

ypkpathway

Ypkpathway is a software tool for the automated planning of metabolic pathway assemblies using the Yeast Pathway Kit (Pereira et. al. 2014).

Installation

Python 2.7 and the python packages **pydna**, **networkx**, **biopython**, and **docutils** are required to run ypkpathway. The easiest and most efficient way to install ypkpathway is by first installing the free Anaconda Scientific Python distribution from Continuum analytics. It is a large download, but it installs cleanly in one folder in the users directory (regardless of the operating system) and is easily removable if necessary. Anaconda is available for Windows, Mac and Linux.

Anaconda installation

Go to the website of Anaconda at <https://store.continuum.io/cshop/anaconda/> (Fig1).

Anaconda Scientific Python Distribution - Google Chrome

https://store.continuum.io/cshop/anaconda/

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Anaconda

Completely free enterprise-ready Python distribution for large-scale data processing, predictive analytics, and scientific computing

- 125+ of the most popular Python packages for science, math, engineering, data analysis
- Completely free - including for commercial use and even redistribution
- Cross platform on Linux, Windows, Mac
- Installs into a single directory and doesn't affect other Python installations on your system. Doesn't require root or local administrator privileges
- Stay up-to-date by easily updating packages from our free, online repository
- Easily switch between Python 2.6, 2.7, 3.3, and experiment with multiple versions of libraries, using our conda package manager and its great support for virtual environments
- Comes with xlwings, openpyxl, xlswriter, xlrd, and xlwt for Excel integration and works well with DataNitro and PyXLL

Why Are We Just Giving This Away?

- We want to ensure that Python, NumPy, SciPy, Pandas, IPython, Matplotlib, Numba, Blaze, Bokeh, and other great Python data analysis tools can be used everywhere.
- We want to make it easier for Python evangelists and teachers to promote the use of Python.
- We want to give back to the Python community that we love being a part of.

Download Anaconda

Full Version is Completely Free



Anaconda

Anaconda Add-Ons

Accelerate	\$129.00	Free Trial
IOPro	\$79.00	Free Trial

Fig 1

Download Anaconda installation file for your operating system and follow the installation instructions. Once Anaconda is installed, find and start the Anaconda Command Prompt (see Fig2)

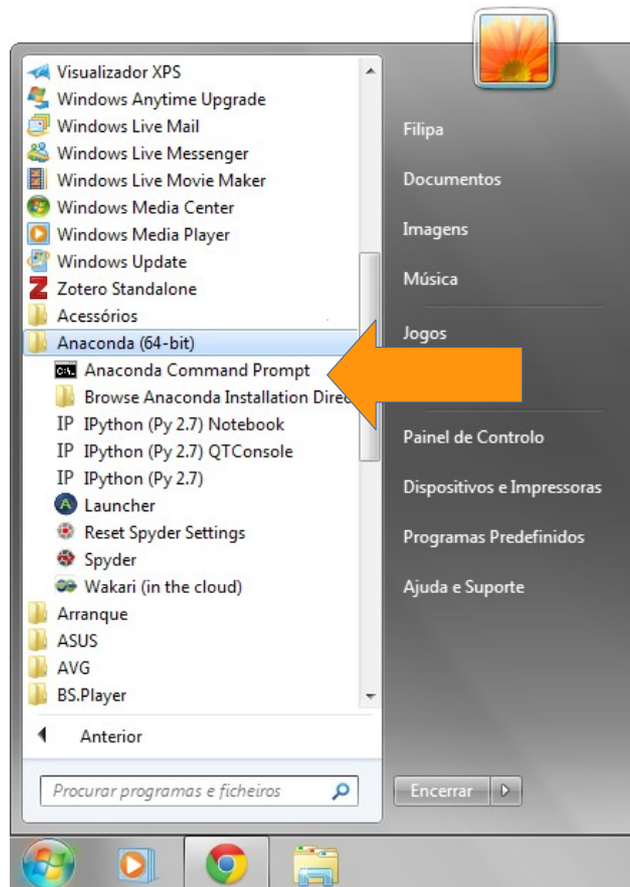


Fig 2 Finding the Anaconda Command Prompt on MS Windows

The Anaconda Command Prompt starts a terminal window. Write “pip install ypkpathway” at the command prompt followed by return (Fig 3). This command will download ypkpathway and install it and all needed dependencies in one go, so make sure you are connected to the internet.

