Exploratory Activities for Understanding Evolutionary Relationships Depicted by Phylogenetic Trees: United but Diverse

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ABSTRACT

Evolution explains both the unity and the diversity of all organisms, and developing students’ ability to represent and communicate evolutionary relationships is an important component of a complete biology education. We present a series of student-centered, exploratory activities to help students develop their tree-thinking skills. In these activities, students use complementary phenotypic and molecular data to explore how to build phylogenetic trees and interpret the evolutionary relationships they represent. This learning module is designed to engage students in the process of science, provide them with active learning experiences using online bioinformatics tools, and foster their appreciation for the evolutionary connections across the tree of life.

Key Words: tree-thinking; cladogram; DNA sequences; evolution; bioinformatics; inquiry.

Introduction

Understanding the evolutionary relationships among organisms is a central goal of biology (Darwin, 1859; Dobzhansky, 1973; Schluter, 2000; Grant & Grant, 2011) and a key component of a complete biology education (Baum et al., 2005; Gregory, 2008; Meisel, 2010; Young et al., 2013). Phylogenetic trees are one of the most important tools that evolutionary biologists use to depict the hypothesized relationships among a set of species. Because phylogenies illustrate the key evolutionary concept that all living organisms are connected through common ancestry, understanding phylogenies is critical for understanding evolution (Baum & Offner, 2008).

Numerous studies have shown that students have difficulty interpreting phylogenetic trees and understanding the evolutionary relationships they represent (Meir et al., 2007; Gregory, 2008; Halverson, 2011; Catley et al., 2013; Dees et al., 2014; Blacquiere & Hoese, 2016; Kummer et al., 2016). For example, many students misinterpret phylogenetic trees by reading the order of species along the tips (usually from left to right) to infer relatedness, and therefore make the common mistake of assuming that “primitive” species on the left give rise to “advanced” species on the right (Meir et al., 2007; Omland et al., 2008; Meisel, 2010). Similarly, many students get distracted by the shape of the tree and erroneously assume that species that appear in close proximity at the tips of the tree are more closely related than those further apart (Baum et al., 2005; Meir et al., 2007; Baum & Offner, 2008). The correct way to interpret a phylogenetic tree is as a set of nested hierarchies: trees can be rotated around any node and still depict the same evolutionary relationships. Teaching students to focus on the relative branching pattern of the tree helps emphasize that evolution is indeed a branching process, and that there is no single, linear narrative to describe the divergence of species (Baum et al., 2005; Meisel, 2010).

Science educators have argued that developing students’ tree-thinking skills should be a priority for biology curricula (Baum et al., 2005; Catley, 2006; Meisel, 2010), and more educational resources are needed to train students to accurately read phylogenetic trees and understand what they communicate (O’Hara, 1997; Baum et al., 2005; Meisel, 2010; Halverson, 2011). Additionally, research indicates that student performance and conceptual understanding increases in classes that incorporate inquiry-based instruction (Minner et al., 2005; Freeman et al., 2014). In particular, the integration of web-based resources allows students to gather up-to-date information in virtually any subject area and become actively engaged in the learning process through exploration (Kuzoff et al., 2009; Satterthwait, 2010). Particularly among contemporary students who have grown up in a technology-driven world, the use of online resources has the potential to enhance classroom activities and invigorate science education (Mistler-Jackson & Songer, 2000; Jang, 2009; Kuzoff et al., 2009). We capitalized on the wealth of online bioinformatics resources to develop a series of inquiry-based activities aimed at improving students’ tree-thinking skills. The learning module described here will help students recognize the evolutionary relationships among organisms, understand how these relationships are determined, and appreciate that phylogenetic trees are hypotheses that are subject to revision following new discoveries and analyses.
Disciplinary Core Ideas that reflect unifying principles in life sciences, including structure and function, variation of traits, evidence of common ancestry and diversity, natural selection, and adaptation. These core ideas are essential for understanding the central organizing principle that evolution explains both the unity and diversity of life on Earth. We list the specific student activities in the module that are aligned with each NGSS dimension in Table 1.

Details of the Module

| NGSS Dimension | NGSS Topics & Codes | Student Activities |
|----------------|---------------------|--------------------|
| Science and Engineering Practices | Developing and using models, Analyzing and interpreting data, Constructing explanations, Obtaining, evaluating, and communicating information | Construct phylogenetic trees as a visual representation of the evolutionary relationships among organisms. Search online bioinformatic resources (NCBI website) to collect genetic data. Use online phylogenetics software (Phylogeny.fr) to align DNA sequences and construct molecular phylogenies. Make a prediction about the evolutionary relationships among present-day animals, and then test it using genetic data and phylogenetic analyses. Evaluate the impact of new data by adding a “Discovery” species to the phylogenetic tree. Compare the advantages of different types of data (morphological vs. molecular) for determining the evolutionary relationships among organisms. |
| Crosscutting Concepts | Patterns, Systems and system models, Structure and function, Stability and change | Group organisms based on their shared characteristics. Construct phylogenetic trees to model evolutionary relationships. Explain the relationship between DNA and protein production. Compare if/how the phylogenetic tree changes with the addition of new data (the “Discovery” species). |
| Disciplinary Core Ideas | Structure and function (LS1.A), Variation of traits (LS3.B), Evidence of common ancestry and diversity (LS4.A), Natural selection (LS4.B), Adaptation (LS4.C) | Explain the relationship between DNA and protein production for the hemoglobin molecule used in the phylogenetic analyses. Evaluate the variation in nucleotide sequences across the different animals on the Alignment page of Phylogeny.fr. Observe that the nucleotide sequences for the hemoglobin gene vary among species, but that there are also many overlaps. Identify the shared, derived characteristics among a group of organisms that make them adapted to their particular environments. Consider how similar environments and lifestyles may lead to the evolution of similar phenotypic characteristics among distantly related species through the process of convergent evolution. |
relationships among organisms, and that new evidence continues to improve our understanding of the tree of life.

