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ABSTRACT

RNA GENOME ANNOTATION WITH A FOCUS ON T. BRUCEI

by Brett Bucci

The goal of this project is to identify untranslated regions (UTRs) and UTR-indicating patterns in the genome of *T. brucei. T. brucei* is an interesting organism, and as the cause of African sleeping sickness—which infects 300,000-500,000 people and a significant number of cattle annually—is currently the subject of considerable research. Using existing algorithms, several patterns have been found that may lead to more complete UTR annotations in the *T. brucei* genome. The most encouraging sequence is the 11-base sequence GAGGG[CG]TGGGG, which appears in five hypothetical genes near the tail. Discovery of several such sequences could guide laboratory experimentation toward more useful results and a better allocation of time and resources.

by Brett Bucci

A Thesis Submitted to the Faculty of New Jersey Institute of Technology in Partial Fulfillment of the Requirements for the Degree of Master of Science in Computational Biology

Department of Computer Science

January 2008

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APPROVAL PAGE

RNA GENOME ANNOTATION WITH A FOCUS ON T. BRUCEI

Brett Bucci

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INTRODUCTION

The goal of this project is to identify UTRs and UTR-indicating patterns in *T. brucei*. Current UTR annotations are limited, and are mostly focused on chromosome I. Several algorithms exist to predict UTRs, and many have been predicted by sequence homology and other methods, but without experimental evidence functionality cannot be verified. One of the aims of this project is to determine the best UTR candidates, and perhaps guide laboratory experimentation toward more useful results. One of the sequences found to recur in putative UTR regions also seems to be present toward the end of several hypothetical proteins, and may be a good indication of where to direct laboratory resources.

METHODS

The two main sources of annotated UTR sequences were GeneDB [1] and NCBI [5]. To find UTRs in NCBI, the author performed the following steps. This search will produce approximately 34 results.

- 1. Enter trypanosoma brucei in the search field
- 2. Select Organisms on the Limits tab and click Go
- 3. Enter 5'UTR in the search field
- 4. Click on the Limits tab and change to All Fields
- 5. Click History
- 6. Click the numbered link next to Search trypanosoma brucei Field: Organism
- 7. Click AND in the pop-up menu, and then click Go

To find UTRs in GeneDB, the following will yield about 15 results.

- 1. Select T. brucei from the Protozoa menu on the right
- 2. Enter *UTR* under Full Content Search , and then click the Full Content Search button

PUTATIVE UTRS

The sequences in Appendix A have been annotated as UTRs by either GeneDB or NCBI. Where possible, some subsequent sequence information has been provided. The key to the annotations is as follows:

The **bold** portion of the sequence is what's annotated as a UTR by GeneDB or NCBI. The <u>underlined</u> portion is one of the following highly conserved sequences that appears four times in this data A[AT]AG[CT]AGAGG), or twice GAGGG[CG]TGGGGG (see note below).

The sequence GAGGG[CG]TGGGG appears 11 times in the *T. brucei* genome according to a BLAST search using *The T. Brucei Genome Project* (8) website:

*Tb10.389.1530 (741 bp) positions 621-631
Tb927.2.2070 (474 bp) positions 132-142
Tb11.22.0002 (486 bp) positions 264-274
Tb10.329.0010 (513 bp) positions 281-291
Tb927.8.3080 (3915 bp) positions 1143-1153
*Tb927.3.2780 (3309 bp) positions 3146-3156
*Tb927.3.3050 (3096 bp) positions 2926-2936
Tb10.05.0160 (1569 bp) positions 528-538
*Tb927.3.1910 (1776 bp) positions 1738-1748
Tb11.01.6770 (2172 bp) positions 1536-1546
*Tb11.02.0020 (1941 bp) positions 1719-1729
Figure 3.1 Locations of GAGGG[CG]TGGGG sequence in *T. brucei* genome.

The above sequences marked with asterisks (*) are good candidates for further exploration because the likely UTR indicator appears in the last 20% of the sequence. There are five such sequences. Although the sample is small, this is noticeably more than the statistically expected number of appearances, which is approximately two. It is important to note that each of these sequences is currently a hypothetical protein, and that laboratory experimentation would be required to confirm functionality. With further UTR information, the five sequences with potential UTR regions might be good targets. This could be a good indicator of 5' UTRs.

UTR SEARCH

To search for coding regions in unannotated sequences, two main tools were used. The first tool, UTRscan [9], was developed by researchers at Istituto di Tecnologie Biomediche in Italy. UTRscan searches for approximately 30 patterns that are believed to indicate 3' or 5' UTR regions. More information about the patterns, including descriptions and sequence permutations, can be found at UTRsite [10]. These descriptions include functionality, mentions of conservation in other species, references, and historical information.

Another resource, BlastUTR [7], is maintained by the same researchers and looked promising, but has not been functioning properly.

MEME [3], the second tool used to analyze sequences for UTRs, was developed by three researchers at the University of California, San Diego. It searches input sequences for motifs and provides detailed output including locations, regular expressions, and p-values. MEME has the ability to find quite a few motifs depending upon the input parameters. These motifs nearly always have quite a bit of variability in the actual sequence, with only certain sequence locations being fixed. The sequences below show some of the MEME hits with the least variability.

The sequences in Appendix B were obtained from NCBI and run through UTRscan. Each sequence included a UTR in the annotation. The <u>underlined</u> regions are hits from UTRscan. The **bold** regions are identified as UTRs in the sequence's annotation. The *blue italicized* regions are motifs found by MEME.

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The sequence CACACATACAC (which appeared twice in the UTR of AM168497) appears 24 times in the *T. brucei* genome according to a BLAST search using *The T. Brucei Genome Project* website:

Tb09.v1.0620 (117 bp) 74-84 *Tb927.1.1320 (231 bp) 11-21 *Tb927.1.3440 (246 bp) 3-13 Tb09.160.3650 (297 bp) 202-212 Tb927.1.4060 (306 bp) 245-255 Tb927.1.4510 (306 bp) 59-69 Tb09.211.2690 (312 bp) 106-116 *Tb927.1.1250 (312 bp) 30-40 Tb927.5.2330 (13254 bp) 3048-3058 Tb927.4.4800 (393 bp) 264-274 Tb09.160.4060 (456 bp) 159-169 Tb09.211.3260 (510 bp) 417-427, 409-419 Tb09.211.4260 (537 bp) 275-285 tmp.1.100 (8300 bp) 3217-3227 Tb09.160.1410 (543 bp) 258-268 Tb927.3.1190 (6984 bp) 6671-6681 *Tb927.6.3210 (678 bp) 35-45 *Tb927.2.4440 (714 bp) 53-63 Tb11.02.4490 (714 bp) 454-464 *Tb927.4.3550 (1029 bp) 5-15 *Tb927.4.4810 (1095 bp) 162-172 Tb927.4.3280 (1233 bp) 566-576 Tb11.01.3740 (2637 bp) 813-823 *Tb11.01.6760 (1788 bp) 19-29 Figure 4.1 Locations of CACACATACAC sequence in *T. brucei* genome.

The above sequences marked with asterisks (*) are good candidates for further exploration because the likely UTR indicator appears in the first 20% of the sequence. There are eight such sequences. The sample is small as above, but again this is noticeably more than the statistically expected number of appearances, which is approximately five. Since these appear in the front of the sequences, these are more likely to indicate 3' UTRs. To get an idea how common the sequences in UTRsite are (these are the sequences that UTRscan searches for), the author submitted the first 330,000 bases of chromosome II as input into UTRscan. The results are shown in Appendix C.

The most common UTRsite sequence found in this section of Chromsome II was 15-LOX-DICE, with 153 occurrences. The following is a histogram of the 15-LOX-DICE locations as output by UTRscan above. Each bucket represents 27,500 bases. Therefore, bucket 1 counted sequence locations 1-27,500, bucket 2 counted locations 27,501-55,000, etc. The distribution is fairly even, with occurrences an average of 2,170 bases apart in this sample. Submitting each segment of the genome sequentially (in roughly 330,000 base sections, since the limit imposed by UTRscan is 350kb) could yield more interesting patterns.



Figure 4.2 Distribution of 15-LOX-DICE locations on chromosome II.

The next most common UTRsite sequence was K-Box with 34 occurrences. Since the 15-LOX-DICE appear to distributed relatively uniformly, the author became curious about how some of the other output sequences line up with respect to these. The locations and distances of the 34 K-Box sequences were compared to the 15-LOX-DICE sequences and an interesting relationship was found.

