# Poseidon – A framework for archaeogenetic human genotype data <sup>2</sup> management

Clemens Schmid <sup>1,2</sup>, Ayshin Ghalichi <sup>1</sup>, Thiseas C. Lamnidis <sup>1</sup>, Dhananjaya B. A. Mudiyanselage <sup>1,3</sup>, Wolfgang Haak <sup>1</sup>, and Stephan Schiffels <sup>1</sup>

<sup>1</sup>Max Planck Institute for Evolutionary Anthropology, Leipzig, Germany

<sup>2</sup>International Max Planck Research School for the Science of Human History, Max Planck Institute for Geoanthropology, Jena, Germany

<sup>3</sup>Saarland University, Saarbrücken, Germany

## • 1 Abstract

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The study of ancient human genomes, archaeo- or palaeogenetics, has accelerated in the last ten years, with 10 now thousands of new ancient genomes being released each year. Operating at the interface of genetics, anthro-11 pology and archaeology, this data includes features from all three fields, including rich meta- and context-data, 12 for example regarding spatiotemporal provenience. While archives and standards for genetic sequencing data al-13 ready exist, no such infrastructure exists for combined genetic and meta-data that could ensure FAIR principles 14 across the field. Here, we present Poseidon, a framework for open and FAIR data handling in archaeogenetics, 15 including a specified package format, software tools, and public, community-maintained online archives. Posei-16 don emphasises human- and machine-readable data storage, the development of convenient and interoperable 17 command line software, and a high degree of source granularity to elevate the original data publication to the 18 main unit of long-term curation. 19 20

<sup>21</sup> Keywords: ancient DNA, data management, FAIR data

## $_{22}$ 2 Introduction

The technology to extract and sequence DNA from human remains thousands of years old has revolutionised 23 the study of the human past. This is documented by groundbreaking new insights, from our evolutionary 24 relationships to distant relatives like Neanderthals [1, 2] to prehistoric [3, 4] and historic migrations [5, 6]. 25 Since the sequencing of the first ancient modern human genome in 2010 [7], hundreds of studies have been 26 published, accompanied by massive datasets of ancient human DNA sequences. A drop in sequencing costs and 27 new technologies like hybridisation capture [8, 4, 9] have in fact lead to an acceleration of new published ancient 28 genomes, with data now coming out faster than individual researchers typically can keep track of and co-analyze. 29 Recently, the threshold of genome-wide data for 10,000 ancient human individuals has been surpassed [10]. 30

To make all this new data publicly available, researchers can partly rely on existing infrastructure for the archival and distribution of modern genetic data, such as the Sequence Read Archive (SRA) [11], the European Nucleotide Archive (ENA) [12] or other INSDC databases (https://www.insdc.org). However,

<sup>34</sup> this infrastructure has not been prepared to also capture the rich context-data ranging from archaeological field

observations to radiocarbon dating that accompanies ancient samples. Nor is there a standardised archive yet for
 derived genotype data that is routinely used to substantiate most if not all of the conclusions in archaeogenetic
 papers. This raises multiple concrete issues:

- Ancient individuals only constitute meaningful observations if their spatiotemporal provenience is known.
   Current practice renders it difficult to maintain the connection between archaeological context and sampled
   genomic data, as this information is generally kept separately.
- Specific results of typical archaeogenetic analyses (e.g. PCA, F-Statistics, kinship estimation) can only
   be fully reproduced with genotype-level data. But current practice is not to include this data with a
   publication, be it for its unwieldy size or the lack of a central repository to easily share it.

Meta-analyses involving large amounts of data require tedious curation. Despite the fact that archaeoge netic genotype data is largely standardized, and common practices for reporting processing steps, data
 quality indicators, as well as spatial and temporal sample origin have emerged, combining information
 from different papers is still hindered by severe structural variation.

A major project addressing some of these problems in human archaeogenetics is the Allen Ancient DNA 48 Resource (AADR), which is a curated dataset of public ancient DNA data assembled by the Reich Lab at 49 Harvard University [13]. While the AADR clearly fills a gap in the field, there continues to be a need for open 50 standardization and the creation of a community-maintained archive. Both standard and archive should be 51 well-specified to be human and machine-readable, fully open and transparent, and permanently downloadable 52 with their entire version history. They should be geared towards scientific practice and include well-documented 53 interfaces for research software. To ensure fairness, i.e. equal access for users and contributors from different 54 backgrounds, and long-term reliability, they should be as independent as possible from specific institutions, key 55 persons [14] or infrastructure providers, and ideally controlled by the community-of-practice as a whole. 56 Here, we present 'Poseidon' (https://www.poseidon-adna.org), a computational framework including 57

<sup>58</sup> an open data format, software, and community-maintained archives, to enable this standardised and FAIR <sup>59</sup> [15] handling of archaeogenetic data. The name *Poseidon* is inspired by the notion of a *sea of data* benefiting <sup>60</sup> from structure and governance. We imagine Poseidon to simplify a variety of concrete workflows and mitigate <sup>61</sup> technical challenges practitioners in the field of human archaeogenetics regularly face:

• Data storage: Archaeogenetic samples/ancient individuals (we often use these terms synonymously below, see Supplementary Text 8 for a definition) can only be effectively analysed with context data. The Poseidon package format allows one to store archaeogenetic genotype data with arbitrary spatiotemporal or archaeological information on a per-sample level. The package format enforces human- and machinereadability for the context data to enable its computational co-analysis with the genomic data, while also maintaining a high level of flexibility and extensibility to accommodate arbitrary additional variables.

