**Introduction**

Post-transcriptional regulation (PTR) of gene expression is now recognized as a major determinant of cell phenotypes. The recent availability of methods to map protein-RNA interactions in entire transcriptomes such as RIP, CLIP and their variants, together with global polysomal and ribosome profiling techniques, are driving the exponential accumulation of vast amounts of data on mRNA contacts in cells, and of corresponding predictions of PTR events. However, this exceptional quantity of information cannot be exploited at its best to reconstruct potential PTR networks, as it still lies scattered throughout several databases and in isolated reports of single interactions. To address this issue, we developed the second and vastly enhanced version of the Atlas of UTR Regulatory Activity (AURA 2), a meta-database centered on mapping interaction of trans-factors with human and mouse UTRs. AURA 2 includes experimentally demonstrated binding sites for RBPs, ncRNAs, thousands of cis-elements, variations, RNA epigenetics data and more. Its user-friendly interface offers various data-mining features including co-regulation search, network generation and regulatory enrichment testing. Gene expression profiles for many tissues and cell lines can be also combined with these analyses to display only the interactions possible in the system under study. AURA 2 aims at becoming a valuable toolbox for PTR studies and at tracing the road for how PTR network-building tools should be designed. AURA 2 is available at http://aura.science.unitn.it.
of a single gene or a collection of functionally related genes. This wealth of data is complemented by a set of data-mining tools providing novel views on PTR of either individual or groups of genes. Particular attention was given to the interface, which is designed to be easy to use but flexible enough to accommodate advanced usage and complex queries. AURA 2 is intended to be a one-stop broad resource for PTR, targeted at both command-line averse biologists and bioinformaticians.

**Results**

The strength of a resource such as AURA 2 lies in its ability to be constantly updated and include, in a timely manner, as many as possible data types and records concerning its area of focus. On the other hand, clever exploitation of such a data collection must be empowered by various adequate retrieval and mining features, eventually allowing users to effectively reach the data they need. We will now describe what can be found in AURA 2, both in terms of data types and quantities therein contained and in terms of website features such as search modes, analytical tools and data download options. A graphical summary of AURA 2 capabilities is displayed by Figure 1.

**Data types and figures**

AURA 2 currently includes data about two model species, namely *Homo sapiens* and *Mus musculus*. However, its structure allows for any number of model species to be inserted and other, such as *Danio rerio*, *Caenorhabditis elegans* and *Saccharomyces cerevisiae*, will follow later on. Both included species share a layer of basic annotation that goes from a gene, transcript and UTR model obtained from the UCSC Genome annotation\(^{15}\) to phylogenetic conservation, secondary structure folding and Gene Ontology term associations. Human data also includes two different data sets of transcript half-life measurements, obtained by microarray\(^{16}\) and high-throughput sequencing\(^{17}\) respectively. Finally, uncoupling between translatome and transcriptome gene expression variations, suggesting the occurrence of post-transcriptional regulation, is quantified and displayed for both human and mouse genes based on a meta-analysis of data sets containing such profiles.\(^3\) On top of these annotations, a number of different PTR databases and data sets of various kinds were collected and integrated. We decided to consider only experimentally derived data, thus excluding predictions. This was done with the notable exception of AU-Rich Elements, obtained from AREsite,\(^{18}\) and of the secondary structure of mRNAs, because
In particular, five search modes are available, enriched by various filters and parameters: along with the classic gene search, providing an overview of all PTR events mediated by the UTRs of a single gene whose name corresponds to the query term, the user can also search for a trans-factor (e.g., an RBP or a ncRNA) in order to obtain all the UTRs which share a regulatory event by this factor; these UTRs can be grouped by Gene Ontology terms in order to provide an immediate clue to functions and processes controlled by this trans-factor through its UTR targets. The other three search methods are named co-regulation, sequence and batch searches. Co-regulation search allows users to retrieve all UTRs controlled at the PTR level by two or more trans-factors: this feature can be extremely useful, for instance, to identify different factors controlling a common process or to detect competing regulation phenomena. Sequence search performs a BLAST query against 5’ and 3’ UTRs of the selected organism, permitting to display and browse all matching UTRs and to download results. The last search mode is the batch search, which includes two analytical tools and is offered in three flavors. Users can input a list of genes in which they are interested: the first option consists in selecting and browsing all UTRs of these genes together; secondarily they can build a PTR network in which edges are regulatory relationships (factor F is controlling an mRNA M and so on) and which outlines all factors regulating and possibly shared by the genes in the input list. The last option consists of performing a Fisher’s exact test in order to understand which trans-factors or cis-element categories are significantly enriched in the provided gene set. When combined, these options will provide both a graphical and intuitive way of producing hypotheses on the regulation of thousands of human and mouse mRNAs and a sound statistics indicating which factors may be interesting to pursue for subsequent investigations.

