UniFrag and GenomePrimer: selection of primers for genome-wide production of unique amplicons

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ABSTRACT

Summary: the complementary programs UniFrag and GenomePrimer were developed to provide a reliable high-throughput method to select the most unique regions within genomic DNA sequence(s) and design primers therein, involving minimal user intervention and maximum flexibility.

Availability: freely available for educational and research purposes by non-profit institutions at http://molgen.biol.rug.nl/molgen/research/molgensoftware.php

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Supplementary information: http://molgen.biol.rug.nl/publication/primer_data/

Various programs have recently become available for the selection from coding sequences of single primers (e.g. OligoArray, Rouillard et al., 2002, and ROCK, Strain and Chmielewski, 2001) or primer pairs (e.g. PrimeArray, Raddatz et al., 2001) for the production of DNA microarrays using oligonucleotides or amplicons. With GST-PRIME (Varotto et al., 2001), a large number of primer pairs can be generated starting from a list of accession numbers (GIs). PrimoUnique (Chang Biosciences; web-site: http://www.changbioscience.com/primo/primou.html) designs primer pairs for each member of a list of DNA sequences with high similarity (a family). Those primer pairs that might aspecifically amplify a different family member are eliminated.

We use a high-throughput approach for amplicon design and production for DNA-microarrays of bacterial genomes. As some specific features, such as high throughput selection of unique regions and subsequent primer design, were not present in available software packages, the complementary programs UniFrag and GenomePrimer were developed. Together, these programs allow selecting unique regions within a DNA sequence and automatically designing primers, with minimal user intervention and maximum flexibility. UniFrag and GenomePrimer were not developed for primer design on genes containing introns. By using unique regions in DNA sequences for primer design, cross-hybridization during DNA microarray experiments is minimized. Also, as aspecific priming during PCR is minimized, PCR is more successful. Moreover, a unified primer design will allow better robotization of PCR.

UniFrag runs on a Unix platform and only requires a locally installed ‘Blastall’ program (the version used was 2.2.3; http://research.nhgri.nih.gov/blastall/). UniFrag consists of five sub-programs written in Pascal and compiled by FreePascal 1.0.4 (http://www.freepascal.org/) under Red Hat Linux release 7.2 (http://www.redhat.com). GenomePrimer was written in Borland Delphi 5.0 and runs on any Microsoft Windows platform.

Figure 1 presents a flow scheme of UniFrag and GenomePrimer. Input files for UniFrag are: (i) a FASTA file containing the DNA sequences from which the unique regions have to be selected; (ii) a FASTA file containing a reference set of DNA sequences formatted with Formatdb for Blastall; and (iii) a configuration file containing the options set by the user. A reference set should consist of, at least, the DNA sequences that are used as input for the UniFrag program but other DNA sequences (such as genome sequences) can also be added. In microarray experiments, RNA originating from an organism other than the one represented on the slide might be used. By using the genomic sequence (if available) of the other organism in the reference set of UniFrag, possible cross-hybridization will be prevented.

In a typical UniFrag run (Fig. 1), overlapping fragments are generated by using a ‘window’ (a window corresponds to a fragment with a size set by the user) that ‘slides’ over each of the input sequences. The ‘windows’ overlap each other with an overlap size that can be set by the user (Fig. 1A and B). By generating overlapping fragments, the chance increases to find a fragment that is unique within a DNA sequence. Sequences that are smaller than the minimum fragment size are saved in a list of ‘leftovers’

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design program, which selects thousands of primers within seconds (Fig. 1F). Primers are examined for: (i) equal distribution of G and C; (ii) occurrence of palindromic sequences; and (iii) homology between primers of a primer pair. The melting temperature can be set according to either one of two rules: (i) \[ T_m = 4 \times (GC) + 2 \times (AT) \] (Suggs et al., 1981); or (ii) \[ T_m = 62.3 + 0.41 \times (GC) - (500/\text{length}) \] (Sugimoto et al., 1996). Desired tags, for instance for the re-amplification of all amplicons using a tag-specific oligonucleotide pair, can be easily introduced.

General result statistics such as success rate, average primer length, number of nucleotides to be synthesized and specific characteristics such as the number of (too short and/or too low GC-content) amplicons are displayed by the Report Generator (Fig. 1G). If primer design failed in too many cases, the settings can be easily adjusted and a new design can be performed until all primers/amplicons meet the desired characteristics. Those DNA sequences on which no primer pair could be designed by the GenomePrimer selection criteria can be selected for primer design using other criteria.