• Data acquisition: Research in archaeogenetics is strongly dependent on incorporating published reference data. Poseidon features public archives with per-article packages that can be downloaded through an open web API. The packages include genotypes, context data and machine-readable citation information. To ensure computational reproducibility, Poseidon supports version tracking for these packages, with old versions directly available through the web interface. Beyond hosting data, the web infrastructure also provides options to report issues and suggest transparent data updates.

Data analysis: Working with common software tools in human archaeogenetics (e.g. smartpca [16, 17],
 ADMIXTURE [18], qpAdm [4, 19]) requires frequent merging, sub-setting and file format conversion for the samples of interest. The Poseidon core team develops the software tool *trident* to simplify these operations

both with Poseidon packages and unpackaged genotype data. The unified package format allows for new 77 analysis tools with flexible data detection, low-memory stream processing and on-the-fly aggregation of 78 widely used genome-wide statistics as demonstrated in the *xerxes* software tool also developed by the core 79 team.

- 80
- Data publication For each sample, human archaeogenetic papers should include genotypes, context 81 information and quality-control data, so e.g. estimates for the aDNA damage or contamination. Poseidon 82 offers a standardized and reusable way to share this data directly with the publication and/or through 83
- our public archives.

#### 3 **Overview** 85



Figure 1: Schematic overview of the core components of the Poseidon framework. The Poseidon package specification forms the foundation on which various open source software tools and the Minotaur workflow are based. They, in turn, underpin and enable the public Poseidon data archives.

Poseidon consists of three major components: A data format, software, and public archives (see Figure 1). 86 At the foundation stands a data format to store genotype data together with context information. The software 87 implements this specification, relies on and validates its promises, and builds convenient, useful functionality 88 on top of it, both for individual users and for our cloud infrastructure. Finally, the public archives store data 89 using the data format and employ the software for validation, modification and book-keeping. 90

The Poseidon schema (Supplementary Text 1) defines the structure and format of a Poseidon package (Figure 91

2). This includes the general layout and purpose of the main files, the POSEIDON.yml, the janno and the .ssf 92

file, and detailed definitions of the variables within them, both regarding semantics and syntax (Supplementary 93

Text 2). The short definitions in the schema are explained in more detail on the Poseidon website, which also 94

serves as a central hub for all components of the framework. 95

The Poseidon software suite enables users to create, download, inspect, subset, merge and analyse Poseidon 96 packages (Supplementary Text 3-7). It is mostly implemented in Haskell [20], a purely functional programming 97 language, and split over multiple command line tools openly available as statically compiled executables for 98



Figure 2: Schematic overview of the Poseidon package structure. The POSEIDON.yml file defines the package and interlinks the additional data files for genotype, context- and bibliography information in a relational structure.

<sup>99</sup> the major desktop and server operating systems. The main tool, *trident*, provides data handling functionality, <sup>100</sup> including the code necessary for the client-server infrastructure of the public archives. The *xerxes* software <sup>101</sup> implements commonly used genome-wide allele-frequency statistics, *qjanno* allows for SQL-like queries on .janno <sup>102</sup> files and the *janno* R package offers a .janno file interface for the R programming language.

Using these tools, we conceived and implemented three public data archives for sharing and maintaining 103 published (ancient) human DNA data packaged in the Poseidon format: The Poseidon Community Archive 104 (PCA), the Poseidon Minotaur Archive (PMA) and the Poseidon AADR Archive (PAA). They already store 105 considerable amounts of public genotype data (Figure 4, Figure 5 and Supplementary Text 8). Each of these 106 archives is at their base represented by a public repository on GitHub, versioned with Git and capable of holding 107 large genotype data files through an integration with Git's Large File Storage (LFS) functionality. This setup 108 allows for community driven maintenance through Git pull requests (Figure 3). A custom webserver mirrors the 109 data from GitHub with an explicit version history, and allows to query the archive data and download packages 110 in the browser, from the command line and through web-frontends. While the PCA focuses on author-submitted 111 genotype data, the PMA only accepts data that went through a specific standardized genotype data processing 112 pipeline, the Minotaur workflow, with a semi-automatic interface on GitHub and a processing queue currently 113 hosted on computational infrastructure at the Max Planck Institute for Evolutionary Anthropology (MPI-EVA). 114 Finally, the PAA stores releases of the AADR dataset in Poseidon format, after a light-weight curation step to 115 ensure format compatibility. 116

### <sup>117</sup> 4 Structure

The following sections explain the different components of Poseidon in detail: The Poseidon package, the software tools, and the public archives including the Minotaur processing workflow. The underlying repositories with specifications and code are available on GitHub (https://github.com/poseidon-framework) and, each in a

current version, at a long-term archive: https://doi.org/10.17605/OSF.IO/ZUQGB



Figure 3: Schematic overview of the most common interaction pathways between the archaeogenetic community of practice and the Poseidon infrastructure. Community members share data either by preparing .ssf files as build-instructions for the Minotaur workflow or by directly submitting packages to the Community Archive. The data in the archives can then in turn be downloaded from the archives through the Poseidon webserver and its API.

