Rfam 14: expanded coverage of metagenomic, viral and microRNA families

Ioanna Kalvari 1, Eric P. Nawrocki 2, Nancy Ontiveros-Palacios 1, Joanna Argasinska 1, Kevin Lamkiewicz 3,4, Manja Marz 3,4, Sam Griffiths-Jones 5, Claire Toffano-Nioche 6, Daniel Gautheret 6, Zasha Weinberg 7, Elena Rivas 8, Sean R. Eddy 8,9,10, Robert D. Finn 1, Alex Bateman 1 and Anton I. Petrov 1,*

1European Molecular Biology Laboratory, European Bioinformatics Institute, Wellcome Genome Campus, Hinxton, Cambridge CB10 1SD, UK, 2National Center for Biotechnology Information, National Library of Medicine, National Institutes of Health, Bethesda, MD 20894, USA, 3RNA Bioinformatics and High-Throughput Analysis, Friedrich Schiller University Jena, Leutragraben 1, 07743 Jena, Germany, 4European Virus Bioinformatics Center, Leutragraben 1, 07743 Jena, Germany, 5Faculty of Biology, Medicine and Health, University of Manchester, Oxford Road, Manchester, M13 9PT, UK, 6Université Paris-Saclay, CEA, CNRS, Institute for Integrative Biology of the Cell (I2BC), 91198, Gif-sur-Yvette, France, 7Bioinformatics Group, Department of Computer Science and Interdisciplinary Centre for Bioinformatics, Leipzig University, 04107 Leipzig, Germany, 8Department of Molecular and Cellular Biology, Harvard University, Cambridge, MA 02138, USA, 9Howard Hughes Medical Institute, Harvard University, Cambridge, MA 02138, USA and 10John A. Paulson School of Engineering and Applied Science, Harvard University, Cambridge, MA 02138, USA

Received September 15, 2020; Revised October 14, 2020; Editorial Decision October 15, 2020; Accepted October 21, 2020

ABSTRACT

Rfam is a database of RNA families where each of the 3444 families is represented by a multiple sequence alignment of known RNA sequences and a covariance model that can be used to search for additional members of the family. Recent developments have involved expert collaborations to improve the quality and coverage of Rfam data, focusing on microRNAs, viral and bacterial RNAs. We have completed the first phase of synchronising microRNA families in Rfam and miRBase, creating 356 new Rfam families and updating 40. We established a procedure for comprehensive annotation of viral RNA families starting with Flavivirus and Coronaviridae RNAs. We have also increased the coverage of bacterial and metagenome-based RNA families from the ZWD database. These developments have enabled a significant growth of the database, with the addition of 759 new families in Rfam 14. To facilitate further community contribution to Rfam, expert users are now able to build and submit new families using the newly developed Rfam Cloud family curation system. New Rfam website features include a new sequence similarity search powered by RNAcentral, as well as search and visualisation of families with pseudoknots. Rfam is freely available at https://rfam.org.

INTRODUCTION

Rfam is the database of non-coding RNA (ncRNA) families (1), each represented by a multiple sequence alignment (known as the seed), a consensus secondary structure, and a covariance model to annotate non-coding RNAs in nucleotide datasets using the Infernal software (2). Rfam and Infernal are commonly used to annotate ncRNAs in newly sequenced genomes (3,4), and are the core components of the genome annotation pipelines in Ensembl (5), Ensembl Genomes (6), NCBI Prokaryotic and Eukaryotic Gene Annotation (7,8) and other resources. For example, PDBe uses Rfam to enable searching for RNA chains, such as tRNA or rRNA, in 3D structures (9), while RNAcentral uses Rfam to detect incomplete sequences, RNA type annotations errors, and provide other quality controls (10). The manually curated multiple sequence alignments from Rfam are also used for training and benchmarking new software, such as secondary structure prediction algorithms (11,12).

