Sequence analysis

RBR: library-less repeat detection for ESTs

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ABSTRACT

Motivation: Repeat sequences in ESTs are a source of problems, in particular for clustering. ESTs are therefore commonly masked against a library of known repeats. High quality repeat libraries are available for the widely studied organisms, but for most other organisms the lack of such libraries is likely to compromise the quality of EST analysis.

Results: We present a fast, flexible and library-less method for masking repeats in EST sequences, based on match statistics within the EST collection. The method is not linked to a particular clustering algorithm. Extensive testing on datasets using different clustering methods and a genomic mapping as reference shows that this method gives results that are better than or as good as those obtained using RepeatMasker with a repeat library.

Availability: The implementation of RBR is available under the terms of the GPL from http://www.ii.uib.no/~ketil/bioinformatics

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Supplementary Information: Supplementary data are available at Bioinformatics online.

1 INTRODUCTION

Expressed Sequence Tags (ESTs) are a valuable source of information that allows identification of genes in sequenced genomes. In addition, they provide a low-cost approach to mapping the gene complement of organisms for which genome projects are still too expensive. Clustering ESTs and assembling consensus sequences can reveal important information about the transcriptome, including splice variants and putative single nucleotide polymorphisms (SNPs).

The clustering procedure is based on identifying matches between the ESTs and grouping together sequences with sufficient overlaps. Repeats, in the form of sequencing artifacts, contamination, low quality sequence and genomic repeats, represent a serious problem for clustering, as they can occur in otherwise unrelated sequences. Matches resulting from repeats may lead to erroneously grouping ESTs that arise from different genes, and it is therefore important to mask repeats prior to clustering.

We have previously developed RepeatBeater (Schneeberger et al., 2005) and shown that it successfully masks repeats and improves the accuracy of clustering. In this paper we present a new method named RBR that is conceptually simpler than RepeatBeater, and while RepeatBeater was tightly coupled to xsact (Malde et al., 2003), RBR works independently of clustering method.

To analyze the effectiveness of RBR, we use several representative datasets of realistic sizes. We construct reference clusterings by mapping all ESTs to the genomic sequence, retaining only sequences that can be mapped unambiguously and grouping ESTs whose genomic matches overlap.

By comparing with the reference clusterings, we show that RBR produces results that are at least as good as those produced by the most widely used library-dependent repeat masking methods. Even for the set of human ESTs it gives results superior to those produced by methods using human repeat libraries. For some organisms, the results are dramatically improved over those produced by existing methods.

1.1 Existing methods

There are currently two main approaches to masking EST data. Library-based masking identifies undesired sequence parts by comparing with a library of known repeats. Autonomous masking identifies undesired sequence from properties of the sequence itself.

Repetitive regions often constitute large parts of a genome, and some classes of complex genomic repeats, like those originating from transposon and retrotransposon activity or retroviral elements, are collected in libraries which can then be used for masking.

While libraries derived from the genome are commonly used for masking ESTs, they are not targeted specifically at transcripts. For instance, LINE transposons normally contain genes, and for the purpose of EST clustering, the copies will either be treated as a single gene or as multiple homologues. Some transposons do cause problems, for instance the short SINE/Alu repeat is found in human ESTs (Schneeberger et al., 2005).

In EST data, there are also vector sequences and genomic contamination from the host organism, typically Escherichia coli. Similar to genomic repeats, these can be masked using a library.

Genomic repeats are in general organism specific. For novel organisms, where the genome is not yet available for constructing a good repeat library, effective masking of genomic repeats is difficult.

Autonomous masking typically deals with simple repeats, which consist of contiguously repeated short subsequences (2–4 nt) often caused by polymerase slippage, and low complexity regions (LCRs), which are regions containing an overabundance of some nucleotides. In addition to poly-A tails, which can be considered a special case of either type, these repeats are common in the UTR regions of genes.
The accuracy of base calling tends to deteriorate towards the ends of the ESTs (Liang et al., 2000; Ewing and Green, 1998), and the error rate increases until at some point the remaining sequence is noisy. These low-quality sequence parts are commonly trimmed. Usually a cut-off is set from quality values determined by the base-calling software, but heuristics using only the sequence exist.

