Topological Trait Estimation for Phylogenetic Reconstruction of Fossil Evolution

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1 Introduction

Topological methods have emerged as powerful tools in data analysis, particularly for complex, high-dimensional datasets such as those found in paleontology. The study of dinosaur evolution, traditionally reliant on morphological analysis of fossilized bones, is an area ripe for innovation through these topological techniques. Figure 1 shows how a dinosaur fossil can be scanned and turned into a 3D image. Estimating morphological traits from 3D scans of bones can provide insights into the evolutionary relationships between different species and offer a clearer picture of their lineage and adaptations over time.

Current approaches to this problem involve manually entering and comparing the morphological data, a process that



Figure 1: A 3D scan of the left maxilla of an Edmontosaurus.

is not only time-consuming but also prone to human error and subjectivity [1]. One computational approach, topological data analysis (TDA), offers a mathematical framework that captures the intrinsic geometric and topological properties of shapes, allowing for a more robust and efficient method of trait estimation. Unlike traditional methods, TDA can handle variations in shape, scale, and orientation, which enables more accurate comparisons of

complex structures [6].

In analyzing the traits of dinosaur bones, TDA would allow researchers to quantify differences and similarities by representing the shapes as high-dimensional data points. By treating these shapes as topological spaces, it becomes possible to construct a phylogenetic tree that reflects the true evolutionary history of a given species with greater precision and less manual intervention [5]. A phylogenetic tree is a diagram that displays the lines of evolutionary descent of different species. By using the traits found on certain dinosaur fossils, I will draw my own tree with good resolution by identifying key morphological features that suggest specific evolutionary branches or relationships. This is the area I wish to further study, particularly for its potential to enhance our understanding of evolutionary relationships.

2 Background

Recent studies have demonstrated the power of TDA in fields such as neuroscience and anatomy. Notably, Bendich et al. [2] utilized TDA to analyze brain artery structures, revealing correlations between age, sex, and the bending of arteries that were not present in previous studies. In Figure 2, we see how an artery is constructed and then studied using various techniques. TDA methods provide new insights into complex data by capturing changes in the connectedness of structures and enabling the analysis of high-dimensional characteristics, such as shape and form, often overlooked by traditional metrics.

With advancements in 3D imaging and data acquisition technologies, applying TDA to 3D data feature simplification has gained significant traction in recent years. A study pro-

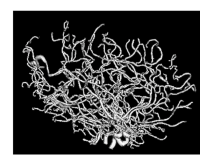


Figure 2: By using tools from TDA, [2] uncovered previously unknown traits of the complex branching structure of arteries, as seen here

duced by Ou Li [4] introduced a method that leverages TDA for segmenting and simplifying 3D point cloud data, a representation commonly only obtained through low-cost 3D sensors. The study demonstrated that TDA can effectively partition complex, disorganized 3D data into subsets with consistent characteristics, using various techniques that simplify the data structure while retaining critical geometric information [4].

Dornbusch et al. [3], developed a method that utilizes 3D point clouds to extract and parameterize morphological traits of plant organs, which are then used to create a comprehensive architectural model of the plant. Fitting these 3D point clouds into a structural model, the method enables a precise portrayal of the leaf and stem shape and their spatial orientations [3]. By applying this technique to 3D scans of fossils, one can extract and parameterize various morphological traits, such as bone structure and dimensions, with high accuracy. This detailed characterization allows for reconstructing more exact evolutionary trees and a better understanding of the anatomical variations among different species. Integrating this approach will significantly enhance the ability to trace evolutionary relationships and track the development of specific traits over time.

While computational methods for analyzing 3D scans have advanced significantly in fields like medical imaging and industrial design, paleontology has not yet fully leveraged these tools. Current research in paleontology has a noticeable lack of specialized algorithms and methods tailored for interpreting the complex, irregular shapes of fossilized remains, such as dinosaur bones. The gap lies in the adaptation and development of these advanced computational methods to suit the specific challenges of paleontological data.

Building on existing research in 3D scan analysis and computational methods, my goal is to advance these techniques more specifically for the field of paleontology. By applying and adapting algorithms such as the one used by Dornbusch et al. [3], I aim to develop new methods for analyzing and characterizing dinosaur bones from 3D scans. This work seeks to fill the current gap in applying these advanced techniques to paleontological data, ultimately improving the accuracy of evolutionary trees and our understanding of prehistoric life. I will

ensure the accuracy of my phylogenetic tree through comparissons of known fossil relatives that have been hand caluclated and use those results to validate the methods I choose.

I look forward to expanding my research experience by taking on the challenge of an independent project for the first time. With a solid background in coding with Python, Java, and C++, I have the skills that will be instrumental in developing and implementing the new algorithms necessary for this research. This project will allow me to deepen my knowledge of computational topology and enhance my skills in algorithm development and data analysis. I am eager to contribute to the field of paleontology with a tool that not only advances our capability to analyze fossil data but also has the potential to influence broader applications in scientific research.

3 Methods

To begin this project, I will preprocess 3D scans of dinosaur bones to clean and normalize the data. Specifically, I will ensure consistency in scale, orientation, and resolution by removing noise, filling holes, and aligning scans using tools from Python, such as trimesh and pygltflib. Next, I will adapt existing algorithms, such as point cloud registration and surface fitting to identify key morphological features of the bones. This will involve implementing and customizing these algorithms in Python, utilizing libraries like NumPy, SciPy, and Py-Torch. Following this, I will develop new algorithms specifically tailored to paleontology, using techniques like principal component analysis (PCA) and singular value decomposition (SVD) to analyze and compare features across specimens. Certain machine learning models, such as convolutional neural networks (CNN) may also be utilized to enhance automated characterization. Experiments will validate the effectiveness of these methods by comparing them to known data, using metrics like shape similarity scores and classification accuracy to ensure robustness across different types of bones. The final stage will involve documenting the findings in an abstract or research paper, which will highlight the potential applications

of these methods in paleontology and beyond.

4 Timeline

An outline of the estimated timeline for this project is listed below.

- 1. Clean, normalize, and align 3D scans using trimesh and pygltflib: 2 weeks
- 2. Implement and adapt existing algorithms to identify key morphological features of the bones: 3 weeks
- 3. Develop and customize new algorithms tailored to paleontology using PCA, SVD and CNNs to enhance automated characterization: 5 weeks
- 4. Conduct experiments to validate methods, comparing them to known data: 3 weeks
- 5. Analyze the results, refine methods as needed, and compile findings for documentation:

 3 weeks

5 Collaboration with Faculty Sponsor

My primary faculty sponsor for this project is Dr. Brittany Fasy, whose ongoing research in 3D shape analysis aligns with the methodological aspects of this project. I plan to meet with Dr. Fasy regularly as her expertise in computational geometry and 3D data analysis will be invaluable in guiding the development and refinement of algorithms for this project. Additionally, I will be collaborating with Dr. Christopher Organ, whose work in evolutionary biology and paleontology will provide crucial insights into the biological relevance of the morphological features I am going to analyze. I am eager to engage in this research project, which not only supports the work of my mentors but also aims to make a meaningful contribution to the fields of computational topology and paleontology.

References

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