The K-Box sequences tend to precede the 15-LOX-DICE sequences. In 23 of 34 instances (62%), the nearest 15-LOX-DICE sequence was "behind" the K-Box in question. In other words, from the K-Box's starting position, it was usually more likely to find a nearby 15-LOX-DICE sequence in the forward direction. This could mean that the combination of a K-Box followed closely by a 15-LOX-DICE provides a stronger indication of a potential UTR segment than either sequence alone. The average distance from a K-Box to the next 15-LOX-DICE ahead of it is 1457 bases, while the average distance from a K-Box to the previous 15-LOX-DICE sequence is 6080 bases. Although the sample is small, this is more than a four-fold increase.

To confuse matters, the average forward distance from a K-Box to a 15-LOX-DICE is 3,225 bases, while only 2,619 bases in the backward direction. This might suggest that in the 62% of instances in which the forward 15-LOX-DICE is closer the sequences are in some way correlated. Since the average distance between 15-LOX-DICE sequences in this sample is 2,170 bases, it would also appear that the K-Box segments are occurring in the larger gaps between 15-LOX-DICE hits. This makes sense probabilistically, since if one assumes a uniform K-Box distribution, longer spans for the K-Box sequences to fall in would yield more hits. Otherwise, the expected average distance from a K-Box to the nearest 15-LOX-DICE would be about 1,000 bases. The actual average of 1,733 bases is greater, but not alarmingly so.

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DR. GOPAL'S RESEARCH

ORBIT [6], one of the tools used here to help predict whether an RNA sequence is coding or non-coding, was developed by Dr. Shuba Gopal, currently at the Rochester Institute of Technology. Dr. Gopal's work has focused on creating an alternative to annotation by sequence homology because organisms that are long since evolutionarily diverged tend to yield many false positive coding regions [2]. *T. brucei*, for example, are thought to be more than 800 million years diverged from *S. cerevisiae*, its nearest evolutionary neighbor.

Her paper (An organism-specific method to rank predicted coding regions in Trypanosoma brucei) describes a method that separates coding and non-coding regions based on nucleotide composition. Using standard sequence homology, more than 500 coding regions have been noted on *T. brucei* chromosome I, yet barely one-fourth of these have assigned functions. The reason so many regions remain unassigned is because there is little evidence for function besides homology, and experimental determinations of function for so many regions is unfeasible. However, if educated guesses could be made as to which regions to look at first (i.e., that were the most likely to be true coding regions), then the effort might be worthwhile. This is what ORBIT attempts to accomplish.

ORBIT identifies differences in nucleotide composition between coding regions and the region immediately upstream. This upstream region is rich in thymine and cytosine; an abundance of these pyrimidines appears to signal a trans-splice site. These

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trans-splicing signals are assumed to indicate non-coding regions because it is very unlikely that they will occur in the middle of a coding sequence.

To determine whether or not a region codes for a protein, ORBIT uses linear discriminant analysis (LDA). While LDA may not be as sophisticated as other pattern recognition methods, it was the optimal classifier for this simple coding vs. non-coding decision. Transition probabilities at the dinucleotide level were calculated using maximum likelihood estimation. The codon level, which comprises groups of three nucleotides and has three potential reading frames in each direction, did not provide useful classification information for the coding vs. non-coding decision.

Dr. Gopal's other tool is Motif-er. Motif-er's primary use is for genome visualization. *T. brucei* chromosomes I and III are currently mapped with coding regions predicted by ORBIT as well as current public annotations. Sequence information can be downloaded along with the coding likelihood score as predicted by ORBIT's LDA classifier.

DISCUSSION

The results are encouraging if not entirely concrete. With a relatively limited data set, enough clues and motifs have emerged to continue searching using similar methods. The motifs found by MEME in the UTR-only data set have yielded clues as to other possible UTRs, as shown by the five hypothetical genes in which the sequence GAGGG[CG]TGGGG appears near the tail.

The small number of UTRs that are currently annotated leaves a lot of room for improvement in this area. The current techniques are a good start, and some more advanced techniques could be a decisive step in better UTR predictions. While ORBIT's use of LDA may be optimal for a two-pattern classifier, more advanced techniques such as Support Vector Machines (SVMs) may be able to better learn the sequences and identify untranslated regions. Another advantage of using SVMs may be that there is more information than just that contained in dinucleotide transitions.

There are several sequences that show Internal Ribosome Entry Sites at their tails. There are also sequences whose annotated UTR does not agree with UTRscan's results. The 330kb section of chromosome II against which UTRscan was run gives an indication of the tool's sensitivity. In this section of bases UTRscan found 267 hits from the UTRsite list. This amounts to a potential UTR-indicating sequence every 1,236 bases. This might be slightly more than expected, however, each UTR might be composed of several different sequences, and thus this inter-UTR spacing would increase. The sequence CACACATACAC was found by MEME to appear twice in the same UTR, and may be a promising key to other UTRs. MEME is a very valuable tool, but it will be easier to use without the 60,000 base restriction. Being able to submit an entire chromosome's sequence at a time, for example, will allow motifs that appear farther apart than 60,000 bases to be elucidated. For example, motifs that appear infrequently—perhaps only once every 100,000 bases—could be stronger indicators of UTRs than more common sequences.

If motifs could be generated by MEME and shown graphically on a map of the genome similar to the one used in Motif-er, the location of these motifs could be compared with the predicted coding regions. This could provide very valuable insight.

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APPENDIX A

UTR ANNOTATIONS

This appendix contains sequences annotated as UTRs by either GeneDB or NCBI. The **bold** portion of the sequence is what's annotated as a UTR by GeneDB or NCBI. The <u>underlined</u> portion is one of the following highly conserved sequences that appears four times in this data A[AT]AG[CT]AGAGG), or twice GAGGG[CG]TGGGGG (see note below).

Tb927.1.1000 5' UTR Source: GeneDB Chromosome 1 289,877 ... 289,892 ORBIT: -16 bp UTR non-coding (.840) -1263 bp gene coding (.969) UTRscan: -1 IRES in 80 bp sequence centered at UTR GAATGAAGGTAGTACTATGCGTCGCTTATTGTGTCT...

Tb927.1.700 3' UTR Source: GeneDB Chromosome 1 231,710 ... 232,503 **ORBIT:** -794 bp UTR non-coding (.999) -1323 bp gene coding (.963) ACTTCCAGAAAAAATATATTTCTGCAAAATACTTTTGGAAGTTTGTCTTG TCTTTATAGATGAAGGATTTGTTTTTTTTGTGATGTTTTCAAGGTTAAT <u>AAGCAGAGG</u>TAAATTTTTTGGTGACACAAAAATTGGGAAGCTTCGTGTT CTTACTTGTTCAACTGAAAAATGCCTTTTCAGGAATTCATATTTGGGAGT TATTGTGGTGTAGAAGGACTGAGGAACAGAAGAAAGCAGAGGTTATTTG TGATACTGGGAAATGGACTCTAAAAATGAGAAATAAAGGGAAAGAGAAA GGAAGAGTGATATATATTTATTTTGGAAAAAAACACCTTTCGTTTGCTT

Tb927.1.700 5' UTR Source: GeneDB Chromosome 1 233,825 ... 233,904 ORBIT: -80 bp UTR non-coding (.918) GTTCAGCTCTTTGGTGATATCAAAGCATAATTGCTGCGGAGATACGTTTT TCCACCTAATAAGTAATTGTGATACAAGATCAAATCGTTTGGACTGTAGG ...

Source: GeneDB Chromosome 1 862,869 ... 863,029 ORBIT: -161 bp UTR non-coding (.999) -1062 bp gene coding (1.000)

Tb927.1.720 3' UTR Source: GeneDB Chromosome 1 235,665 ... 235,769 ORBIT: -105 bp UTR non-coding (1.000) -1530 bp gene coding (.972) TGTACATCAGGCGAAGGGTTTGTTTTTTTTTTTTTCTCCTGCCCTATGTTTTT CTGATGTCGTGGGAGGTTTTGAATACTTTTAGTATATCGTTTATTATTTGT GAACATTGGATGATAAGGAGTAAT...

DQ826505 3' UTR Source: NCBI Chromosome 1 58 bp ORBIT: non-coding (.996) ACTAGTTTCTGTACTATATTGTGAGTAGCCAGCTTTGACCAAAATATAAC TGACTGCTATGTATTCGAAAAGCA...

