Homology, Orthology, and Trees

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Outline

• Systematics as a unifying principle

• Basics of phylogenetic trees

• Homology, orthology, paralogy, xenology…

• Inference of trees and modern phylogenetics
Part I - Systematics

• Biology: *why are there so many things?*

Systematics

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• Diversity is a fundamental fact of biology
  – It is created by a process: Evolution
Systematics

• Biology: *why are there so many things?*

• Diversity is a fundamental fact of biology
  – It is created by a process: Evolution

• All organisms are the way they are because they evolved to be that way

Systematics

• Biology: *why are there so many things?*

• Diversity is a fundamental fact of biology
  – It is created by a process: Evolution

• “Nothing makes sense except in the light of evolution” – Theodosius Dobzhansky
Systematics

- Diversity is a challenge and an opportunity

- All life shares a common origin
  - Any organism can be used to understand any other organism
Systematics

- Diversity is a challenge and an opportunity

- All life shares a common origin
  - Any organism can be used to understand any other organism

- But life forms are radically different
  - Evolution is the key to comparison

~ 6 orders of magnitude difference in genome size across organisms!!
Systematics

• In order to compare organisms, you must *systematize* (group) them.
  – Same goes for parts of organisms.

• Modern systematics uses phylogenetic trees

Basics of Phylogenetics

• Only figure in Darwin’s “Origin of Species”
Basics of Phylogenetics

• Trees show the relationship between pattern and process
Basics of Phylogenetics

• Phylogenetic systematics (cladistics)
  – Organisms should be grouped by phylogenetic relationships

• Key terms:
  – Clade
  – Monophyly
  – Paraphyly

Willi Hennig (1913 – 1976)

Basics of Phylogenetics

• Clade: an ancestor and all of its descendants
Basics of Phylogenetics

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Basics of Phylogenetics

• Clade: an ancestor and all of its descendants

Not A Clade!!
Basics of Phylogenetics

• Monophyletic group: organisms in a clade
Basics of Phylogenetics

- Monophyletic group: organisms in a clade

Basics of Phylogenetics

- Monophyletic group: organisms in a clade

- A group is *not* monophyletic if their most recent common ancestor has descendants that are not in the group
Basics of Phylogenetics

- Tree nomenclature: nodes
Basics of Phylogenetics

- Tree nomenclature: nodes

- Node

- Leaf or Tip

- Root
Basics of Phylogenetics

• Tree nomenclature: branches
  – Measures of evolutionary rate

Branch or Edge

Basics of Phylogenetics

• Tree nomenclature: branches
  – Measures of evolutionary rate

Branch or Edge

Nothing!!
Basics of Phylogenetics

- Topology

Any node can be rotated without changing topology
Basics of Phylogenetics

• The tree can be unrooted without changing topology

```
C       A
   \    /  \\
  B     D
```

