A better way to find RNA virus needles in database haystacks

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Graphical overview of the pipeline starting with the RNA Virus MetaTranscriptomes (RVMT) database to uncover the expansion in RNA virus diversity. Credit: Simon Roux

A zoo once offered a coloring book featuring polar bears in winter scenes that came with crayons in various shades of white. To researchers searching for sequences of RNA viruses in large data sets, their work may be akin to finding a single snowflake on a colored-in page of that book.

Published online September 28, 2022, in Cell, a team led by researchers at Tel Aviv University in Israel, the National Center for Biotechnology Information, and the U.S. Department of Energy (DOE) Joint Genome Institute (JGI), a DOE Office of Science User Facility located at Lawrence Berkeley National Laboratory (Berkeley Lab) describe a computational pipeline that can specifically scan for those snowflakes, or RNA virus sequences. Using this workflow, the team combed through more than 5,000 data sets of RNA sequences (metatranscriptomes) generated from diverse environmental samples around the world, resulting in a five-fold increase of RNA virus diversity.

"The world of viruses around us is vast, and we now have the means to explore it," said Eugene Koonin, a senior investigator at the NCBI and one of the senior authors on the paper, of the uncovered viral diversity. "Although the technical challenges of data analysis at this scale are formidable."

Computational sieves to filter sequences

There are more microbes on the planet than particles in a handful of dirt, and viruses vastly outnumber the microbes. Advances in sequencing technologies and computational tools have uncovered a diversity of viruses that infect not just crops, animals and humans, but also microbes whose presence or absence can impact the planet's nutrient cycles.

While most organism's genetic information is encoded in DNA, with RNA delivering the instructions inside DNA to the cell, RNA viruses store their genetic information in RNA without a DNA stage. "I would argue RNA viruses globally are even less known than DNA viruses," said Simon Roux, a JGI scientist and one of the project co-leads. "But same as DNA viruses, RNA viruses infect microbes all across the world and lead to cell death and/or profound changes in the cell physiology during infection."

While all RNA viruses have a gene that encodes for an enzyme called RNS-directed RNA Polymerase (RdRP), necessary for replicating the RNA genome replication, detecting it has been a challenge. Finding the RNA virus snowflakes in the snowstorm of genomic data involved developing special computational sieves to filter out sequences that were unlikely to contain the RdRP sequence.
The work resulted from a three-way collaboration that began in 2019, recalled Uri Neri of Tel Aviv University, one of the project co-leads and first author of the study. Members of the Tel Aviv and NCBI teams, who were already working on mining prokaryotic viruses together, learned from JGI’s Nikos Kyrpides that his Microbiome Data Science group was also working on RNA virus mining. After a couple of virtual meetings of the three teams it was clear that a larger collaborative effort would be far more effective in achieving higher quality results compared to smaller individual efforts. This is also the type of synergistic and collaborative community spirit that the JGI advocates for and actively promotes.

The team used all the publicly available metatranscriptome datasets from the JGI’s Integrated Microbial Genomes & Microbiomes (IMG/M) system. “We then looked into many more samples and refined our methodology,” Neri said. “Our team grew and so did the scope of the project.” To this end, Kyrpides emphasized, the contributions of the numerous JGI science users in collecting and submitting their microbiome samples for sequencing at the JGI cannot be overstated. Their cooperation and support, he said, and in several cases, their permission to use yet unpublished sequence data, was absolutely critical for the success of this effort and so was the acknowledgement of their contribution.

Both Roux and Koonin both noted that other research in parallel has reported similar “dramatic expansions” of the global RNA virome. “We now need to compare and reconcile the findings, coming up with a single, non-redundant dataset,” said Koonin. “Hopefully, relatively soon we will be able to estimate the actual size of the RNA virome. However, this is now real Big Data, we are dealing with billions of sequences, and soon, with trillions. The development of efficient, automated approaches to analyze and classify sequence data at this scale is essential.”