DQ826504 3' UTR

Source: NCBI Chromosome 1 19 bp ORBIT: coding (1.000) AGAAAAGACACGACCAGAAATGGCCAACACATCG...

N45755 5' UTR Source: NCBI Chromosome 1 376 bp ORBIT: non-coding (.999) TGTNCACCCGCTGTCGNCCGCTCTAGAACTAGTNNTTCCNCTGTGNCTGC AGGNTTTCNGNACGAGGTTGGTCGCCGCGAAGTTATNCCATACAAGGGC GTTTTTAGGCAGCAAAANCCAAGCAA<u>ATAGCAGAGG</u>CAAGGNGCTTCCN CGTAAGTNTAGTTAGTGGAGCGGTTTTCTNATGCNAACAGNCGTNGCTN TCCTGTTGNTNNTTTACAGNGGCAGTNNTTTTNTNGTNCAGTNTTTGGGG GCCATTTNGGANAAATGCCNTTTTACAAATAACNNTGGTAAGTAGCTTGT NTGTNGTGTTTNAGNNNACGTTGCTTCTANNGAANGTTTNNAAATTGGTN AATGTCCCTNNTTTNTTGGTGTTGGGAGT

APPENDIX B

UTRSCAN OUTPUT FROM NCBI SEQUENCES

The sequences in this appendix were obtained from NCBI and run through UTRscan.

Each sequence included a UTR in the annotation. The underlined regions are hits from

UTRscan. The **bold** regions are identified as UTRs in the sequence's annotation. The

blue italicized regions are motifs found by MEME.

AJ243568

ATCATCCTATATCCTCTGTGCCGCGGCCTCGCTCCAGGCGCTTTACCGCCACA AGAGGAATTCCCTCAATGAGGGTCTCCGCTTGTTCACTTTAGGAAGGCCACA AACCATCCGTTCCCGCAACGGGTGGAGACCCCAGCGTTCCCCAAACGCCGCTT CTCCAACTCCCGAAGAACCATCACCGCTTTTCGGGCGTCACGACTCGCCATCC AACGACAGCTTTTTTCGCCACATTGGAAGGAAGGTGGACAAAACACCCATCC ACCACGTGCCTTTTTCCCGTTTCGGTGAAGCCGGTGGGTAAGGAAATTGG GCGCCCAGAAAAGGGCCGTTACGGGAATTGAACCCGTGACCTCCTGCACCCA AAGCAGGAATCATACCACTAGACCAAACGGCCACACCGGCGGGGCACCAGGTC CAACTTATGCACCTATGCTGGAGTAGATTGGAGATAGCGCCGGGTCCCCGAA GCACCGTGGCGCAGGGGAAGCGCGATGGGCTCATAACCCATAGGACGTTGG ATCGAAACCAACCGGTGCTAAGTTTTCACATCCACCCTTTTTTCTCCAAAGGA AAATAAAGGTCGCCGGTTCGCAAAAAGTTGACGAGAGTGGGGTTTGAACCCCA CGCCCTCGGAAGGATTGGAACCTTAATCCAACGTCTTAGACCACTCGACCAT CTCGCCACGGGACACCGCTACAGCACAAAACATCGACACACCGCAATGAGC TCGAACCCTCGACAGACGAAAAACCACGTGGTATGGAAGCGGCTGAACAAG CGCCGTCTCGACTTTTACTAGGCGGCGCGCGCGCGCGATTAGCGTTTAAACTTTGGG GTGTCGGCGGTTGTTTCCGTCCGGTGTCAATATTTTTTCGCTTTTCCCACGGA AGGAAAGGTAGCAATTGGGTCCGCTGGAACTCGGCTTCGCGACTGCCTTCTG TGCCAAAGTGGCCAGAGACCCTAATAAGAGACATAAAGTTGAGTCCAGCAAC CGACTGCCGTCGCGCTTCGTCCAACAGCAAACTACGAACAAAATCCCACGGG CGGCGAGCACATTTCTGCTAACTAAGAGTCTGCCCGACAGAAACGAAATAAG ATGCCATAGTCCTTGCCACCTGATACTGGTCACTGGTGAAGCCGCGTCGTCAC CCACGTCGTCGCTGTACCTGTGACAGCCAATTCATCACTCTCAGGTGCTTCG ACCAGGAGGATCAACATCGATTGTGCATCAGTCCCATGTGGCCGGGAAGCGG

GCTTGTCTGCGATCCGGCTAGCTCTAGTCCAATCGATTTTGTCGGGGGGCAAAT TTGGAGTAAAGCATGCTACCTGTTCCAAGGGATGGCCACTCCATACTAACTG TCTCGGCTAGACCCGCGGAGTTAACACGCAACGACCGGGTAATTCATCCCAG CAAATCGGTCAGAGCCGCCATGGCACACTCACCGCGGGCGCTTGTGTTTTC CAGAACTGAGGAAACAACGAGCGTCGCTATTAAGGCGCAGCACTAAACAGC ATCAGTCACGCGCCGCTAAAAGAGCGAGTCCCAGTGGAGAGGTATTAAATAA TTAAAGTAAGGTCTATCGGGTTATGAACAGTTCAGTTGGTTATAATATCCCGG GTGGAATCGGAATATTGAGGTCCTTTACTTTTAACACTGAGCATATATTTGCC GACATGAAAATTGCGGGCGCGTACGCTGGAGGAAAATGCTGCGCTGAAGGG TGCGACTCGGAAGAAGAGAGTACTCAGCCACAGCGGCCAGTATCGAACCAA ATGCAGTGAGGCCACGGCCGGCTGCAAACAGAAAACAGGGTTCCCCATACA ACAAACGGGTGTTCACAGGATTCACAATCCCAAAAATCTAGTACCAAACACT GTCAGCCTTCTCGCATCCACTAAGGATCTAAAATGTAACTTTTACAAGTAAAT ATTTGCAACTAAGAGAATTTTAACGCAGCGGAAATATAGACACAAGACAAAT CCAAATAGCACCTGCACGCATAACCGAAGGTATACAGGAAGATTCCTCCCAT AAATGTACAGCTCGCGCCGATCCAGATAGGCAATTGAAGTACGTATGACCTT CATAATTACCTTTTGCGATTGCATAGAGGCAACAAAGGGGGTCTTGGGAAATG AGGATGGAGATGGGATTTAAATGGTACGAGTAGTATAGTCGATTGCGTTGCA CAAAAAATCCGCTAACGCTGCTGCATTTCTCATTTGACTCGACCATTGGGGGAG AGTCGTGAAACGCGCGGGGTAGTAGGAAGGGGAAAGAGCGTGATACCGGCCG CCGGAAAGCTTGCAGCTGCACCACTCGCTATTCCACCTCCACGCCGTGGTATG CGTCTTGCCTCGCTCTCTGCGGTCGACTGCGAATTATGCACAGGGAACCCAAT ATGTCAGTTATAACACCCACAAAGGGGGACCGGGAAGAGGTGGTGAACTTATC GAAGGGTTTATTTGGATCAAGCGAATATCGTCACCTTAGGGAGGCGATTGGG GGCGCAGAAGCTATTGCTTCCACCAACTTATGGGTTTCTATTTTCCGATGGTG CAGGGATTTATAGGGGTGAGTGTGTCTTCCTCGATACTTCGGTCAAACGTTAA AATGTCCTCCAACTGTGGCATCCGTAAATTCGTTACGATCTTCCACCACGGAA TAACACATACGCGGGCTTGTGGCAATCAAGATGCGGATTTTGCTGAAACATG CATTTTTGCAGCCTCAGAAAAAGATCCCAACGATGCACTTTTATGGTATATCC ACCACTCGTCAGTTCCCAGGCCCTCTGCGTGCTTGCTGAAAGGGTGAGGGAG TTGCAAGGGTAATGTTAGAAAATCGCTGCGCGTACTTTCGGCGCGATTTGTCT ATTTACTTTAGCTTTCGATTGGTTAGGACCTCTATTTTCGGGTAAGCAATAAC TCATTTCATTCTCGTAGAAACGGTGGCTACACGAAATTCCAATGTTCCCT TTCCGCGGAAAAGTCGCTTGTACGAGAAACTTGTGAAGCAGACGAGGGCGCC CACTCGTTATGGGGGGCATGGGGAAAGGGTGGTGGTGAGAAACCATAGGACGC GATCAAATGGCGTTTCTTTTGCCTTTTCCTTTCTCCGAAAAAAAGTGAATACC GGAGAAGGACGTAATGCCGGATAAAATTACATCTCTCGTGGATACGAGGGGA AACTTTCGTTCCCCTACGCAATGCTAAAGATAAATCAGTGGATTGCGAAAAA GGTAACGGCAACTGGTCGATAAATTAGGAATAAAACTAAAACATAATGGCAC TGCACCAAAAATTTTTAGCGTTTTCCTTGCTGTGGTGATGCCAAAAATGGGCG TCGCCTCAACGAATAAAAAAAAATTGACCGATAAAAAGTGAAGTTGGCCAC TTTTGAGGGTGGGGGGGGGGGGGGGGGGGGGGGGCCCTCACTTGCACGGCACT ATTCAGAGGTTCCATTCATGTGTGTTTTTAAGTTGTGATAGATTTACAAAGGA