#### <sup>122</sup> 4.1 The Poseidon Package specification (v2.7.1)

The core idea of Poseidon is to organize genotype data together with relevant meta- and context data in a 123 structured yet flexible, human- and machine-readable format. This format is the Poseidon package, defined in 124 a semantically versioned [21] specification, openly available online (https://github.com/poseidon-framewo 125 rk/poseidon-schema), and also part of this publication as Supplementary Text 1. A Poseidon package must 126 contain a POSEIDON.yml file and genotype data in PLINK or EIGENSTRAT format. It should additionally 127 contain a .janno file to store sample-wise context information and a .bib file for literature references. Option-128 ally, it can also contain an .ssf file with information on the underlying raw sequencing data, an unstructured 129 README.md file for arbitrary meta information and a CHANGELOG.md file to document changes to the 130 131 package.

```
poseidonVersion: 2.7.1
132 1
    title: Switzerland_LNBA
133 2
    description: Genetic data from Late Neolithic and Bronze Age Switzerland
134 3
    contributor:
135 4
        name: Roswita Malone
136 5
        email: roswita.malone@example.org
137 6
    packageVersion: 1.1.2
138 7
    lastModified: 2024-03-15
139 8
    genotypeData:
140 9
      format: PLINK
14110
      genoFile: Switzerland_LNBA.bed
142 1
      snpFile: Switzerland_LNBA.bim
143.2
      indFile: Switzerland_LNBA.fam
144.3
      snpSet: 1240K
1454
1465 jannoFile: Switzerland_LNBA.janno
```

#### 147.6 bibFile: Switzerland\_LNBA.bib

Example 1: A typical, neither minimal nor maximal, POSEIDON.yml file.

A Poseidon package is defined by a POSEIDON.yml file, using the YAML markup language (https://ya ml.org) with a set of predefined fields, which stores some general information and, most importantly, relative paths to the other files in the package (see Example 1). A full list of the specified fields is provided as part of the schema, but only poseidonVersion (the schema version the package adheres to), title (a short package name), and genotypeData (context information for and relative paths to the genotype data files) are mandatory. Besides its function to define the package, the POSEIDON.yml also enables software to easily crawl for packages in a file system.

At the time of writing, Poseidon supports genotype data for single nucleotide polymorphisms (SNPs) in two 155 common file formats: a binary-encoded format as introduced by the plink software package (PLINK) [22] and a 156 plain-text, human-readable format defined for the eigensoft software tools (EIGENSTRAT) [17]. Both formats 157 are structurally similar, and split into three files: A genotype table (.bed/.geno); a SNP file (.bim/.snp), defining 158 the SNPs in the genotype table; and an individual file (.fam/.ind) for the respective samples. In the future, 159 Poseidon may add support for other formats, e.g. the more flexible vcf file format [23]. Arbitrary SNP sets are 160 allowed and supported, but special keywords are reserved for the commonly used Affymetrix HumanOrigins 161 array [24], and the so-called 1240k in-solution hybridization capture reagent [25]. 162

The .janno file accompanies the genotype data to provide context for each sample. It is designed as a tabular, 163 tab-separated text file with a set of predefined columns. Each row corresponds to one entry in the individual file 164 (.fam/.ind), featuring at least the following mandatory columns: Poseidon\_ID (a sample identifier), Genetic\_Sex 165 (Female, Male, Unknown) and Group\_Name (one or multiple group identifiers). Other, optional columns include 166 standard information on the spatiotemporal provenience of a given sample, its genetic data quality metrics, and 167 which major laboratory procedures it was subjected to. Supplementary Text 2 includes documentation for all 168 specified columns of the .janno file. One particular column documents the scientific publication(s) within which 169 the genetic or contextual data for a given sample was originally reported. This is a list column with BibTeX 170 keys, which, in turn, must be specified in the .bib file of the Poseidon package, thus ensuring that each sample 171 can be properly cited. 172

The .ssf file is similarly structured as the .janno file and stores sequencing source data, i.e. meta-information about the raw sequencing data behind the genotypes in a Poseidon package. The rows in this table correspond to sequencing entities, typically the set of unprocessed reads sequenced from DNA libraries or even multiple runs/lanes of the same library. At the time of writing, the specified columns mostly focus on how to download a given dataset from the INSDC databases such as ENA or SRA (https://www.insdc.org). The .ssf file does not have any mandatory variables, but entries can be linked to the package with the list column poseidon\_IDs in a many-to-many relationship.

#### <sup>180</sup> 4.2 Software tools

The following software tools were developed to work with Poseidon packages and other infrastructure of the Poseidon ecosystem. They can be considered reference implementations of the Poseidon schema, but are by no means exclusive nor comprehensive. The Poseidon schema is defined as an independent, versioned entity, and software developers can easily implement other tools in addition to what we as the Poseidon core team currently offer.

All Poseidon software is open-source (MIT-License) and available on GitHub (https://github.com/posei don-framework). That is also where users should report issues.

