Computational Approach for Mining Simple Sequence Repeats in Expressed Sequence Tags

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Abstract

Expressed sequence tags, the short sequences of cDNA, are mined for identifying and characterizing simple sequence repeats for studying genetic variations. Web-based tools due to lack of server maintenance, become unusable; also few available stand-alone tools lack processing adequateness. Therefore with the intent to process multiple expressed sequence tag files without size limitations, proper validations, and the ability to retrieve more genome-related features, a simple to use, speed-efficient portable stand-alone tool has been developed. The algorithm is implemented in Java using a microsatellite search algorithm, with a dictionary-based approach MISA – Perl script, called via command line for data mining. Another parallel module retrieves additional information from GenBank files. In the pipeline primer, three were invoked for designing batch primers. This algorithm with an extended interface in Java Net Beans provides naïve users with a simple interactive tool for mining microsatellites, statistical analysis, and primer designing on one platform in the form of a stand-alone application. The number of repeats/ interruptions parameters can be reset through the graphical interface. This tool has interactive modules with proper validations, batch processing, and cost-effective analysis of tandem repeats as compared to peers; the source code can be upgraded in the future.

Keywords: BLAST, Data mining, EST sequence, Java pipeline, Microsatellites, Primer Design.

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INTRODUCTION

Expressed sequence tags (ESTs) or "single-pass" cDNA sequences are short sequences (<1000bp) of cDNA and are generated in large batches. These are tags of expression for the cDNA library of interest. More than 74.2 million ESTs from all species are available in public databases. The explosive growth of next-generation sequencing data has resulted in ultra-large-scale datasets. These datasets are a source of prime importance for gene discovery, gene transcripts, and gene-sequence determination; also, EST containing simple sequence repeats (SSRs) have become an attractive choice for the development of SSR markers. Simple sequence repeats (SSRs) or "microsatellites" ¹ are short tandem repeats (motifs) of length 1–6 nucleotides² and are found in genomes of both prokaryotes and eukaryotes.³ Microsatellites can be genomic if developed from genomic DNAs (gSSRs), or can be expressed, referred to as EST-SSRs if developed from an expressed sequence database.^{4, 5} EST- SSRs have high power because of their associations with expressed genes, directly contributing to a phenotype,⁶ also beneficial, being more conserved used as functional markers.⁷ Massive information can be churned from these databases by comparing ESTs from multiple species. They are likely to be more capable of cross**Corresponding Author:** Umang, School of Computer Science, Shri Venkateshwara University, Gajraula, Uttar Pradesh, India, E-mail: anilumang@yahoo.co.in

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genera transferability and are used for associating conserved genomic regions amid species, and genus, for comparative genomics studies, and for establishing evolutionary relationships.⁸⁻¹⁰ EST-SSR markers are also useful in markerassisted selection studies and facilitate in establishing genetic linkage maps.¹¹⁻¹² The SSRs found in the coding region affect gene activation, resulting in the expression of protein and reflects lesser polymorphism in the coding part.¹³ EST-SSR are directly associated with genes affecting a particular trait; therefore, they are considered a better resource for their use in breeding improvements,¹⁴ and SSRs present in the non-coding region affect gene regulation.¹⁵ In eukaryotes,

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EST-SSR markers are more profusely found in coding regions than in non-coding regions.^{2, 16} EST sequences are likely to be conserved evolutionarily; therefore, the expected rate of success is likely to be more in cross-species polymerase chain reaction amplification of EST-SSRs than in cross-species amplification of SSRs developed from genomic DNA.¹⁷ The rapid discovery of known or unknown genes from ESTs may contribute to the understanding of complex adaptive mechanisms as sorghum EST were obtained to identify and analyze genes that could respond to biotic stress.¹⁸

Considering the importance of EST analysis in studying genetic similarity/dissimilarity, various studies have been made to identify and characterize them in vitro, which is labor-intensive and time-consuming. Therefore with the advancement in next-generation sequencing technology and the easy availability of ESTs in public databases, many bioinformatics tools have been developed to analyze ESTs in silico. These bio computational tools aided the development of EST-SSR markers on a large scale in a cost-effective manner.2, 19 In a survey from the literature, it was revealed that most of the studies made by researchers using expressed sequence tags in species all over made the use of MISA20 based software for microsatellite analysis, Primer 321 to design primers and BLAST for homology search as observed in bioinformatics analysis of the ESTs from Rhizophora stylosa Griff. Genomic Library22, Pomegranate,23 Cacao, 24 Euphorbia esula, 25 characterizations of simple sequence repeats in watermelon genome, 26 Development of EST-SSRs in the genus Rubus13, and many more. Speciesspecific databases using different technologies were developed as conifer EST;27 a comprehensive annotated transcriptome data set in yam28 and garlic; the GarlicESTdb29 was developed to enrich the EST information in public databases. SpicEST, a comprehensive database, was developed for two spices plants, ginger and turmeric.³⁰

These researchers made use of software that fulfills the particular objective; these tools were found using web-based technology, species-specific databases, and stand-alone software. All these tools have different features that cater to different needs as per study or objectives.