Here we describe community-driven improvements and updates that are available in Rfam 14 (releases 14.0–14.3), including new RNA families, new features on the Rfam website, and new tools for expert users to contribute families to the database. Rfam 14.3 contains 3444 families, rep-
representing a 28% increase since version 13.0. The majority of the new families have been generated in collaboration with other RNA resources and experts, in particular ZWD, miRBase and European Virus Bioinformatics Center (EVBC), supplemented by in-house literature curation. Following the transition from annotating a subset of ENA (13) to annotation of a collection of non-redundant and complete genomes in Rfam 13.0 (1), we have expanded the set of genomes in the Rfam sequence database (Rfamseq) by 76% in Rfam 14.0 to reflect the increase in number of genomes. Rfamseq now includes 14,772 genomes from all domains of life. Here, we also describe the newly-developed Rfam Cloud curation pipeline, which marks a further major step towards ensuring that Rfam remains a core open and sustainable resource for the whole RNA community.

NEW FAMILIES FROM ZWD

Rfam releases 14.1 and 14.3 included 253 new families from the Zasha Weinberg Database (ZWD) that were discovered by a systematic computational analysis of intergenic regions in Bacteria and metagenomic samples (14), as well as newly identified ribosomal leaders (r-leaders) (15). ZWD stores the original manually curated multiple sequence alignments produced in the Breaker and Weinberg groups over the last decade. ZWD is a git-based resource that currently includes 417 alignments and is available at https://bitbucket.org/zashaw/zashawinbergdata. Examples of the new and updated ZWD-based families are shown in Figure 1.

Prior to version 14, Rfam had incorporated 180 families from ZWD, but in Rfam 14 a new process was developed to automate the addition of ZWD alignments. Since the inception of Rfam in 2003, it has been a strict requirement that sequences in seed alignments were derived from Rfamseq, the underlying sequence database that is searched by covariance models for all families with each release. The Rfamseq database sequences must also exist in public databases like GenBank or ENA and each seed sequence is automatically checked to ensure that it is a valid sequence of a publicly available sequence. However, many of the ZWD families come from environmental samples, so the sequences were not found in the INSDC archives. Previously such sequences were replaced with closely related ones from INSDC or removed, which required modifying the user-submitted alignments and could result in smaller alignments missing covariation compared to the originals. In order to preserve the manually curated alignments as much as possible and avoid the error-prone manual steps, we imported all ZWD sequences into RNAcentral (10) to create stable accessions and allowed for the RNAcentral identifiers to appear in Rfam seed alignments. Furthermore, we have updated the Rfam pipeline to allow seed alignment sequences to derive from any valid ENA or GenBank accession. This added flexibility not only allowed us to import the ZWD families, but will facilitate the construction of seed alignments, especially using the new cloud-based Rfam family building pipeline (see below).

To confirm the completeness of the import, we systematically compared Rfam families with ZWD using the Infernal cmscan program (2) to search all ZWD sequences with Rfam covariance models. Ninety eight percent of ZWD alignments have now been imported into Rfam except for seven families lacking covariation support as determined by R-scape (16) and 43 families marked ‘Not for Rfam’ in ZWD. The mapping between ZWD alignments and Rfam accessions is provided in Supplementary Information.

We also used CaCoFold (17) to identify the pre-existing ZWD-based Rfam families requiring an update using the new RNAcentral import mechanism. For example, the manA alignment from Rfam (RF01745) had only 29 statistically significant basepairs while a CaCoFold structure based on the corresponding alignment in ZWD contained 48 significant basepairs. A comparison between the Rfam and ZWD versions of the alignment revealed a mistake in the Rfam secondary structure introduced during manual import. As a result, the manA Rfam alignment was updated to its original version from ZWD (Figure 1A, right). We are systematically reviewing other ZWD-based families that may require an update and will release them in future Rfam versions (17).

NEW WORKFLOW FOR VIRAL RNA FAMILIES

Despite the small size (3400–41,000 nucleotides) of RNA virus genomes, they contain several conserved RNA structures vital for protection against exonucleases, genome diversification, and play a crucial role in various stages of their viral life cycle (18). Many viruses rely on RNAs to infect and replicate inside a host: for example, the cis-acting element of coronaviruses is essential for replication (18), while in Dengue viruses replication depends on RNA structure-mediated circularization (19,20). RNA viruses are often highly contagious and can lead to fast-emerging severe diseases (21). Being able to identify and understand viral RNAs is essential for the scientific community to develop novel drugs and treatments in response to pandemics like COVID-19.

