

May 15, 2020 Version 1

## Installation instructions for RNA-seq analysis using a conda environment V.1

DOI

[dx.doi.org/10.17504/protocols.io.bgffjtjn](https://dx.doi.org/10.17504/protocols.io.bgffjtjn)

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**Protocol Citation:** Laise Moraes 2020. Installation instructions for RNA-seq analysis using a conda environment. [protocols.io  
https://dx.doi.org/10.17504/protocols.io.bgffjtjn](https://dx.doi.org/10.17504/protocols.io.bgffjtjn)

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**Protocol status:** Working

**Created:** May 15, 2020

**Last Modified:** May 15, 2020

**Protocol Integer ID:** 37063

### Abstract

The purpose of this protocol is to define and provide instructions for creating a conda environment for phylogenetic analysis

## Installing Miniconda

- 1 Open a terminal (Ctrl + Alt + T).

Create a directory called softwares to your HOME directory, switch to it and then download the 64-bit Python 3 Miniconda installer.

Install Miniconda quietly, accepting defaults.

After installation, remove the Miniconda installer from directory.

Set the Miniconda permanent PATH and update conda packages to the latest compatible version.

### Command

```
mkdir $HOME/softwares
cd $HOME/softwares
wget https://repo.continuum.io/miniconda/Miniconda3-latest-Linux-
x86_64.sh
bash Miniconda3-latest-Linux-x86_64.sh -bfp $HOME/softwares/conda
rm Miniconda3-latest-Linux-x86_64.sh
export
PATH="$HOME/softwares/conda/bin:/usr/local/share/rsi/idl/bin:$PATH"
echo 'export
PATH="$HOME/softwares/conda/bin:/usr/local/share/rsi/idl/bin:$PATH"' >> $HOME/.bashrc
conda update -y -n base conda
```

## Creating conda environment

- 2 Create a YML file to build a phylogenetic analysis environment called rna-seq.

### Command

```
cat > rna-seq.yml <<EOF
name: rna-seq
channels:
- bioconda
dependencies:
- bedtools
- bioconductor-rdavidwebservice
- bowtie2
- bwa
- cutadapt
- fastqc
- igv
- multiqc
- tophat
- trimmomatic
- samtools
- star
- subread
EOF
```

### 3 Create the environment from the phy.yml file

### Command

```
conda env create -f rna-seq.yml
```