UTRdb and UTRsite (RELEASE 2010): a collection of sequences and regulatory motifs of the untranslated regions of eukaryotic mRNAs

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ABSTRACT

The 5’ and 3’ untranslated regions of eukaryotic mRNAs (UTRs) play crucial roles in the post-transcriptional regulation of gene expression through the modulation of nucleo-cytoplasmic mRNA transport, translation efficiency, subcellular localization and message stability. UTRdb is a curated database of 5’ and 3’ untranslated sequences of eukaryotic mRNAs, derived from several sources of primary data. Experimentally validated functional motifs are annotated and also collated as the UTRsite database where more specific information on the functional motifs and cross-links to interacting regulatory protein are provided. In the current update, the UTR entries have been organized in a gene-centric structure to better visualize and retrieve 5’ and 3’ UTR variants generated by alternative initiation and termination of transcription and alternative splicing. Experimentally validated miRNA targets and conserved sequence elements are also annotated. The integration of UTRdb with genomic data has allowed the implementation of an efficient annotation system and a powerful retrieval resource for the selection and extraction of specific UTR subsets. All internet resources implemented for retrieval and functional analysis of 5’ and 3’ untranslated regions of eukaryotic mRNAs are accessible at http://utrdb.ba.itb.cnr.it/.

INTRODUCTION

One of the main challenges of the post-genomic era is the understanding of the mechanisms that control the spatio-temporal regulation of gene expression. The fate of newly synthesized mRNA with respect to its nucleo-cytoplasmic transport, stability, translation efficiency and subcellular localization is determined at the post-transcriptional level. Such regulation is mostly mediated by cis-acting elements located in the 5’ and 3’ untranslated regions of mRNAs (5’UTR and 3’UTR) (1) and miRNAs interacting with their specific targets in 3’UTR (2,3).

Various specific functional sequence elements and miRNA targets have been identified and characterized in mRNA UTRs. These elements usually correspond to short oligonucleotide tracts whose biological activity relies on a combination of their primary sequence and specific secondary structure. These motifs act either as target sites for RNA binding factors or interact directly with the translation machinery. Additionally, miRNA targets, usually located in the 3’UTR, present a very degenerate complementarity with the miRNAs, tolerating several mismatches, gaps and G–U pairings, outside of 6–8 bp continuous seed region at the 5’-end of the miRNA. Additionally, some UTRs may be targeted by complementary natural antisense transcripts masking RNA binding protein or miRNA binding sites (4).

Notably, it is now clear that the same gene may generate several transcript variants, through the use of alternative sites for the initiation and termination of transcription and through alternative splicing. Alternative transcripts can differ both in the coding and in the untranslated regions
(5). Specifically, alternative 5' and 3' UTRs may differentially modulate the gene expression due to the presence of different combinations of functional motifs and miRNA targets.

The availability of a large collection of functionally related sequences—such as UTRs—is invaluable for structural and functional analyses and for a better understanding of the specific role of different variants. To address this issue we have developed a new version of UTRdb, a collection of 5' and 3' UTR sequences derived from eukaryotic mRNAs, where the entries have been organized in a gene-centric structure in order to provide relevant information about splicing variants. Sequences collated in UTRdb were recovered from the National Center for Biotechnology Information (NCBI) RefSeq transcripts (6) using custom software. For human genes, a more comprehensive collection of UTRs is available [derived from the full set of over 300,000 alternative full-length transcripts collected in ASPicDB (7)] generated by a thorough analysis of all available EST/mRNA data.

All UTRdb entries are further annotated for the occurrence of validated regulatory elements, conserved elements and structured RNAs, and miRNA targets (see below). Furthermore, the completeness of 5'UTRs is assessed by the occurrence of mapping CAGE tags (22) (if available) and/or a polyA tail. We have also further expanded UTRsite, a collection of regulatory elements located in 5' and 3' UTRs and whose function and structure have been experimentally determined and published. The UTRsite collection may prove useful in automatic annotation projects of unknown expressed sequences as well as for finding previously undetected signals in known sequences. In the present release, the information for each UTRsite entry has been further enriched including data on functional interacting RNA-binding proteins.

The gene-centric structure of UTRdb facilitates a full integration with all possible gene attributes collected in the NCBI Gene database (8) or other genomic resources such as the UCSC genome browser (9). In this way, the retrieval of specific UTR subsets is possible based on the features associated with each gene, for example a GO term (10), a MIM identifier (11) or a Unigene accession (12).

**GENERATION OF UTRDB AND ITS INTEGRATION WITH OTHER DATABASES**

UTRdb entries are automatically generated through the accurate parsing of the feature table of NCBI RefSeq and ASPicDB transcripts for the UTRref and UTRfull sections of UTRdb database, respectively. ASPicDB contains all possible transcript isoforms for a gene reconstructed by using all available transcript and EST sequences as described in (13). UTR entries are then annotated for the occurrence of tandem and interspersed repetitive elements by using RepeatMasker (v3.2.8, March 2009; A.F.A. Smit, R. Hubley & P. Green RepeatMasker at http://repeatmasker.org), and known regulatory motifs collected in the UTRsite database, as detailed in (14). Each UTRsite entry (Figure 1) is prepared/reviewed/updated by expert scientists (in many cases, those who performed the experimental analysis) by using a suitably developed submission tool (15).

