MCB 5472

Intro to Trees

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OLD ASSIGNMENTS Turn your script that calculates the reverse complement of a sequence into a subroutine

```
use warnings;
#modified from Dustin
MAIN: {
        my $seq='';
        my $revcomp='';
        print "Please enter sequence to be reverse complementedn":
        chomp(seq = \diamond);
        $revcomp=&Reverse(&Complement($seq)); #Next line would work just as well
        # $revcomp=Reverse(Complement($seq));
        print "\n".
        print "The reverse complement of \$seq=$seq"." is"." $revcomp\n\n";
exit.
sub Reverse {
        use strict;
        use warnings;
        my string = [0];
        my $rev= reverse $string;
        return $rev;
}
sub Complement{
        # you can use a local variable with the same name as in the main program e.g.
        my $seq= $_[0];
        $seq =~ tr/ATGCYRatcqyr/TACGRYTAGCRY/;
        return ($seq);
        #$seq in the calling program is not impacted.
        #my$comp= $_[0];
        #$comp =~ tr/ATGCYRatcqyr/TACGRYTAGCRY/;
        #return $comp;
L D
```

OLD ASSIGNMENTS

Write a script that takes all files with the extension .fa (containing a single fasta formated sequence) and writes their contents in a single multiple sequence file.

Solution 1:

```
use warnings;
system ("cat *.fa > all_fa_files.faa");
```

OLD ASSIGNMENTS

Solution 2:

use warnings;

open (OUT, ">temp.out") || die "cannot open: \$!";#open a file for output # as the names of the infiles have not been read yet, I assign this a temporary name to be renamed later
#
while(defined(\$file=glob("*.fa"))){ #loop opens one file ending in .fa after the other. stores the name in \$file
 open (IN, "<\$file") || die "cannot open: \$!"; #reports error if file cannot be opened
 #and assigns \$file to IN
 @filename_parts=split(/\./,\$file); # writes first part of filename into \$filename_parts[0]
 \$outname .= "\$filename_parts[0]".'.'; ### concatenated \$filename_part[0] into new outname ###
 #repeats this for every cycle
 while (defined(\$line=<IN>)){print OUT \$line;} #writes contents of \$file line by line into filehandle OUT
} #no more *.fa files in directory
close OUT; #closes filehandle OUT and temp.out
\$outname="\$outname".'fa.multiple';#adds useful extension to outfile
system ("mv temp.out \$outname"); # uses unix command mv (move)to rename temp.out into new \$outname

assignments continued (use class 5 as sample)

Assume that you have the following non-aligned multiple sequence files in a directory:

<u>A.fa</u> : vacuolar/archaeal ATPase catalytic subunits ; <u>B.fa</u> : vacuolar/archaeal ATPase non-catalytic subunits; <u>alpha.fa</u> : F-ATPases non-catalytic subunits, <u>beta.fa</u> : F-ATPases catalytic subunits, <u>F.fa</u> : ATPases involved in the assembly of the besterial flow

<u>F.fa</u>: ATPase involved in the assembly of the bacterial flagella.

Write a perl script that executes muscle or clustalw and

- 1) aligns the sequences within each file
- 2) successively calculates profile alignments between all aligned sequences.

Hints:

```
system (command) ; # executes "command" as if you had typed command in the command line
```

challenge:

Often one wants to build families of homologous proteins extracted from genomes. One way to do so is to find reciprocal best hits.

Tools:

The script *<u>blastall.pl</u>* takes the genomes indicted in the first line and calculates all possible genome against genome searches.

This script <u>simple_rbh_pairs.pl</u> takes two blastall searches (genome A versus genome B) in -m8 format and listing only the top scoring blast hit for each query) and writes the GI numbers of reciprocal best hits into a table.

The script <u>run_pairs.pl</u> runs all possible pairwise extractions of RBHs

Task: write a script that combines the pairwise tables keeping only those families that have a strict reciprocal best blast hit relationship in all genomes.

New assignment

Check chapter U40 in the Unix/Perl primer – sad ⊗

Read chapters P17 and P18 on regular expressions

(these are really useful, if you repeatedly run a program that generates complex output, and you want to extract a single value into a table, e.g. a program that calculates theoretical isoelectric points ...)

