Analysis of Genomic Sequence Using DSP Techniques in Labview

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ABSTRACT: A cognitive radio (CR) is a transceiver which automatically detects available channels in wireless spectrum and accordingly changes its transmission or reception parameters. In this paper, it proposes an algorithm for the energy-efficient and spectrum-aware communications requirements in CR network. It enables each node to determine and regulate its transmission strategy to provide minimum energy consumption without sacrificing end-to-end delay performance and also maximizes overall spectrum utilization. Spectrum sensing is one of the essential parameter to be considered in CR networks. Therefore, the security aspect of spectrum sensing should be addressed well. Using a Trust-Worthy algorithm, it improves the trustworthiness of the Spectrum sensing in CR-Networks. It implemented using Network Simulator-2.

KEYWORDS: Rheumatic Arthritis, palindrome sequence, LabVIEW, Genomic analysis, Digital Signal Processing.

INTRODUCTION

The analysis of the genomic sequence is done using LabVIEW. In this project, Rheumatic Arthritis (RA) which is a chronic systemic inflammatory disease involving primarily the peripheral synovial joints is the disease taken for analysis. Many genes which are responsible for RA disease were found out and also the genomic sequence for each of these genes were found using databases such as KEGG (Kyoto Encyclopedia of Genes and Genomes) and National center of Biotechnology Information (NCBI). Along with normal genes few normal genes were also taken. Both normal and abnormal genes were then compared using the Digital Signal Processing techniques (DSP). Here, Fast Fourier Transform (FFT) is applied to achieve the comparison. The FFT tool is available in LabVIEW Software 2011 version. Appropriate code was written to extract a string sequence, convert this string sequence into numeric sequence and then apply FFT both the normal and abnormal. Gene sequence was given as inputs to the code. Analysis of the spectrum obtained for both normal and abnormal sequence was done by computing the mean amplitude. Separate code were written and implement for calculating the mean amplitude.

II. LITERATURE SURVEY

Digital Signal Processing (DSP) applications in genomic sequence analysis have received great attention in recent years. DSP principles are used to analyse genomic and proteomic sequences. [1]This paper has describe a method of generating Finite Impulse Response (FIR) of the genomic sequence. The same DNA sequence is used to convert into proteomic sequence using transcription and translation, and also digital filtering technique such as FIR filter applied to know the frequency response. The frequency response is same for both gene and proteomic sequence.[2]In this paper new methods are being developed to analyse DNA sequences, the DNA sequences should be converted into numeric sequences. Then the DSP algorithms are used in DNA analysis. This method is being used in our paper.[3]They have review the role of digital filtering techniques in gene identification. Long-range correlation between base pairs in DNA sequences has been discussed in brief which corresponds to a 1/f type of power spectrum. They have also described some of the recent applications of Fourier methods in the study of proteins. Finally they have mentioned the role of Karhunen-Loeve like transforms in the interpretation of DNA microarray data for gene expression. The role of signal processing in genomics and more generally biological sciences has been quite impressive.
SOFTWARE IMPLEMENTATION

The method to implement this paper is based on database of genomic sequence, LabVIEW and digital signal processing techniques. The acquired DNA sequence is compared with the standard DNA sequence structure. The analysis of these two sequences are done using DSP tool that are available in LabVIEW. A genomic sequence is accessed from the standard database and pasted in a text document. This text file is called in program through file input output function. Genomic sequence are generally in the form of string (mainly A,G,T and C). This string is converted into numerical form using select function along with comparison. Array of this, string converted into its corresponding numerical form, is obtained. FFT is applied and PSD coefficients are obtained. Using unbundle by name function, value of amplitude at each point can be accessed, mean amplitude is hence calculated. Also the given sequence is palindrome or not can be verified.

Fig. 1 Block diagram

In fig 1 the block diagram for the proposed work is shown

Fig. 2 Flowchart showing the 4 major steps

The implementation of this system is done in four steps. these are shown in the form of a flowchart in figure 2.

Fig. 3 Accessing the genomic sequence (courtesy: KEGG)
Step 1. Genomic sequence Extraction

The genomic sequence of the genes responsible for Rheumatic Arthritis (RA) is taken from a standard database. There are many websites available for the extraction of genomic sequences. The National center of Biotechnology Information (NCBI) database is the most popular one. Others are Kyoto Encyclopaedia of Genes and Genomes or KEGG, PubMed, etc. The procedure followed for the same is that we need to enter the official gene name or the gene number to access the genomic sequence. They are two types of genomic sequences one is the AA sequences which stands for the amino acid sequencing and the other is the NT sequence which stands for nucleotide sequences. Here, NT sequence has been used. A snapshot for KEGG website is showed in the figure 3.

Fig.4 Conversion of string to numeric form.

Step 2. Conversion of string to numeric form

The accessed genomic sequences which is in the form of NT sequence is converted to a complex format. As already mention earlier, the sequence obtained is in a string format and hence to apply FFT on it we need a discrete form. In a DNA sequence we have to assign numbers to the characters A, T, C, G, respectively. A proper choice of the numbers can provide potentially useful properties to the numerical sequence. For example, if we choose complex conjugate pairs T = A* and G = C*, then the complementary DNA strand is represented conjugate, symmetric numerical sequences which have interesting mathematical properties, including generalized linear phase. In this work complex conversion is taken as below and the code for the same is as shown in figure 4.

A → 1 + j
G → -1 + j
T → +1 - j
C → -1 - j

There are other conversions like binary where presence and absence is represented by 1 or 0 and other one is representing using electron ion interaction potential (EIIP) values.