In releases 14.2 and 14.3, Rfam created 22 new families, focusing on Coronaviridae and Flavivirus structured RNAs. These releases are the first of many planned extensions for viral RNA families in Rfam. We will continually update the functional RNA structures of viral clades in Rfam and aim to provide a comprehensive database for virologists interested in RNA secondary structures. This effort is the first of its kind to bring bioinformaticians and virologists together to publish detailed and specific RNA alignments and secondary structures for a broad range of RNA viruses, and thus complements protein-based databases like vFam (22) and RVDB (23).

Coronaviridae families

In response to the SARS-CoV-2 outbreak, the Rfam team prepared a special release 14.2 dedicated to the Coronaviridae RNA families (24), including ten new and four revised families that can be used to annotate SARS-CoV-2 and other Coronaviridae genomes with RNA families. The new RNA families represent the entire 5′- and 3′-untranslated regions (UTR) for Alpha-, Beta-, Gamma- and Deltacoronavirus subfamilies (Figure 2A–D). The families were built...
Figure 1. Example ZWD-based Rfam families. (A) Metagenomic-based RNAs Drum (RF02958), LOOT (RF03000), and manA (RF01745) RNAs; (B) S4-Clostridia (RF03140), L4-Archaeoglobi (RF03135) and L13-Bacteroidia (RF03127) r-leader families. All families have been created in releases 14.1 and 14.3, except for manA, which was updated in 14.3. Purines (adenine or guanine) are shown as ‘R’, while pyrimidines (cytosine or uracil) are shown as ‘Y’.

based on a set of high-quality alignments produced with LocARNA (25) and reviewed by expert virologists from the Marz group (University of Jena) and the EVBC. In addition, a set of alignments was built for the Sarbecovirus sub-genus UTRs, including the SARS-CoV-1 and SARS-CoV-2 UTRs (Figure 2E). While the Alpha-, Beta- and Delta-coronavirus alignments and structures were refined based on the literature (26–28), the Gammacoronavirus families are based on prediction alone due to the lack of experimental data.

We also reviewed and updated the existing Coronaviridae Rfam families, including Coronavirus packaging signal (RF00182), Coronavirus frameshifting stimulation element (RF00507), Coronavirus s2m RNA (RF00164), and Coronavirus 3′-UTR pseudoknot (RF00165). Two Rfam families were superseded by the new whole-UTR alignments and removed from the database: the Coronavirus SL-III cis-acting replication element (RF00496) representing a single stem that is now found in Alpha- and Betacoronavirus 5′-UTR families (aCoV-5UTR and bCoV-5UTR) and Coronavirus_5p_sl1_2 (RF02910) representing two stems from aCoV-5UTR. The new and updated Coronaviridae Rfam families are available at https://rfam.org/covid-19.

Flavivirus families

In order to create new Flavivirus families, we identified a set of full-length, non-redundant Flavivirus genomes to serve as a source of sequences. All Flavivirus genomes marked as ‘complete’ were downloaded from the ViPR database (10 443 genomes as of August 3rd, 2020) (29). Since most of the known flaviviral genomes are represented by the Dengue virus genomes (∼58%), the ViPR-based Dengue sequences were excluded and the curated high-quality RefSeq sequences were used instead. The genomes were scanned with the existing Rfam models for Dengue virus SLA (RF02340), Flavivirus capsid hairpin cHP (RF00617), and Flavivirus 3′ UTR CRE (RF00185) to identify the genomes with complete 5′ and 3′ UTRs. The resulting 2661 genomes were used to refine the existing Flavivirus Rfam families as well as previously published models (30) to produce a set of 12 new and 2 updated Rfam families (Table 1).