#### <sup>188</sup> 4.2.1 trident (v1.4.1.0)

<sup>189</sup> trident is a command line software tool to create, download, inspect, merge, and subset Poseidon packages – <sup>190</sup> and therefore the central tool of the Poseidon framework. It is implemented in Haskell as one executable for <sup>191</sup> the poseidon-hs library and can be installed from source with the build systems cabal [26] or stack [27]. To <sup>192</sup> ease installation we provide pre-compiled, static binary executables for Linux, Windows and macOS. We also <sup>193</sup> maintain a package recipe under poseidon-trident on the bioconda channel [28], enabling installation of trident <sup>194</sup> through the conda (https://docs.anaconda.com) package management system.

trident includes multiple sub-commands for different operations on and with Poseidon packages – Supple-195 mentary Text 3 gives a detailed overview including all specific arguments. It supports two mechanisms to obtain 196 Poseidon packages: A user can create them from genotype data with trident init or download them from our 197 community-maintained archives with trident fetch. The most involved and technically complex sub-command 198 in trident is trident forge, which allows users to both subset and merge Poseidon packages. To simplify the 199 process of package maintenance, trident genoconvert converts the genotype data between the formats Po-200 seidon supports and trident rectify automates some common tasks such as checksum and version updates. 201 For package inspection, trident includes trident list, which returns lists of packages, groups, or samples for 202 a given package selection (either locally, or remotely by accessing the web API for the community-maintained 203 online archives). trident summarise compiles some basic summary statistics about a package collection and 204 trident survey gives an overview to which degree .janno files in a package collection are filled with data, i.e. 205 the level of completeness for context information. The last and arguably the most important inspection sub-206 command is trident validate, which parses all components of a package to report violations of the Poseidon 207 schema. This ensures the structural integrity of Poseidon packages and maintains machine-readability. 208

```
209 1 trident list --remote --individuals --raw | grep "Finland\|VolgaOka"
210 2 trident fetch -d . -f "*2018_Lamnidis_Fennoscandia*,*2023_Peltola_VolgaOka*"
211 3 trident forge -d . -d ../myData/ -f "Finland_Levanluhta,VolgaOka_IA,<mySample>" -o newPac
```

Example 2: A basic trident command line workflow to explore the community archive, download relevant packages and create a new package from the downloaded data, as well as a local/private data collection. trident list queries the webserver (--remote), and returns a tab separated (--raw) table of individuals/samples (--individuals) available in the public Community Archive (PCA). This list can then be filtered with standard command line tools (here grep). With trident fetch two packages are selected for download (-f ...) into the current working directory (-d .). These two packages as well as an additional, local package collection (-d ../myData/) can then be read into forge to create a new package newPac, specifying (-f ...) the groups Finland\_Levanluhta, VolgaOka\_IA and the local sample mySample.

A simple *trident* workflow could look like Example 2. The first two commands here require interaction with the Poseidon webserver. This server software is, in fact, also implemented in Haskell as a part of *poseidon-hs* and can be started with a hidden (and for most end users irrelevant) *trident* sub-command trident serve. On the client side trident list and trident fetch use this API and interact, by default, with the endpoints at https://server.poseidon-adna.org. A different server can be set with --remoteURL. More about the available endpoints below.

The forge sub-command is the technically most involved operation in *trident*. It discovers all Poseidon packages under a list of base directories (-d), reads them and their components into dedicated in-memory data structures, parses the query language in -f (or from a file with --forgeFile) to decide which entities (individuals, groups, packages) should be selected, and ultimately generates the new package including genotypes and context data in a single, low-memory stream processing run. All components of the Poseidon package format (i.e. the POSEIDON.yml file, the genotype data, the .janno, the .ssf, and the .bib file) are modelled in *poseidon*hs as algebraic data types with dedicated parsers. Corrupted files are rejected upon reading, including cross-file

compatibility checks within each package (e.g. bibtex keys mentioned in the .janno file must be specified in the .bib file). The query language supported by trident forge in -f is a flexible domain-specific language (DSL) to describe arbitrary positive and negative entity selection scenarios. For the stream processing of genotype data in binary PLINK and EIGENSTRAT format, *poseidon-hs* relies on the the sequencing-formats library (https://github.com/stschiff/sequence-formats), which allows *trident* to read, filter and write data from a large amount of files at once, while also unifying SNPs to a consensus by recoding alleles on-the-fly. Varying

231 SNP sets in multiple packages can be merged using either an intersection or a union operation.

#### 232 4.2.2 xerxes (v1.0.1.0)

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xerxes is another command line software tool based on the poseidon-hs Haskell library. While trident is meant 233 for data management, *xerxes* is intended for basic, every-day data analysis operations. Its most advanced and 234 stable sub-command xerxes fstats makes extensive use of the genotype data stream processing introduced 235 above to implement commonly used genome-wide statistics. These are F-Statistics ( $F_2$ ,  $F_3$  and  $F_4$ , see Example 236 3, in several variants differing in bias-correction and normalisation) [24],  $F_{\rm ST}$  [29], heterozygosity, and pairwise 237 nucleotide mismatch rates for assessing pairwise relatedness. All statistics are evaluated with error-estimation 238 using weighted block-Jackknife [30]. See Supplementary Text 4 for the *xerxes* user guide and Supplementary 239 Text 5 for a whitepaper documenting the mathematical underpinnings of the implemented algorithms. 240

```
2411 xerxes fstats -d . \
242 2 --stat "F4(Mbuti,Nganasan,Lithuanian,Finland_Levanluhta)" \
```

```
243 3 -- stat "F4(Mbuti, Nganasan, Lithuanian, Russia_Bolshoy)"
```