RepeatMasker (Smit et al., 2004) is probably the most widely used tool for masking against a library (typically interspersed repeats and E.coli and vector sequences), and by default uses cross_match (P. Green, unpublished data) to identify matches. It also masks LCRs. RepeatMasker is distributed (and normally used) with the RepBase (Jurka et al., 2005) repeat library.

One popular utility for autonomous masking is mdust from TIGR, which implements the DUST algorithm (R. L. Tatusov and D. J. Lipman, unpublished data). Other alternatives include DustMasker (Morgulis et al., 2006) and SeqClean (G. Pertia, unpublished data), the latter combines autonomous masking with optional masking against a library. Search and alignment programs, like BLAST (Altschul et al., 1990) and BLAT (Kent, 2002), often incorporate autonomous masking as well.

There appears to be little consensus on a ‘best practice’ of steps to apply. TGICL (Pertia et al., 2003), a widely used EST analysis pipeline from TIGR, uses mdust to mask sequences for low complexity before clustering and assembly, but does not mask against genomic repeats.

Other pipelines (e.g. Krause et al., 2002; HarvESTer from BioMax Informatics AG, unpublished data) use either RepeatMasker or similar library-based masking (e.g. Pontius et al., 2003; Miller et al., 1999; D’Agostino et al., 2005) in addition to autonomous masking.

2 METHODS

RBR identifies repeats by first calculating the frequencies of word occurrences in the entire dataset. Then, for each sequence the frequencies of the words it contains are collected, and a threshold is determined based on these frequencies. Sequence positions corresponding to frequency peaks above this threshold are then masked. RepeatBeater (Schneeberger et al., 2005) used a similar approach, but its algorithm is more complex, and as the implementation relies on pre-processing the sequences with xsact, it is slower and more complicated to use.

2.1 The RBR algorithm

Given a word size \( k \), RBR first calculates the frequency of every \( k \)-word in the full dataset. For each sequence, the distribution of word frequencies is collected, and the baseline for the sequence is estimated. Words with frequencies significantly above this baseline are then masked. RepeatMasker (Smit et al., 2004) is probably the most widely used tool for masking against a library (typically interspersed repeats and E.coli and vector sequences), and by default uses cross_match (P. Green, unpublished data) to identify matches. It also masks LCRs. RepeatMasker is distributed (and normally used) with the RepBase (Jurka et al., 2005) repeat library.

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In the absence of repeats or read errors in the sequences, and with the sequences uniformly distributed over the originating gene, we expect the distribution of word counts to approximate a binomial distribution. To see this, let \( n \) be the number of sequences whose origin in the genome overlaps the origin of a given sequence \( S \), and \( p \) be the average fraction of this overlap. For each position in \( S \), the probability of an individual sequence overlapping it in that position is \( p \), and as there are \( n \) sequences, we get a distribution of \( B(n, p) \). Furthermore, from the assumed uniform distribution, and with the added assumption that sequences have similar lengths, we expect \( p \) to be 0.5.

In practice sequences vary in length and quality, and the distribution is non-uniform. Therefore the assumptions above lead to only an approximation of the actual distribution. Instead of assuming a fixed \( p \), we estimate the distribution empirically.

The variance \( \sigma^2 \) of the binomial distribution \( B(n, p) \) is \( np(1 - p) \), and thus the standard deviation grows with the square root of the mean. We therefore start by identifying the modal interval \( (m - \sqrt{m}/2, m + \sqrt{m}/2) \) containing the maximum number of word occurrences.

For a binomial distribution, we can estimate \( 1 - p \) from the empirical variance and mean as \( \hat{q} = \sigma^2/\mu \). We adjust the interval around \( m \) so that estimated \( \hat{q} \) for the frequencies in the interval is equal to the stringency parameter \( s \). This distribution is used as the baseline to mask all words in the sequence with frequencies above \( \mu + \sigma \hat{q} \).