A single Rfam family was generated for the complete Flavivirus 5′ UTR (RF03546) to respect the conserved order of the structural elements SLA and SLB. However, a single model could not represent the variability of the 3′ UTR as it can vary both in terms of structural composition and length (from 400 to 900 nucleotides). Therefore
we produced three general 3' UTR models that are valid for all viruses of the *Flavivirus* genus and represent the xRNA, CRE, and DB elements (RF03547, RF00185, and RF00525), as well as a set of specialised models for the viral clades based on their host-range (31,32). For insect-specific flaviviruses (ISFV), we created two specialised models: CRE and xRNA (RF03545 and RF03541) with increased sensitivity for ISFVs, as well as three families unique to ISFV: the repeated elements (Ra and Rb) individually (RF03544 and RF03543) and as a combined element (RF03542) (30,33). In most instances, Ra and Rb co-occur. However, in some ISFVs (e.g. Quang Binh virus (QBV) and Mosquito flavivirus (MSFV)), the Rb element is missing from some repeats (30). Furthermore, there may be novel as yet undescribed *Flavivirus* species with another repeat pattern. Therefore, the individual repeated elements and the co-occurring model were created and integrated into the Rfam.

We provide three tick-borne flavivirus (TBFV) models, enabling more sensitive searches for the CRE (RF03538) and xRNA (RF03536) elements, as well as the SL6 stem (RF03537) that appears in some 3' UTRs in TBFVs (30,34). Although there is no known vector (NKV) for some flaviviruses, they can still be grouped and annotated using the specialised CRE and xRNA models (RF03540 and RF03539). The remaining host-specific group, the mosquito-borne flaviviruses (MBFV), will be imported in a future Rfam release, once the detailed analysis becomes available in the literature. Finally, two *Flavivirus* families (RF00465 and RF02549) have been removed from Rfam since they were superseded by the new families.

Figure 2. New Rfam *Coronaviridae* 5' and 3' UTR families are depicted schematically within the complete viral genomes. (A) *Alphacoronavirus* UTRs (RF03116 and RF03121); (B) *Betacoronavirus* UTRs (RF03117 and RF03122); (C) *Gammacoronavirus* UTRs (RF03118 and RF03123); (D) *Deltacoronavirus* UTRs (RF03119 and RF03124); (E) *Sarbecovirus* UTRs (RF03120 and RF03125); (F) visualising SARS-CoV-2 UTRs using the *Sarbecovirus* Rfam models.

Generalising the workflow for annotating viral RNAs in Rfam

We are generalising the procedure established for the *Coronaviridae* and *Flavivirus* families to annotate structured RNAs in all viruses, starting with the human pathogens like *Hepacivirus* (Hepatitis C viruses), *Filoviridae* (e.g. *Ebolavirus*) and *Rhabdoviridae* (e.g. Rabies viruses). The
Table 1. New and updated Flavivirus RNA families from release 14.3. The two updated families are marked with an asterisk

| Rfam accession | Rfam ID | Rfam description |
|----------------|---------|------------------|
| RF03546        | Flavivirus-5UTR | Flavivirus 5’ UTR |
| RF00185*       | Flavi_CRE | General Flavivirus exoribonuclease-resistant RNA element |
| RF00525*       | Flavivirus_DB | Flavivirus DB element |
| RF03547        | Flavi_xrRNA | Insect-specific Flavivirus 3’ UTR cis-acting replication element (CRE) |
| RF03549        | Flavi_JSFV_CRE | Insect-specific Flavivirus 3’ UTR cis-acting replication element (CRE) |
| RF03544        | Flavi_JSFV_repeat_Ra | Insect-specific Flavivirus 3’ UTR repeats Ra |
| RF03543        | Flavi_JSFV_repeat_Rb | Insect-specific Flavivirus 3’ UTR repeats Rb |
| RF03542        | Flavi_JSFV_repeat_Ra_Rb | Tick-borne Flavivirus short stem-loop SL6 |
| RF03541        | Flavi_JSFV_xrRNA | Tick-borne Flavivirus exoribonuclease-resistant RNA element |
| RF03540        | Flavi_NKV_CRE | No-Known-Vector Flavivirus 3’ UTR cis-acting replication element (CRE) |
| RF03539        | Flavi_NKV_xrRNA | No-known vector Flavivirus exoribonuclease-resistant RNA element |
| RF03538        | Flavi_TBFV_CRE | Tick-borne Flavivirus 3’ UTR cis-acting replication element (CRE) |
| RF03537        | Flavi_TBFV_xrRNA | Tick-borne Flavivirus short stem-loop SL6 |

