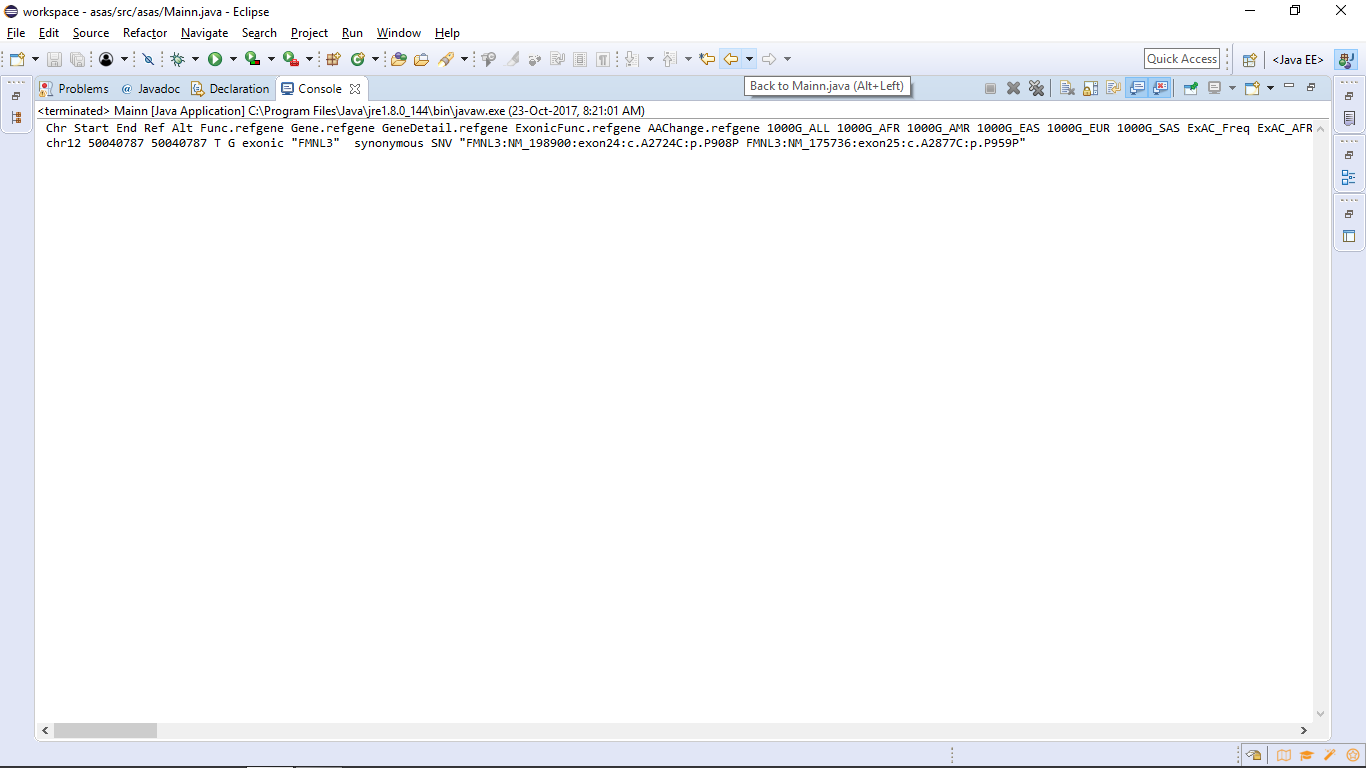


Fig. 16 – Output of abnormality detection

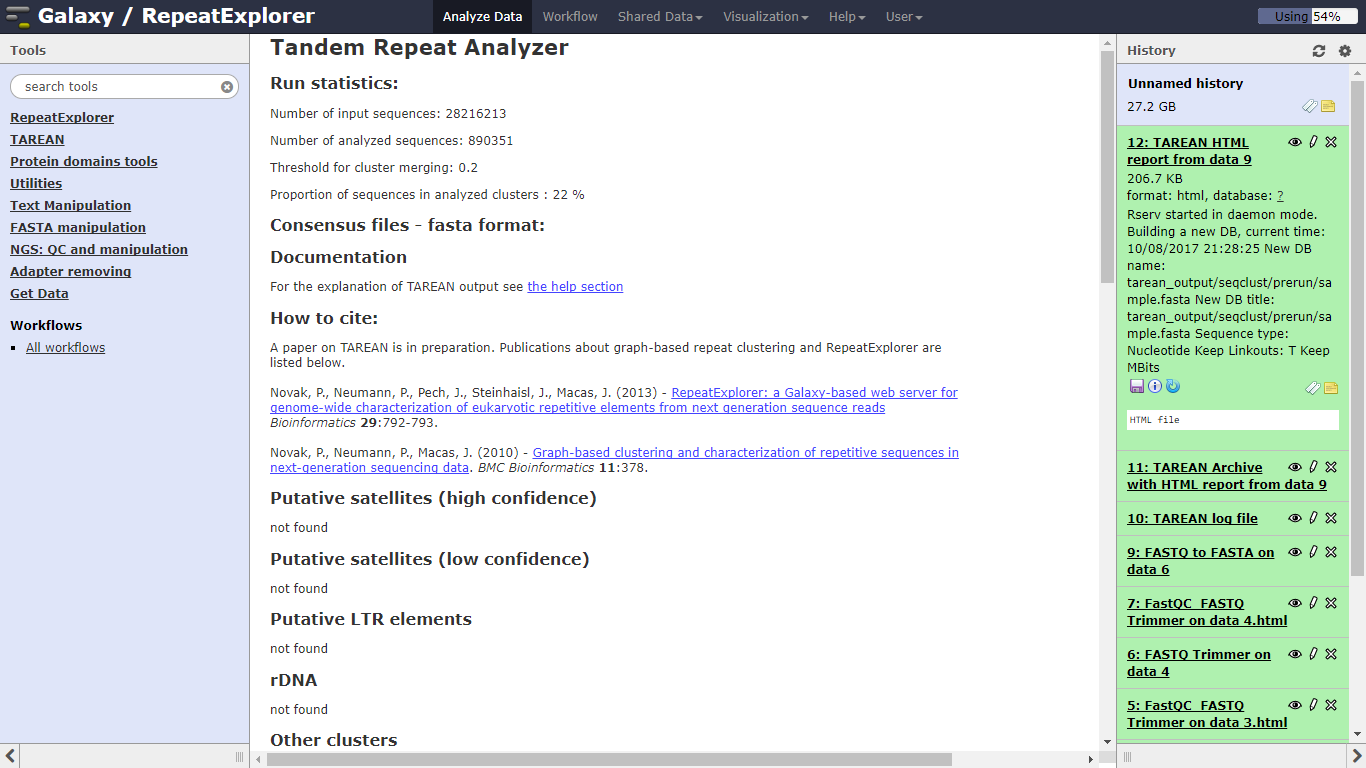


Fig. 17(a) – Result of tandem repeats

