# UTRdb 2.0: a comprehensive, expert curated catalog of eukaryotic mRNAs untranslated regions

Claudio Lo Giudice<sup>1</sup>, Federico Zambelli<sup>2,3</sup>, Matteo Chiara<sup>(b2,3)</sup>, Giulio Pavesi<sup>2,3</sup>, Marco Antonio Tangaro<sup>3</sup>, Ernesto Picardi <sup>(b1,3)</sup> and Graziano Pesole <sup>(b1,3,\*</sup>

<sup>1</sup>Department of Biosciences, Biotechnology and Environment, University of Bari A. Moro, 70126 Bari, Italy, <sup>2</sup>Department of Biosciences, University of Milan, 20133 Milan, Italy and <sup>3</sup>Institute of Biomembranes, Bioenergetics and Molecular Biotechnology, Consiglio Nazionale delle Ricerche, 70126 Bari, Italy

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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. Since 1996, we have developed and maintained UTRdb, a specialized database of UTR sequences. Here we present UTRdb 2.0, a major update of UTRdb featuring an extensive collection of eukaryotic 5' and 3' UTR sequences, including over 26 million entries from over 6 million genes and 573 species, enriched with a curated set of functional annotations. Annotations include CAGE tags and polyA signals to label the completeness of 5' and 3'UTRs, respectively. In addition, uORFs and IRES are annotated in 5'UTRs as well as experimentally validated miRNA targets in 3'UTRs. Further annotations include evolutionarily conserved blocks, Rfam motifs, ADAR-mediated RNA editing events, and m6A modifications. A web interface allowing a flexible selection and retrieval of specific subsets of UTRs, selected according to a combination of criteria, has been implemented which also provides comprehensive download facilities. UTRdb 2.0 is accessible at http://utrdb.cloud.ba.infn.it/utrdb/

# INTRODUCTION

Even after over two decades since the completion of the first draft assembly of the human genome, and the compilation of a complete catalog of all the human genes, we are still far from a detailed appreciation of the molecular mechanisms that regulate the spatio-temporal expression of protein coding genes and their translation into their 'final' gene products. While technological advances such as the massive, parallel sequencing of RNAs have enabled the quantitative and precise measure of RNA expression levels in a wide range of human cells and tissues (1), several independent lines of evidence confirmed that mRNAs and proteins levels are only partly correlated, and in general mRNA levels alone are not predictive of the concentration or activity of proteins in vivo (2-4). Several biological mechanisms that regulate post-transcriptional mRNAs stability and translation are at the base of this observation. For example, the nucleo-cytoplasmic transport of the newly synthesized RNAs (5), their stability (6), translation efficiency (7) and subcellular localization (5) are regulated at the posttranscriptional level. In protein-coding mRNAs, such regulation is mainly mediated by cis-acting elements located in the 5' and 3' untranslated regions (5'UTRs, 3'UTRs) also through the interaction with complementary RNAs (e.g. miRNAs) or enzymatically driven chemical modifications (e.g. m6A) or editing processes (e.g. A to I). Evolutionarily free from the constraint of encoding proteins, UTRs may modulate gene expression through various functional elements which have been characterized and experimentally validated in numerous mRNAs (8,9). These elements may correspond to short oligonucleotide tracts whose biological activity relies on both their sequence and secondary structures (10). For example, some specific sequence motifs function as target sites for RNA binding proteins, modification or editing enzymes or interact directly with the translational machinery.

The 5'UTRs may contain structural motifs and upstream Open Reading Frames (uORFs) that regulate translation efficiency or promote cap-independent translation in specific physiological conditions (11). uORFs, are found in nearly half of human 5'UTR mRNA transcripts (12), and can suppress translation through various mechanisms, including translation reinitiation, ribosome leaky scanning or the functional activity of the uORF-encoded peptide. 3'UTRs may contain, as well, sequence and structural motifs modulating mRNA stability, translational efficiency and subcellular localization in several ways. In particular, their degen-

\*To whom correspondence should be addressed. Tel: +39 0805443588; Email: graziano.pesole@uniba.it

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erate complementary interaction with miRNAs has been associated with translation downregulation (13).