method is based on clustering the set of available complete genomes in ViPR (29) and RefSeq (8) and calculating high-quality genome alignments of representative viruses. These alignments are refined by local RNA secondary structure information using LocARNAs analyses (35). The structurally conserved elements, as identified by RNAz (36), are extracted from the alignment, compared with the literature and manually curated by expert virologists from the EVBC before submission to Rfam. The new viral families will enable researchers to rapidly annotate viral genomes with conserved RNA structures using the Infernal software and Rfam covariation models, detect structured non-coding viral regions in metagenomic data, and gain insight into the recombination of the conserved RNA structures (37).

SYNCHRONISING MICRORNA FAMILIES BETWEEN RFAM AND MIRBASE

MicroRNAs are a class of ∼22 nt ncRNA that regulate gene expression at the post-transcriptional level. Animal and plant genomes contain hundreds to thousands of microRNA genes, many of which have been implicated in processes such as development and disease (38). For example, the mir-17-92 cluster and mir-155 have been shown to act as oncogenes [oncomirs; reviewed in (39)]. Understanding the evolutionary relationships between microRNAs of different species allows the transfer of gene annotation and functional information, for example from model organisms to human. MicroRNA sequences and annotations are aggregated in miRBase (40), the authoritative resource for published microRNA genes.

mirBase is primarily a sequence database, but both Rfam and mirBase contain classifications of microRNA families. However, before Rfam 14.3 the two databases have not been coordinated or synchronised. Previously, miRBase used a semi-automated, clustering method relying on BLAST (41). These sequence-only mirBase families have higher coverage but lower quality than the Rfam microRNA families. In release 14.2, Rfam contained 529 microRNA families, while miRBase v22 annotated 1,983 microRNA families. Only 28% of the miRBase families matched one or more of the Rfam 14.2 families. There was therefore an opportunity to create up to 1500 new families to increase the coverage of microRNAs in Rfam, as well as investigate and rationalise the entries that are unique to each database. Here, we present the first phase of a comprehensive review and classification of microRNA gene families in collaboration with miRBase.

Based on miRBase v22, we have manually curated an initial set of 1678 multiple sequence alignments. The remaining ∼300 families require more detailed consideration and curation, including merging and splitting, and will be revisited in a second phase. The manually curated alignments were used as seeds for building the covariance models, with which we searched the Rfam sequence database for homologs of these new microRNA families. At the time of writing, we have created and submitted 356 new microRNA families to Rfam and updated 40 existing families. Work is underway to create and review the remaining microRNA families, which will be made available in the subsequent Rfam release.

The workflow established here will rationalise microRNA families in the key RNA database resources, and ensure consistency between Rfam, mirBase, and RNAcentral. mirBase will retain its focus on sequences, and Rfam will be the primary resource of microRNA family classifications. These Rfam microRNA family classifications will be made available in both miRBase and RNAcentral. The relationship between the databases will create a cycle of improvement. All Rfam microRNA seed alignments will contain only sequences that are validated as microRNAs by miRBase. Rfam microRNA families will be used to identify new member sequences, and these new sequences will be reviewed by miRBase for inclusion in both miRBase and RNAcentral. The validated microRNA sequences will then be added to an updated version of the Rfam seed alignment. Since miRBase and Rfam are both RNAcentral member databases, the synchronisation of miRBase and Rfam is facilitated by consistent use of RNAcentral sequence identifiers for all sequences in microRNA seed alignments. The new Rfam models coupled with Infernal will enable other resources, including the key genome browsers and model organism databases, to annotate microRNA sequences in genome sequences in a rigorous and sustainable way.

CLOUD-BASED FAMILY CURATION SYSTEM

Creating new Rfam families is a computationally expensive process that depends on searching the Rfam non-redundant collection of complete genomes (360Gb as of release 14.3)
using Infernal (2). Such searches are impractical without access to storage, memory, and CPU resources that can parallelise the execution and reduce the running time. To enable the scientific community to build Rfam families without setting up their own computational infrastructure, the Rfam curation pipeline was packaged in software containers using Docker and deployed using Kubernetes, a container orchestration engine that automates cloud deployment and manages containerised applications. The new cloud-based family curation pipeline, Rfam Cloud, is hosted at the Embassy Cloud platform provided by EMBL-EBI.