Overview

This module was designed as a story in which students are applying for an internship at a local college. Stories help capture students’ attention and keep them engaged in activities because they appeal to the human desire for goals (Wilson, 2002; Schultheis & Kjelvik, 2015). The module is presented in two sections: (1) a “Diagnostic Assessment” in which students investigate the relationships among fictitious alien species using solely morphological data; and (2) an “Application” in which students use online resources to gather real DNA sequences on present-day animals, construct a phylogenetic tree based on these genetic data, and then revise their tree after the addition of a new species. An educator’s guide with step-by-step instructions for the entire module (Supplemental File 1), ready-made student worksheets and activity cards (Supplemental Files 2–4), handouts for students on how to use the online resources (Supplemental Files 5 and 6), and a complete educator’s key (Supplemental File 7) are all provided as Supplemental Material with the online version of this article.

Section 1: Comparing Shared Characteristics among Fictitious Species (Days 1–2)

In the “Diagnostic Assessment” section, students compare the characteristics (i.e., phenotypes) among nine fictitious alien species using a character matrix, group the species on the basis of their shared characteristics, and then build a phylogenetic tree based on these groupings (Figure 1). The tree contains multiple gains and losses of traits across two major lineages, which are designed to challenge students’ misconceptions that evolution is a linear, progressive process toward increasing complexity. This activity is designed to help students understand how shared, derived characteristics are used to infer evolutionary relationships, and that, in general, species with recent common ancestry will have more phenotypic characteristics in common.

Figure 1. Alien species used in the “Diagnostic Assessment” section of the module. Students are asked to group species on the basis of shared characteristics to infer their evolutionary relationships.

Section 2: Using Online Tools to Construct a Real Phylogenetic Tree (Days 2–3)

In the “Application” section, students apply the tree-thinking skills they developed in the alien species activity to analyze the evolutionary relationships among 10 present-day animals: nine mammals and a bird (Figure 2). Students are asked to construct a phylogenetic tree for the 10 species based on their knowledge of shared phenotypic characteristics. Students then search the National Center for Biotechnology Information website (https://www.ncbi.nlm.nih.gov) to collect DNA sequences for the hemoglobin alpha protein for all 10 species, and use these molecular data to construct a phylogenetic tree with the web-based phylogenetic alignment and analysis software Phylogeny.fr (http://www.phylogeny.fr; Dereeper et al., 2008). Allowing students to explore computational resources used by scientists to collect genetic data, align DNA sequences, and construct molecular phylogenies helps them appreciate the importance of computational technologies for evolutionary analyses and scientific research.

Lastly, to mirror the discovery of a new species or the incorporation of new data, students are asked to select one of six possible “discovery” mammals to add to their phylogenetic tree: bat, platypus, hedgehog, tarsier, walrus, or whale. These organisms were intentionally selected to challenge students’ understanding of evolutionary relationships among the animals used in the “Application” section of the module. The relative branching pattern was established using the online phylogenetic software Phylogeny.fr, based on the DNA sequences for the hemoglobin alpha protein. The original 10 species are indicated in black, and the addition of two “discovery” species is illustrated in red. All images are from Wikimedia Commons. Photo credits (top to bottom): Hasitha Tudugalle, Anbeast25, mtoz, Hans Hillewaert, George Shuklin, CK Subramanya.

Figure 2. Phylogenetic tree depicting the evolutionary relationships among the animals used in the “Application” section of the module. The relative branching pattern was established using the online phylogenetic software Phylogeny.fr, based on the DNA sequences for the hemoglobin alpha protein. The original 10 species are indicated in black, and the addition of two “discovery” species is illustrated in red. All images are from Wikimedia Commons. Photo credits (top to bottom): Hasitha Tudugalle, Anbeast25, mtoz, Hans Hillewaert, George Shuklin, CK Subramanya.
phylogenies after the inclusion of new data led to fruitful discussions processed with modern computers. Finally, the process of revising the data, and for the speed at which large amounts of information can be appreciated for scientists who regularly analyze large-scale molecular exceptions. Students also found it exciting to use the online resources on the tree revealed common misconceptions about phylogenies, able to predict and test their hypotheses about the evolutionary relationships for High School Students Supplements to S.D. • Supplemental File 3: Card Set A – Alien Species • Supplemental File 4: Card Set B – Present-Day Animals (including the six Discovery Species) • Supplemental File 5: Resource no. 1 – How to Use NCBI • Supplemental File 6: Resource no. 2 – How to Use Phylogeny.fr • Supplemental File 7: Educator’s Key to Student Handout • Supplemental File 8: DNA Sequences for 10 Original Species (optional handout for lower level) • Supplemental File 9: DNA Sequences for 6 Discovery Species (optional handout for lower level) **Acknowledgments**

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