The freely available web-based version of PrimoUnique allows designing one unique primer pair at a time. In contrast to UniFrag and GenomePrimer, PrimoUnique does not allow constraints in amplicon length. As only the primers and not the amplicon sequences are unique, PrimoUnique is not usable for effective genome-wide primer design in contrast to the combination of UniFrag and GenomePrimer. The PrimeArray program allows using a predetermined amplicon length cut-off whereas in GenomePrimer a window of amplicon sizes is used. The variation in amplicon size allows primers to be chosen more flexibly in a location on the DNA sequence. This allows using more stringent primer characteristics during the design, which results in a more efficient (and successful) high-throughput approach because subsequent PCR conditions can be standardized. GenomePrimer provides the user with a number of criteria for primer design: (i) preferred location of primers in a sequence (for instance to obtain 3′-, 5′- or central amplicons of a gene); (ii) preferred primer length and reduction of primer length, while maintaining annealing properties, to reduce production costs; (iii) 3′- G or C on primers to improve the PCR extension step, which is especially important for low GC organisms such as Lactococcus lactis; and (iv) preferred GC content of the primer.

UniFrag was used to select the unique regions in all predicted open reading frames (ORFs) of Lactis IL1403 and Streptococcus pneumoniae TIGR4 (Table 1, supplementary information). More unique ORF fragments could be selected from Lactis IL1403 (1516; 68% of the total amount of 2214 ORFs) than from S.pneumoniae TIGR4 (1390; 56% of the total amount of 2495 ORFs) because the genome sequence of the latter contained more

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**Fig. 1.** Flow chart of UniFrag and GenomePrimer. Arrows with a plus (+) or minus (-) sign indicate data that met or did not meet the selected criteria (in case of steps A and D) or signify a successful or not successful primer design (step G), respectively.

(Fig. 1E). The remaining overlapping DNA fragments are compared using BlastN against the reference set and the results are filtered using an expectancy value cutoff parameter (e-value) commonly used in blast searches (Fig. 1C). From the filtered blast output, the fragment with the highest expectancy value (most discriminating fragment), is selected for each input DNA sequence (Fig. 1D). If the selected fragment has an e-value higher than the ‘cutoff unique’ parameter, the input DNA sequence is used for primer design; if not, the input DNA sequence is saved in the ‘leftover’ list (Fig. 1E). UniFrag can be run for any number of passes (n in Fig. 1), decreasing fragment and overlap sizes until the maximum number of unique fragments is obtained (Table 1, supplementary information). Other sequences can be added to the resulting list of fragments (i.e. sequences of the ‘leftover’ list). The FASTA file generated can then be used for subsequent GenomePrimer processing.

Via user-friendly interface frames, various selection criteria can be chosen before running the GenomePrimer program, which selects thousands of primers within seconds (Fig. 1F). Primers are examined for: (i) equal distribution of G and C; (ii) occurrence of palindromic sequences; and (iii) homology between primers of a primer pair. The melting temperature can be set according to either one of two rules: (i) \[ T_m = 4 \times (GC) + 2 \times (AT) \] (Suggs et al., 1981); or (ii) \[ T_m = 62.3 + 0.41 \times (GC) - (500/\text{length}) \] (Sugimoto et al., 1996). Desired tags, for instance for the re-amplification of all amplicons using a tag-specific oligonucleotide pair, can be easily introduced. General result statistics such as success rate, average primer length, number of nucleotides to be synthesized and specific characteristics such as the number of (too short and/or too low GC-content) amplicons are displayed by the Report Generator (Fig. 1G). If primer design failed in too many cases, the settings can be easily adjusted and a new design can be performed until all primers/amplicons meet the desired characteristics. Those DNA sequences on which no primer pair could be designed by the GenomePrimer selection criteria can be selected for primer design using other criteria.

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ORFs smaller than 500 bp, which were ignored in the analysis. The \textit{S.pneumoniae} TIGR4 genome contained less ORFs (69; 2.8\%) in which no unique fragment could be identified than \textit{L.lactis} IL1403 (80; 3.6\%). In most cases, the ORFs of which no unique fragment could be identified were specified by insertion elements or transposon sequences. Table 1 (supplementary information) clearly illustrates that decreasing the fragment size results in a significantly higher amount of unique ORF fragments. By additionally decreasing the overlap size, the probability increases that a certain fragment is unique.

\textit{GenomePrimer} was initially tested on the predicted ORFs of the \textit{L.lactis} bacteriophage r1t (Table 2, supplementary information). A 100\% success rate for primer design and amplicon production of the 51 r1t ORFs was obtained. For the selection of primers for ORFs larger than 90 bp in the complete genomes of \textit{S.pneumoniae} TIGR4 and \textit{L.lactis} IL1403, a smaller range of amplicon length was used (80–800 bp) than for r1t (89–1842 bp). Success rates of both primer sets (100\% and 99.5\% for \textit{L.lactis} and \textit{S.pneumoniae} TIGR4, respectively) were high in the first round of PCR. PCR reactions were carried out under standard conditions in a Bio-Rad iCycler 1 PCR machine (Bio-Rad, Hercules, CA) in 96-wells format with a non-proofreading DNA polymerase.

Reliable high-throughput genome-wide primer design on unique fragments of ORFs is the strength of using \textit{UniFrag} and \textit{GenomePrimer} software. Using unique fragments in microarray studies should reduce cross-hybridizations, which will improve data quality and make data analysis more straightforward.

\textbf{REFERENCES}