Rfam Cloud provides access to a command line environment that enables users to create or modify Rfam families. The Rfam family building process involves several steps: starting with a single sequence or a seed alignment containing known examples of a family, the user can search for similar sequences using a covariance model, refine the family by adding more sequences into the seed alignment, and identify a bit score cutoff (the gathering threshold) that separates homologous sequences that constitute an Rfam family from non-homologous sequences. This process can be iteratively repeated to improve the seed alignment and the associated covariance model. The user can also perform quality control checks to verify the format and ensure that there are no overlaps with existing families. Upon successful completion of quality control steps, new families can be submitted to the Rfam team for review and inclusion in the main Rfam database. The Rfam Cloud documentation describes the curation tools and provides guidelines and tips for common tasks, such as selecting the gathering threshold for a family and is available at https://rfam.org/cloud.

Rfam Cloud enables RNA family building using Rfam’s Infernal-based search pipeline and requires some manual intervention between iterations. Two alternative RNA family building tools with more automation than the Rfam Cloud pipeline are GraphClust2 (42) and RNAliken (43), both of which also use Infernal amongst other tools. However, Rfam Cloud is directly linked to the Rfam database where the nascent families can be deposited and accessed by the larger community.

Following a testing period in 2019, Rfam Cloud was successfully used in a higher education setting in collaboration with the University of Paris-Saclay. A three-month masters level RNA bioinformatics course was held from October 2019 to January 2020, where eight teams of three graduate students used Rfam Cloud to build RNA families. Each team was assigned one candidate sequence to initiate the family building process. The students used Rfam Cloud at their own pace and were provided support using a Slack workspace where the Rfam team answered questions and helped with troubleshooting. The students produced 6 new Rfam families (RF03530-RF03535) that are listed in Table 2.

Table 2. New Rfam families created during an Rfam course at University of Paris-Saclay

| Rfam accession | Rfam ID | Description                                                                 | Reference |
|----------------|--------|------------------------------------------------------------------------------|-----------|
| RF03530        | bgG-cis-reg | cis-regulatory element of the bgG/LicT operon                              | (44)      |
| RF03531        | n00280_RNA | Clostridioides difficile sRNA included into helicase gene                   | (44)      |
| RF03532        | SQ2397_RNA | cis-regulator of HTH transcription factor                                   | (44)      |
| RF03533        | SQ1002_RNA | Clostridioides difficile sRNA SQ1002                                      | (44)      |
| RF03534        | sRNA71   | Staphylococcus aureus small RNA                                             | (45)      |
| RF03535        | TEG147   | Staphylococcus aureus small RNA TEG147                                     | (45)      |

The Rfam Cloud pipeline is currently used by a group of 15 external users contributing to family curation. Following approved requests, users are provisioned with a private cloud account where they can access the Rfam curation tools through a command line interface. New accounts can be requested at https://rfam.org/cloud.

Getting credit for authoring Rfam families using ORCID

The ORCID registry (https://orcid.org) provides unique identifiers that unambiguously link researchers with their scientific papers and other outputs. The Rfam database is now integrated with the ORCID system, allowing authors of Rfam families to get credit for their contributions by adding Rfam family accessions to their ORCID profiles. The ORCID identifiers have been manually associated with existing Rfam families and the new families. This feature was enabled by the ‘Claim to ORCID’ functionality provided by the EBI Search (46). The process includes three steps: (a) search for an ORCID identifier on the EMBL-EBI website; (b) manually select all or a subset of listed entries and click ‘Claim to ORCID’; (c) login to ORCID using the same ORCID identifier and agree to add the Rfam entries to the ORCID record. As the RNA community begins using the Rfam Cloud infrastructure, this integration will allow the growing number of Rfam contributors to get credit for their work.