DQ246439

CGCTATTATTAGAACAGTTTCTGTACTATATTGAATCCACTACAAGACAG CAGGCACAAGCTTCGATACCATCCAAATTAACAACAATTATGAGCATCAC TTTTCATAGTTTATGGCTACTTCTGACAGTGTTGTGCACTGCAGGTATTCGTG **GTGATCGAGTCTGGTACGATTGTCCAGAAAAAGGTGTGGACACATCGCGGGA** CGGCATACAAGCTTTGTGCCGTGCGGCGGAGCAGTTCAGAGGCCTATCACAA ACAGTAACATCTGCTGTGGAAACTTCCGCTACTGCTTCGAGTAAAGCATTTGA AGCAAAGGTACAAGCAGAGGAAGCTGTGGAACTTGCCGAGTCAAAAGGCCT AAACGTTACGAAAGCGAAGGAAGCTGCTGTGAGAGCAACACTCGCTGCTGA AGCTGCGGCTACGGCTGCAAGTAATGTGGAAATTAACGCTGCAAATATTGCT GCGGTGCCGTGGTCACAACCAAGCAGTGATGCAGGTTTACAGAAGCTGGCAC TATGTGAAAACATCGACAAGAGTTTACGACAGTTGGCATCAGAGTGCTCGAA GAGAGCGGAAAACGTGACAGCCCAATCGCTCAGTGAGGCGTTGGAGGGACT AGATGTTGAGTTCCACAAAGAGTTTATGTGGTTGCAGCACCACCTCCGGGAG **<u>GCAGTTCATGCACGAAAAACAGGCTGAGGATGCAGCTGCCGAGGCGAATGAA</u>** ATTGCCGGCACAAATACGGGACCAGTTGGAAGTTCCGTTGCATCACCTGAAG **GGTCAGTGCTGCTACTGATGGCTGGACTGTTTCTCAGTTCTCTACTGTAAGAG** GTTACAACTCCATGAATTGTGATCCCAGCCATTCCACTTGTCTCCACACA GTTATGTGCACATACAAGTGGGGGCAGCAAACATTCTTTCATATTAACCTT ATTTTCTATCAGTGACTATTAATATTGTTATTTTATTTGTTTCCTTTATA **TTTCGAACAGTTGTTGTAACTTTTATCCTTCATTTTTATTACTTCTTAACT** AGTGTGTCTATTGTCCTACTTTTTAGCTATATTTTTTCCTTTATTACCTTT GTTATTTCATTATTTATTCATGTAATGCACTGAGGTGATTGCGTGTAAG CATGGCCACCTATGCGTTACAATAGTGACACTTTTTAAATCAAAATCTAT **CATGTCTACGCATCCATAATGTCCTATTGTATGCGCAAGGCTAACTGATT CTGAGGGTTTTTTATTTATCATGAAAAAA**

Z15031

TTTCTGTACTATATTGCGTCCTTTCCCACAACGGTAAACATCTAAGAAATA ATGGCAGAGGAGACATCGTTGGTTGCAGATAAGGTTCCAGAGCCAGCGGTGA TTGATGCCGTCGCAGATGCAATGCCGGACAGCCTCGAAGACGCTCTCCGGAT TGTGTTGATGAAAGCTCGTGAGACGAATGGCCTCATTTGCGGCCTTTCAGAG GTCACACGGGCCTTGGACCGCCGCACGGCTCACCTTTGTGTACTTGCTGATGA AACATTGACCTCGTAAGCATGGACGAGCGCGAGAAACTCGCTCAATGGGCAG GACTCACCAGAATGGCCGCCGACGGTTCGGTACGGAAGACGTTGAAGTGTTC CTGCCTTGCTGTAAGGGATTTTGGCGAGCGCACAAAGGCTCTTGACTACCTTC **TGTCGCAACTGCAGTAATGTAGTGAACTGTCGCGGCACCGACATCAGCAC** GGAGTACGAGGGCGGGGACTGTCATGTTTGTGTTTGCATTGTTGAGGTGG TGTAGCATGTTGGAGGAAACGTTTTCCTATGTCCTCCATTACAGCTCGTT CCATTTGGACGTTGCCCCCTCTTCTGAGCTTATTGATATTTCGTTGTTTA AACAATGAAATTGATACTCCCCTTTTTCCCTTGTTGAACCCTCGATTCCT CTGTGATTTTGCTCTTCTTACTTGTCTGCCGACCTTTCTCTTTGAGGACG TCGATCTGTGGGAGGGGGGGGGGGCCACTTCTCTCTTAAAATGGTTTACTTTATT CAAAAGCTATCAGGAGTGACAAGGGTCTACGATGCATTACTTCAGTAAA

AM295303

CGATCCCTAGAGTGATTGCACCTTGAGAACTGGTGGCAGCAGAGTTGTG AGCTTTTACTAAAAGAAAAAAGGCCACGTGATCGACTTGTAATCAACCG CTTTTGACAGGACGCGCGTGCAATAATAAGGGGGGTTCGTATTCTTTTG **TTTTTTTTCACCAGGCAGTCCGTTGGATGCTAGCTCTTACATCACGCCG** GTTGCTGCAGCAAACATTTATGCGATGCTGTAAGAGTGTAAACAGTGTG ACCCTTGTTGGCGTCGTTCACGATATTCAGAGCGGCTTTGTATATGAAGATGC CGTCACGCAGTTTACACTAACAACGACGAGCATTGACACGACACATCCAACT CAGGAGGTTGTCGTCGAAAAGGACCACCACACGATCCGCTGCTTCGGCGAGC GAATGGGAGGCTTCGTCTTACCCCTCAACTTGAGCCTTCCTGCAACAACACT TTTACTTTCCATACATTCAGGTTCAGCCACCTCATGGCCAGGTGGCGGTCATC CATGGCGACAGGCGGACGGTTCCCGCGGCAGTGAACCCTGCGGTGGAGGAT ATAAAGTCGGAAAAGGAGGGCGCTGGTGGTGGCCAGAGTGGTGGCGTCGT AATGCCCTTGTGTTTCCGCAGTGTGAAGGAACTGTGATGGTTTTCTTCCT CACAGATGTGGAGGTTTCGTGAAGTGAATGGATTGGTGGGGGAAATCGCT TTCCTCTGCTTTGTTTCGCGCTTCGGCGACCTTCGGTGTGTTTGGAATGC **CGGCGTTGTCAACTAAATTAATGGTGTGTGACTTTGTGGTGCTTCGGTTG GTGAGTTTAATCGCAGGTCTCTGTGTTTC**

