

#### A bioinformatic pipeline to elucidate the links between viruses and their hosts in microbial communities, applied to viruses in anaerobic digestion processes

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# A bioinformatic pipeline to elucidate the links between viruses and their hosts in microbial communities, applied to viruses in anaerobic digestion processes

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## Context

- $\succ$  Viruses are key-players in microbial ecosystems. However, predicting hosts from viruses is still a major challenge in microbial ecology.
- > We developed a bioinformatic pipeline including the detection of CRISPR protospacers in viral contigs, a method previously used to predict hosts from marine viruses [1].
- > We applied our pipeline to anaerobic digestion (AD) ecosystems, in the context of organic waste treatment. We focused on the diversity of viruses infecting methanogens, the latter being the key actors of methane production during AD process.

# Data

- Samples originating from 2 distinct AD microcosms
- > Cellular and viral DNA extracted and sequenced by Illumina NextSeq (2 \* 150bp): ≈40M read pairs per sample for cellular metagenomes and  $\approx$ 20M for each metavirome.

# **Pipeline**

- > The most generic steps were scripted as a snakemake workflow (https://forgemia.inra.fr/cedric.midoux/workflow\_metagenomics) to favor reproducible and scalable data analysis. It was run on the cluster of the INRAE MIGALE bioinformatics platform.
- > Several steps specifically dedicated to the prediction of hosts from viral contigs were performed using bash and python scripts.
- > For the cellular metagenomes, a non-redundant spacer database was built from the spacer sequences obtained from Spacer detection step. In addition, a metagenome-assembled genome (MAG) database was constructed from cellular metagenomic data with *Metabat2*.
- $\succ$  Host prediction was performed by using alignment-free (WiSH) or alignment-dependent (*Blastn* or *SpacePharer*) methods.

### References

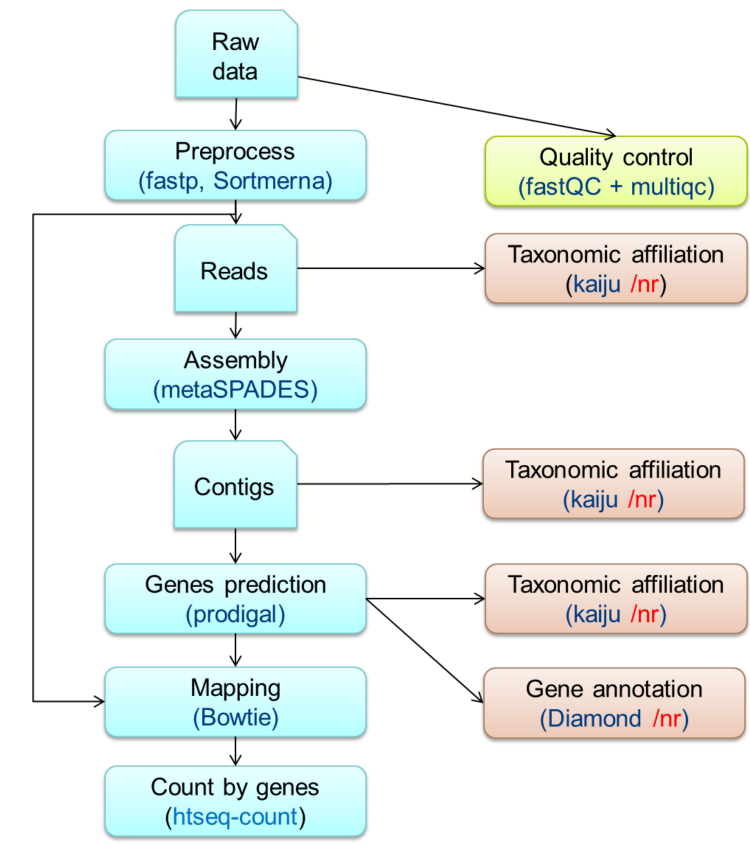
Felipe H Coutinho, et al. Marine viruses discovered via metagenomics shed light on viral strategies throughout the oceans. *Nature communications* 8, no. 1: 1-12, 2017

CRISPRCasdb spacer (<u>https://crisprcas.i2bc.paris-saclay.fr/Home/DownloadFile?filename=spacer\_34.zip</u>)