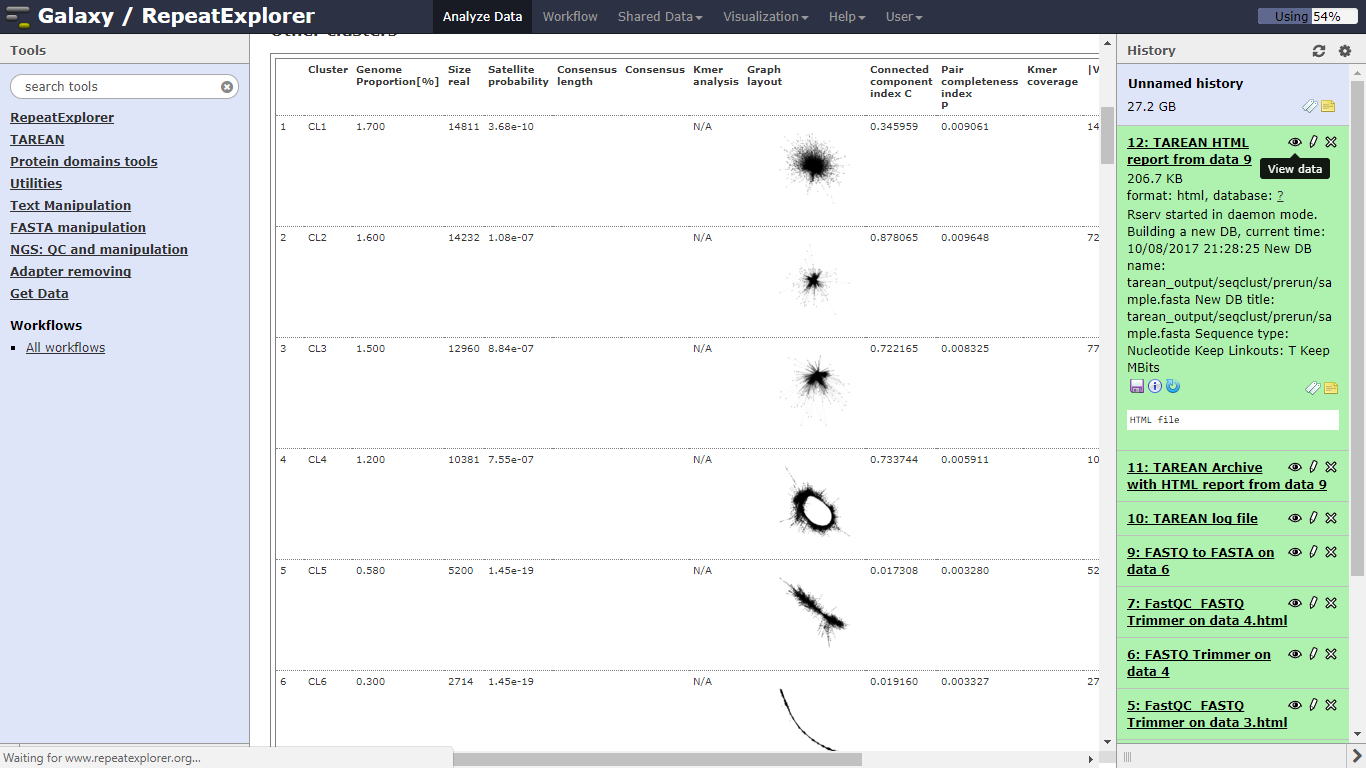


Fig. 17(b) – Result of tandem repeats

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