New assignment

Write a script that takes a genome (fna file) and calculates the GC content.

Modify the script to count tetra- and penta-nucleotide frequencies.

Modify the script so that it creates a table that gives the GC, tetra- and penta-nucleotide content in a sliding window moving through the genome.

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(For which of these programs, and for which problems might you want to consider, or correct for strand bias?)

Steps of the phylogenetic analysis

Phylogenetic analysis is an inference of evolutionary relationships between organisms. Phylogenetics tries to answer the question "How did groups of organisms come into existence?"

Those relationships are usually represented by tree-like diagrams.

Note: the assumption of a tree-like process of evolution is controversial!



Phylogenetic reconstruction - How

Distance analyses

calculate pairwise distances (different distance measures, correction for multiple hits, correction for codon bias)

make distance matrix (table of pairwise corrected distances)

calculate tree from distance matrix

i) using optimality criterion(e.g.: smallest error between distance matrixand distances in tree, or useii) algorithmic approaches (UPGMA or neighbor joining) B)

Phylogenetic reconstruction - How

Parsimony analyses

find that tree that explains sequence data with minimum number of substitutions

(tree includes hypothesis of sequence at each of the nodes)

Maximum Likelihood analyses

given a model for sequence evolution, find the tree that has the highest probability under this model.

This approach can also be used to successively refine the model.

Bayesian statistics use ML analyses to calculate posterior probabilities for trees, clades and evolutionary parameters. Especially MCMC approaches have become very popular in the last year, because they allow to estimate evolutionary parameters (e.g., which site in a virus protein is under positive selection), without assuming that one actually knows the "true" phylogeny.

What is in a tree?

Trees form molecular data are usually calculated as unrooted trees (at least they should be - if they are not this is usually a mistake).

To root a tree you either can assume a molecular clock (substitutions occur at a constant rate, again this assumption is usually not warranted and needs to be tested),

or you can use an **outgroup** (i.e. something that you know forms the deepest branch). For example, to root a phylogeny of birds, you could use the homologous characters from a reptile as outgroup; to find the root in a tree depicting the relations between different human mitochondria, you could use the mitochondria from chimpanzees or from Neanderthals as an outgroup; to root a phylogeny of alpha hemoglobins you could use a beta hemoglobin sequence, or a myoglobin sequence as outgroup.

Trees have a branching pattern (also called the topology), and branch lengths.

Often the branch lengths are ignored in depicting trees (these trees often are referred to as cladograms - note that cladograms should be considered rooted). You can swap branches attached to a node, and in an unrooted you can depict the tree as rooted in any branch you like without changing the tree.

Test: Which of these trees is different?



More tests <u>here</u>

Terminology

•Branches, splits, bipartitions

In a rooted tree: clades (for urooted trees sometimes the term clann is used)
Mono-, Para-, polyphyletic groups, cladists and a natural taxonomy

The term cladogram refers to a strictly bifurcating diagram, where each clade is defined by a common ancestor that only gives rise to members of this clade. I.e., a clade is **monophyletic** (derived from one ancestor) as opposed to **polyphyletic** (derived from many ancestors). (Note: you do need to know where the root is!)

A clade is recognized and defined by shared derived characters (= synapomorphies). Shared primitive characters (= sympleisiomorphies, aternativie spelling is symplesiomorphies) do not define a clade. (see in class example drawing ala Hennig).

To use these terms you need to have **polarized characters**; for most molecular characters you don't know which state is primitive and which is derived (exceptions:....).

Terminology

Related terms:

autapomorphy = a derived character that is only present in one group; an autapomorphic character does not tell us anything about the relationship of the group that has this character of other groups.

homoplasy = a derived character that was derived twice independently (convergent evolution). *Note that the characters in question might still be homologous (e.g. a position in a sequence alignment, frontlimbs turned into wings in birds and bats).*

paraphyletic = a taxonomic group that is defined by a common ancestor, however, the common ancestor of this group also has decendants that do not belong to this taxonomic group. Many systematists despise paraphyletic groups (and consider them to be polyphyletic). Examples for paraphyletic groups are reptiles and protists. Many consider the archaea to be paraphyletic as well.

holophyletic = same as above, but the common ancestor gave rise only to members of the group.

homology

Two sequences are homologous, if there existed an ancestral molecule in the past that is ancestral to both of the sequences

Types of Homology

Orthologs: "deepest" bifurcation in molecular tree reflects speciation. These are the molecules people interested in the taxonomic classification of organisms want to study.

