#### **PHYLIP**

#### Joe Felsenstein

Depts. of Genome Sciences and of Biology, University of Washington

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#### Software for this lab

This lab is intended to introduce the PHYLIP package and a number of major phylogeny methods.

1. You should have downloaded the PHYLIP version 3.695 package from the main PHYLIP web site:

http://evolution.gs.washington.edu/phylip.html
(see the link to "Get me PHYLIP") there. Follow the installation instructions linked to
there.

2. You may want to try a pre-release version of the 4.0 version of PHYLIP instead. You will find it here:

http://evolution.gs.washington.edu/phylip/download/sisg2013/ It and some of the v3.695 programs need you to have a recent version of Oracle Java installed, for the Java front ends. If you have a Windows machine or a Linux system, it can be downloaded and installed free from http://java.com (Mac OS X Java should be good enough).

3. If you have come with a tablet with the iPad or Android operating systems, there is no version of PHYLIP available for that. Instead we will try to give you a URL allowing you to log in to one of our local machines so that, if you have an SSH client, you can open a terminal window and run PHYLIP on our system through that.

#### **PHYLIP**

- Distributed since 1980
- Originally in Pascal, now in C
- Intended to provide "basic transportation"
- Intended to provide a wide variety of methods
- Freely available (unless you try to charge others for it)

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## **Advantages of PHYLIP**

- 1. Free (in the sense of "free beer"), easily obtainable
- 2. Runs on all major platforms
- 3. Very good documentation
- 4. Lots of people around who know how to use it
- 5. Often used in teaching about phylogenies.
- 6. Runs can be automated by using input redirection and command files
- 7. Support for PHYLIP-format files by many other programs such as ClustalW, MacClade and PAUP\*

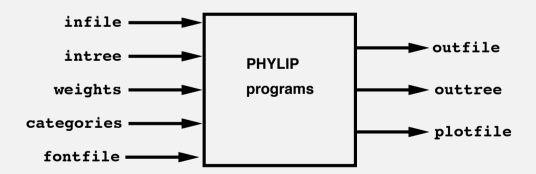
Over 30,000 registered users in over 50 countries including: Fiji, Cuba, Papua New Guinea, Iran, Iceland. Large numbers of users in countries such as India, Brazil, Argentina, Russia, and China where even modest cash prices for software can be a major burden.

# **Disadvantages of PHYLIP**

- 1. Tree search less thorough than some other packages such as PAUP\*.
- 2. Much, much slower than packages such as PAUP\* and RAXML
- 3. Character-mode interface is not mouse/windows GUI
- 4. Manual steps such as renaming file names can be tedious
- 5. Still no: codon model (but coming very soon), Bayesian inference.
- 6. Not as many options available as in other programs
- 7. Cannot read NEXUS standard files

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# **PHYLIP** programs



These are the default file names. If the input files do not exist (or if the output files exist and you choose not to overwrite them), you will be asked for the file name. This is not a bug.

## Input format for PHYLIP (DNA, Interleaved)

7 112 CCAAACCTGT CCCCACCATC TAACACCAAC CCACATATAC AAGCTAAACC AAAAATACCA Bovine CCAAAAAAC ATCCAAACAC CAACCCCAGC CCTTACGCAA TAGCCATACA AAGAATATTA Mouse Gibbon CTATACCCAC CCAACTCGAC CTACACCAAT CCCCACATAG CACACAGACC AACAACCTCC CCCCACCGT CTACACCAGC CAACACCAAC CCCCACCTAC TATACCAACC AATAACCTCT Orang CCCCATTTAT CCATAAAAAC CAACACCAAC CCCCATCTAA CACACAAACT AATGACCCCC Gorilla Chimp Human CCCCACTCAC CCATACAAAC CAACACCACT CTCCACCTAA TATACAAATT AATAACCTCC CCCCAGCCCA ACACCCTTCC ACAAATCCTT AATATACGCA CCATAAATAA CA TCCCACCAAA TCACCCTCCA TCAAATCCAC AAATTACACA ACCATTAACC CA GCACGCCAAG CTCTCTACCA TCAAACGCAC AACTTACACA TACAGAACCA CA ACACCCTAAG CCACCTTCCT CAAAATCCAA AACCCACACA ACCGAAACAA CA ACACCTCAAT CCACCTCCCC CCAAATACAC AATTCACACA AACAATACCA CA ACATCTTGAC TCGCCTCTCT CCAAACACAC AATTCACGCA AACAACGCCA CA ACACCTTAAC TCACCTTCTC CCAAACGCAC AATTCGCACA CACAACGCCA CA