Additionally, alternative cleavage and polyadenylation (APA), a widespread mechanism to generate mRNA isoforms with alternative 3' UTRs, has been implicated in the modulation of protein abundance, as well as in the regulation of mRNA localization and the spatial organization of protein synthesis (14). Finally, all the regulatory mechanisms described above can be further modulated by specific RNA editing or chemical modifications.

Although UTRs comprise only <1% of the human genome, about 3.7% of all currently known human genetic variants are located in UTRs, suggesting that they might represent a hotspot of genetic variation in human protein coding genes, a consideration that further emphasizes the importance of performing/unraveling accurate functional annotation of UTR elements (15). Indeed, GWAS investigations identified several genetic variants in UTR regions. These variants, potentially associated with complex disease risks, may affect gene expression, thus playing a critical functional role, but are usually more tricky to functionally assess with respect to variants affecting coding sequences. Moreover, 599 variants (0.5% of the total number) in the UTRs of protein coding genes are known to be implicated in the pathogenesis of human disorders according to the Clin-Var database (16).

Therefore, the availability of an extensive collection of highly curated annotations of functional elements in human UTRs could represent an invaluable resource for a better understanding of gene expression regulation and its role in health and disease conditions. Importantly, similar considerations could be applied to several model and non-model organisms, for which a reference genome assembly and annotation is currently available. At present >500 complete genomes are available through the dedicated resources curated by EMBL/Ensembl (17). Accurate and comprehensive annotations of UTR regions could provide a useful resource for advancing functional genomics also in these species, and for the execution of complex and detailed comparative genomic analyses.

To address these issues, we have developed a new version of UTRdb (18), a collection of 5' and 3'UTR sequences derived from eukaryotic mRNA collected in the Ensembl (17) and Ensembl Genomes (19) databases with over 500 different species represented.

UTRdb is based on a gene and transcript-centric annotation model, which facilitates the full integration with other specialized databases and resources, and, at the same time, permits the compilation of detailed reports providing comprehensive information on alternative 5' and 3'UTRs. UTRdb entries boast a comprehensive collection of annotations, including: regulatory elements collected in Rfam (20), conserved elements in PhastCons (21), ADAR-mediated RNA editing (22) and m6A modifications (23). In addition, upstream open reading frames, complemented with mass spectrometry proteomic data (when available) (24), and Internal Ribosome entry sites (25) are provided for 5' UTRs, while experimentally validated miRNA targets are reported for 3'UTRs (26,27). Finally, to provide information concerning the completeness of 5' and 3' UTR sequences and/or alternative TSS/TTS, UTRdb integrates a large collection of CAGE experiments (28) and provides annotations of experimentally validated poly-A signals (29).

We believe that by providing a comprehensive and annotation-rich collection of annotations UTRdb may be regarded as a highly useful resource for investigations on the regulatory roles of UTRs, their role in the maintenance of cellular homeostasis and in the onset and progression of human disorders, including cancer (30). Finally, UTRdb, providing useful information for the functional assessment of UTRs, may represent a valuable resource for the development of mRNA vaccine technology, whose recent impressive breakthrough greatly supported the fight against the COVID-19 pandemic (31).

# DATA COLLECTION AND PROCESSING

#### **Data collection**

Gene annotations in GTF format and the corresponding FASTA files for 615 organisms were downloaded from Ensembl, including: 313 organisms from Ensembl rel 107 (http://ftp.ensembl.org/pub/release-107/), 125 from Ensembl plants rel 54 (http://ftp.ensemblgenomes.org/pub/ plants/release-54/) and 177 from Ensembl metazoans (http: //ftp.ensemblgenomes.org/pub/metazoa/release-54/) rel 54 (17). All GTF files were preliminarily evaluated and organisms lacking UTR annotations were discarded. In particular, a total of 42 species (19 from Ensembl rel 107, 13 from Ensembl plants and 10 from Ensembl metazoans) were not included in UTRdb for this reason, thus reducing the total number of organisms incorporated in our database to 573. Lists of orthologous genes, annotations of repeated elements, low complexity DNA regions and small genetic variants were downloaded from Ensembl through their Representational State Transfer (REST) API (32) (https://rest. ensembl.org/) by means of custom Python scripts. Collections of human genetic variants significantly associated with genotypic traits or disorders according to a compendium of GWAS were downloaded from the GWAS Catalog (33) (https://www.ebi.ac.uk/gwas/).