Paralogs: "deepest" bifurcation in molecular tree reflects gene duplication. The study of paralogs and their distribution in genomes provides clues on the way genomes evolved. Gen and genome duplication have emerged as the most important pathway to molecular innovation, including the evolution of developmental pathways.

Xenologs: gene was obtained by organism through horizontal transfer. The classic example for Xenologs are antibiotic resistance genes, but the history of many other molecules also fits into this category: inteins, selfsplicing introns, transposable elements, ion pumps, other transporters,

Synologs: genes ended up in one organism through fusion of lineages. The paradigm are genes that were transferred into the eukaryotic cell together with the endosymbionts that evolved into mitochondria and plastids (the -logs are often spelled with "ue" like in orthologues)

see Fitch's article in $\underline{\text{TIG 2000}}$ for more discussion.

Trees – what might they mean?

Calculating a tree is comparatively easy, figuring out what it might mean is much more difficult.

If this is the probable organismal tree:



what could be the reason for obtaining this gene tree:



lack of resolution



e.g., 60% bootstrap support for bipartition (AD)(CB)

long branch attraction artifact



e.g., 100% bootstrap support for bipartition (AD)(CB)

What could you do to investigate if this is a possible explanation? use only slow positions, use an algorithm that corrects for ASRV

Gene transfer

Organismal tree:



molecular tree:





Gene duplication and gene transfer are equivalent explanations.



Horizontal or lateral Gene

Ancient duplication followed by gene loss

Note that scenario B involves many more individual events than A

1 HGT with orthologous replacement gene duplication followed by
 independent gene loss events

What is it good for?

Gene duplication events can provide an outgroup that allows rooting a molecular phylogeny.

Most famously this principle was applied in case of the tree of life – the only outgroup available in this case are ancient paralogs (see

http://gogarten.uconn.edu/cvs/Publ_Pres.htm for more info).

However, the same principle also is applicable to any group of organisms, where a duplication preceded the radiation (<u>example</u>).

Lineage specific duplications also provide insights into which traits were important during evolution of a lineage.

Function, ortho- and paralogy

molecular tree:



The presence of the duplication is a taxonomic character (shared derived character in species B C D).

The phylogeny suggests that seq' and seq have similar function, and that this function was important in the evolution of the clade BCD.

seq' in B and seq' in C and D are orthologs and probably have the same function, whereas seq and seq' in BCD probably have different function (the difference might be in subfunctionalization of functions that seq had in A. – e.g. organ specific expression)





SSU-rRNA Tree of Life

Ochiai, K., Yamanaka, T., Kimura, K., and Sawada, O. (1959) Inheritance of drug resistance (and its tranfer) between Shigella strains and Between Shigella and E.coli strains. Hihon Iji Shimpor 1861: 34 (in Japanese)

Gray GS, Fitch WM (1983):

Evolution of antibiotic resistance genes: the DNA sequence of a kanamycin resistance gene from *Staphylococcus aureus*. *Mol Biol Evol 1983, 1(1):57-66.*

Sorin Sonea (1988): The global organism: A new view of bacteria. The Sciences, 28:38-45.



Robert Natkin, Tiepolo Turn, 1984





BioSystems 31 (1993) 111-119

Bio Systems

Horizontal transfer of ATPase genes — the tree of life becomes a net of life

Elena Hilario, Johann Peter Gogarten*

Department of Molecular and Cell Biology, University of Connecticut, 75 North Eagleville Rd., Storrs, CT 06269-3044, USA

1995:

GOGARTEN, J.P. (1995): The early evolution of cellular life, Trends in Ecology and Evolution, 10, 147-151.