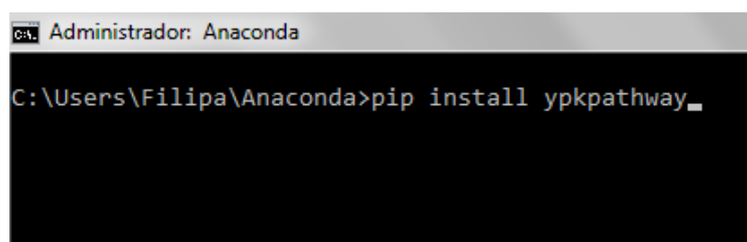


Fig 3

The installation process will generate some text output, important is that it ends with “Successfully

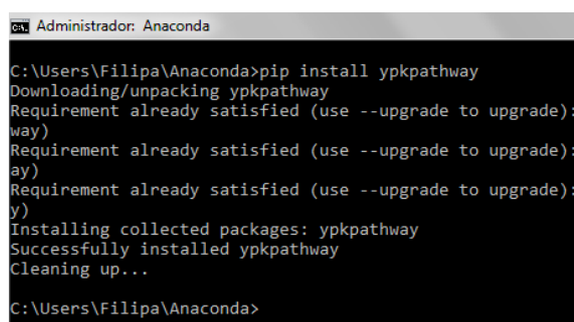
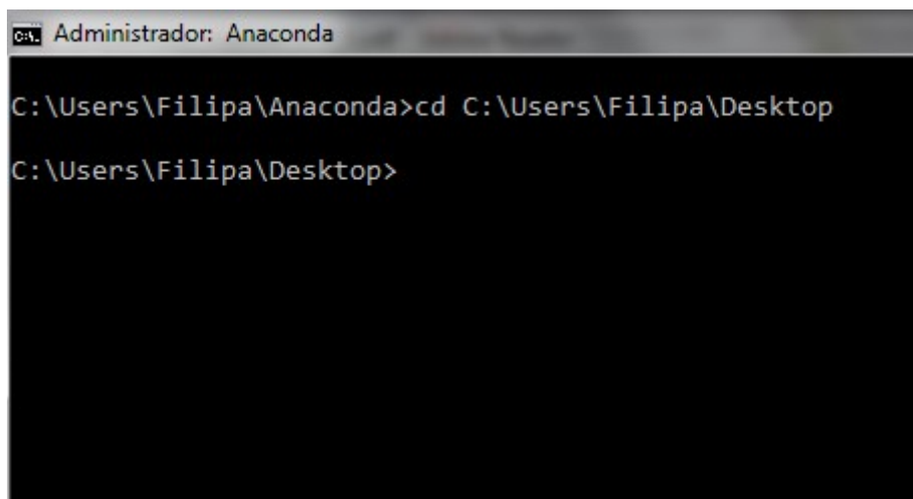


Fig 4

installed ypkpathway Cleaning up...” (Fig 4).

Use

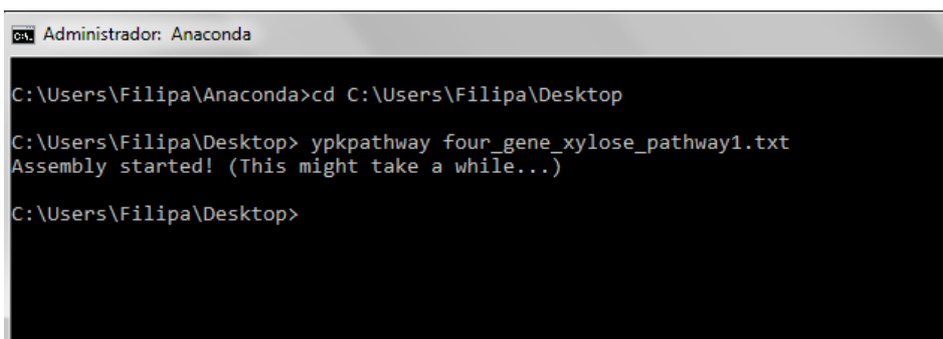
Ypkpathway is meant to be used in the terminal. It is most practical to navigate the terminal to where you have the data file that ypkpathway should process. Issue the commands (“cd” means change directory) in Fig 5 to specify the working directory where the program will read the data files and write the results. In this case we chose the Desktop (cd C:\Users\Filipa\Desktop), the actual name of the folder will be different on another computer.



```
Administrator: Anaconda
C:\Users\Filipa\Anaconda>cd C:\Users\Filipa\Desktop
C:\Users\Filipa\Desktop>
```