Evolutionary conserved elements identified by Phast-Cons were obtained from the UCSC Genome Browser database (https://hgdownload.soe.ucsc.edu/downloads. html) (34).

When multiple PhastCons annotations for an organism, were available, the one incorporating the largest number of alignments/species was selected (e.g. 'phast-ConsElements135way.txt.gz' instead of 'phastConsElements26way.txt.gz' in *Caenorhabditis elegans* entries).

Annotations of miRNA targets for the Homo sapiens GRCh38 and Mus musculus GRCm39 reference genome assemblies were downloaded from the Ensembl genome browser, through their public mysql databases, using the following commands: 'mysql -h ensembldb.ensembl.org -u anonymous -P 5306 homo\_sapiens\_funcgen\_107\_38 'select \* from mirna\_target\_feature' and 'mysql -e -h ensembldb.ensembl.org -u anonymous -P 5306 mus\_musculus\_funcgen\_107\_39 -е 'select from mirna\_target\_feature', respectively. A custom script was applied to retain only target sites supported by experimental evidence according to Tarbase v8 (https://dianalab. e-ce.uth.gr/html/diana/web/index.php?r=tarbasev8) (27).



**Figure 1.** UTRdb data processing and database organization. A total of 26 098 657 UTR sequences were extracted from Ensembl and Ensembl Genomes included in UTRdb. Functional annotations were retrieved from their respective repositories (e.g. D1, D2, D3, ..., Dn) and integrated by means of custom Python scripts. All data was organized in MySQL tables. The web-interface was developed using Bootstrap and HTML5.

Alternative polyadenylation sites (in BED format) were obtained from APADB (http://tools.genxpro.net:9000/ apadb/download/) version v2 (29) and the UCSC Liftover tool (35) (available at https://genome-store.ucsc.edu/) was used to convert genomic coordinates to different reference genome assemblies when required (e.g. hg19 to hg38 for the human genome). Additionally, miRNA target sites predicted by TargetScan (32) and miRanda (36) and included in APADB were incorporated in UTRdb as well. Annotations of IRES (Internal Ribosome Entry Site) motifs were collected from IRESite (http://iresite.org/ IRESite\_web.php?page=browse\_cellular\_transcripts) (25), a database of experimentally validated IRES structures.

Epitranscriptome modifications, including RNA editing events due to the deamination of the adenosine (A) in inosine (I) and m6A sites were obtained from the specialized databases REDIportal (http://srv00.recas.ba.infn. it/atlas/download.html) and RMVar (https://rmvar.renlab. org/download.html) (22,23), respectively. Table 1. Distribution of UTR annotations collected in UTRdb (full values are reported for the five most represented organisms in each Ensembl group)

Species	5'UTR		3'UTR	
	N	Average L	N	Average L
Vertebrates				
Homo_sapiens.GRCh38.107	93 457	231	87 932	1241
Mus_musculus.GRCm39.107	56 214	222	52 526	1135
Mus_spretus.SPRET_EiJ_v1.107	46 627	194	44 162	975
Danio_rerio.GRCz11.107	33 326	226	30 928	953
Sus_scrofa.Sscrofa11.1.107	36 098	338	36 757	1434
Metazoans				
Crassostrea_gigas.GCA902806645v1.54	72 508	211	72 800	613
Ascaris_suum.ASM18702v3.54	52 374	688	55 488	1618
Pomacea_canaliculata_gca003073045v1.GCA003073045v1.54	39 028	425	39 051	1686
Limulus_polyphemus_gca000517525v1.Limulus_polyphemus_2.1.2.54	34 796	240	33 600	1281
Drosophila_melanogaster.BDGP6.32.54	30 091	316	30 211	600
Plants				
Triticum_dicoccoides.WEWSeq_v.1.0.54	214 610	350	240 626	476
Aegilops_tauschii.Aet_v4.0.54	201 304	407	222 362	553
Hordeum_vulgare.IBSC_v2.42	184 618	365	203 233	549
Zea_mays.B73_RefGen_v4.42	112 659	534	113 636	729
Triticum_aestivum.IWGSC.54	88 479	192	92 320	345