OTHER IMPROVEMENTS

Pseudoknot search and visualisation

The R-scape software analyses covariation support for RNA secondary structure based on multiple sequence alignments (16). Starting with version 1.2.0, R-scape systematically identifies pseudoknots and other non-nested interactions provided that they have covariation support (17) and displays them using R2R (47). We analysed all Rfam seed alignments with R-scape and added pseudoknot visualisation to the Rfam website (Figure 3A). In addition, we updated the Rfam text search to enable searching for families with or without pseudoknots and filter the results by whether the pseudoknots have covariation support (Figure 3B).

New sequence search integrated with RNAcentral

The Rfam sequence search has been updated to use the RNAcentral reusable sequence search component. The search is executed on the RNAcentral cloud infrastructure.
Figure 3. (A) R-scape visualisation of the skipping-rope RNA (RF02924). The nucleotides forming the pseudoknot are labelled pk1 and are shown as a separate stem. The basepairs with significant covariation, according to R-scape, are colored green. (B) Pseudoknot facets from the Rfam text search.

Figure 4. Rfam sequence search using the RNAcentral sequence search component. (A) Query sequence. (B) A sequence identical to the query found in RNAcentral. (C) Rfam classification using Infernal. (D) Alignment between the query and the Rfam covariance model. (E) Secondary structure visualised using R2DT displayed using the consensus secondary structure from the corresponding Rfam model. Clicking the link opens a new window with the detailed secondary structure diagram. (F) Similar sequences from RNAcentral.
and, in addition to annotating the query sequence with Rfam families using Infernal, identifies similar sequences in the RNAcentral sequence database using nhmmer (48).

The new search also performs the clan competition procedure (49) which selects the longest and highest scoring hit if several Rfam families from the same clan match the query sequence. The new search can show related RNAs even if a query sequence does not match any Rfam families (e.g. most IncRNA queries will not have matches in Rfam). The search is also integrated with R2DT (50) to visualise RNA secondary structure in standard, reproducible, and recognisable layouts (Figure 4E).

The user interface features facets that enable filtering similar sequences by RNA type, organism, or the source database (Figure 4F). The results can also be filtered with any keyword and exported for further processing. The search is available under the search sequence tab at https://rfam.org/search.

CONCLUSIONS

After eighteen years of development, Rfam still continues to grow as new RNA families are regularly reported in the literature. The recent innovations and improvements described here are focused on collaboration with the RNA community and other RNA resources to share data and tools to classify RNA families. Furthermore, the new Rfam Cloud pipeline is designed to involve more RNA experts in the creation of new families and narrow the gap between cutting edge research and manual database curation. Our future plans, in addition to completing the microRNA and viral projects, include the integration of experimentally determined 3D structure information into seed alignments and connecting the existing RNA families with the latest literature using text mining. The new Rfam Cloud and ongoing collaborations with resources such as ZWD, EVBC and miRBase cements Rfam’s position as the community resource for RNA families. We invite new data submissions and feedback at https://docs.rfam.org/page/contact-us.html.

DATA AVAILABILITY

Rfam is available under the Creative Commons Zero (CC0) license at https://rfam.org. The data can be accessed in the FTP archive, as well as through an API and a public MySQL database (see https://docs.rfam.org and (51) for instructions). All code is available on GitHub under the Apache 2.0 license at https://github.com/Rfam.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

ACKNOWLEDGEMENTS

The authors would like to thank the organisers of the Be-nasque 2018 meeting where several collaborations described in this paper have been established. We would like to thank Michael T. Wolfinger (University of Vienna) for providing the ISFV, TBFV and NKV Flavivirus alignments. We also thank Ramakanth Madhugiri (Justus Liebig University Giessen) for reviewing the Coronaviridae alignments. We thank the Scientific Advisory Board members for useful feedback.

FUNDING

Biotechnology and Biological Sciences Research Council (BBSRC) [BB/M011690/1, BB/S020462/1]; European Union’s Horizon 2020 research and innovation programme [654039]; Intramural Research Program of the National Library of Medicine at the NIH; Carl Zeiss Foundation [0563-2.8/7382/2]; NIH National Human Genome Research Institute grant [R01-HG009116]; DFG [MA5082/7-1]. Funding for open access charge: Research Councils UK (RCUK), Conflict of interest statement. None declared.