UTRdb entries are also annotated for the occurrence of validated miRNA targets, collected in miRecords (16), a large, high-quality database of experimentally validated miRNA targets resulting from meticulous literature curation. Furthermore, we annotated a set of 3'UTR sequences that have a high likelihood to represent *bona fide* miRNA target recognition sites, as predicted by the HOCTOR tool (17).

For a subset of seven organisms, namely human, mouse, rat, cow, dog, chicken and *Arabidopsis*, for which a suitable genome assembly is available, we also determined the genomic coordinates of UTRs. For such species we were able to clean all redundancies based on the observation of coincident UTRs coordinates, arising from alternative mRNA isoforms.

Additional annotations are specifically provided for genome-linked UTRs. These include: (i) highly conserved sequence blocks from the 17-way PhastCons vertebrate conserved elements (18); (ii) significantly conserved tracts detected by EvoFold (19); and (iii) structural conserved non-coding RNAs detected by RNAz (20).

PhastCons detects evolutionarily conserved elements using a genome-wide multiple alignment based on a phylogenetic hidden Markov model (21). EvoFold is a general comparative genomics method based on phylogenetic stochastic context-free grammars for identifying RNA secondary structures encoded in the human genome and conserved in an eight-way genome-wide alignment of the human, chimpanzee, mouse, rat, dog, chicken, zebrafish and pufferfish genomes (19). RNAz evaluates conserved genomic DNA sequences for signatures of structural conservation of base pairing patterns and exceptional thermodynamic stability. We employed three sets retrieved from (20), with regions conserved with *P*-value >0.9 in human, mouse, rat and dog (Set 1); human, mouse, rat, dog and chicken (Set 2); and human, mouse, rat, dog, chicken and either fugu or zebrafish (Set 3).

To assess the completeness of 5'UTRs in human and mouse we used the mapping data of the CAGE tags indicating the location of the transcription start site (22). The 5'-end of a 5'UTR has been considered as complete when at least five CAGE tags map in a nearby position (a window of 5bp around the mapping position of the 5'-end of the 5'UTR). Analogously, a 3'UTR is considered complete at its 3'-end if a polyA signal and/or a polyA tail is detected in the original transcript sequence.

The UTRdb and UTRSite data have been organized into relational databases using MySQL as the Database Management System. A novel implementation detail in this new release is that several physical databases (containing UTR sequences and annotations from Refseq and ASPicDB transcripts, chromosome coordinates of source transcripts for the seven model organism, taxonomic data, etc.) are used to store all the information on UTRs and their annotations. The new search and retrieval system retrieves and integrates data contained in these different relational databases to give out the requested data on
Figure 1. Sample UTRsite entry. The general information section includes the pattern syntax of the regulatory motif in a format suitable for PatSearch software (23) and the number of hits/kb expected in a sequence collection of randomly generated sequences of the same nucleotide composition of UTRdb. The cross-link to the RFAM database (24), transcripts, genes and RNA binding proteins, if available are also provided, as well as all relevant references (not shown here).
UTRs and related annotations [such as the database from which a UTR was recovered (Refseq or ASPicDB), genomic coordinates and structure, miRNA targets and conserved elements localization, functional elements, etc.]. An exemplar entry of UTRdb is shown in Figure 2.

**UTRdb CONTENT**

UTRdb (UTRef section, release 2010) contains a total of 473,330 5' UTR and 527,323 3' UTR entries, respectively, from 483,605 genes in 79 species (see the Supplementary Data for more information).
A total of 788,370 UTRsite motifs are annotated (317,767 in the 5′UTRs and 470,603 in the 3′UTRs), 20,191 experimentally validated miRNA targets, and 242,773 conserved regions.

For human, the UTRfull section is also available, including UTRs derived from full-length transcripts collected in ASPicDB (7). Overall, UTRfull contains 124,345 and 194,503 5′ and 3′UTRs respectively (3.7 gene) and 3′UTRs (5.18 gene), with 348,412 annotated UTRsite motifs, 649,679 conserved elements and 105,209 experimentally validated miRNA targets.

AVAILABILITY OF UTRdb

UTRdb and UTRsite are accessible through a newly developed retrieval system where simple and advanced search forms are available. UTRs can be retrieved by several accession IDs, GO terms and MIM identifiers. Additionally, the advanced form permits a further refinement of the UT subset to be retrieved using several criteria including the number of CAGE mapping tags (for 5′UTRs), the length of the UTR, the number of spanning exons, the occurrence of UTRsite motifs, conserved elements and miRNA targets.

A download facility for selected UTR entries in FASTA format is also available.

Further online utilities are UTRscan and UTRblast. The UTRscan feature allows the enquirer to search user-submitted sequences for any of the motifs collected in UTRsite. The UTRblast utility allows database searches against any of the UTRdb sections.

UTRdb, UTRsite and other related resources are publicly available at http://utrdb.ba.ibt.cnr.it/.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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