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# Format for trees in tree files (Newick standard)

(Mouse:0.87231,Bovine:0.49807,(Gibbon:0.25930,(Orang:0.24166, (Gorilla:0.12322,(Chimp:0.13846, Human:0.08571):0.06026):0.04405):0.10815):0.39538);

More than such tree can be placed end-to-end in the same tree file.

The Newick standard was defined by an informal standards committee in 1986. It is described on this web page:

http://evolution.gs.washington.edu/phylip/newicktree.html

# **PHYLIP** guide

A useful guide to using PHYLIP with molecular sequences has been produced by Jarno Tuimala. It can be downloaded as a PDF from

http://koti.mbnet.fi/tuimala/oppaat/phylip2.pdf or using the link to it on the main PHYLIP web page.

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# For more information on many other programs

... at my PHYLIP web site there is a master list of over 390 phylogeny programs, with descriptions and links.

To find it simply put the phrase "Phylogeny Programs" into your favorite search engine.

However, it is not really up-to-date. I have had to stop work on it as I have no one to help me on that.

### What to do in the PHYLIP likelihood lab exercise

- Get a DNA or protein sequence data set of aligned sequences. You
  can use one of the ones provided by the course if you wish. They
  are also at
- http://evolution.gs.washington.edu/sisg/2013/data/
- 2. Make a copy the data file as file infile, and then run either Dnaml or Proml, whichever is appropriate. Use the R to do a "Gamma distributed rates" analysis and then the A options to set it to a mean block length of about 3. After you accept the menu settings, you will be asked for a coefficient of variation of rates (you could set this at 2.0) and for the number of rate categories used to approximate the Gamma distribution (about 5-6 would be good).
- 3. Look at the tree by looking at the output file outfile (when you examine that file, you will need to make sure the font is a fixed-width one such as Courier) and also by renaming outtree to intree and then using Drawgram (perhaps with font file font1). You can also try Drawtree. (In using these, when you get a preview of the graph, use the File menu to choose whether you want to change settings. The final plot will be called plotfile.

#### More to do: the PHYLIP distance lab exercise

Use your data set and analyze it by the Neighbor-Joining method:

- 1. Make a copy of your sequences and call that file infile
- 2. Run Dnadist or Protdist, whichever is appropriate.
- 3. The distance matrix is in file outfile
- 4. Rename that infile
- 5. Run Neighbor, using the default options except maybe the outgroup-rooting option.
- 6. The output file outfile will show your tree, and the output tree file treefile has the Newick-format representation of it. Save them by renaming them. When examining the output file, use a constant-width font to avoid distortion of the tree.

### More to do: the PHYLIP bootstrap lab exercise

Use that distance matrix method to do a bootstrap analysis:

- 1. (use Seqboot, then renaming outfile to infile, (You can use 1000 replicates if you have DNA sequences (use menu option R), but don't do 1000 replicates for a protein data set as this will be too slow). When asked for the random number seed, provide any odd number whose last two digits give a remainder of 1 when divided by 4 (for example, they might be 45).
- 2. Use that infile of many data sets as an input for Dnadist or Protdist, using the M (Multiple input data sets) option (with multiple data sets, not weights).
- 3. The multiple distance matrices are now in file outfile. Rename that to infile.
- 4. Now run program Neighbor, making sure to set the multiple data sets option M and provide the number of the bootstrap replicate distance matrices.
- 5. Rename the output file outtree (which will contain multiple bootstrap estimates of the tree) to intree.
- 6. Run program Consense which makes an Extended Majority-Rule Consensus Tree from these trees.
- 7. Look at the consensus tree by examining outfile, or renaming outtree to intree and running either Drawgram or Drawtree.
- 8. The branch lengths of this consensus tree are weird (they reflect levels of bootstrap support rather than amounts of change. Can you figure out a way, using the original sequences and the consensus tree and menu option U (User-defined tree) in the likelihood program, to get more reasonable branch lengths in that tree?