Table 2. Distribution of UTR functional annotations available in UTRdb 2.0

Annotation	5'L	JTR	3′UTR	TR
	Human	Others	Human	Others
uORFs	78 276	10 677 649	-	-
IRES	239	102	-	-
CAGEs	2885	77 673	-	-
Alternative PolyA sites	-	-	124 015	56 180
miRNAs targets	51 428	7972	741 407	205 379
Conserved blocks	431 218	1 048 459	2 095 576	2 731 101
RNA editing	15 293	253	336 959	6998
m6A modifications	8050	6517	28	34 445
Variants	5 890 887	8 789 358	22 821 251	21 051 930
Rfam motifs	159	24 922	528	54 478

Known RNA functional secondary structure elements and other sequence motifs associated with UTRs were annotated according to Rfam (20) (https://rfam.xfam.org/), while mass spectrometrybased proteomic evidences of potential upstream ORFs translation were obtained from PRIDE (24) (https://www.ebi.ac.uk/pride/). Cap Analysis of Gene Expression (CAGE) tags were downloaded from fantom5 (28) (https://fantom.gsc.riken.jp/5/) in BED format for the following species: *Homo sapiens* (GRCh38), *Mus musculus* (GRCm39), *Canis lupus* (Dog10K\_Boxer\_Tasha), *Gallus gallus* (gca000002315v5.GRCg6a), *Macaca mulatta* (Mmul\_10) and *Rattus norvegicus* (mRatBN7.2).

#### Construction of UTRdb relational database tables

A compendium of highly curated and accurate resources for functional annotation of UTRs were collected (see Data Collection and Processing section) and automatically processed by custom software in order to generate *ad hoc* tables containing UTRdb entries and attributes (Figure 1). The sequence, and corresponding annotation of gene models, in the form of a GTF file, were retrieved for every eukaryotic genome reference assembly available from the Ensembl genome browser. *Ad-hoc* Python scripts were applied to retrieve genomic coordinates of UTRs and extract (by means of the pysam (37) Python module) the corresponding sequences.

Functional annotations, including polyA signals, RNA editing sites, SNPs, miRNA binding sites, PhastCons evolutionarily conserved elements and many others (see see Data Collection and Processing section), were retrieved from their respective repositories (as detailed above), formatted according to the requirements of UTRdb and integrated in the database, again by applying custom Python scripts (Figure 1).

Annotations of uORFs were obtained by performing in silico translation of 5' UTR sequences, ORFs formed by an AUGs without an in-frame downstream stop codon (38) were also included.

# DATABASE CONTENT AND WEB INTERFACE

#### UTRdb content and structure

The knowledge base incorporated in UTRdb 2.0 represents a more than 25-fold increase in terms of the number of UTRs and 7-fold increase in terms of the number of species represented in the database, compared to previous release (15). UTRdb 2.0 includes functional annotation for a total of 26 098 657 UTRs—12 908 478 5'UTRs (with a mean



Figure 2. The advanced search front-end of UTRdb and an example of the tabular output as well as a sample entry.

length of 300 nt) and 13 190 179 3'UTRs (with a mean length of 753 nt) (Table 1)-from 6 688 161 genes in 573 organisms. Approximately, 0.7% of the UTRs in the database are from H. sapiens and collectively 8.8% are associated with model organisms. Importantly in UTRdb 2.0 the plant repertoire of UTRs derived from plants is greatly expanded compared to previous versions; a total of 8 452 197 UTRs from 112 plant species is included in our database, while only six species were available in previous releases. An average of 22 527 5'UTRs and 23 019 3'UTRs are available for every species incorporated in the database, however the number and size of UTRs associated with different species vary greatly; probably this is due to a series of factors including: evolutionary constraints/history, the quality of the available genome annotations, the total number of genes in the genome (Table 1).