Fig.5 A part of the code resulting in a FFT spectrum
Step 3. Applying DSP techniques

After the conversion is done, the next step is to create a waveform. To build this waveform, the LabVIEW tool called the "build waveform" tool is used after which the FFT is applied. Hence the resultant spectrum obtained is ready for analysis. The build waveform icon and the FFT tool can be seen in figure 5.

![Image of build waveform and FFT](image)

Fig. 5. Finding mean amplitude

Step 4. Analysis (comparison)

We compare the two waveforms obtained for normality and abnormality by finding out their mean amplitude, whose code is given in figure 6.

![Image of palindrome detection](image)

Fig. 6. Identification of a palindrome sequence

The code is shown in figure 7 where if the 5' to 3' sequence and 3' to 5' sequence is entered, it will show whether it's a palindrome sequence or not. Many restriction endonucleases (restriction enzymes) recognize specific palindromic sequences and cut them. The restriction enzyme EcoR1 recognizes the following palindromic sequence:

5' - G A A T T C - 3'
3' - C T T A A G - 3'

IV. PALINDROME SEQUENCE

Detecting palindromes in DNA sequences is a central problem in computational biology. Identifying palindromes could help scientists advance the understanding of genomic instability. DNA sequences containing long adjacent inverted repeats (palindromes) are inherently unstable and are associated with many types of chromosomal rearrangements. In this paper, we present a simple tool to assist biologists detecting palindromes in DNA sequences. A palindrome is a sequence of letters or words which reads the same in forward as well as backward directions. DNA palindromes are words from the nucleotide base alphabets A, C, G, T that are symmetrical in the sense that they read exactly the same as their complementary sequences in the reverse direction. DNA palindromes are crucial for gene regulation, DNA replication, and initiation of gene amplification.

![Image of palindrome detection](image)

Fig. 7. Identification of a palindrome sequence

The code is shown in figure 7 where if the 5' to 3' sequence and 3' to 5' sequence is entered, it will show whether it's a palindrome sequence or not. Many restriction endonucleases (restriction enzymes) recognize specific palindromic sequences and cut them. The restriction enzyme EcoR1 recognizes the following palindromic sequence:

5' - G A A T T C - 3'
3' - C T T A A G - 3'

V. RESULT

In a Normal sequence, we observe that the mean amplitude is less than 1, and there is one clear peak obtained. For an abnormal sequence (with respect to RA disease), the mean amplitude is more than 1. Also, no clear peak is obtained in this case. Hence, we have successfully obtained power spectra of both normal and abnormal sequence and compared them. Snapshots of the various normal and abnormal genes are shown in figure 8, 9, and 10 respectively.
In Fig 8, the figure on the left shows the spectrum obtained from a sequence of TLR9 and the figure on the right shows an sequence of BLK.

In Fig 8, the figure on the left shows the spectrum obtained from a normal sequence (TLR9) and the figure on the right shows an abnormal sequence (BLK), hence we clearly observe distortions in the abnormal sequence.

In Fig 9, the figure on the top shows the spectrum obtained from a sequence of HBA2 and figure on the bottom shows an sequence of CD5.

In Fig 9, the figure on the top shows the spectrum obtained from a normal sequence (HBA2) and figure on the bottom shows an abnormal sequence (CD5). For normal sequence we observe that we obtain only one peak.

In Fig 10, the figure on the left shows the spectrum obtained from a sequence of ABO and figure on the right shows a sequence of PTPN22.
In the Fig 10, The figure on the left shows the spectrum obtained from a normal sequence (ABO) and figure on the right shows an abnormal sequence (PTPN22). Hence you can distinguish a normal sequence from an abnormal one through the graph obtained.

In fig 11, The effectiveness of the approach has been brought out in predicting the gene F56F11.4 with five exons. The snap shot of this gene is shown, where the five peaks show the five coding regions of this gene.

<table>
<thead>
<tr>
<th>Gene name</th>
<th>Mean amplitude obtained</th>
</tr>
</thead>
<tbody>
<tr>
<td>TLR9(normal)</td>
<td>0.89</td>
</tr>
<tr>
<td>HBA2(normal)</td>
<td>0.95</td>
</tr>
<tr>
<td>ABO(normal)</td>
<td>0.98</td>
</tr>
<tr>
<td>BLK(abnormal)</td>
<td>1.04</td>
</tr>
<tr>
<td>CD5(abnormal)</td>
<td>1.02</td>
</tr>
<tr>
<td>PTPN22(abnormal)</td>
<td>1.01</td>
</tr>
</tbody>
</table>

An observation table has been made by noting down the mean amplitude values for various genes which is given in the table 1.

VI. CONCLUSION

The application of DSP methods to genomic data have begun to make important contributions to genomic research. In this system, complex based values-based approach, as an alternative to the binary sequence method, has been suggested for genomic signal processing. Open access to raw genomic data makes it easy for DSP experts to get involved in genomic research. With the huge amount of powerful techniques developed over the years being applied to genomics, we can hope to see rapid advances in specialized areas such as customized drug design and genetic remedies, which will greatly benefit humankind.

VII. FUTURE SCOPE

Further efforts can be made to improve the accuracy of the system since it is of utmost importance in this case. Likewise, the same algorithm can be applied to various other diseases like cancer etc. Also, we can make this algorithm available online so that it can be accessed as an open source code. We can also predict the in a eukaryotic genome, the introns and exons, start codon and stop codon, donor splice sites (transition from an exon to an intron or vice versa), and a CpG island (a region rich in CG pairs that may promote gene function) using DSP techniques.
Efficiency of the developed module can be still improved by detecting the stage of disease. The proposed algorithm can be made as universal standard and also can be used to predict the other disease.

REFERENCES