Example 3: Calculating two F<sub>4</sub> statistics with *xerxes*, reproducing an analysis of Lamnidis et al. 2018 [31]. We assume the base directory (-d .) to include Poseidon packages with the groups Mbuti, Nganasan, Lithuanian, Finland\_Levanluhta and Russia\_Bolshoy. These happen to be described in the Community Archive packages 2018\_Lamnidis\_Fennoscandia [31], 2014\_LazaridisNature [3] and 2012\_PattersonGenetics [24], but xerxes will find those automatically if they are stored below the base directory. The individual statistics are specified with --stats and a dedicated domain specific language. For complex analyses *xerxes* offers a more powerful configuration file interface with the --statConfig argument.

```
groupDefs:
244 1
      CEU2: ["CEU.SG", "-<NA12889.SG>", "-<NA12890.SG>"]
245 2
      FIN2: ["FIN.SG", "-<HG00383.SG>", "-<HG00384.SG>"]
246 3
247 4
    fstats:
      type: FST # this will create 2x2 = 4 statistics
248 5
      a: ["French", "Spanish"]
249.6
      b: ["Han", "CEU2"]
250 7
251 8
      type: F4 # this will create 2x2x2x1 = 8 statistics
      a:
         ["French", "Spanish"]
252 9
      b:
         ["Han", "CEU2"]
253.0
      c: ["Nganasan", "Lithuanian"]
2541
      d: ["Mbuti"]
25512
```

**Example 4:** An example of a configuration file for *xerxes*, specifying a total of 12 statistics of  $F_{ST}$  and  $F_4$  to be computed for all combinations of listed populations. In addition, the example features adhoc group definitions in the **groupDefs** section, which shows how to exclude individuals from a group using the minus-sign. More complex features of these configuration files include frequency-ascertainment, and selection of entire packages to act as groups.

We highlight three *xerxes* features that stand out: First, the user interface provides for a powerful way to specify families of statistics using combinatorical expansions. For example, users can specify in a configuration file (Example 4) that  $F_{\rm ST}$  should be computed between all combinations of two lists of populations, and multiple

such families can be requested simultaneously. Second, statistics can be easily adapted with custom group-259 definitions, in which individuals, groups or entire packages (similar to the trident forge selection language) 260 can be excluded or included. If a specific individual turns out to be an outlier that should not be included in 261 the computation of allele frequencies of its group, one can define a custom group and exclude that individual on 262 the fly, without the need to change the underlying genotype definition files. Third, users do not have to specify 263 the exact packages that their populations or individuals reside in, but can pass a base directory with dozens or 264 hundreds of packages, and *xerxes* will automatically select only the relevant packages, involving the groups or 265 individuals needed, thereby reducing memory- and run-time overhead: Similar to trident forge, xerxes uses 266 stream-processing to merge the relevant genotype-files without the need to load large genotype matrices into 267 memory. This is particularly important for dense genotyping data sets, for example involving tens of millions 268 of SNPs as with the 1000 Genomes dataset [32]. 269

#### 270 4.2.3 qjanno (v1.0.0.0)

qjanno is a command line tool implemented in Haskell to run SQL queries on .janno files (or arbitrary delimiterseparated text files) and therefore to conveniently subject the context data provided with Poseidon packages
to a wide range of data transformations. It was built on top of a hard fork of v0.3.3 of the qsh package
(https://github.com/itchyny/qhs, MIT-License). See Supplementary Text 6 for the *qjanno* user guide.

On startup, *qjanno* creates an SQLite [33] database in memory. It then reads the requested, well-structured text files, attributes each column a type and writes the contents of the files to tables in the in-memory database.

277 It finally sends the user-provided SQL query to the database, waits for the result, parses it and returns it on the

<sup>278</sup> command line. The query gets pre-parsed to extract file names and then forwarded to an SQLite database server

via the Haskell library sqlite-simple (https://github.com/nurpax/sqlite-simple). That means qjanno can

<sup>280</sup> parse and understand most SQLite3 syntax (see Example 5 and 6) with some minor exceptions (e.g. PRAGMA

<sup>281</sup> functions).

2821 qjanno "SELECT Poseidon\_ID FROM 2018\_Lamnidis\_Fennoscandia.janno WHERE Country <> 'Finland'"

Example 5: A simple *qjanno* query on the command line. It extracts the columns Poseidon\_ID and Country of the file 2018 Lamnidis Fennoscandia.janno and returns all rows where Country is not 'Finland'.

qjanno does not have a complete understanding of the .janno-file structure, and mostly treats it like a normal .tsv file. But .janno files are still given special consideration with a number of pseudo-functions, which allow to search Poseidon packages and .janno files recursively to load them together into one database table (see Evenuela 6)

Example 6).