Eventually, interactions displayed by the trans-factor, co-regulation, network generation and regulatory enrichment modes can be filtered for the expression of involved genes in a set of 45 different tissues and 3 cell lines (see Materials and Methods). This powerful feature allows the user to investigate only the potential interactions which are relevant to the system under study. 

The UTR Browser

UTRs selected for detailed visualization by any of the search modes described above are then displayed in individual page segments (UTR Browser) including a basic annotation section (showing, for instance, genomic coordinates, length and overall conservation) and a dedicated genome browser. This component, based on JBrowse41, allows the user to zoom and move throughout the UTR sequence, add or remove tracks and rearrange their order to best suit the user’s focus. Along with sequence and conservation tracks, regulatory sites of RBPs, microRNAs, variations and each category of cis-elements are each laid out on independent tracks, for the maximum flexibility in choosing which data to visualize and with which arrangement. Further features accessible through the UTR Browser are gene expression measurements for the gene and the trans-factor at hand, and annotated UTR secondary structure visualization. The former feature takes advantage of several hundred gene expression profiles performed on more than 20 different tissues and stored at
the Genotype-Tissue Expression project (GTEx)\textsuperscript{32}: this database is queried on-the-fly for the displayed gene and for the related trans-factor (only when the trans-factor search is used); relevant gene expression measurements are displayed in a pop-up window. The latter feature provides the UTR secondary structure folding (drawn by the VARNA plugin\textsuperscript{33}), annotated with a green gradient for phylogenetic conservation and three different colors for SNPs, RBPs and ncRNA binding sites: users can thus readily understand whether a regulatory event relies on or can be influenced by a particular predicted secondary structure element.

Data download
All of the information displayed in the various UTR Browser parts is also downloadable through the UTR card feature, which consists of a textual, human-readable (but also machine-parseable thanks to its structured layout) collection of all the data regarding the investigated UTR. The card feature is also available for trans-factors such as RBPs and microRNAs and includes, along with basic annotation and descriptions, all the binding sites for the factor and related target UTR data. The complete AURA 2 data set can also be retrieved by downloading the database dump (to replicate the complete database on a local machine), by downloading a reduced version of the database, called AURAlight, which includes only data concerning cis-elements and trans-factors interactions with UTRs in tab-delimited format, or by accessing the AURA 2 Mart. This latter feature is an instance of the federated database system BioMart\textsuperscript{34} containing all of AURA 2 data. BioMart provides an unique query interface shared by all data from the Mart offered by AURA 2 without additional learning efforts. A comparison of AURA 2 features, data content and mining tools with respect to its previous version and other available resources is in Supplementary Materials (Supplementary Text, Supplementary Table 1–2). Furthermore, a user guide to AURA 2 features is also available in Supplementary Materials (Supplementary Text).

Discussion
We think that AURA 2 is a considerable step forward for PTR data integration and PTR network-building capabilities. Nevertheless, interaction data are still available only for a limited number of RBPs and non-coding RNAs: this aspect represents a limit on the possibility to accurately reconstitute regulatory networks. While this issue will be progressively solved with the predictable increase of available data sets (from CLIP and CLIP variants), it must as of now be taken into account when building such networks. It is to be stressed that all data sets present in AURA 2 were included ‘as-is’, without reprocessing data: this choice has been made on the basis of the heterogeneity of the techniques employed to obtain such data. Even if we decided to reprocess all these data sets, we would have needed to employ several analysis protocols, with their inherent biases and limitations. Furthermore, we decided not to exclude any data set on the basis of quality judgments: references to the original publication are available for every and each binding site throughout the database, allowing users to further investigate and eventually decide on the value of the data they wish to exploit. A quality measure for the high-throughput-derived data sets (RIP, CLIPs, etc.) considering properties such as presence of replicates and filtering stringency will likely be subject of future developments in AURA 2. Evidence has been accumulating in the last years to suggest that interactions, both in terms of cooperation and competition, do happen between RBPs and microRNAs.\textsuperscript{35,36} A small but increasing number of publications are also now indicating that lncRNAs may be involved as well.\textsuperscript{37} We think that these phenomena will likely gain more importance as they are elucidated and characterized, emphasizing the opportunity of exploiting them to discover biological implications and to promote better network-building approaches in PTR. We plan to integrate this kind of data in AURA 2 and possibly develop some dedicated search and visualization modes to allow the user for seamless and effective study of this interaction processes. Another emerging topic in PTR is RNA epigenetics. As can be seen in AURA 2, genome-wide transcriptome methylation and editing data sets are beginning to appear and the next years will most likely start to reveal the impact of these regulatory processes and the potential interplay with other PTR actors. Integrating from the start these data into the PTR networks we are building will help to better understand the functional effects of RNA epigenetics and improve its characterization. We also aim at expanding AURA 2 by including annotation and data for other model organisms which are relevant in RNA biology, such as \textit{C. elegans}, \textit{S. cerevisiae}, \textit{D. rerio} and \textit{D. melanogaster}. This will require collaboration with experts of these species to achieve effective and useful data integration. On a broader perspective, being able to involve the PTR research community in maintaining AURA 2 updated and in determining the directions to be taken in its development would be of great value to make it as effective and useful as possible: we indeed are available for establishing new and thriving collaborations toward this aim.