UTRdb provides a rich set of functional annotations for UTRs. These include, but are not limited to: 6 306 354 evolutionarily conserved elements, 359 503 A-to-I RNA editing events, 49 040 m6A modifications and 58 553 426 nucleotide variants. Additionally, 10 755 925 candidate uORFs, 341 IRESs and 80 558 CAGEs are annotated in 5'UTR entries, while 180 195 alternative polyAdenylation sites are associated with 3'UTR entries (Table 2). UTRdb 2.0 has been structured as a relational database, built on the MySQL framework and easily accessible through an *ad hoc* webinterface developed in Bootstrap and HTML5.

# Web interface

Server-side operations including MySQL querying and retrieval are handled by Python scripts with the support of dedicated modules (e.g. pyMySQL, python CGI). UTR sequences and their associated annotations are displayed through dynamic and sortable tables generated by means of the DataTables jQuery module.

The overall architecture of the database has been completely redesigned with respect to previous releases. Indeed, annotation categories (i.e. miRNA targets, Rfam motifs, RNA editing sites and so on) are now organized in multiple databases, each including a set of organism-specific tables (generally linked by the transcript id as the unique primary key). This organization increases the scalability of the database allowing straightforward updates at organism level (i.e. when new genomic assemblies or annotations are released) without the need to shut down the entire service or reindex the full database.

Queries to UTRdb 2.0 can be performed using either a basic or an advanced interface. The former has been designed for quick queries, and supports limited metadata fields/filters: the type of UTR (5', 3', or both), the organism and a gene identifier, which can be specified in the form of a gene symbol, Ensembl gene or transcript IDs. To facilitate the user experience, an autocomplete function for gene symbols and gene and transcript ids has been designed to allow more complex queries enabling different filtering schemes through several parameters, which can be used individually or in combination. For example, users can filter entries according to UTRs size and/or the number of exons they span, the presence of Rfam motifs, miRNA target sites or epitranscriptomics modifications (A-to-I edit-

ing or m6A). In case of 5'UTRs, users could additionally filter the results for the presence of upstream ORFs (uORFs), Internal Ribosome Entry sites (IREs) or CAGE tags. For 3'UTR queries, instead, users can extract entries containing annotations of alternative polyadenylation sites. Since each UTRdb entry is identified by a unique identifier in the format {UTR\_type\_genome\_assembly\_transcript\_id}, the database can also be queried by entering the unique id of the entry. Annotations of low quality for example UTRs of <10 nt in size, are flagged by a warning at the beginning of the entry. An overview of the search interfaces, as well as a sample entry, are displayed in Figure 2. Additionally, UTRdb 2.0 features explicative, interactive graphical representations of the annotations. The various functional annotations associated with any UTR can be downloaded in the form of standard JSON tables, while all UTR and related functional annotations for an organism can be retrieved in bulk through a dedicated download web page.

# **UTRdb 2.0 UPDATE AND FUTURE PLANS**

One of the key improvements of UTRdb 2.0 has been the redesign and re-implementation of the service's backend that, while less visible for users than the complete frontend overhaul, will provide solid ground for the future of UTRdb in terms of easier maintenance, more straightforward integration of new or updated annotations as they become available, and development of novel functionalities.

Considering the biological relevance of UTRs, we plan to regularly update UTRdb, in correspondence with the new releases of Ensembl, providing researchers an accurate, sustainable and accessible resource through the inclusion of annotations from novel organisms, the interoperability with other resources through crosslinks and a dedicated API. Additionally, we will work to enhance the quality of UTR annotations as new data (e.g. from long reads technologies such as Oxford Nanopore and PacBio) will become available and allow the referencing of data in both a locationindependent and resource-dependent manner through resolvable identifiers (URLs) (by means of the identifiers.org registry) and constantly improving the compliance with FAIR principles (Findability, Accessibility, Interoperability and Reusability).

#### DATA AVAILABILITY

All the Python scripts to extract UTR sequences from Ensembl Fasta files and retrieve their associated annotations from multiple databases are available on our dedicated github repository at https://github.com/BioinfoUNIBA/ UTRdb.

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