AM168497

AJ879575

ACAGTTTCTGTACTATATTGTATGAAATAGGTCCGGCGTGTTGGTACAAT AATGCGTAGGAACAATCGCACACGTTGCACTATTGATGATGTGAATGGAATG CTTGCCAGAAACGCGCAGCTACGCAACGCCTTACAGGAAAGGTACAAGCAGT TGAAAATGCGATATGAACAGCTGGCAGCTCTGAGAGCGGCCCTCTATCCTTC GCGGGGGGTGACTCTGAGGAAACTAGGTGTAAGACAGGAAACTGCTGACGG TGCTGAGGAGGTACGTTTCCTTGACGATTATACAACGGGAGGTGTGGGTAAT CCCCCCTTTCGTGACGCTGGTATTTATTCAGCCAAGAATATTGTATGTTACGC ACCGGTTCCACCTACACGAGACGAGTTGCGATGGAAAGGAGTGACGTTGGCA TTTCCACAGTTGGCATACGTACATTCTTTGGCAATGCTGCCGGAAACAGCGAA CAGTTTTTCTAAAGCTCTCCAGTGGAGTCGAGAAGAAGATAGTGCACTCAGA GAGCAGGTTCACGCCTACAAGGGCGCAAGGTGTGGACCTTCTTTTGGAAAGC ACTTGGTGCACCCGGTCAATCACGATTCGAAGTGGCGAGTCATTATATGGT TGCAGCAGCTTGGACTCATCGAACCGGGGGAAAAACGACAAAACGATGATGT GGATCCCAGAGGAACAACGCGATATTGCTTTGCGGGGAGGCAGTGTGGCGTCA TTTAGGTGATGAGGGTGGCATTATGGCGGCTTACGTGGAAATTATTAGCGTTG CAGCACGGAAATGTGTGTGCCTTTCGGAGGTAACATGTAATGAATCACTTGT GTTCCCTCCGTATGTGTGGGTGAAGCGTACCACCTTCAATTGGCTAACTTCAC TTCTCATACAGAAAGCGAAAGCTCCTCTCGTGCGATGTGAAGGAGAATTTTC AGATGATACACCTTTGTGCATGCACTTTAAGAGTGACATTTCCCTAAAACTTT CTGCAGAAGATATGATGGCGTGTTTGTTGGCTTTCAAGGGTGAGGTGTTCGG AGAGGTGGGTGGACTGCGGTTTATTGAGCGCGCGTTTTTGCCGAAGAATAGG ACCTATGCACTGAAGGCCACAGACTTCATGGCTAAAAGAGAGGGCAAGAAA CATAAACTGGATGAAGAAGTTTGAGGTGCCTGGGGGGGGAGATCAAAAATAAA **GGAGCATAGTGCCAAGTAATGTGATTATCATGCACAATATATCACTGTTC** CTCACCTTGCGCGCACTCGCATATGAACTTGCTGTTGCGCCATATCTATG TGTGCGTGGGTGTTACTGAGAACCCTCGCTTTCCTTTGCTTGTTTGCCGA TGCTGTTGCTGCGGGTTCCTCCTGTCCATTTGTGTGAGTACTGCGTAT AAATTAATGCGTAAATATATACGGCACTGAGGAGTTTGTATGTTTGTCTC

L30155

TATTATTAGAACAGTTTCTGTACTATATTGGGTGTCAAACTACTGCCGCA **TAAACTACGGTTATCCCAAATTTAAGAGAAAGCAATAAAGCATCA**ATGTG GTCCCGGAGGTAACAGATATCACGCTGGAGGCCGCCCGCAAGCAGAAAATTC ACAACCTGAAGTTGAAGACCGCCTGCCTTTCGAATGAGGAATATGTCCAGGA CCTGCACGTATCCGAGTGGAGTGAGACGCAGAAGCAGAAGCTGCAGGCTGC ACACGAGAAAGCGCATGAATTGCTTGCCTCAGTGGAGGGTGGGACGAAGTG GAGCCTGACAGAGGCGTATGACATCAAGAAGCTGATGCGCGTCTGTGGTCTT GAGATGTCTGTGCGTGAACTGTACAAGCCGGAGGACAAGCCACAGTTCATGG AGATTGTTGCACTCAAGAAGACAATGAACGAACTGAAGCAACATCACAACA AGACTCGCACGGTGTCTTTCACCGGCATGATCGACAATGCCATCGCCAAACT GGAGAAAATCCAAGACGAACTGCGCCGGTCCCAGCTCGACGCTTCTGAGATG GCGCAAGTTCCTGTGGCTGCACTGAAGAATATTGAGGACACGATGAACGTGG CTGTTGTGCAGACGGCTCTTCTTGGGAACGAGGAGCAGATCAAAGCCCAACT TGCAGCCGTTGAGAAGGCGAACGAAATCCGTAATGTTGCCATTGCCGATGGT GAGATGGCGATTGCTGAGGAACAGTATTACATTAAGGCGCAGCTGTTGGAGC ACCTTGTGGAGCTTGTGGCCGACAAGTTTCGCATCATTGGGCAAACTGAGGA TGAGAATAAGAGCTTCAGTAAGATCCACGAGGTACAGAAGAAGTCATTTCAG GAATCTGCCTCAATCAAGGACGCGAAGCGCCGCCTTAAGCAACACTGCGAGG ACGACCTACGTAACCTTCACGATGCCATCCAGAAAGCTGACTTGGAGGACGC CGAAGCCATGAAACGGTTCGCCACGCAGAAGGAGAAGTCGGAGCGGTTCAT CCACGAGAACCTCGACAAACAGGACGAGGCATGGCGTCGCATTCAGGAACT GGAGCGCGTGTTGCAGCGCCTTGGGACGGAGCGTTTTGAAGAGGTGAAGCGC CGTATTGAGGAGAACGACCGCGAGGAGAAGCGTAAGGTGGAGTACCAACAG CTGCGACCTTGCGCTTCGCTGCATGGGTATGCTGGAGGAGATCGTAGCCGAG ACCTTCGGCTGCAGGTGCACCAGGAGTACCTGGAGGCATTCCGTCGCCTGTA CAAAACTCTTGGCCAGCTTGTGTACAAGAAAGAAAGCGCCTGGAGGAGATT GATCGCAACATCCGCACCACACACATTCAACTGGAGTTTGCCATTGAGACCT TTGACCCCAACGCGAAACTACACTCCGATAAGAAGAAGACCTATACAAACT TCGTGCGCAGGTGGAGGAAGAGTTGGAGATGCTGAAGGACAAGATGGCGCA GGCGTTGGAGATGTTTGGACCTACTGAGGATGCGCTGAACCAGGCTGGTATC GATTTTGTTCACCCTGCTGAGGAGGTTGAGTCCGGCAACATGGATCGCCGCA GCAAGATGGTGGAGTACCGTGCACACCTGGCGAAGGAGGAGGAGGAGGTGAAGA TTGCCGCGGAGCGCGAGGAGCTGAAACGATCTAAGATGCTCCTGAGCCAGCA **GTACCGCGGCCGCACGATGCCCGAGATCACTCAGTAGCGCTGCGCTTAAAT GTCTTTCATTATAATCAATGTATAACCTTTATGTAGTATTTCAATCTATGC** CGCTGTGTACGTGCACTGCGGTGCCTATCCTTCGGCATTAGAGAGTCAC **TGTTTGTGTAGATCGTAGCTGCATGTCTG**