Materials and Methods
Website implementation
The database was designed and implemented on a MySQL Community Server 5.5 (Oracle, Santa Clara, CA, USA). The website is implemented with Python programming language\textsuperscript{38} and the Django web framework,\textsuperscript{39} exploiting JQueryUI graphical components.\textsuperscript{40} The UTR browser employs jBrowse\textsuperscript{33} as genome browser to display sequence, and data tracks: UTRs secondary structures are visualized through the VARNA plugin,\textsuperscript{33} while CytoscapeWeb\textsuperscript{41} is used to display PTR networks. Sequence search is performed by means of a local installation of BLAST\textsuperscript{50} on a custom database containing 5’ and 3’ UTR sequences. The BioMart site was implemented on BioMart 0.8.34 by producing a copy of the AURA database containing all the data and employing a database schema adequate for BioMart.

Basic annotation data retrieval
Gene symbols, synonyms and annotations were retrieved from HGNC\textsuperscript{42} and MGI\textsuperscript{43} for human and mouse respectively.
Transcripts and UTRs models, UTR secondary structure folding and phylogenetic data were retrieved from various tracks of the UCSC Genome Browser for both species. Displayed half-life measurements were retrieved from two genome-wide measurements, obtained through microarray and high throughput sequencing respectively. Uncoupling data was obtained by computing differences in expression variations between translatome and transcriptome profiles in 20 human and mouse data sets, as presented in. Links to Human Protein Atlas and miRBase for additional trans-factor information, and to PAZAR for transcriptional regulation data, were composed by plugging-in the trans-factor name in the URL structure of these two websites. Genes associations with Gene Ontology terms were retrieved from the Gene Ontology website for both organisms.

**Databases integration**

The latest version of integrated databases (listed in Supplementary Table 2) were downloaded in full from their respective websites. When needed, genomic coordinates were converted to the most recent genome assembly version (hg19 for human and mm10 for mouse) and mapped to UTRs. One database, GTEx was integrated by retrieving expression plot images when requested for a given gene and/or trans-factor.

**High-throughput data sets collection**

A literature search was performed to retrieve high-throughput data sets for RBPs and ncRNAs, deriving from techniques such as the many variations of RIP and CLIP. Given the wide number of techniques covered by these data sets (thus inherently containing a certain degree of heterogeneity), we decided to insert the data as processed by the authors of each publication: details for each experiment can be accessed through links to the original publications, available on the website. A parallel search focused on identifying studies profiling RNA modifications such as RNA methylation m5C and m6A, alternative polyadenylation and translation initiation sites at a genome-wide level. Collected and inserted data sets are listed in Supplementary Table 3. As for the integrated databases, listed in Supplementary Table 1, genomic coordinates were converted to the most recent genome assembly version and mapped to UTRs.

**Low-throughput data set building**

An extensive, manually curated, RBP-by-RBP literature search was performed in order to collect and extract all RBP-mRNA interactions deriving from mechanistic or single-gene experiments, in papers published from 1994 to 2013. Data obtained from assays producing various degrees of information (protein binding to the mRNA, protein binding to a specific UTR or binding to a defined location in a given UTR) was included; the associated uncertainty of precise binding location was indicated in the UTR browser by graphically coding it with colors and drawing patterns.

**Gene expression data set building**

HTS-derived gene expression profiles for 45 tissues were retrieved from GTEx, and three further profiles for HeLa, HEK293 and MCF7 cell lines were retrieved from published RNA-seq studies. RPKMs were computed for these last three data sets by means of Cufflinks. Expressed genes lists were obtained by filtering these profiles for genes having a RPKM value greater than 0.1.

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

**Acknowledgments**

We would like to thank Luca Andreotti for reviewing the website graphics and providing helpful advices toward its improvement.

**Supplementary Material**

Supplementary material may be found here: www.landesbioscience.com/journals/translation/article/27738/
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