RSAT 2018: regulatory sequence analysis tools 20th anniversary

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ABSTRACT

RSAT (Regulatory Sequence Analysis Tools) is a suite of modular tools for the detection and the analysis of cis-regulatory elements in genome sequences. Its main applications are (i) motif discovery, including from genome-wide datasets like ChIP-seq/ATAC-seq, (ii) motif scanning, (iii) motif analysis (quality assessment, comparisons and clustering), (iv) analysis of regulatory variations, (v) comparative genomics. Six public servers jointly support 10 000 genomes from all kingdoms. Six novel or refactored programs have been added since the 2015 NAR Web Software Issue, including updated programs to analyse regulatory variants (retrieve-variation-seq, variation-scan, convert-variations), along with tools to extract sequences from a list of coordinates (retrieve-seqbed), to select motifs from motif collections (retrieve-matrix), and to extract orthologs based on Ensembl Compara (get-orthologs-compara). Three use cases illustrate the integration of new and refactored tools to the suite. This Anniversary update gives a 20-year perspective on the software suite. RSAT is well-documented and available through Web sites, SOAP/WSDL (Simple Object Access Protocol/Web Services Description Language) web services, virtual machines and stand-alone programs at http://www.rsat.eu/.

INTRODUCTION

Initiated in 1998 (1,2), the Regulatory Sequences Analysis Tools (RSAT) project aims at deploying software tools to detect cis-regulatory elements in genomic sequences, via a Web interface. RSAT functionalities include de novo motif discovery, analyses of motif quality, motif comparisons and clustering, motif scanning to predict transcription factor (TF) binding sites (TFBSs), detection and analysis of regulatory variants, and comparative genomics to discover motifs based on cross-species conservation (Figure 1). Over the last 20 years, the RSAT team has maintained uninterrupted service, while extending developments prompted by the advances in the field of regulatory genomics (Supplementary Figure S1). This Anniversary article gives a 20-year perspective on the software suite, describes its main functionalities, focusing on the novelties since the previous NAR Web server issues (3–6), and presents the various access and training modalities.

RSAT OVER THE LAST 20 YEARS

From yeast-tools to RSAT

The development of RSAT (initially named yeast-tools) (1,2) was prompted by the sequencing of the yeast genome
The motivation was to ease the extraction of non-coding sequences upstream of genes, and the prediction of TF binding sites (TFBSs). The programs for \textit{ab initio} motif discovery, still today at the core of the suite, are based on a variety of criteria: over-represented oligonucleotides (\textit{oligo-analysis} (1)), of spaced pairs (\textit{dyad-analysis} (8)), or positionally biased oligonucleotides (\textit{position-analysis} (9)). In 2000, a series of bacterial genomes were added to the suite, leading to the first RSAT release in 2003, supporting 100 genomes (6) (Supplementary Figure S1).

At that time RSAT started to support Position Specific Scoring Matrices (PSSMs) motif representation, in particular within the core tool \textit{matrix-scan}, which scans sequences to locate putative TFBSs (10). As more and more prokaryotic genomes were sequenced, it became possible to use cross-species conservation to detect putative regulatory signals in non-coding sequences (phylogenetic footprinting) with \textit{footprint-discovery} (11,12) (available for Prokaryotes and Fungi). In the 2008 server update (5), the number of tools almost doubled, programmatic access was offered as SOAP Web Service, and almost 700 genomes were supported, with six mirror servers across Europe and Mexico being available.

\textbf{RSAT in the high-throughput sequencing era}

The microarray technology that enabled measuring full transcriptomes and its ChIP-on-chip application opened the field of regulatory genomics to genome-wide analyses. In 2007, the ChIP-seq approach (13,14), taking advantage of high-throughput sequencing technology, revolutionised the field with an unprecedented level of precision and quantity of experimentally-detected functional TF binding regions. In contrast with alternative tools, RSAT programs coped well with datasets obtained from this technology, prompting the development of the user-friendly integrated pipeline \textit{peak-motifs} (4,15,16), which enabled the online analysis of full datasets without size restriction. To better cope with the increasing size of datasets, multiple RSAT tools have been optimized over the years to reduce execution time. The 2011 RSAT server update (4) comprised new tools to retrieve sequences on-the-fly from Ensembl (retrieve-ensembl-seq (17)), to generate control sets, to discover motifs (\textit{info-gibbs} (18)), to evaluate the quality of PSSMs (\textit{matrix-quality} (19)) and to compare them (\textit{compare-matrices}).