#### 287 1 qjanno "SELECT Date\_Type,count(\*) AS n FROM d(.) GROUP BY Date\_Type"

Example 6: Another *qjanno* query. With 'FROM d(.)' *qjanno* searches all latest versions of Poseidon packages under the current working directory, reads their .janno files and appends them to a common database table. This table is then grouped by the Date\_Type column (*C14, contextual, modern*). The output features a row-count for each group in a new summary column n.

#### <sup>288</sup> 4.2.4 janno R package (v1.0.0)

<sup>289</sup> The *janno* R package simplifies loading and handling .janno files in R and the popular tidyverse [34] R package

 $_{290}$  ecosystem. It provides a dedicated R S3 class janno that inherits from the tibble class to allow tidy reading and

<sup>291</sup> manipulating the context information in a Poseidon package (see Example 7). Supplementary Text 7 features <sup>292</sup> its user guide.

```
293 1 janno::read_janno(".") |>
```

#### 294 2 dplyr::group\_by(Date\_Type) |>

#### 295 3 dplyr::summarise(n = dplyr::n())

Example 7: A small R workflow enabled by the *janno* R package. The code is equivalent to the *qjanno* Example 6. read\_janno() discovers and reads all .janno files under the current working directory into an object of classs janno. The dplyr verbs group\_by and summarise can then be applied to this object to count the rows per Date\_Type group.

As trident and qjanno (with -d .../d(...)), the package's read\_janno() function searches janno files re-296 cursively under a given directory and loads them into a single data frame. The reading process includes validation 297 according to the Poseidon schema. String list columns in .janno files are translated to true list columns in R. 298 Beyond basic functionality (print(), write\_janno()), the package features one more major function for janno 299 objects: process\_age(). This function processes the age information in the Date\_\* columns of the .janno file to 300 derive a set of new columns useful for further chronological analysis via radiocarbon calibration: Date\_BC\_AD\_Prob 301 is a list column with the complete post-calibration probability distribution, Date\_BC\_AD\_Median\_Derived is the 302 median of this distribution and  $Date_BC_AD_Sample$  stores n random samples drawn from it. The radiocarbon 303 calibration is implemented with the *Bchron* R package [35]. 304

#### 305 4.3 Public archives

With a standardized, versioned data format, Poseidon does not require central infrastructure. Its software tools work independently and users can apply them locally on their own data. But Poseidon was also developed to mitigate the issue of increasingly tedious data preparation for future research projects. To that end, the Poseidon ecosystem includes three public, openly curated archives, which share implementation and infrastructure, but differ in their goals, mode of maintenance, and data content. See the Figures 4 and 5 for an overview of the current data content in the archives. Supplementary Text 8 explains this comparison in more detail.



Figure 4: Spatiotemporal distribution of ancient individuals in the PCA and PAA (AADR v54.1.p1) public Poseidon archives. See Supplementary Text 8 for an explanation of how individuals were counted. The map shows the qualitative presence and absence of samples from both archives in a 5°-resolution grid. Especially highlighted are areas and time periods for which the PCA includes samples from currently 13 author-submitted Poseidon packages.



Figure 5: Multiple charts to compare the current content of the PCA and PAA (AADR v54.1.p1) public Poseidon archives. See Supplementary Text 8 for an explanation of how individuals were counted and more in-depth descriptions of each chart. A) Stacked barchart of publications and how they are distributed across packages in PCA and PAA. Publications represented in multiple packages are counted towards the shaded area to get a correct total, B) Barcode plot of individuals available in each archive per publication through time, C) Stacked barchart of individuals and how they were added to the archives, D) Sankey diagram of individuals matching across PCA and PAA, highlighting individuals unique to each archive on the right, E) Stacked barchart of dating information per individual available in the archives, F) Stacked barchart of spatial coordinate coverage in the archives.

#### 312 4.3.1 Technical infrastructure

All archives are hosted in dedicated Git repositories on GitHub (e.g. https://github.com/poseidon-frame 313 work/community-archive for the PCA), where each individual package is stored in a directory named after 314 the package. This setup has a number of advantages: Git provides version control down to the individual lines 315 of each meta- and context data file, Git and GitHub together include co-working features that allow users to 316 submit new packages and suggest concrete changes to existing ones, and GitHub is a comparatively affordable 317 host with advanced automation features. Git is by default not suitable for large, binary files, but GitHub offers 318 large file storage with the Git LFS extension. The archives make use of this for all .bed and .bim files. Each 319 change in the public archives is automatically validated using GitHub Actions on their cloud infrastructure, 320 running a number of scripts on the new state, including trident validate, to ensure continuous structural 321 integrity. 322

As already introduced above for trident list --remote and trident fetch, the data in the public 323 archives is not just available on GitHub, but also and more conveniently via a web API provided by an open web 324 server running trident serve. This service is hosted by the scientific IT service provider of the Max Planck 325 Society (Gesellschaft für wissenschaftliche Datenverarbeitung mbH Göttingen, https://gwdg.de). Once per 326 day the server fetches the latest changes to the archives on GitHub and incorporates new packages and package 327 versions. It does so through a Git-integrated bookkeeping mechanism using the hidden trident subcommands 328 chronicle and timetravel. Note that the server also provides outdated package versions (for the community 329 archive starting from 2023-06-12) to maintain computational reproducibility. Here are the endpoints the server 330 supports: 331

• https://server.poseidon-adna.org/packages returns a JSON list of all packages

- https://server.poseidon-adna.org/groups returns a JSON list of all groups
- https://server.poseidon-adna.org/individuals returns a JSON list of all samples/individuals

https://server.poseidon-adna.org/zip\_file/<package\_name> returns a complete zip file of the
 package with the given name

The most important arguments for these are ?archive to select the archive that should be queried, ?additionalJannoColumns to add more detailed information to the /individuals response and ?package\_version to select a specific package version with /zip\_file.