AACGCTATTATTAGAACAGTTTCTGTACTATATTGGCAAGACATACTGGG GAAAAAGAAAGAAACCTGCACTGACCAAAAAGACTGAACGAAACGAAA TTGAGGCGATTGAGACGCCCCTTTCTGTAAGCGGGGTTTAGTTCATATTC GAACGGAATGGGAGGGTGTACCTCACGTGGGCTCTCAGAAGAGAAACTCGC ATGTTACTCCCACCGTACCGGCAACCTCGTTGATGAGCATCTTTCGACAGGGG CAGTTGAGGCACACGAGCTTCAGCCCTTCTTTTCTTCTCTGCTGGGAGCCATC ACGGACCTTTTGAAGTGCAGCCGGGAGGATGCCGTCGAGTTCCTTGCATGCT CAAGTAGTGCAAACCCACGGGCGGCAGAGTTGTTTACTTCCTTTTGTGCTGCA AATCCGCTGAATCTCATAAAATGGGATGTGAATCACGCGAAGTTCATGATGA TATGGATAAAGTACGACGACGACAACAGCGGGGGACATATGCGTTCGTGAATT GAGGAAAATCTTGAAGGGTTTGAGCTTCCCTGAAAGGCTCTCACAAAAGATG GGGACATTTATGTCTCTGACAAGACTTAATGAACTGACATATGCAATGCGAA ATGTCGTGGGTCCCGATCGGGACGTGATGACAAAGGCTGAGTTTGTTACCTTC TAGACGCTATTGGCTGTACCGAAGAGCATCCTCTACATTTGGACGCATTCTTA TCATTTCTCAGCGACAGGCGCTTTAACTCCATTGTGAACAACAGAAAGGTGTC TAGTGTTTACCACGATATGACTCGCCCGATATGTGAGTATTTCATCAATTCCT CGCACAATACCTACCTTACGGGTGATCAACTCTTGAGCAAATCTTCCACGGAT ATGTACAAGAGGGTTCTACTGGATGGCTGCCTCTGCGTTGAACTTGATTGTTG **GGATGGTCGCAAGGGTCAGCCTGTAGTTTATCATGGTTACACAAGGACTTCC** AAGCTTTGGTTCCGGGACTGTATTAGCACGATCAAGAAGTATGCTTTCGTTAA TTCAATATACCCTGTCATTTTGTCGCTTGAGGTTCACACTAGCCTCCGCCAAC AGGATCGAATGGCGGAAATTTTGTGTGAAACGCTCGGAGATATGCTATTCTG CAGTCCTTGGGGTGCTGGTGAACAGACTTCTTTCACGTTCTCGCCGGAAGCGC TAAAGGGAAAAATTCTGCTAAAGAGCAAACGGGGCTACTACACCTACCGATGG GGTACAGGTTGATGATGACGACGATGAGGATGAGGAAGCCGATGGTGTGGT GGAAAATTTCGTACCACCTGAAACTGCTCGGCGTTGTCGCGGTGGTGGAAAA ACGAATTCAAGGGGTGCTGAAAAGAAAGAAGGGTTTCAAAGGTTTCAGAG AAACTATCTCGTCTTATTTCAATCGAATCCATTGGTTATAAGGGTGTTGAGGA TCTAAGTTATCTTGAAACGCGTCAACCATATCACTGCAGCTCTTTTACTGAAG GGAAAGCGGGGAAAATTGCCTCTTCTAACCAGGAGGAGTTTGTTGCCGTCAA TAATCGGTGTTTGAGTCGCATATACCCCACAGGAACTCGCATCGGTAGCAGT AACTTTCATCCTCAAACGTTTTGGAATTGTGGGTGCCAACTAGTTGCACTCAA TTGGCAGAATTACAAGTCATACCAGCTTAGGCTAAATAGGGGGGTTCTTCAGC GACAATGGTAACTGCGGCTACCTTCTCAAACCGACTGCTGTGGACATTGCAC GTGCAAGGGGGCCAAAACGGCAGTCACGGTTGCTCACAATAGAAATTATATC AGCTTTCTGTCTTCCCAGGCGGAAAAATGCATCCGGTAGCAGTATTGTGGATT CTCGCGTAGCCGCCTTGATTGAGGGCCCCGGCATGGAGAAAAGCCAACGAAA TAAACAACGAGTTCTGCTGGAAGGTGTACGAGTGGGAACTGTCCACCCTTGT CATGCAGGTGTATGACGAGGATACCAAAAGCAATAACCTTCTGGGTGAATAT GTTGTGCCATTACGTGCCCTAAAAGAAGGAATTCGCCAGGTCCCCCTTCGAG ACCTCAAAGGATCTATTATACATGGCTCTTTTTTAATGGTTCAAGTATCTTATC

AGTAGGAGTTTGAGAATATCGTGTTCTTCAATTGGGTTACAAGTGTGCGT TGCATGTACCCGACTTTATGGCACCATGTTGTGCCATGGTCTTCACAGCG GACACTGTTATTTTCATTTTGTTGGCCGTTCTATCGTGCCTTCGGTGAG TATTACACTCATCACGACATCATATATGAACACACCGACGGTGTTGTTTT GTTTTGTTTTTTTTTGCGTGTGGGTAAAATCACCAGGGGATTCCCAATG **TTTCGACCGAGGATAATGTTTCATTCAGTATCTACTTTTCCAATTCATCA** CTTCATTTGAGTTTATTTCTTTGTTGTCTCCGCTTACTTTCATGTCCTCAC ACCGTGAGGCAGGATAGGTGAGCTGAACGATTTTTTTTCCTTTGTGTGT GGTGGTTAGAAGGAAGGAACGTAATATGGTAATTGGGCTTATATGGGTC TGATGCTTCATTTTGTCCTTCCCTTCGCACACGTGCTTCCAGCTACGTGT TCGTATTTCTGTAAATGTAAATTCGTTTCGTAGAACGGATGCCTGTTTCC TCTGCCGATTGTTTCGCTGCTCAAGGGACTGCGGCGGTCGCAGTTAGGT AACTGATTTTCCGTTCCACATCCCTACTTTCTAGTTGATCGAATATACGG AAATTAAGCTACAAAAGATATCGGAAGTCAATGATGGTCCAGACGAAAA AGTGCATCGCCACTTTCCGGGGCCACAGGCAAGATATGTGGGATGGGCAA

Z54338

CTGTTTCCACATTG*TCGTCGGG*GGTTGTGTCGCTTCTGTAGACTTTCCATTTTC TATTTGGGGACTCTTTTGAAACCTCTCTGACCATAATTTGCTTTCATTCCTTCT TCTGTTTCGTCTTTTCCAATACTCGATGGCAGCCTCCCTTTCCAGTAGCACAAT GGCAAAGAAGGTCAAGTCGAAGGTGGACACCATCAA*CACCAAGATCC*AACTG GTGATGAAATCCGGCAAATACGTTCTCGGGACGCAGCAGTCACTCAAGACAC TTCGTCAGGGCCGCAGTAAACTCGTTGTCATTTCCGCTAACTGCCCGCCGATC CGCAAGGCGGAGATTGAGTACTACTGCACTTTAAGCAAGACGCCAATTCACC ACTACAGCGGCAACAACCTTGACCTTGGAA*CGGCATGCGGA*AGGCATTTCCG TGCTTGCGTACTTTCCATTACGGATGTTGGTGACTCTGACATCACTTCTGCAT AATCGCAACGGTGTAGGTGTGTGCCGTATGTCCTTACCGAGAGTCGTTCAAGT GATTT

M81386

ACTGTGCACTGGACGAGTGCTCATAGGTCTCGGCGTTGGTATTTTATGCTCTG TTTGCCCCATGTATGTGAATGAGAACGCACATCCCAAACTCTGCAAGATGGA CGGTGTGTTGTTCAGGTGTTCACAACACTTGGCATTATGCTTGCCGCGATGC TGGGTCTGATTTTGGACAAAACAGGAGCTAGTAAAGAAGAGGCAAACATGG CTGGGCGGTTACACGTTTTTTCAGCGGTACCGCTTGGATTGTCCGTCGCCATG TTCCTAGTGGGCATGTTCCTCCGCGAAAGCACTGCAACATTTGCCCAAGACG ATGATGGTAAGGCTGATGGCGGAATGGACCCCAACGAGTATGGCTGGGGGGCA GATGTTGTGGCCACTGTTCATGGGCGCTGTAACCGCTGGTACGCTGCAGCTGA CTGGGATCAACGCGGTAATGAACTATGCGCCGAAGATTACAGAGAACCTCGG AATGGATCCATCACTTGGCAACTTTCTGGTTATGGCATGGAATTTTGTGACAT CCCTTGTGGCTATTCCACTTGCGTCACGCTTTACGATGCGTCAAATGTTTATC ACCTGTTCCTTTGTTGCGTCATGTATGTGCTTGTTCCTATGTGGAATCCCAGTG TTCCCCGGTGTTGCAGGAAAAGAGGTGAAGAATGGTGTGGCAACTACTGGTA TCGCCCTGTTCATAGCTGCATTTGAGTTTGGTGTTGGGTCGTGCTTCTTCGTGC TTGCACAGGACCTTTTCCCACCATCATTCCGACCTAAGGGCGGTTCGTTTGTT GTCATGATGCAGTTTATCTTTAACATCCTTATTAACCTATTGTATCCCATTACA ACTGAAGCTATATCTGGTGGGCCAACTGCCAACCAGGACAAGGGACAGGCC GTTGCATTCATACTGTTTGGTTTAATTGGCTTGATTTGTTCCGTTCTGCAGTTC TTTTACTTGTATCCATATGATGCCAATCAGGACCATGAGAATGACCATGGTGG TGAGCCTGTGGAACAGAAGACATATCCCGTTGAAGCATCTCCGCGGAACTAA **CTTGCATATTTCTCGCGCATCTGATGAATTAAAAAAAACGATTATTGCA** TAACATGATTATCTGACCACAAAACGTTTTGTAGTTTGAAGGAGGTAATT GGGTAATGTTTTAGAGGTCGTCAATATTAGTGGCGTTAATGAAAACGGA TTTTAAAATTTACTTCTTTTTGCTGTTTTATGTTGTCTATATACTTTTG TTTTTCCATCAAGTCGACTGTGCCTATTATTATCTGCTCGGTTTTGTAGC AGCGGATGGACAGATGGATGAAGTGATATATGAGGGCAGTATGCTGTTA GTGTGTATGTGCACTCTAAAGCTGCTGCTGTGTCGGGATAGTGATTTAC TGAAGACGTTTCTTAATAGTTTTTTGAAAAAAAACATTGTGTTGTTTTTT ATTGGTGTACAGGGGGACAACTTGTTTATT

APPENDIX C

UTRSCAN OUTPUT FROM CHROMOSOME II

This appendix shows the results of submitting the first 330,000 bases of chromosome II as input into UTRscan.

