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PaleoProPhyler: a reproducible pipeline for phylogenetic inference using ancient proteins

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Summary

Ancient proteins from fossilized or semi-fossilized remains can yield phylogenetic information at broad temporal horizons, in some cases even millions of years into the past. In recent years, peptides extracted from archaic hominins and long-extinct mega-fauna have enabled unprecedented insights into their evolutionary history [1, 2, 3, 4, 5, 6]. In contrast to the field of ancient DNA - where several computational methods exist to process and analyze sequencing data - few tools exist for handling ancient protein sequence data. Instead, most studies rely on loosely combined custom scripts, which makes it difficult to reproduce results or share methodologies across research groups. Here, we present PaleoProPhyler: a new fully reproducible pipeline for aligning ancient peptide data and subsequently performing phylogenetic analyses. The pipeline can not only process various forms of proteomic data, but also easily harness genetic data in different formats (CRAM, BAM, VCF) and translate it, allowing the user to create reference panels for phyloproteomic analyses. We describe the various steps of the pipeline and its many functionalities, and provide some examples of how to use it. PaleoProPhyler allows researchers with little bioinformatics experience to efficiently analyze palaeoproteomic sequences, so as to derive insights from this valuable source of evolutionary data.

Statement of Need

Recent advances in protein extraction and mass spectrometry [7, 8, 9] have made it possible to isolate ancient peptides from organisms that lived thousands or even millions of years ago. Certain ancient proteins have a lower degradation rate and can be preserved for longer than ancient DNA [10, 11, 12, 13]. The sequences of these proteins contain evolutionary information and thus have the potential to resolve important scientific questions about the deep past, which are not approachable via other methods. Tooth enamel proteins in particular have been successfully extracted from multiple extinct species, in order to resolve their relationships to other species [1, 2, 4, 6, 14, 5].

Ancient proteomic studies typically use combinations of custom scripts and repurposed software, which require extensive in-house knowledge and phylogenetic expertise, and are not easily reproducible. Barriers to newcomers in the field include difficulties in properly aligning the fractured peptides with present-day sequences, translating available genomic data for comparison, and porting proteomic data into standard phylogenetic packages. The creation of automated pipelines like PALEOMIX [15] and EAGER [16] have facilitated the streamlining and reproducibility of ancient DNA analyses, which has been particularly helpful for emerging research groups around the world. This has undoubtedly contributed to the growth of the field [17]. Yet, the field of palaeoproteomics still lacks a “democratizing” tool that is approachable to researchers of different backgrounds and expertises.

Another important issue in phyloproteomics is the rel-
ative scarcity of proteomic datasets [18, 19]. There are currently tens of thousands of publicly available whole genome sequences, covering hundreds of species [20, 21, 22, 23, 24]. The amount of publicly available proteome sequences is much smaller in comparison. For most vertebrate species, lab-generated protein data does not even exist and phyloproteomic research is reliant on sequences translated in silico from genomic data. These, more often than not, are not sufficiently validated or curated [25]. As a result, assembling a proper reference dataset for phyloproteomics can be challenging. Given how important rigorous taxon sampling is in performing proper phylogenetic reconstruction [26, 27], having a complete and reliable reference dataset is crucial. In the case of proteins, the typically short sequence length and the low amounts of sequence diversity - due to the strong influence of purifying selection - means that absence of knowledge about even a single amino acid polymorphism (SAP) can strongly affect downstream inferences [3, 28, 29, 30].

To address all of the above issues, we present “PaleoProPhyler”: a fully reproducible and easily deployable pipeline for assisting researchers in phyloproteomic analyses of ancient peptides. “PaleoProPhyler” is based on the workflows developed in earlier ancient protein studies [2, 4, 6], with some additional functionalities. It allows for the search and access of available reference proteomes, bulk translation of CRAM, BAM or VCF files into amino acid sequences in FASTA format, and various forms of phylogenetic tree reconstruction.

Description of the Pipeline

To maximize reproducibility, accessibility and scalability, we have built our pipeline using Snakemake [31] and Conda [32]. The Snakemake format provides the workflow with tools for automation and computational optimization, while Conda enables the pipeline to operate on different platforms, granting it ease of access and portability. The pipeline is divided into three distinct but interacting modules (Modules 1, 2 and 3), each of which is composed of a Snakemake script and a Conda environment 1. The modules are intended to synergize with each other, but can also be used independently. An in-depth explanation of each step of each module, as well as the code being run in the background, is provided on the software’s Github page as well as in the supplementary material.

Application

As proof of principle, we deploy this pipeline in the reconstruction of ancient hominid history using the publicly available enamel proteomes of Homo antecessor and Gigantopithecus blacki, in combination with translated genomes from hundreds of present-day and ancient hominid samples. In the process, we have generated the most complete and up to date, molecular hominid phyloproteomic tree 2. The process of generating the reference dataset and its phyloproteomic tree using PaleoProPhyler is covered in detail in the step-by-step Github Tutorial. The dataset used as input for the creation of the phylogenetic tree is available at Zenodo'

Protein Reference Dataset

In order to facilitate future analyses of ancient protein data, we also generated a publicly-available palaeoproteomic hominid reference dataset, using Modules 1 and 2. We translated 204 publicly available whole genomes from all 4 extant Hominid genera [21, 22, 33]. Details on the preparation of the translated samples can be found in the supplementary materials. We also translated multiple ancient genomes from VCF files, including those of several Neanderthals and one Denisovan [34, 35]. Since the dataset is tailored for palaeoproteomic tree sequence reconstruction, we chose to translate proteins that have previously been reported as present in either teeth or bone tissue. We compiled a list of 1696 proteins from previous studies [36, 37, 38, 39, 40, 41] and successfully translated 1,543 of them. For each protein, we translated the canonical isoform as well as all alternative isoforms, leading to a total of 10,058 protein sequences for each individual in the dataset. Details on the creation of the protein list can be found in the supplementary materials. The palaeoproteomic hominid reference dataset is publicly available online at Zenodo, under the name ‘Hominid Palaeoproteomic Reference Dataset’

Availability and Community Guidelines

PaleoProPhyler is publicly available on github: The software requires the prior installation of Conda. The github repository contains a tutorial for using the workflow presented here, with the proteins recovered from the Homo antecessor and Gigantopithecus blacki as examples. We welcome code contributions, feature requests, and bug
reports via Github. The software is released under a CC-BY license.

**Author Contributions**

- **Ioannis Patramanis**: Conceptualization, manuscript writing, code writing for the Snakemake scripts, compilation of the Conda environments and application of the pipelines to produce the results described in the 'Application' and 'Protein Reference Dataset' section.

- **Jazmin Ramos Madrigal**: Manuscript review, conceptualization and code for multiple R and bash scripts utilised by the Snakemake script as steps of the pipeline.

- **Enrico Cappellini**: Manuscript review and editing

- **Fernando Racimo**: Conceptualization, manuscript writing, review and editing

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Figure 2: Phyloproteomic tree generated using Paleo-ProPhyler’s Module 3. The tree was constructed using 9 protein sequences obtained from enamel and includes more than 100 hominid individuals translated from genomic data, two individuals from published palaeoproteomic datasets as well as sequence data from a Macaca and a Hylobates individual, which are used to root the tree.
Bibliography


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