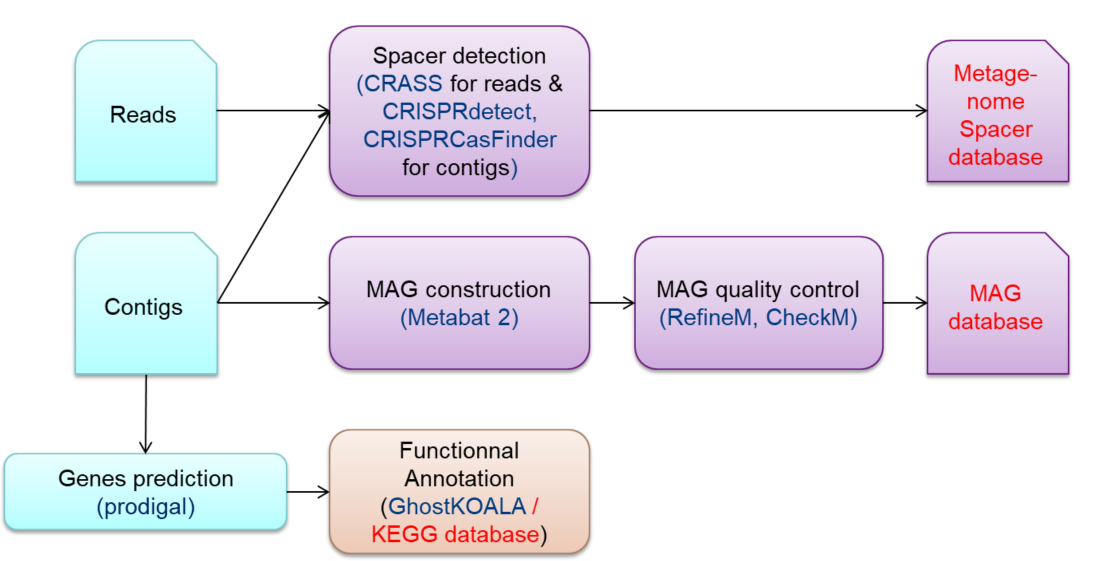
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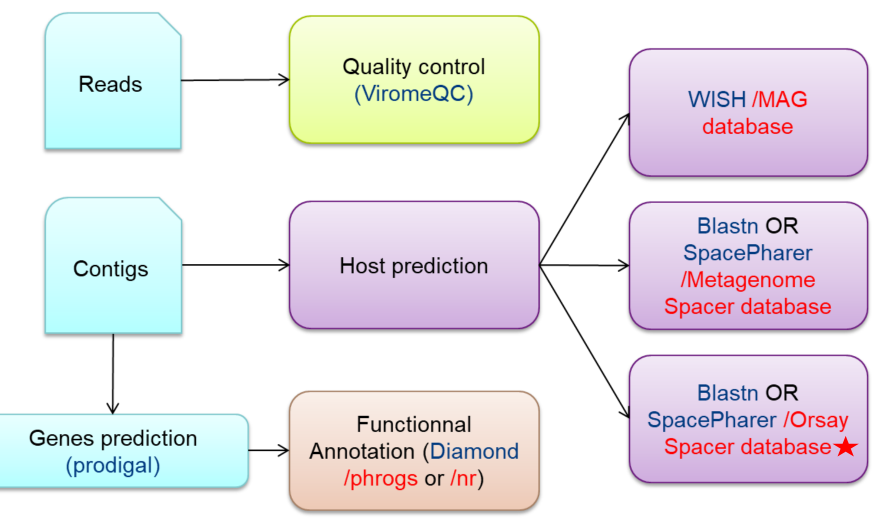
Substrate = Formate



### Generic pipeline for both cellular and viral metagenomes (snakemake)



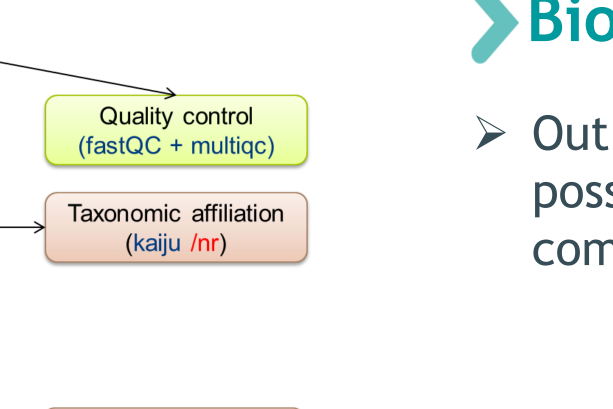
### Steps specific for cellular metagenomes



### Steps specific for metaviromes

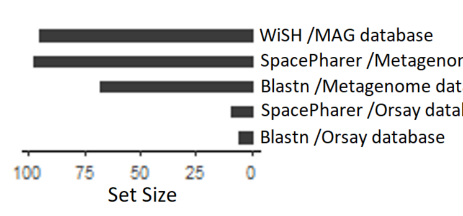


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# **Biological results**

possibly originating complementary approaches.



Sets of contigs identified by different approaches

 $\succ$  One of them is likely a novel spindle-shaped virus, a morphotype specific for archaeal viruses. Viruslike particles with this morphotype were also observed by Transmission Electron Microscopy.

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1	500	1k	1.5k	2k	2.5k	3k	

 $\succ$  Using the developed pipe-line, we were able to discover new viral diversity and to identify some host-virus links. > The pipeline will be used in future metagenomics projects and will continue to be improved by the addition of new features.





 $\succ$  Out of 5570 viral contigs longer than 3 kb, 199 were identified as from archaeal thanks to viruses