In 2015, after a drastic increase in available genomes in RSAT (~3300 at that time, with support for Ensembl...
Genomes), it became necessary to reorganise the public mirrors into taxon-specific servers, concomitantly better accommodating the specific needs of user communities. The NAR server update (3) further presented novel tools, such as matrix-clustering (3,20), a clustering tool to regroup similar PSSMs and offers a dynamic visualisation of aligned PSSMs. We also introduced a first series of tools (variation-seq, retrieve-variation-seq) to predict the impact of non-coding variants on cis-regulatory elements. To facilitate a local installation of the suite, Virtual Machines were made available for download.

Currently, the RSAT Website includes six novel or refactored programs (tagged with asterisks in Table 1) for a total of 52 programs. As of January 2018, RSAT public servers support 10 032 locally-installed genomes (including 9 451 Prokaryotes, 238 Fungi, 91 Metazoa, 66 Plants and 186 Prokaryotes). For Prokaryotes, we now only support NCBI genome assemblies classified as ‘Complete Genome’ and ‘Chromosome’, leaving out any genome project classified as ‘Contig’ or ‘Scaffold’. Nevertheless, additional genomes can be installed on request.

**RSAT 2018 NOVELTIES**

Many RSAT functionalities are described in the previous 2015 NAR update (3). We focus on the main novelties below (Table 1), situating the new tools in the global context of the suite.

**Obtaining sequences and homologous genes**

RSAT maintains locally-installed genomes and integrates on-the-fly access to external databases (Ensembl and UCSC). It offers tools to retrieve sequences relative to annotated genomic features (retrieve-seq for promoter sequences of local genomes, retrieve-ensembl-seq (17) for Ensembl vertebrate species). For genome-wide epigenomic datasets where genomic coordinates are usually specified in BED format, corresponding sequences can be extracted from UCSC (fetch-sequences from UCSC) and from local genomes with a new program supporting repeat-masking (retrieve-segbed). In addition, retrieve-ensembl-seq supports the retrieval of sequences from homologous genes. A new tool also relies on Ensembl Compara (21) to return detailed information on homologous genes in a set of reference organisms (get-orthologs-compara, currently only for Plants). In Fungi and Prokaryotes, lists of orthologous genes can be obtained with get-orthologs.

**Obtaining motifs (PSSMs)**

The ChIP-seq ‘revolution’ gave rise to a dramatic increase in the number of PSSMs stored in established motif databases, such as JASPAR (22), and a multiplication of independent motif collections. To facilitate the access to motifs, RSAT now locally hosts 50 external motif databases (JASPAR (22), Cis-Bp (23), FootprintDB (24), etc.) (Supplementary Table S1), covering DNA and RNA binding motifs in a wide range of organisms (Metazoa, Prokaryotes, Fungi, Plants). These collections have been homogenised in TRANSFAC format to alleviate format conversion. A new tool enables the extraction of particular motifs from these collections, based on identifiers or names (retrieve-matrix). The selection menu for motif collections is organised by color-coded taxons and is searchable to simplify access. This menu is also integrated in the tools using PSSMs as input, so that motifs can now be directly selected, rather than copy/pasted from external databases. To cope with the problem of motif redundancy within and across collections, we have established non-redundant motif collections by automatically clustering all the motifs from these collections (20).

**Detecting regulatory variations**

Population genomics and Genome-Wide Association Studies (GWAS) projects produce information on genetic variants (SNPs, indels), many of which are located in non-coding regions of the genome, and may thus affect cis-regulatory elements. To predict the impact of sequence variations on TF binding, variants and their flanking sequences can be extracted (retrieve-variation-seq) and scanned with a collection of motifs (variation-scan). This tool has been refactored to support multiallelic variants and indels, and optimized for time efficiency. A new tool further eases file format conversion between VCF, GVF and varBed (convert-variations). Altogether, RSAT sequence variation analysis tools (variation-info, retrieve-variation-seq, convert-variations and variation-scan) enable users to input any motif collection, and either retrieve Ensembl annotated variants, using either IDs or genomic coordinates, or input their own variants of interest (manuscript in prep).