Limited web-based tools are available that are costly to deploy, maintain, and also have processing inadequateness as they deal only with limited range of genome sequences. Web-based EST analysis pipeline ESTPiper streamlines typical large-scale EST analysis components.³¹ EST mining tools like EST2uni³² ParPEST³³ and ESAP plus³⁴ literature is available, but the registration process for direct access fails, it seems to server maintenance problem. Bioinformatics Tool Kit for EST Analysis³⁵ was developed by using a combination of five software that performs preprocessing/cleaning and clustering and can be called via command line on Linux platform is non-interactive, lacks batch processing, and is difficult for non-programmers to use. None of the stand-alone software was found to provide microsatellite identification/ analysis with more genome information, batch primer designing, and BLAST options on one platform.

Therefore considering the importance of *in silico* mining of ESTs and the privilege of server independence for desktop applications, this stand-alone java package for mining expressed sequence tags has been designed with a rich interactive graphic user interface that bridges the gap between existing technologies for the benefit of learners and researchers by enabling them to save intensive labor and cost.

MATERIALS AND METHOD

Material

Publicly available Batch files of EST sequences were downloaded in FASTA format from GenBank at the National Center for Biotechnology Information- website (http://www. ncbi.nlm.nih.gov/).

Graphic User Interface (GUI)

An interactive, user-friendly graphical user interface is implemented using Java Net Beans IDE 8.0.2 swing components, Perl and Java. It displays the tutorials to use the software. It allows MISA parameters and interruption numbers to set and reset as per requirements and saves them in the configuration file MISA.ini; the users are allowed to select multiple files for batch processing. The validation checks for EST files, name, path and count in the repository are displayed after submission. Users can clear the text box and exit from the system when required. Upon batch submission, users can simply click the Mine SSRs button to identify microsatellites and then press the Design Primers button or BLAST to choose BLAST options as per requirements. BLAST algorithm parameters can be changed by selecting them from the graphical interface.

The Algorithm with Regular Expressions

Pseudocode

Module 1

//Input -

- Reset repeat numbers
- Reset interruptions /parameters in misa.ini via interface
- Save.

Module 2

//Input -

• upload EST files in FASTA format as a file stream;

open each file in read mode

check for">" FASTA FILE

if FASTA format

then Upload to fasta files directory / create directories if not present/ delete old files if present.

else

skip



- Start process builder via CMD to call perl.exe misa.pl // a Perl script having dictionary based
- // approach data mining algorithm to detect microsatellites
- Execute each file serially on misa.pl
- Close files

Create directories if not present/ delete old files if present. //Output

mined text files saved to different directory and statistics text files saved to another directory

Module 3

//Input

- Upload Genbank file
- Check for extension
- Open file containing motifs
- Match accession Id from file containing motifs of each motif from GenBank file
- copy start position and end position of simple sequence repeats /motifs
- find coding /non-coding or overlapping region
- Fetch flanking region of 200 nucleotides from an upstream and downstream position for similarity search using BLAST//(200[motif]200)//output files with coding /non-coding information of motifs and sequence information to generate primers

Module 4

- install Primer3 software
- With default parameters, run output files with coding /non-coding information of motifs and sequence information to generate left and right primers, left and right primer length, primer TM, primer GC %, product size

Module 5

Use BLAST to find common, unique, and polymorphic simple sequence repeats by running files with flanking regions of motifs



Figure 1: Diagrammatic representation of Java based tool.



Figure 2: Screenshot representation of Java tool showing ESTs files deposited to repositories; Identification, Primer Design and BLAST Options.

Pipeline

The entire algorithm was implemented using Java programming language that performs a call to configuration file misa.ini, then misa.pl. Misa.pl (MISA, http://pgrc.ipkgatersleben.de/misa/download/misa.pl) is a Perl script that was used for mining perfect and compound simple sequence repeats. Motif coding / non-coding region from corresponding GenBank files was mined using guava-18.0.jar. Primer 3 software (http://primer3.org/releases.html) with default parameters was used to design primers. The flanking regions of 200 nucleotides were fetched in the pipeline from corresponding FASTA files to design batch primers for the identified microsatellites. BLAST GUI has been added for similarity search of flanking regions and provides Blast options like BlastN, BlastX, TBlastX, BlastP, and TBlastN. The entire workflow of the algorithm has been diagrammatically represented in Figure 1.

Transfer of Mined Data

Outputs are written in tab-delimited text files and saved in designated folders. These files can be imported to any relational database management system for downstream analysis.