Pattern = Histone 3'UTR stem-loop structure

Pattern not found

Pattern = IRE

Pattern not found

Pattern = SECIS-1

Pattern not found

Pattern = SECIS-2

Found 3 matches in 1 sequences

seq :[152066,152127] :GATA ATGTATGGA A TGAA AGTGTGGA AAC AAGGTTGAGAGAAA TCATGTG TGAG ACTG TATT seq :[171238,171301] :ATGA CAAAT A TGAT GTTACCAT TTA AAA AATAAGTAGCAATA GTAGTAG GGAG AAAGCTA TTGT seq :[313819,313877] :CGCT GACAACT A TGAT GTTTTAGA AAG CAGCGGCATGGTCG GTGAAAC GGAC TAC AGCG

......

Pattern = APP

Pattern not found

Pattern = CPE

Pattern not found

Pattern = TGE

Found 3 matches in 1 sequences

seq :[16493,16527] :GTCA ATTGAATATCT CA TTTC TT GTATGTT TTTCT seq :[53295,53331] :CTCA CACTGAGGCCGCA CA TTTC TT TCAATTG TATCT seq :[138215,138251] :CTCA CACTGAGGCTGCA CA TTTC TT CCAATTA TATCT

Pattern = NANOS TCE

 $I allelli = IAAAOS_ICE$

Pattern not found

Pattern = 15-LOX-DICE

Found 153 matches in 1 sequences

seq :[6,21] :CCCTATCGCT CAA ATG seq :[266,281] :CTCCACCCCT TTC AGG seq :[2743,2757] :CCCCACCTCG AT ATG seq :[3062,3078] :CCTTACCCCT CACA ACG seq :[5563,5581] :CCCTTCCTTC TCCATT AAG seq :[9871,9889] :CCCTCACTCT GAGTAA ATG seq :[11047,11060] :CCCCGCCCGT A ACG seq :[18933,18952] :CCCTTCATCC TCTGCGC ACG seq :[22569,22582] :CTCTGTCTCT G AGG seq :[23118,23137] :CCTCGCCCTT CCCCGGG ACG seq :[23191,23209] :ACCGTCCTCC TTCACT ATG seq :[24782,24801] :CCCCTTCCCT GCCAGTA AAG seq :[26891,26907] :CCCTAACTCT GCCA ACG seq :[32386,32405] :CCCCGTACCC TTCCTAG AAG seq :[32735,32750] :CCCCACCTCT TCG ACG seq :[34158,34176] :CCCTTCCTTC TCCATT AAG seq :[35291,35306] :CACACCCTCC GAG AGG seq :[37159,37179] :CCCTACCCTTC ACAAAAT AAG seq :[37460,37476] :CCAACCCTCC TGCG AGG seq :[39542,39560] :CCCTTCCTTC TCCATT AAG seg : [42814,42833] : TCCCGTCTCC ATTCAAA AAG

seq :[45668,45682] :CCCTATCTCT AC ACG seq :[45933,45952] :CCTCACCTCC GGCACTC AAG seq :[46083,46097] :CCGCTCCTCC TA ATG seq :[46104,46118] :TCCCTCCTCT TT AAG seq :[47343,47357] :CGCCTCCTCC GA AAG seq :[48263,48281] :CTCCATCTCT CAGTCC ACG seq :[48712,48728] :CCCCTTCCTT CAAT AGG seq :[51793,51807] :CACATCCTCC CT AAG seq :[52928,52943] :CCCATCCTGT CGT ATG seq :[54150,54165] :CCCTATCGCT CAA ATG seq :[54412,54427] :CCCCACCCCT TTC AGG seq :[56896,56910] :CCCCATCTCG AT ATG seq :[57215,57231] :CCTTACCCCT CACA ACG seq :[59739,59757] :CCCTTCCTTC TCCATT AAG seq :[60892,60907] :CACACCCTCC GAG AGG seq :[65131,65149] :CCCTTCCTTC TCCATT AAG seq :[66259,66274] :CACACCCTCC GAG AGG seq :[68127,68147] :CCCTACCCTTC ACAAAAT AAG seq :[68428,68444] :CCAACCCTCC TGCG AGG seq :[70510,70528] :CCCTTCCTTC TCCATT AAG seq :[73798,73817] :TCCCGTCTCC ATTCAAA AAG seq :[76652,76666] :CCTTATCTCT AC ACG seq :[76917,76936] :CCTCACCTCC GGCACTC AAG seg : [77088,77102] : TCCCTCCTCT TT AAG seq :[78327,78341] :CACCTCCTCC GA AAG seq :[79244,79262] :CTCCATCTCT CAGTCC ACG seq :[79331,79344] :CCTCATCCTC A AAG seq :[79693,79709] :CCCCTTCCTT CAAT AGG seg :[86382,86400] :TCCTATCTCT ACACAG ATG seq :[86647,86666] :CCTCACCTCC GGCACTC AAG seq :[86818,86832] :TCCCTCCTCT TT AAG seq :[88057,88071] :CACCTCCTCC GA AAG seq :[88980,88998] :CTCCATCTCT CTGTCC ACG seq :[89432,89448] :CCCCTTCCTT CAAT AGG seq :[91140,91155] :CACACCCTCC GAG AGG seq : [92989,93009] : CCCTACCCTTC ACAAAAT AAG seq :[94052,94068] :CCCCTTCCTT CAAT AGG seq :[97485,97503] :CCCTATCTCT ACACTG ATG seq :[99759,99772] :ACCTTCCTCC G AAG seq :[101976,101994] :CCCTTCCTTC TCCATT AAG seq :[104542,104556] :CACATCCTCC CT AAG seq :[108116,108130] :CCCTATCTCT AC ACG seq :[108381,108400] :CCTCACCTCC GGCACTC AAG seg :[112664,112682] :CCCTTCCTTC TCCATT AAG seq :[116949,116967] :CCCTCACTCT GAGTAG ATG

seq :[118166,118184] :CACCGCCCTT GCCAAC ACG seq :[122748,122761] :CGCTGTCCCC A ACG seq :[122769,122785] :CCTCACCCCC GCAC AAG seq :[124499,124512] :CCCCGCCCGT A ACG

seq :[137843,137858] :CCCATCCTGT CGT ATG seq :[139317,139332] :CCCCACCCCT TTC AGG seq :[141788,141802] :CCCCACCTCG AT ATG seq :[142107,142123] :CCTCGCCCTCC ACA ACG seq :[143895,143912] :CCCCACTCCT CTCTT ATG seq :[144749,144764] :CTCCACCTCT TCG ACG seq :[146166,146184] :CCCTTCCTTC TCCATT AAG seq :[150463,150481] :CCCTCACTCT GAGTAG ATG seq :[151680,151698] :CACCACCCTT GCCAAC ATG seq :[156217,156230] :CGCTGTCCCC A ACG seq :[156238,156254] :CCTCACCCCC GCAC AAG seq :[157969,157982] :CCCCGCCCGT A ACG seq :[172656,172671] :CCCTTTCCTCC CG AGG seq :[173468,173483] :CCCCACCCCT TTC AGG seq :[175945,175959] :CCCCACCTCG AT ATG seq :[176660,176673] :CTCTGTCTCT A AGG seq :[177209,177228] :CCCCGCCCTTC CCCGGG ACG seq :[177282,177300] :ACCGTCCTCC GTCACT ATG seq :[178874,178893] :CCCCTTCCCT GCCAGTA AAG seq :[180982,180998] :CCCTAACTCT GCCA ACG seq :[186813,186829] :CCCCTTCCTT CAAT AGG seq :[190228,190242] :TCCTATCTCT AC ACG seq :[190493,190512] :CCTCACCTCC GGCACTC AAG seq :[190664,190678] :TCCCTCCTCT TT AAG seq :[191401,191415] :CCCGGCCTCT GC AGG seq :[191901,191915] :CGCCTCCTCC GA AAG seq :[192597,192615] :CTCCTCCTCC GTCCAT ATG seq :[197810,197823] :ACCTTCCTCC G AAG seq :[198065,198083] :CCCCACTTCT TTGGCA ATG seq :[200000,200018] :CCCTTCCTTC TCCATT AAG seq :[201083,201096] :ACCCACCTCC T AAG seq :[201099,201114] :CACACCCTCC GAG AGG seq :[204477,204491] :CTCTTCCTCC TC ACG seq :[204774,204793] :CCCCTTCCCT GCCAGTA AAG seg : [206490,206508] : CCCCGTCCACT GCAGT AAG seq :[206886,206902] :CCCTAACTCT GCCA ACG seq :[212206,212224] :CTCCATCTCT CAGTCC ACG seq :[212293,212306] :CCTCATCCTC A AAG seq :[212655,212671] :CCCCTTCCTT CAAT AGG seq :[219344,219358] :CCCTATCTCT AC ACG seq :[219609,219628] :CCTCACCTCC GACAACT ATG