2.2 Constructing reference clusterings

When each EST contains enough non-repeat sequence to determine its position in the genome, the ESTs can be clustered by location, which greatly reduces the effect of any repeats. The result is similar to a clustering based on EST sequence similarity alone when the ESTs are masked optimally, and we use this to serve as a reference for comparing maskings (Kalyanaraman et al., 2003; Wang et al., 2004; Wu et al., 2004; Wu and Watanabe, 2005). We emphasize that this is only a coarse approximation to the biological transcripts, for example, incompleteness, sense-antisense transcription (Shendure and Church, 2002; Yelin et al., 2003) and trans-splicing events (Huang and Hirsh, 1992) will still contribute to inaccuracies in the clustering.

To assess RBR’s performance, we developed GEM, a system for clustering ESTs based on overlap in genomic position. It uses BLAT (Kent, 2002) to determine a spliced alignment to the genome, and has several parameters to set criteria for the clustering.

To verify that RBR is able to identify and eliminate repeats, we used a set of 1355 human ESTs taken from a cluster known to contain multiple genes SINE/Alu repeat (Schneeberger et al., 2005). The sequences were mapped to the human genome using GEM, resulting in 339 distinct clusters. We also mapped 50 000 ESTs from Oryza sativa, and 100 000 from each of Caenorhabditis elegans and Arabidopsis thaliana to their respective genomes using GEM.

In order not to end up with artificially clean datasets, we used relatively lax parameters (90% sequence similarity and 75% of the sequence length) to match ESTs against the genome and discarded sequences that matched multiple locations equally well. The ESTs were then clustered if they overlapped 20 nt or more in the matches against the genome.

The resulting data sets contained 47 302 ESTs from O.sativa, 84 655 ESTs from C.elegans, and 97 919 from Arabidopsis.

2.3 Comparing clusterings

Many of the commonly used indices for comparing clusterings are based on categorizing and counting pairs of clustered objects. Specifically, for a set of objects \( \{x_1, x_2, \ldots, x_n\} \), and clusterings \( K \) and \( C \), we denote by \( C(x_i) \) and \( K(x_i) \) the clusters containing \( x_i \) in \( C \) and \( K \), respectively. We then define four variables

\[
a = \left| \{(i, j) \mid C(x_i) = C(x_j), K(x_i) = K(x_j)\} \right|
\]

\[
b = \left| \{(i, j) \mid C(x_i) = C(x_j), K(x_i) \neq K(x_j)\} \right|
\]

\[
c = \left| \{(i, j) \mid C(x_i) \neq C(x_j), K(x_i) = K(x_j)\} \right|
\]

\[
d = \left| \{(i, j) \mid C(x_i) \neq C(x_j), K(x_i) \neq K(x_j)\} \right|
\]

where \( 1 \leq i < j \leq n \).

Many cluster indices can then be calculated from these variables. In particular, the Jaccard index (Jain and Dubes, 1988) is defined as

\[
J = \frac{a}{a + b + c}.
\]
One problem with Jaccard, and indeed with all the measures based on counting pairs, is that modifications to large clusters will affect the score much more than modifications to small clusters. A different approach that suffers less from this drawback is based on entropy. Meila (2005) introduces the variation of information VI to compare clusterings. For two clusterings K and C, VI is defined as

\[ VI(K, C) = H(C, K) - I(C, K) \]

where \( H(C, K) \) is the total entropy of the clusterings, and \( I(C, K) \) denotes the mutual information, defined as

\[ I(C, K) = H(C) + H(K) - H(C, K) \]

The confusion matrix \( M \) is the \( c \times k \) matrix defined by \( M_{i,j} = |C_i \cap K_j| \). If we define \( M_i = \sum_j M_{i,j} \) and similarly, \( M_j = \sum_i M_{i,j} \), we have the following definitions

\[ H(C) = \sum_i \frac{M_i}{n} \log \frac{M_i}{n} \]

\[ H(K) = \sum_j \frac{M_j}{n} \log \frac{M_j}{n} \]

\[ H(C, K) = \sum_{i,j} \frac{M_{i,j}}{n} \log \frac{M_{i,j}}{n} \]

VI can now be calculated directly from \( M \) as

\[ VI = \frac{1}{n} \left( \sum_i M_i \log M_i + \sum_j M_j \log M_j - 2 \sum_{i,j} M_{i,j} \log M_{i,j} \right) \]

We believe VI is a more appropriate measure, but for consistency with previous work, we will also provide the Jaccard index.