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Genome Data Shake Tree of Life

RESEARCH NEWS

New genome sequences are mystifying evolutionary biologists by revealing unexpected connections between microbes thought to have diverged hundreds of millions of years ago



Science, 280, p.672ff (1998)

rRNA-based tree of life (top). One puts Aquifex close to archaea (left); another splits archaea (right).

31 orthologous protein sequences present in 191 genomes each, concatenated, 8,089 sites

Ciccarelli *et al.* 2006 Toward automatic reconstruction of a highly resolved tree of life.



*Dagan and Martin, Genome Biology, 2006

Tree, Web, or Coral of Life?



Charles Darwin Photo by J. Cameron, 1869

"The tree of life should perhaps be called the coral of life, base of branches dead"

The tree of life Show perhaps he called the conte & life, base of hander dead, to that pepagers count

Page B26 from Charles Darwin's (1809-1882) notebook (1837/38)

Which Type of Coral ?



Darwin's Coral a Red Algae?

(Bossea orbignyana)



According to **Horst Bredekamp**, parts of the diagram in Darwin's origin of species (center) may reflect the branching properties of a specimen Darwin collected himself.

From Florian Maderspacher: "The captivating coral--the origins of early evolutionary imagery." Current Biology 16: R476-8 2006 "As buds give rise by growth to fresh buds, and these, if vigorous, branch out and overtop on all sides many a feebler branch, so by generation I believe it has been with the great Tree of Life, which fills with its dead and broken branches the crust of the earth, and covers the surface with its everbranching and beautiful ramifications."



Charles Darwin in "On the Origin of Species by Means of Natural Selection or the Preservation of Favoured Races in the Struggle for Life", p 162 ff, London, John Murray, 1859

THE POTATO OF LIFE

Successive independent crystallization of the three domains of life from a population of pre-cells

Glansdorff N, Xu Y, Labedan B: *The Last Universal Common Ancestor: emergence, constitution and genetic legacy of an elusive forerunner. Biol Direct* 2008, **3**:29.

Kandler O (1994) *The early diversification of life*. In *Early Life on Earth, Nobel Symposium*, vol. 84, pp. 152-509: Columbia Univ. Press.

Woese CR (2002) On the evolution of cells. PNAS 99: 8742-7



The Coral of Life (Darwin)



Coalescence – the process of tracing lineages backwards in time to their common ancestors. Every two extant lineages coalesce to their most recent common ancestor. Eventually, all lineages coalesce to the cenancestor.



Illustration is from J. Felsenstein, "Inferring Phylogenies", Sinauer, 2003

Simulations of organismal evolution by coalescence



Y chromosome Adam

Lived approximately 50,000 years ago

Thomson, R. *et al.* (2000) *Proc Natl Acad Sci* U S A 97, 7360-5

Underhill, P.A. *et al.* (2000) *Nat Genet* 26, 358-61



Mitochondrial Eve

Lived 166,000-249,000 years ago

Cann, R.L. *et al.* (1987) *Nature* 325, 31-6

Vigilant, L. *et al.* (1991) *Science* 253, 1503-7

Albrecht Dürer, The Fall of Man, 1504

Adam and Eve never met ⊗ The same is true for ancestral rRNAs, EF, ATPases!







Lineages Through Time Plot



red: molecular lineages (with gene transfer)



Bacterial 16SrRNA based phylogeny (from P. D. Schloss and J. Handelsman, Microbiology and Molecular Biology Reviews, December 2004.) The deviation from the "long branches at the base" pattern could be due to

- under sampling
- an actual radiation
 - due to an invention that was not transferred
 - following a mass extinction



Sequence editors and viewers

Jalview <u>Homepage</u>, <u>Description</u> Jalview as <u>Java Web Start Application</u> (other JAVA applications are <u>here</u>)

Jalview is easy to install and run. Test file is <u>here</u> (ATPase subunits) (Intro to ATPases: 1bmf in spdbv) (gif of rotation <u>here</u>, movies of the rotation are <u>here</u> and <u>here</u>) (Load all.txt into Jalview, colo**u**r options,

mouse use,

PID tree,

Principle component analysis -> sequence space)

More on sequence space <u>here</u>