Basics of Phylogenetics

• The tree can be unrooted without changing topology

```
C       A
   \    /  \\
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```
Basics of Phylogenetics

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Basics of Phylogenetics

• Interpreting trees
  – Trees tell us the relative relatedness of leaf nodes
Basics of Phylogenetics

• Interpreting trees
  – Trees tell us the relative relatedness of leaf nodes

• Common misperceptions:
  – D is not the “ancestor” of any other leaf and is not necessarily an older lineage
  – The tree does not tell us that these tips are “related” (all organisms are related)

Part I - Summary

• Phylogenetics gives us a way to organize biological diversity in a rational way

• Trees are powerful representations of the evolutionary process

• Trees hold two kinds of information:
  – Hierarchical relationships
  – Evolutionary rate
Part II – The comparative method

Homology

• When comparing parts of organisms, you need a criterion of “sameness”
  – Evolutionary “sameness” is called “homology”

“Homologue…The same organ in different animals under every variety of form and function…Analogue…A part or organ in one animal which has the same function as another part or organ in a different animal”
  - Richard Owen (1843)
Homology

• Evolutionary or phylogenetic homology
  – Organs (or genes, or…) in two or more species that
    are similar due to common descent
  – I.e. they are descended from a similar organ in their
    most recent common ancestor

• Note
  – This means homology is binary
  – No such thing as % homology
Homology

This is trickier than it sounds!!

Organs are not monolithic entities!!

Bat wings and bird wings are homologous as vertebrate forelimbs

But they are analogous as wings

askabiologist.asu.edu/human-bird-and-bat-bone-comparison

Homology

• Homology also applies to genes
  – How can we tell whether genes are homologous?
Homology

• Homology also applies to genes
  – How can we tell whether genes are homologous?

• Sequence matching scores derived from alignment
  – Null distributions of scores are easily derivable
  – Sequence space is HUGE!
    • Non-homologous gene scores are not distinguishable from random

Homology

• Homologous gene families are very large

• How do we find one-to-one correspondence across species?
Orthology

Walter Fitch

1929 - 2011

What is an Ortholog?

• Need a basis for comparing genes across species
  – Orthology is used nearly universally

• “Same gene in a different species”

• “Homologous genes are related by speciation, whereas paralogs are related by duplication”
What is an Ortholog?

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What is an Ortholog?

“Two genes whose common ancestor resides at a Y junction (speciation) are orthologous. Two genes whose common ancestor resides at a horizontal bar junction (gene duplications) are paralogous.” – Walter Fitch (2000) *Trends in Genetics*
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Deceptive simplicity:

Orthology is a pairwise relationship

It is not transitive

Co-orthology is often abstracted from, with confusing results

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What is an Orthologous Group?
What is an Orthologous Group?

Part II - Summary

• Homology and Orthology give us a way to compare genes in different species
  – Homology contains whole families
  – Orthology is a direct comparison, usually denoting more functional similarity

• Both concepts are potentially slippery
Part III – Inferring Phylogenies

Inferring Phylogenies

• A doctor’s girlfriend accuses him of injecting her with HIV. He said it was vitamin B12.
  — Who’s right?

• Phylogenetics to the rescue!!
Inferring Phylogenies

Gravitropism defects

Significantly overlapping sets of orthologs

McGary et al. (2013)

Waardenburg syndrome
Inferring Phylogenies

How do we infer relatedness between genes?

Algorithms:

1. Random starting tree
2. Measure fit of data to tree under optimality criterion
3. Choose another tree
4. Iterate (for how long?)
### Number of Possible Trees

<table>
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<tr>
<th>Number of Taxa</th>
<th>Number of unrooted trees</th>
<th>Number of rooted trees</th>
</tr>
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<td>3</td>
</tr>
<tr>
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<td>5.86E+096</td>
</tr>
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<td>70</td>
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</tr>
<tr>
<td>80</td>
<td>2.18E+137</td>
<td>3.43E+139</td>
</tr>
</tbody>
</table>

For comparison the universe contains only about $10^{89}$ protons. (http://www.pagines.ma1.upc.edu/~casanellas/eaca/tree_number.html)

### Inferring Phylogenies

#### Algorithms:
- Random starting tree
- Measure fit of data to tree under optimality criterion
- Choose another tree

#### Heuristic Search:
Search a sub-space of trees with a well-defined stopping criterion
Inferring Phylogenies

- Optimality criteria for inferring trees
  - Pairwise distance methods
  - Maximum parsimony
  - Likelihood/Bayesian methods

Inferring Phylogenies

- Phylogenies are based on alignments
  - Taxa are represented row-wise
  - Columns are sites in the genome
  - Can be nucleotides or amino acids
Parsimony Score

Downpass (postorder traversal) Length = 4

Figure courtesy of David Hillis
Parsimony Score

Figure courtesy of David Hillis
Parsimony Score

Figure courtesy of David Hillis

Parsimony Score

Figure courtesy of David Hillis
Parametric Methods

- Maximum likelihood methods find the tree that maximizes the probability of a model
  \[ \text{arg max } P(M | T) \]
- Bayesian methods calculate the probability of a tree given a model
  \[ P(T | M) \]
  Bayes’ theorem:
  \[ P(T | M) = \frac{P(M | T) P(T)}{P(M)} \]

Models of Evolution

- Parametric criteria are evaluated analogously to parsimony
  - One tree is tried at a time!!
Performance

• Parametric methods (likelihood, Bayesian) perform best except in cases of egregious model violation
  — Con: they are much slower

• Distance methods are the norm in prepackaged software. Parsimony still used widely
  — Con: both are statistically inconsistent when internal branch lengths get longer

Part III - Summary

• Phylogenetics is a powerful tool for
  — Bioinformatics
  — Evolutionary biology
  — Virology and medicine

• Numerous methods exist
  — Parametric methods perform the best but are slower
Parametric Methods

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Models of Evolution

- Continuous time Markov models
  \[
P(t) = \begin{pmatrix}
p_{AA}(t) & p_{GA}(t) & p_{CA}(t) & p_{TA}(t) \\
p_{AG}(t) & p_{GG}(t) & p_{CG}(t) & p_{TG}(t) \\
p_{AC}(t) & p_{GC}(t) & p_{CC}(t) & p_{TC}(t) \\
p_{AT}(t) & p_{GT}(t) & p_{CT}(t) & p_{TT}(t) \\
\end{pmatrix}
\]
Models of Evolution

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Probability of changing from cytosine to thymine
In time \( t \) along a branch

Wait times are exponentially functions
If transition probabilities are equal (\( =.25 \)),
all probabilities approach .25