Fig 5

The syntax is very simple as ypkpathway takes only one argument which is the name of the data file to be processed, in this case `four_gene_xylose_pathway1.txt` (Fig 6) which is a datafile accompanying this document.



```
Administrator: Anaconda
C:\Users\Filipa\Anaconda>cd C:\Users\Filipa\Desktop
C:\Users\Filipa\Desktop> ypkpathway four_gene_xylose_pathway1.txt
Assembly started! (This might take a while...)
C:\Users\Filipa\Desktop>
```

Fig 7



Fig 6

The data file can have any name as long as it is a text file containing the sequences to be assembled. See next section “Indata” for proper formatting of this file. The result is a new folder created in the same directory called `ypk_assembly` containing the assembly report (Fig 7).

Indata

The data file is simply a list of the TPs and genes that should be assembled in a text format (either FASTA or Genbank). The datafile.txt file (which can have a different name) can have the structure depicted in Fig 8. The sequences in Fig 8 are truncated for clarity and could also be given in Genbank format or a mix of FASTA and Genbank formats. The sequences could be linear fragments (as in the example four_gene_xylose_pathway1.txt file accompanying this document).

```
>TEF1tp
ACAATGC...AAA
>gene1
atgatc...taa
>TDH3tp
ATAAAAA...AAA

>TDH3tp
ATAAAAA...AAA
>gene2
atgcac...tag
>TPI1tp
TGTTTAA...AAA
```

Fig 8 Two sets of three sequences, each forming a tp gene tp cassette. Dots symbolize sequence not shown for clarity.

The Yeast Pathway Kit was designed for the reuse of the genetic parts, especially terminator-promoter and genes cloned in the pYPKa. These plasmids are named pYPKa_Z_XXXN and pYPKa_E_XXXN, where XXXN represent the actual identifier of the tp or gene. Once constructed, terminator-promoter plasmids can be reused for other pathways, in which case they do not need to be constructed again.

Sequences can also be given to the ypkpathway program in the form of the entire pYPKa_Z_XXXN, pYPKa_A_XXXN or pYPKa_E_XXXN sequences, typically generated from a previous assembly experiment. These will be recognized by the ypkpathway algorithm and the assembly report will indicate that these were given and not constructed (Fig 9).

```

>pYPKa_Z_TEF1tp
tcgcgcgttt...ACAATGC...AAA...ctttcgtc

>pYPKa_A_gene1
tcgcgcgttt...atgac...taa...ctttcgtc

>pYPKa_E_TDH3tp
tcgcgcgttt...ATAAAAA...AAA...ctttcgtc

>pYPKa_Z_TDH3tp
tcgcgcgttt...ATAAAAA...AAA...ctttcgtc

>pYPKa_A_gene2
tcgcgcgttt...atgcac...tag...ctttcgtc

>pYPKa_E_TPI1tp
tcgcgcgttt...TGTTAA...AAA...ctttcgtc

```

Fig 9 The pYPKa sequences will not be assembled by the algorithm.

In the same way

sequences can be supplied as pYPK0_tp_gene_tp sequences (Fig 10).

```

>pYPK0_TEF1tp_gene1_TDH3tp
tcgcgcgttt...ACAATGC...AAA...atgac...taa...ATAAAAA...AAA...ctttcgtc

>pYPK0_TDH3tp_gene2_TPI1tp
tcgcgcgttt...ATAAAAA...AAA...atgcac...tag...TGTTAA...AAA...ctttcgtc

```

Fig 10

The ypkpathway algorithm also permits the use of data files with any valid combination of the three kinds of sequences (Fig 11). In the example in Fig 10 two pYPKa sequences were supplied, one for the tp1 and one for the gene1. The tp2 was never cloned before, so it was given as a linear

sequence. The pYPK0_TDH3tp_gene2_TPI1tp vector was made in a previous experiment and was also given.

```
>pYPKa_Z_TEF1tp
tcgcgcggttt...ACAATGC...AAA...ctttcgtc

>pYPKa_A_gene1
tcgcgcggttt...atgatc...taa...ctttcgtc

>TDH3tp
ATAAAAA...AAA

>pYPK0_TDH3tp_gene2_TPI1tp
tcgcgcggttt...ATAAAAA...AAA...atgcac...tag...TGTTTAA...AAA...ctttcgtc
```

Fig

11

Output

The ypkpathway program creates a folder in the current working directory (the directory from which ypkpathway was called). The folder is called “ypk_assembly”. This folder will be overwritten by ypkpathway if it already exists.