**Enhanced Web interface and visualisation**

The home page has been extensively redesigned to simplify navigation and facilitate access to the tutorials, training material, and to the question-based menu guiding new users to the appropriate tools depending on their aims. A box to search the tools has been added, while tools that are not available on certain servers now appear deactivated in the menu. The number of organisms supported on each server is now clearly displayed. To accommodate the increasing number of supported organisms (especially in the Prokaryotes server), the selection menu has been replaced by a search engine implemented in Ajax. The visualisation tool feature-map has been re-implemented using modern libraries (d3) (feature-map2). A twitter account @RSATools is now alive with the feed displayed on the main page.

**USE CASES**

We present three use cases that exemplify applications integrating the novel tools into routine analysis (Supplementary Use Cases).

Use Case 1: Identify the binding motif for VRN1 in the promoters of the Flowering Locus T-like 1 orthologous genes. This use case on the Plant server integrates the usage of the novel tools get-orthologs-compara and feature-map2, with matrix-scan and retrieve-sequence.

Use Case 2: Select the motifs of transcription factors that conform AP-1 heterodimers, identify and reduce redundancy within this set of motifs, and detect AP1 binding...
| Application                          | Program name             | Input                          | Output                          | Description                                                                                                                                                                                                 |
|-------------------------------------|--------------------------|-------------------------------|---------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Obtaining sequences (Sequence Tools)| retrieve-seq             | Gene names                    | Sequences                       | Given a set of gene names, returns upstream, downstream (relative to ORF start) or unspliced ORF sequences. Segments overlapping an upstream ORF can be excluded or included. From a set of genomic coordinates (bed file), collects the sequences from the UCSC genome browser. |
|                                     | fetch-sequences (from UCSC) | Genomic coordinates           | Sequences                       | From a set of genomic coordinates (bed file), collects the sequences from the UCSC genome browser. Supports repeat masking option.                                                                           |
|                                     | *retrieve-seq-bed         | Genomic coordinates           | Sequences                       | Returns upstream, downstream, intronic, exonic, UTR, mRNA or CDS for a list of genes from EnsEMBL vertebrates.                                                                                          |
|                                     | retrieve-ensembl-seq      | Gene names                    | Sequences                       | Analyses oligonucleotide occurrences in a set of sequences, and detects over/under-represented oligonucleotides, using various background models and scoring statistics.          |
| Motif discovery                     | oligo-analysis           | Sequences                     | Over/under-represented oligonucleotides + PSSM |                                                                                                                                                                                                          |
|                                     | dyad-analysis            | Sequences                     | Over/under-represented dyads + PSSM | Detects over-represented dyads (spaced pairs of oligonucleotides) within a set of sequences.                                                                                                           |
| NGS ChIP-seq                        | peak-motifs              | Sequences                     | Discovered motifs + predicted sites |                                                                                                                                                                                                          |
| Pattern matching                    | crer-scan                | Transcription factor binding sites | Cis-regulatory enriched regions (CRER) | Given a set of cis-regulatory elements (predicted sites, annotated sites, ChIP-seq peaks), detects regions presenting a significant enrichment in CRERs.                                                                 |
|                                     | matrix-scan (-quick)     | Sequences + PSSMs             | Matching positions in input sequences | Scans sequences with one or several PSSMs to identify instances of the corresponding motifs (putative sites). Supports a variety of background models (Bernoulli, Markov chains of any order). |
| Motif quality and comparisons       | *retrieve–matrix          | Motif collection + motif name/ID | Motif (PSSM)                    | From a chosen motif collection (supported external database), extract the PSSMs specified by the provided name or identifier.                                                                                   |
| (Matrix Tools)                      | matrix-quality           | Motif (PSSM) + sequence set(s) | Score distribution statistics + ROC curves | Evaluates the quality of a PSSM by comparing score distributions obtained with this matrix in control sequence sets,                                                                                                                                                     |
|                                     | compare-matrices         | Two sets of PSSM              | Similarity scores + matrix alignments | Compares two collections of PSSMs, and returns various similarity statistics + matrix alignments.                                                                                                         |
|                                     | matrix-clustering        | One or several sets of PSSM   | Clusters of matrices + similarity trees | Clusters similar PSSMs and builds consensus matrices for each cluster.                                                                                                                                  |
| Comparative genomics                | get-orthologs            | Gene names + taxon             | List of homologous genes with percentage of identity, alignment length, and e-value | Given a list of Ensembl stable gene IDs from one or more query organisms, returns orthologs (optionally paralogs and homologs). Relies on primary data from Ensembl Compara.                                           |
|                                     | *get-orthologs-compara  | Ensembl gene ids              | Ensembl gene ids + homology relation information |                                                                                                                                                                                                          |
|                                     | footprint-discovery      | Sequences                     | Conserved dyads + PSSM          | Detects phylogenetic footprints by applying dyad-analysis in promoters of a set of orthologous genes.                                                                                                      |
|                                     | footprint-scan           | Sequences + PSSM              | Conserved motifs + binding sites | Scans promoters of orthologous genes with one or several PSSMs to detect enriched motifs and predict phylogenetically conserved target genes.                                                            |
| Regulatory variants (GeneticVariation Tools) | retrieve-variation-seq | Identifier of variations | Sequences of the variants | Given a set of IDs for genetic variations, returns the corresponding variants and their flanking sequences. The output file can be scanned with the tool variation-scan. |
Table 1. Continued