3 RESULTS

We have selected two parameter settings for RBR, the default configuration of \( s = 1.5 \) and \( d = 6 \), and a more aggressive configuration, using \( s = 1 \) and \( d = 4 \). We also masked the datasets with RepeatMasker, masking with lower case letters using the -xsmall option, and supplying the appropriate -species parameter.

TGICL pre-processes sequences with mdust, and uses MegaBLAST (Zhang et al., 2000) to identify matches, using exactly matching words of length 18 to seed pairwise alignments. Lower case letters are interpreted as masked sequence and excluded from the seeding, but not from the subsequent alignment. For clustering, we ran TGICL with its default parameters, and xsact with a word size (-k option) of 25 and a threshold (-r) of 60.

3.1 Human dataset

To verify that RBR produces a result similar to RepeatBeater, we first ran RBR on the set human ESTs. Using the default configuration, RBR masks 4.6% of the nucleotides differently from RepeatBeater. Comparing RBR to RepeatMasker results in a difference of 7.8%, while comparing RepeatBeater with RepeatMasker gives 7% differently masked nucleotides (Schneeberger et al., 2005). Thus, RBR and RepeatBeater are more similar to each other than either is to RepeatMasker.

From Figure 1 we see that RepeatMasker masks a relatively large part of the human dataset. A large fraction of the masked nucleotides are masked as SINE/Alu repeats, and these repeats constitute almost all nucleotides that are masked by both methods (6.3% versus 6.4%).

RBR masks a smaller amount of sequence, but with a large degree of overlap, and the differences are mainly owing to RBR only masking the most conserved parts of the repeats. In particular, only two sequences in the dataset are masked by RepeatMasker without being masked by RBR, and they turn out to contain be duplicates of the same sequence. Most of the sequence masked by RBR but not RepeatMasker is low complexity and simple repeats, either too short or too ‘noisy’ to be detected by RepeatMasker.

We also compared the clusterings obtained by running TGICL and xsact on the masked sequences with the clusterings obtained by mapping the ESTs to the genome using GEM. The results are displayed in Figures 2 and 3.

We see that the performance of both TGICL and xsact is poor on the unmasked data. The two clusterings using RBR and the masking using RepeatMasker lead to similar clusterings, indicating that although RBR does not mask all SINE/Alu repeats, it masks it to the extent it adversely affects the clustering.

3.2 Large datasets

To see how RBR performs on more realistic datasets, we processed the larger datasets with RepeatMasker, mdust and RBR. The fraction of masked sequence is given in Figure 4. On these sets, RepeatMasker only masks a small fraction of the sequence data,
comparable with mdust. RBR masks a similar amount of sequence as it does on the human dataset.

Again we clustered the masked datasets with TGICL and xsact, and compared with the clusterings derived from the genomic mapping. The results are given as Variation of Information in Figures 5 and 6, and Jaccard indices in Figures 7 and 8.

We see that TGICL performs well on *C. elegans* and *O. sativa* using only the intrinsic masking. The *Arabidopsis* dataset, however, is markedly improved using RBR. For xsact, the clusterings generally see an even greater improvement.

4 DISCUSSION AND CONCLUSION

Except for the human dataset, which was selected explicitly for containing a known repeat, masking against genomic repeats in ESTs generally appear to have little effect on the clustering. Transcript-specific repeat databases would likely improve this situation, or at least reduce the amount of nucleotides masked unnecessarily.

In our experiments, we find no case where using RBR significantly worsens the results, and often the result is markedly improved. We think RBR, or a similar tool, should become an important element in EST analysis, complementing the standard approaches. In particular for exploring new organisms, where no good repeat library exists, we believe RBR provides the best available alternative for masking ESTs.
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Conflict of Interest: none declared.

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