


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Code

 alessia	Updated containers, bumped version	9a844a7 · last year	49 Commits
assets	Added test data, demo data, and asset data	3 years ago	
bin	Small edits and typos	3 years ago	
conf	Improved testing	last year	
data/test_data	Added test data, demo data, and asset data	3 years ago	
modules	Update to DSL2	last year	
LICENSE	Initial commit	7 years ago	
README.md	Updated containers, bumped version	last year	
YAMP.nf	Update to DSL2	last year	
nextflow.config	Updated containers, bumped version	last year	

[README](#)
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Yet Another Metagenomic Pipeline (YAMP)

Thanks to the increased cost-effectiveness of high-throughput technologies, the number of studies focusing on microorganisms (bacteria, archaea, microbial eukaryotes, fungi, and viruses) and their connections with human health and diseases has surged, and, consequently, a plethora of approaches and software has been made available for their study, making it difficult to select the best methods and tools.

Here we present Yet Another Metagenomic Pipeline (YAMP) that, starting from the raw sequencing data and having a strong focus on quality control, allows, within hours, the data processing up to the functional annotation (please refer to the YAMP [wiki](#) for more information).

CAMP: Core Analysis Metagenomics Pipeline

A modular metagenomics analysis system for integrated multi-step data exploration



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