| Application | Program name       | Input                            | Output                                      | Description                                                                 |
|-------------|--------------------|----------------------------------|---------------------------------------------|-----------------------------------------------------------------------------|
|             | *variation-scan    | Variant sequences                | Regulatory variants                         | Scans variant sequences with PSSM and report variations that affect the binding score, in order to predict regulatory variants. Faster version with novel support for indels. |
|             | *convert-variations| File with genetic variants       | File with genetic variants in the specified format | Converts between different file formats that store genetic variation information. The most commonly used formats are: VCF and GVF. varBed format presents several advantages for scanning variations with matrices using variation-scan. Generates a graphical map of features localized on one or several sequences. Several maps can be drawn in parallel, allowing to detect conserved positions. Exports in svg, png/jpeg. |
| Visualisation| *feature-map2      | Coordinates (relative or absolute) | Image depicting features over lines representing sequences | This table presents a selection of key tools equipped with a Web interface. Connect to the RSAT Web site to obtain the complete list of available tools. Novel tools and major updates since the 2015 Web software issue are emphasized by an asterisk (*). |

sites in JUNB keratinocyte ChIP-seq peaks. This use case introduces the usage of retrieve-matrix, matrix-clustering, sequences-from-bed and feature-map2, along with matrix-scan.

Use Case 3: Identify genetic variants associated with melanoma, which could affect AP1 binding sites. Exemplifies the usage of the complete refactored variation tools on the Metazoa server: retrieve-variation-seq and variation-scan, with convert-variations.

ACCESSING AND LEARNING TO USE RSAT

In addition to the public Web sites, RSAT can be remotely accessed via SOAP Web services. RSAT can also be used via Unix command-line, after installation of the suite on a local server or on a computer cloud, either from source code or with a Virtual Machine (on any operating system supporting VirtualBox, including Windows) (3).

To learn how to use the RSAT suite, extensive documentation material is available (3). The latest protocols (25,26) describe motif discovery in plant genomes, but the approach can be applied to any organisms supported on the other RSAT servers. Although the Web interfaces are being continually updated, most of our previously published protocols (10,15,27) are still usable to gain experience in understanding the underlying algorithms, choosing the relevant parameters and interpreting the results.

CONCLUSIONS

The RSAT suite is unique for its broad range of functionalities and supported organisms from all kingdoms. The main alternative is the MEME suite (28), which mainly focuses on motif analyses. Since the beginning of the project, RSAT strives to facilitate inter-connections with complementary programs (including MEME) and motif databases, thanks to a series of utility tools to convert alternative file formats (convert-background-models, convert-features, convert-matrix, etc.). Celebrating its 20th Anniversary, RSAT is gearing up for more interoperability with REST programmatic standards, better packaging with conda associated with a Docker image, and centralised documentation on GitHub.

DATA AVAILABILITY

All public RSAT servers are accessible from the RSAT portal at http://www.rsat.eu/. RSAT Web servers can be freely accessed by all users without login requirement.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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