#### 340 4.3.2 The Community Archive

The Poseidon Community Archive (PCA) stores author-submitted, article-wise Poseidon packages. It focuses on packages prepared by the authors of the respective publication, containing the exact genotype data used for the paper, to ensure a maximum of computational reproducibility. Author submissions are also ideal for the context data in the .janno file, because the respective domain-experts are generally most knowledgeable on data quality and the spatiotemporal origin of their samples.

For historical reasons the PCA does not only contain author submissions, though. To kickstart the public 346 archive development in 2020, we prepopulated it with packages derived from in-house data and previous versions 347 of the AADR, which have since then been further modified and edited, as is transparent in the version history of 348 these packages. This legacy data will remain in the PCA to maintain established workflows. Authors and other 349 community members can take ownership, update entries if need be, and thus have the possibility to (further) 350 improve the quality of these datasets. A contributing guide on the Poseidon webpage explain the details of 351 how to submit a new paper-associated dataset or suggest changes to an existing one. Each submission passes 352 through a checklist-based review process and is eventually confirmed by the Poseidon core team. Contributors 353 and original and intermediate authors are credited via publication keys and corresponding .bib entries, as well 354 as a dedicated Contributor list in the package-defining POSEIDON.yml file. 355

#### 356 4.3.3 The AADR Archive

The Poseidon AADR Archive (PAA) stores releases of the AADR dataset [13] reworked into Poseidon packages. 357 It thus deviates from the PCA and the PMA in multiple important ways: It is not organized by individual 358 publications, includes the versioning of the original provider on top of our own versioning, and relies to a lesser 359 degree on community contributions. The cleaning and repackaging process is documented in an extra repository 360 (https://github.com/poseidon-framework/aadr2poseidon) and mostly has the following goals: i) Creation 36 of a version of the AADR that follows the Poseidon package standard and is thus directly compatible with 362 trident and other Poseidon tooling, ii) increasing the machine-readability of the AADR, especially regarding 363 the sample age information, and iii) providing clean .bib files with all references of publications in the AADR 364 for convenient citation management. 365

#### 366 4.3.4 The Minotaur Archive

The Poseidon Minotaur Archive (PMA) mirrors the PCA in that it stores publication-wise packages, often the very same as the PCA. However, Packages in the PMA do not rely on author-submitted genotype data, but instead include genotypes consistently reprocessed from raw sequencing data, run through the Minotaur workflow (see below). The motivation for this bioinformatic reprocessing is to generate an internally consistent dataset, which optimises cross-package comparability, rather than per-author reproducibility of individual packages –

372 like the AADR.

The submission of packages to the PMA is less direct as for the PCA and involves the preparation of a perpackage recipe to parameterize the relevant processing run. Package recipes are archived in a dedicated GitHub repository, where targeted GitHub Actions guide users through the necessary steps to create and submit a new recipe.

#### 377 4.4 The Minotaur workflow

The reproducibility of our processing behind the PMA is achieved with a semi-automatic computational workflow to generate Poseidon packages from raw sequencing data: the Minotaur workflow. The entry point for this processing pipeline is a *package recipe*, a collection of files containing all the information required to download, validate, and process the raw reads into a Poseidon package. Each recipe must contain an .ssf file, a .tsv file formatted like a valid input .tsv for nf-core/eager [36], a .config file outlining the nf-core/eager configuration parameters for processing, a .sh script that adapts the .tsv file to the local cluster at the time of processing, and finally a .txt file listing all the versions of scripts used when creating the recipe, to ensure reproducibility.

Contributors are able to request packages via GitHub issues on the dedicated minotaur-recipes repository (https://github.com/poseidon-framework/minotaur-recipes), and actively prepare package recipes by providing a configured .ssf file. This .ssf file gets validated through GitHub Actions, and then complemented to a full recipe with all the required files. Each recipe submission passes through a checklist-based review process and is finally confirmed by the Poseidon core team. This approach is designed to standardise and streamline the processing through the Minotaur workflow, while still allowing enough flexibility in its configuration to account for the heterogeneity present in raw sequencing data.