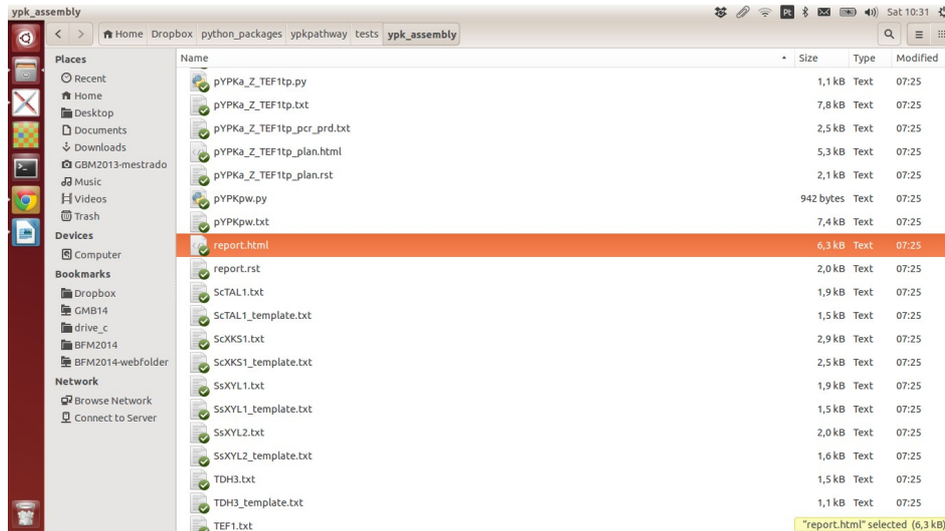


Fig 12

Open this folder (Fig 12) and open the file “report.html” in the web browser (Fig 11). A web page will be created in the browser looking like Fig 13.

Yeast Pathway Kit Assembly Report

Yeast pathway kit assembly 2014-05-03 06:25:54:

1 [pYPK0_TEF1tp_SsXYL1_TDH3tp_SsXYL2_PGItP_ScXKS1_FBA1tp_ScTAL1_PDC1tp_pw \(plan\)](#) 2

List of all [PCR primers](#) needed

[pYPK0_TEF1tp_SsXYL1_TDH3tp \(plan\)](#) 3

- [pYPKa_Z_TEF1tp \(plan\)](#) 4
- [pYPKa_A_SsXYL1 \(plan\)](#)
- [pYPKa_E_TDH3tp \(plan\)](#)

[pYPK0_TDH3tp_SsXYL2_PGItP \(plan\)](#)

- [pYPKa_Z_TDH3tp \(plan\)](#)
- [pYPKa_A_SsXYL2 \(plan\)](#)
- [pYPKa_E_PGItP \(plan\)](#)

[pYPK0_PGItP_ScXKS1_FBA1tp \(plan\)](#)

- [pYPKa_Z_PGItP \(plan\)](#)
- [pYPKa_A_ScXKS1 \(plan\)](#)
- [pYPKa_E_FBA1tp \(plan\)](#)

[pYPK0_FBA1tp_ScTAL1_PDC1tp \(plan\)](#)

- [pYPKa_Z_FBA1tp \(plan\)](#)
- [pYPKa_A_ScTAL1 \(plan\)](#)
- [pYPKa_E_PDC1tp \(plan\)](#)

Fig 13:

Clicking on the first link “pYPK0_TEF1tp_SsXYL1_TDH3tp_SsXYL2_PGItP_ScXKS1_FBA1tp_ScTAL1_PDC1tp_pw” (Fig 13-1) will display the final sequence of the pathway in the browser, a 14800 bp sequence in this case (Fig 14).