RESULTS

This tool has been successfully tested and run on various size EST files for mining and analyzing perfect and compound microsatellites. Batch processing of EST format files results into information such as SSR type (perfect or compound), size, start position, end position, coding and non-coding region, and flanking sequences of 200 nucleotides for both upstream and downstream regions of SSRs are provided with left and right primers, left and right primer length, primer TM, primer GC %, product size. These primers are essential for developing microsatellite-based markers. Details of null primers, i.e., microsatellites for which the primers are not designed due to insufficient flanking regions or poor melting temperature, are stored in separate files. Statistics



Table 1. Comparative reatures of LST processing tool.	Table 1:	Comparative	features of	f EST	processing tools
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Program	Technology	Platform	Features	Remarks
ESAP Plus	PHP,HTML,CSS, Java Script, Apache HTTP server	Web-based	Pre processing, clustering and assembly, SSR Mining and Primer design	Unable to register to access the tool online
PESTAS	JSP	Web-based	Pre processing, clustering and assembling	Link Not available
ParPest	Perl, MySQL database using Red Hat Linux	Web-based	EST cleaning, clustering, assembling and BLAST comparisons.	Link not available
ESTPiper	Perl, JavaScript, JAVA on Linux	Web-based	Base calling, data cleaning, assembly, genome alignment, annotation, analysis of gene ontology.	Link not available
EST2uni	Apache HTTP Server, Perl scripting language, MySQL database management system, and PHP language on Linux platform	Web interface	Pre-processing, clustering, annotation, database creation, and data mining of EST collections	Link for code not available
GMATA	JavaScript, Perl script and R scripts	Stand-alone	No multiple files processing and BLAST options	Downloadable
Krait	Implemented in C and python	Stand-alone	Multiple files can be processed, SSR Mining, individual motif has to be clicked, no BLAST options	Downloadable and Performs very fast
MISA-Web	PHP and UNIX shell monitor server and Perl script	Web-based	No flanking sequences, no coding, and non-coding information.	none
Bioinformatics tool kit	Pherd, LUCY, RepeatsMasker, Cap3 tools ere integrated using Java based pipeline.	Invoked via command line on Linux platform	Pre-processing, clustering, and assembly, EST Nucleotide database similarity searches	Downloadable
JaP-estmining	Java with misa.pl	Stand-alone tool	Multiple files can be processed; SSR Mining with coding and non- coding information and batch primer design with BLAST options and source code is modifiable.	Downloadable

details are displayed in separate files mentioning the total number of sequences examined, the total size of examined sequences (base pair), the total number of identified SSRs, the number of SSR containing sequences, number of sequences containing more than one motif, Number of SSRs present in the compound formation, distribution to different repeat type classes; having unit size with the corresponding number of SSR and frequency of identified SSR motifs. The data is automatically saved to tab-delimited text files in designated folders. BLAST options with different parameters can be used for a homology search.

DISCUSSION

This algorithm implemented via java package uses the dictionary-based approach algorithm misa- a perl script and

regular expressions for data mining in genome sequences; this tool has several unique features. The user can set, reset, and save the number of nucleotides and interruptions in the configuration file. Proper validations are applied to check valid file types and extensions. It checks the FASTA format sequences for processing. For batch processing, the number, names, and location of files sent for mining are displayed (Figure 2). Multiple files are processed with a single click, no need to attach or upload files again and again. The files after mining (simple sequence repeats) are saved to properly designated folders. ESTpiper,31 EST2uni,32 ParPest33 PESTAS36 pipelines are web-based, and we didn't find the link to access them, maybe due to server maintenance problems. However, these pipelines are using different technologies and are limited to cleaning, clustering, assembling, and BLAST comparisons and ESAP plus34 seem challenging to access with a typical registration process. JaP-estmining tool has an advantage over them for being a stand-alone desktop application. We can also use other simple sequence mining stand-alone tools like GMATA.37 But, it cannot process multiple files at one click, so each expressed sequence has to be copied to a different file for processing, thus creating a large number of records, whereas JaP-estmining processes single file containing as many sequences. No file size restrictions for processing the data as it was observed in MISA-web.38The users can either detect microsatellites and related statistics or use corresponding GenBank files; they can identify coding, and non-coding regions; with flanking sequence can go for primer designing set with default parameters. Alternatively, the user can BLAST flanking sequences for homology search with five BLAST options39 as described. The users can either copypaste sequence or upload preformatted ESTs files using the upload file button. The type of query sequence is identified, and then the appropriate BLAST option is enabled; this is an advantage over the Krait tool40 that mines and designs multiple files. Output formats in tab-delimited text files are imported to any relational database management system for downstream analysis.

This Java-based tool varies in terms of file processing features and output, and other utilities, as shown in Table 1.

CONCLUSION

The tool is developed and tested on AMD E-350 processor 1.60 GHz with 2.0 GB RAM and 32-bit operating system and performs much better with upgraded systems. Batch EST files have been processed to identify microsatellites, design primers, and use BLAST options within few seconds. These features altogether are not available in peers. The tool is available on-demand and will be provided on open source platforms soon after adding more functionality.

CONFLICT **O**F **I**NTEREST

The author declares no potential conflicts of interest.

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