The minotaur-recipes repository is mirrored in the computational cluster of MPI-EVA, where the actual 392 processing takes place. In the future, we may outsource this processing to a publicly accessible cloud service 393 to make it fully independent from a particular institution. The raw data is downloaded there, and processed 394 through nf-core/eager with the parameters specified in the .config file of the package recipe. By default, this 395 processing includes adapter trimming and read-pair collapsing, aligning to the human reference, removal or 396 PCR duplicates, masking of the ends of reads to mitigate aDNA damage artefacts, and genotyping. Changes 397 to processing parameters are permitted, and can be specified, explained and recorded in the package recipe. 398 The genotypes generated from this processing are then turned into a Poseidon package, whose .janno file is 399 populated with descriptive statistics generated during the processing. 400

All the code responsible for this pipeline can be found at https://github.com/poseidon-framework/pos eidon-eager. The resulting Poseidon package is finally uploaded to the PMA, where it again undergoes review before being added to the archive. During the review process, missing information for the package is filled in either manually, or pulled from the Community Archive, if appropriate.

## 405 5 Discussion

Archaeogenetic research, as many other fast growing, data-driven fields, is challenged by data heterogeneity, a 406 lack of systematically applied standards, and from difficulties to discover published data. Poseidon addresses 40 these issues on multiple levels: First, Poseidon provides a standardised, yet flexible data format for day-to-day 408 scientific data analysis, large scale automation, and tidy data storage. Our design choice to build Poseidon around 409 the notion of packages, together with integration of bibliography information (the .bib file), emphasises good 410 citation practice, with any downstream merging necessarily resulting in complete bibliography files listing all 411 original source papers. Second, the Poseidon software, such as *trident*, to discover, validate, update, merge, subset 412 and analyse such packages, complements the standard to ease adoption. Third, perhaps the most ambitious 413

component of Poseidon, our public archives, make use of the standard and its versioning feature to host published
data and make it findable and transparently maintainable via GitHub community features.

This multi-layer architecture, with loosely coupled components, allows for a variety of adoption paths or starting points for users and data analysts: The package format alone can serve as a useful storage format for local work, even without using the software or archives. The software can help to create and work with such packages locally, even without the cloud-features and server-access. Finally, our archives can be seen as a transparent hub to download and discover data, even without our software or adoption of the package format.

In light of our developing research field and its position between disciplines, we designed the Poseidon package definition to allow for flexibility, e.g. by allowing arbitrary non-schema columns, as for example used in the PAA to incorporate some fields specific to that data source. Furthermore, we have placed the standard definition itself on GitHub to enable its long-term use as a "living standard", which can be modified and developed further given emerging use-cases and practice from the community, after open discussion and review.

While the Poseidon package format combines various standard formats (e.g. YAML, .tsv, EIGENSTRAT) 426 into its own package specification, opportunities exist to integrate it with larger systems in the Linked Open 427 Data (LOD) world [37]. For example, many of the concepts used in our meta-data definitions exist already in 428 public ontologies, which could be more tightly integrated into our format in the future. Specifically, for example, 429 our Country field in the janno file definition could link to entities in Wikidata [38] or other comparable LOD 430 databases. Our decision for a light-weight flat-file setup has given us leeway to adjust the system exactly to our 431 preferences and the requirements of the field, but integration with the Web of Data thus remains an open tasks 432 for the future. To find partners to establish this uplink in future versions Poseidon is part of the NFDI4Objects 433 initiative (https://www.nfdi4objects.net). 434

Regarding infrastructure, Poseidon is currently very much dependent on GitHub. Following the example of other research data standards [39], all code, data, and issue-tracking is stored there, relying extensively on GitHub's CI system ('GitHub Actions') for automatic code compilation and data validation. The lock-in into GitHub's proprietary platform is slightly mitigated by the open Git format used for code- and data storage, but this is still a serious dependency, including factual drawbacks like a strict bandwidth limitation for large file data downloads.

Beyond these technical questions, finally, Poseidon is also exposed to some of the broader social challenges 441 of scientific data management: Guiding the growth of a healthy community of developers, contributors and 442 maintainers is not trivial. Poseidon currently depends on a small core team consisting of the authors of this 443 paper. A growing number of active collaborators will require committing to a suitable governance system [40]. 444 The work of the core team is currently funded by their employing institution, which also provides computational 445 infrastructure and computing hours. If this commitment gets reduced in the middle- to long-term future, funding 446 may become an increasingly pressing issue – a challenge shared by many research data management projects 447 [41]. 448

Critically the long-term success of Poseidon depends on scientists and generally practitioners in the field 449 of archaeogenetics to reroute resources, not least time, into its development and maintenance, if they consider 450 it valuable for their research and publications. With emerging initiatives like the HAAM Community (https: 451 //haam-community.github.io) and various working groups recently embracing Poseidon-based workflows and 452 first papers referencing it explicitly ([42, 43] and other forthcoming work) we indeed see positive signals towards 453 wider adoption. Regardless of whether this development subsists and a community forms around Poseidon, the 454 Poseidon data format and the software developed for it will remain permanently and openly available for future 455 reference. 456

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## 468 Supplementary Texts

- Supplementary Text 1: Poseidon package specification v2.7.1
- Supplementary Text 2: .janno file details
- Supplementary Text 3: Guide for *trident* v1.4.1.0
- Supplementary Text 4: Guide for *xerxes* v1.0.1.0
- Supplementary Text 5: *xerxes* theoretical background
- Supplementary Text 6: Guide for *qjanno* v1.0.0.0
- Supplementary Text 7: Guide for the *janno* R package v1.0.0
- Supplementary Text 8: Comparison of the public archive content

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