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Mitogenome Assembly with GetOrganelle

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Dakota Betz¹

¹ucsd

Rouse Lab



Dakota Betz

ucsd

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Protocol status: Working

We use this protocol and it's working

Created: July 09, 2021

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Protocol Integer ID: 51443

Abstract

Basic instructions for assembling mitogenomes with the GetOrganelle Program



Installation

1

Note

This protocol includes instructions for a conda install of GetOrganelle, which requires that you have Anaconda or Miniconda already installed on your system.

Detailed installation information and additional installation options for various systems can be found here: https://github.com/Kinggerm/GetOrganelle and here:

https://github.com/Kinggerm/GetOrganelle/wikihttps://github.com/Kinggerm/GetOrganell e/wiki

Before installation, activate your python environment (example code below for python3 environment):

Command

conda activate python3

2 Install GetOrganelle with the following code:

Command

conda install -c bioconda getorganelle

You may need to install updates, be sure to say yes (type y and hit enter) when prompted to update any conda-related programs.

```
Downloading and Extracting Packages
libgfortran-5.0.0
           19 KB
                 100%
libcblas-3.9.0
           11 KB
                                    100%
                 99 KB
                                    100%
urllib3-1.26.6
                 mpc-1.1.0
           100 KB
                 100%
gmpy2-2.1.0b5
           190 KB
                 100%
sympy-1.8
           11.2 MB
                 ######################################
                                    100%
pigz-2.6
           88 KB
                 100%
libgfortran5-9.3.0
           1.7 MB
                 100%
           573 KB
mpfr-4.0.2
                                    100%
                 openssl-1.1.1k
           1.9 MB
                                    100%
                 llvm-openmp-11.1.0
           268 KB
                 100%
           1.3 MB
bowtie2-2.4.4
                                    100%
                 ca-certificates-2021
           136 KB
                 ######################################
                                    100%
perl-5.32.1
           13.4 MB
                 100%
libcxx-11.1.0
           1.0 MB
                 100%
lz4-c-1.9.3
           155 KB
                 100%
tbb-2020.2
           132 KB
                 100%
zstd-1.4.9
           460 KB
                 100%
mpmath-1.2.1
           437 KB
                 100%
liblapack-3.9.0
           11 KB
                                    100%
                 164 KB
getorganelle-1.7.5.0
                 100%
idna-2.10
           52 KB
                 100%
cryptography-3.4.7
           763 KB
                 100%
libopenblas-0.3.15
           8.7 MB
                 100%
scipy-1.7.0
           19.8 MB
                 100%
                                    100%
           113.0 MB
blast-2.6.0
                 spades-3.12.0
           12.3 MB
                 100%
libblas-3.9.0
           11 KB
                                    100%
                 certifi-2021.5.30
           141 KB
                 100%
chardet-4.0.0
           204 KB
                                    100%
                 numpy-1.21.0
           5.8 MB
                 100%
gmp-6.2.1
           774 KB
                                    100%
                 requests-2.25.1
           51 KB
                 100%
brotlipy-0.7.0
           357 KB
                 100%
pyopenssl-20.0.1
           48 KB
                 100%
           27 KB
pysocks-1.7.1
                 100%
Preparing transaction:
Verifying transaction: done
Executing transaction: done
```

Printout of successful installation of GetOrganelle (for reference).

Initialization

3 Next we need to download and initialize the database of your preferred organelle genome type.

Options include: embplant_pt, embplant_mt, embplant_nr, fungus_mt, fungus_nr, animal_mt, and/or other_pt

For animal mitochondrial genomes, use animal_mt:



Command

```
get_organelle_config.py --add animal_mt
```

If you would like to download and initialize multiple organelle genome types, you can list them after the **--add** command above, separated by spaces.

```
[(python3) marina@Lamellibrachia ~ % get_organelle_config.py --add animal_mt
Python 3.8.5 | packaged by conda-forge | (default, Aug 29 2020, 01:18:42) [Clang 10.0.1 ]
DEPENDENCIES: Bowtie2 2.4.4; Blast 2.6.0
WORKING DIR: /Users/marina
/Users/marina/miniconda3/envs/python3/bin/get_organelle_config.py --add animal_mt
Existing databases(s):
Downloading animal_mt.fasta
Downloading animal_mt.fasta [========= ] 100%
Downloaded animal_mt.fasta (30285897 bytes), cost 1.82 s
bowtie2-build animal_mt.fasta ... Making - bowtie2 index ...
Making - bowtie2 index finished.
finished
Downloading animal_mt.fasta
Downloading animal_mt.fasta [======== ] 100%
Downloaded animal_mt.fasta (6674918 bytes), cost 0.72 s
makeblastdb animal_mt.fasta ... finished
```

Printout of successful initialization (for reference). Note that working directories will be different on your system.

4 If you have trouble with the above command (connection to GitHub fails) try a few times more, as the problem generally resolves itself. If you're still having trouble, you can download the repository manually, as in Option 2 available here: https://github.com/Kinggerm/GetOrganelle/wiki/Initialization#default-directory