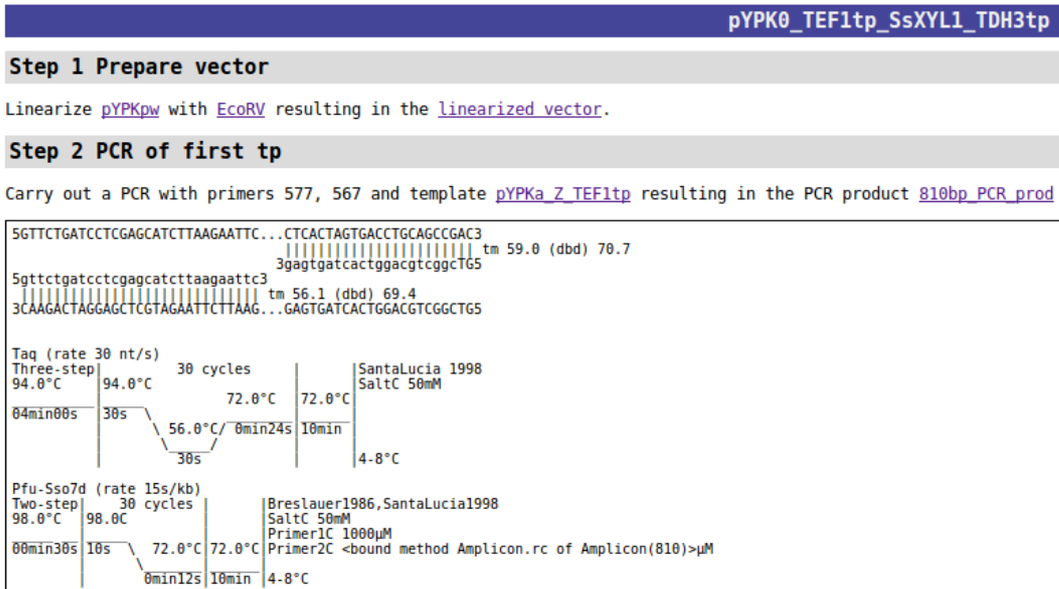


Fig 16

The last “(plan)” link (Fig 12-4) show a plan for the construction of the first pYPKa clone (Fig17). These clones are made from pYPKa vectors linearized with ZraI, AjiI or EcoRV and a linear PCR product.

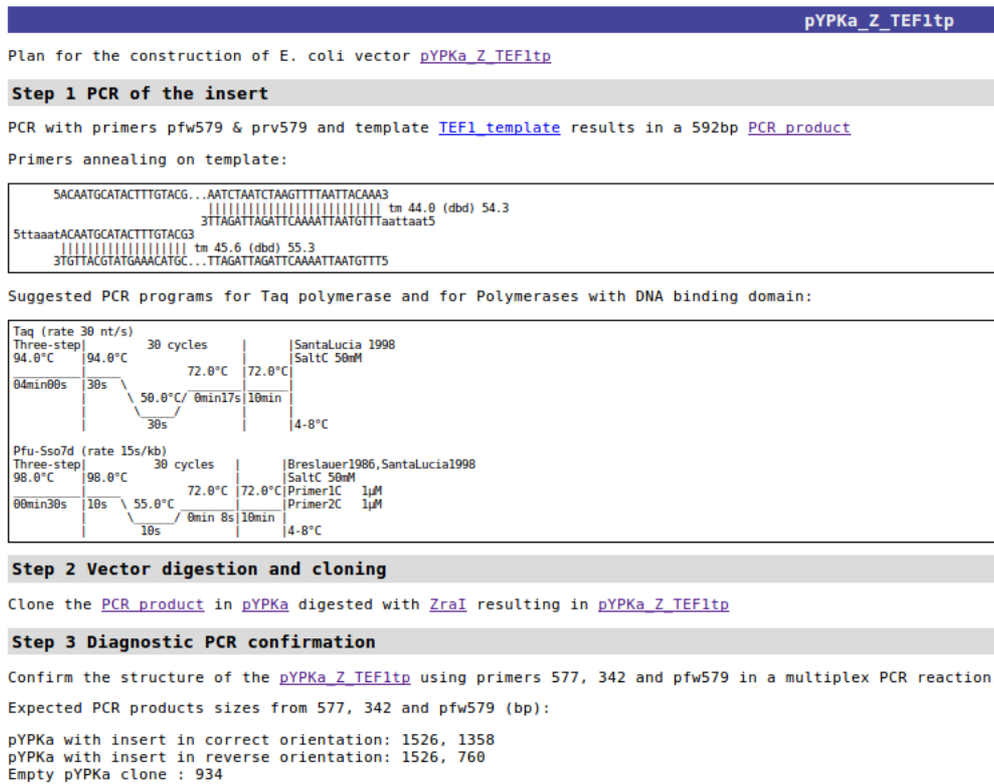


Fig 17

All PCR primers needed for the construction and verification of the pathway can be found under the “[PCR primers](#)” link on the report page (Fig 18). Primers are divided between specific and general