seq :[222123,222142] :CCCCGTACCC TTCCTAG AAG seq :[223882,223900] :CCCTTCCTCC TCCATT AAG

seq :[228156,228174] :CCCTCACTCT GAGTAG ATG seg :[229373,229391] :CACCGCCCTT GCCAAC ATG seg :[233949,233965] :CCTCACCCCC GCAC AAG seg :[257254,257268] :CCCCTGCTCT CA ATG seq :[261219,261236] :CCCCATACTT CCCTC ACG seg :[263044,263061] :CCCCCGCTCC GAGTA ACG seg :[263647,263664] :CCTCACCCCC ACGCC ACG seq :[266171,266184] :CCTCATCTCC C ATG seq :[266572,266590] :CCCCGCTCTT TTGATC AAG seq :[268058,268074] :GCTGCCCTCC GTAG AAG seg :[268119,268136] :CCCCCGCTCC CGTCA ACG seq :[271503,271521] :TCCCGCCTCC CCTCTA ATG seq :[273981,274000] :CCCCGTCTCA TCGGGGG AAG seq :[274202,274221] :CCCTGCCCCCT CCACCG AAG seq :[275600,275614] :CCACTCCTCT GA ATG seq :[276178,276194] :CCCCACCTCT AGAA ATG seq :[276779,276796] :CCCATCATCC TCTTA ATG seg :[279558,279575] :CCCCATACTT CCCTC ACG seq :[281383,281400] :CCCCCGCTCC GAGTA ACG seq :[281986,282003] :CCTCACCCCC ACGCC ACG seq :[284508,284521] :CCTCATCTCC C ATG seq :[284909,284927] :CCCCGCTCTT TTGATC AAG seq :[286395,286411] :GCTGCCCTCC GTAG AAG seq :[286456,286473] :CCCCCGCTCC CGTCA ACG seg :[287840,287858] :CCCCTTCCCT TAACTG AGG seq :[289096,289111] :CCCCACCGCC AGG ATG seq :[299888,299905] :ACCACCCTCC AGAAC ACG seq :[304302,304315] :CCCCCCCC G AAG seq :[305119,305132] :CCACATCCTT G AAG seq :[305596,305613] :CGCTATCCCT TGTGG ATG seq :[309015,309034] :CCCTCCCTGT GCTATCG AAG seq :[315061,315074] :CCCTAACCCT C ATG seq :[316412,316426] :CCCCATCTGC GC AGG seq :[317728,317747] :CCCCGTCCTG AATTGCC ATG seq :[321022,321040] :CCGCACCCCC TTGGGT ATG seq :[322115,322129] :CGCACCCTCC AG ATG seq :[325080,325094] :CACCTCCTCT CA ATG seq :[326850,326865] :CTCCATCCCT TAT AGG seq :[328116,328130] :CACCTCCTCT CA ATG seg :[329943,329959] :TCCTGCCTCC CAAC ATG

----> Checking repeats for 15-LOX-DICE (min: 2) Found 0 matches for pattern 15-LOX-DICE

Pattern = ARE2

Found 0 matches for pattern ARE2

Pattern = TOP

Pattern not found

Pattern = GLUT1

Pattern not found

Pattern = TNF

Pattern not found

Pattern = VIMENTIN

Pattern not found

Pattern = IRES

Pattern not found

Pattern = MSL2-5UTR

Pattern not found

Pattern = MSL2-3UTR

.....

Pattern not found

Pattern = RPMS12_TCE

Pattern not found

Pattern = BRE

Pattern not found

Pattern = ADH_DRE

Found 11 matches in 1 sequences

seq :[12902,12909] :AAGGCTGA seq :[85602,85609] :AAGGCTGA seq :[96691,96698] :AAGGCTGA seq :[107326,107333] :AAGGCTGA seq :[123908,123915] :AAGGCTGA seq :[126344,126351] :AAGGCTGA seq :[159819,159826] :AAGGCTGA seq :[218561,218568] :AAGGCTGA seq :[237528,237535] :AAGGCTGA seq :[319145,319152] :AAGGCTGA seq :[320720,320727] :AAGGCTGA

Pattern = BYDV

Pattern not found

Pattern = Proneural-Box

Pattern not found

Pattern = K-Box

Found 34 matches in 1 sequences

seq :[7894,7901] :ATGTGATA seq :[15701,15708] :GTGTGATA

seq :[18483,18490] :CTGTGATA

seq :[30753,30760] :CTGTGATA seq :[41487,41494] :GTGTGATA seq :[51179,51186] :GTGTGATA seq :[56646,56653] :GTGTGATA

seg :[72471,72478] :GTGTGATA seq :[82204,82211] :GTGTGATA seg :[100710,100717] :ATGTGATA seq :[103920,103927] :GTGTGATA seg :[118548,118555] :ATGTGATA seq :[128291,128298] :GTGTGATA seg :[141538,141545] :GTGTGATA seq :[152062,152069] :ATGTGATA seq :[154474,154481] :CTGTGATT seq :[158999,159006] :CTGTGATA seg :[162618,162625] :GTGTGATA seg :[165402,165409] :CTGTGATA seg :[184835,184842] :CTGTGATA seg :[191776,191783] :TTGTGATA seg :[210685,210692] :CTGTGATA seg :[214120,214127] :ATGTGATA seg :[215167,215174] :GTGTGATA seg :[224794,224801] :ATGTGATA seg :[229755,229762] :ATGTGATA seg :[232168,232175] :CTGTGATT seg :[243103,243110] :CTGTGATA seq :[259844,259851] :TTGTGATA seq :[265300,265307] :CTGTGATA seg :[278192,278199] :TTGTGATA seq :[283639,283646] :CTGTGATA seq :[316371,316378] :CTGTGATC seg :[321231,321238] :CTGTGATT

Pattern = Brd-Box

Found 16 matches in 1 sequences

seq :[11993,11999] :AGCTTTA seq :[20307,20313] :AGCTTTA

seq :[121436,121442] :AGCTTTA seq :[125435,125441] :AGCTTTA seq :[154951,154957] :AGCTTTA seq :[158910,158916] :AGCTTTA seq :[168381,168387] :AGCTTTA seq :[207973,207979] :AGCTTTA seq :[232648,232654] :AGCTTTA seq :[236619,236625] :AGCTTTA

seq :[246732,246738] :AGCTTTA seq :[256155,256161] :AGCTTTA seq :[274599,274605] :AGCTTTA seq :[276022,276028] :AGCTTTA seq :[276853,276859] :AGCTTTA seq :[303252,303258] :AGCTTTA

Pattern = GY-Box

Found 11 matches in 1 sequences

```
seq :[3482,3488] :GTCTTCC
seq :[57635,57641] :GTCTTCC
seq :[64594,64600] :GTCTTCC
seq :[80769,80775] :GTCTTCC
seq :[99110,99116] :GTCTTCC
seq :[99878,99884] :GTCTTCC
seq :[135443,135449] :GTCTTCC
seq :[144077,144083] :GTCTTCC
seq :[221800,221806] :GTCTTCC
seq :[264797,264803] :GTCTTCC
seq :[283136,283142] :GTCTTCC
```

Pattern = Androgen-Receptor

Pattern not found

Pattern = Elastin G3A

Pattern not found

Pattern = Insulin 3'UTR stability

Pattern not found

Pattern = Beta-actin 3'UTR zipcode

Pattern not found

Pattern = Gap-43 stabilization element

Pattern not found

Pattern = Dendritic localization element

Pattern not found

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