REFERENCES

1. Kalvari, I., Argasinska, J., Quinones-Olvera, N., Nawrocki, E.P., Rivas, E., Eddy, S.R., Bateman, A., Finn, R.D. and Petrov, A.L. (2018) Rfam 13.0: shifting to a genome-centric resource for non-coding RNA families. Nucleic Acids Res., 46, D335–D342.

2. Nawrocki, E.P. and Eddy, S.R. (2013) Infernal 1.1: 100-fold faster RNA homology searches. Bioinformatics, 29, 2933–2935.

3. Gemmell, N.J., Rutherford, K., Prost, S., Tullis, M., Winter, D., Macey, J.R., Adelson, D.L., Suh, A., Bertozzi, T., Grau, J.H. et al. (2020) The tuatara genome reveals ancient features of amniote evolution. Nature, 584, 403–409.

4. Kim, B.-M., Kang, S., Ahn, D.-H., Jung, S.-H., Hée, H., Yoo, J.S., Lee, J.-E., Lee, S., Han, Y.-H., Ryu, K.-B. et al. (2018) The genome of common large-arm octopus Octopus minor. Gigascience, 7, giy119.

5. Yates, A.D., Achuthan, P., Akanni, W., Allen, J., Allen, J., Alvarez-Jarreta, J., Amode, M.R., Armean, I.M., Azov, A.G., Bennett, R. et al. (2020) Ensembl 2020. Nucleic Acids Res., 48, D682–D688.

6. Howe, K.L., Contreras-Moreira, B., De Silva, N., Maslen, G., Akanni, W., Allen, J., Alvarez-Jarreta, J., Barba, M., Bolser, D.M., Cambell, L. et al. (2020) Ensembl genomes 2020—enabling non-vertebrate genomic research. Nucleic Acids Res., 48, D689–D695.

7. HafN, D.H., DeCuccio, M., BadrédinN, A., Brover, V., Chetvernin, V., O’Neill, K., Li, W., Chitsaz, F., Derbyshire, M.K., Gonzales, N.R. et al. (2018) RefSeq: an update on prokaryote genome annotation and curation. Nucleic Acids Res., 46, D851–D860.

8. O’Leary, N.A., Wright, M.W., Brister, J.R., Ciufos, S., Haddad, D., McVeigh, R., Rajput, B., Robbertse, B., Smith-White, B., Ako-Adjei, D. et al. (2016) Reference sequence (RefSeq) database at NCBI: current status, taxonomic expansion, and functional annotation. Nucleic Acids Res., 44, D733–D745.

9. Armstrong, D.R., Berriosford, J.M., Conroy, M.J., Gutmanas, A., Anyango, S., Choudhary, P., Clark, A.R., Dana, J.M., Deshpande, M., Dunlop, R. et al. (2020) PDB: improved findability of macromolecular structure data in the PDB. Nucleic Acids Res., 48, D335–D343.

10. The RNAcentral Consortium (2019) RNAcentral: a hub of information for non-coding RNA sequences. Nucleic Acids Res., 47, D221–D229.

11. Puton, T., Kozlowski, L.P., Rother, K.M. and Bujnicki, J.M. (2013) CompaRNA: a server for continuous benchmarking of automated methods for RNA secondary structure prediction. Nucleic Acids Res., 41, 4307–4323.

12. Do, C.B., Woods, D.A. and Batzoglou, S. (2006) CONTRAfold: RNA secondary structure prediction without physics-based models. Bioinformatics, 22, e90–8.

13. Amid, C., Alako, B.T.F., Balavenkataraman Kadhirdavelu, V., Burdett, T., Burgin, J., Fan, J., Harrison, P.W., Holt, S., Hussein, A., Ivanov, E. et al. (2020) The European Nucleotide Archive in 2019. Nucleic Acids Res., 48, D70–D76.

14. Weinberg, Z., Lănșe, C.E., Corbino, K.A., Ames, T.A., Nelson, J.W., Roth, A., Perkins, K.R., Sherlock, M.E. and Breaker, R.R. (2017)
