# Reproducible, portable, and efficient ancient genome reconstruction with nf-core/eager

This manuscript (permalink) was automatically generated from apeltzer/eager2-paper@b50dfde on June 3, 2020.

#### **Authors**

#### James A. Fellows Yates

**(D** 0000-0001-5585-6277 **(Q** ify133 **) y** ify133

Department of Archaeogenetics, Max-Planck-Institute for the Science of Human History · Funded by Max Planck Society

#### Thiseas C. Lamnidis

© 0000-0003-4485-8570 · ○ TCLamnidis · У TCLamnidis

Department of Archaeogenetics, Max-Planck-Institute for the Science of Human History · Funded by Max Planck Society

#### Maxime Borry

© 0000-0001-9140-7559 · ○ maxibor · У notmaxib

Department of Archaeogenetics, Max-Planck-Institute for the Science of Human History · Funded by Max Planck Society

#### Aida Andrades Valtueña

Department of Archaeogenetics, Max-Planck-Institute for the Science of Human History · Funded by Max Planck Society

#### • Zandra Fagernäs

D 0000-0003-2667-3556 · ☐ ZandraFagernas · У ZandraSelina

Department of Archaeogenetics, Max-Planck-Institute for the Science of Human History · Funded by Max Planck Society

#### • Stephen Clayton

Department of Archaeogenetics, Max-Planck-Institute for the Science of Human History · Funded by Max Planck Society

#### Maxime U. Garcia

Department of Oncology-Pathology, Karolinska Institutet  $\cdot$  Funded by Barncancerfonden

#### Judith Neukamm

© 0000-0001-8141-566X · ♥ JudithNeukamm · ¥ JudithNeukamm

Palaeogenetics Group, Institute of Evolutionary Medicine, University of Zurich

#### Alexander Peltzer

Quantitative Biology Center (QBiC), Eberhard-Karls-Universität; Translational Medicine & Clinical Pharmacology, Boehringer Ingelheim Pharma GmbH & CO KG, Biberach an der Riss

#### **Abstract**

The broadening utilization of ancient DNA (aDNA) to address anthropological, palaeontological and biological questions is resulting in a rising diversity in the size of laboratories and scale of analyses being performed. In the context of this heterogeneous landscape, we present nf-core/eager, an advanced and entirely redesigned pipeline for the analysis of ancient DNA genomic data. nf-core/eager builds on existing ideas and concepts introduced in the original EAGER pipeline, and improves various aspects of the analysis procedure by building on computational frameworks such as Nextflow and nf-core. The pipeline aims to address three main points: accessibility and adaptability to different research groups and their computing configurations, reproducibility to ensure robust analytical standards in the field, and updating the EAGER pipeline to the latest routine ancient genomic practises. This new version of EAGER has been developed within the nf-core initiative, to ensure high quality software development and maintenance support; contributing to a long-term lifecycle for the pipeline. nf-core/eager will assist in ensuring that ancient DNA sequencing data can be utilised by a diverse range of research groups and fields.

#### Introduction

Ancient DNA has become a widely accepted source of biological data, helping to provide new perspective for a range of fields including archaeology, ecology, cultural heritage, and palaeontology. The utilisation of next-generation-sequencing has allowed the recovery of ancient DNA from a wide variety of sources, including but not limited to, the skeletal remains of animals (1, 2, 3, 4), modern and archaic humans (5, 6, 7), bacteria (8, 9, 10), viruses (11, 12), plants (13, 14), coprolites (15, 16), dental calculus (17, 18), sediments (19, 20), medical slides (21), parchment (22), and most, recently ancient 'chewing gum' (23, 24). Improvement in laboratory protocols to increase yields of otherwise trace amount of DNA has at the same time led to studies that can total hundreds of ancient individuals (25, 26), spanning single (27) to thousands of organisms (17). These differences of disciplines have led to a heterogeneous landscape in terms of types of analyses, and thus various types of computing resources required by different labs. Particularly, this is true for regions of the world where ancient DNA as a field of research is still emerging.

Two previously published and commonly used pipelines in the field are PALEOMIX (28) and EAGER (29). These two pipelines take a similar approach to link together standard tools used for Illumina NGS data processing (quality control, adapter removal/and or merging, mapping, genotyping etc.), but with a specific focus on tools that are designed for or well-suited for ancient DNA (such as bwa aln (30) for ultra-short reads and mapDamage (31) for aDNA characteristic evaluation). Yet, neither of these pipelines have had major updates to bring them in-line with current routine aDNA analyses. Metagenomic screening of off-target genomic reads for pathogens or microbiomes (17, 18) has become particularly common, given it's role in revealing widespread infectious disease and possible epidemics that had previously been undetected in the archaeological record (32, 33, 11, 12). Without easy access to the latest field-established analytical routines, aDNA studies from groups new to the field risk being published without the necessary quality control checks that ensure aDNA authenticity and without yielding the full range of possibilities from their data.

To address these shortcomings, we have completely re-implemented the latest version of the EAGER pipeline in Nextflow (34) (a domain-specific-language (DSL) designed for the construction of omics analysis pipelines), introduced new features, and more flexible pipeline configurations. In addition, the newly named pipeline - nf-core/eager - has been developed in the context of the nf-core community framework (35), which enforces strict guidelines for best-practises in software development.

### **Results and Discussion**

### Scalability, Portability, and Efficiency

The reimplementation of EAGER into Nextflow offers a range of benefits over the original custom pipeline framework.

Firstly, the new framework provides immediate integration of nf-core/eager into various schedulers in POSIX High-Performance-Cluster (HPC) environments, cloud computing resources, and as well as local workstations. This portability allows both small and big labs to run nf-core/eager regardless of the type of computer or cluster size, with minimal effort or configuration, facilitating reproducibility and therefore maintenance of standards within the field. This is further assisted by the in-built compatibility with software environments and containers such as conda, docker and singularity. This ensures exact versions of software are used by a user, regardless of the setup of their local software environment. Another major change with nf-core/eager is that the GUI input is now replaced with a command-line-interface as the primary user interaction mode. This is more compatible and portable with most HPCs (that may not offer X11 forwarding), and is in line with the vast majority of bioinformatic tools. We therefore believe this will not be a hindrance to new researchers from outside computational biology. However there are plans within the nf-core community to provide multiple alternatives in the near future including a command-line wizard and a web-based input GUI.

Secondly, reproducibility is made easier through the use of 'profiles' that can define configuration parameters. These profiles can be managed at different hierarchical levels. HPC-level profiles can specify parameters for the computing environment (schedulers, cache locations, maximum resource etc.), which can be centrally managed to ensure all users of a group use the same settings. Pipeline-level profiles, specifying default parameters for nf-core/eager itself, allow fast access to setups via a single flag for routine analyses, without having to newly configure each run. Compared to the original EAGER that utilised per-FASTQ XML files with hardcoded filepaths, nf-core/eager allows researchers to publish the specific profile used in their runs alongside their publications, to ensure other groups can generate the same results. Usage of profiles also reduces mistakes caused by insufficient 'prose' based reporting of program settings, which may occur in papers written by researchers unfamiliar with informatics. The default nf-core/eager profile uses parameters evaluated in different aDNA specific contexts (e.g. <u>36</u>), and will be updated in each new release as new studies are published.

nf-core/eager provides improved efficiency over the original EAGER pipeline by replacing the sample-by-sample sequential processing with Nextflow's asynchronous parallelisation. This, combined with pre-defined per-process customisation of resource parameters, reduces unnecessary resource allocation that can occur with new users to each step of an NGS data processing pipeline. This is particularly pertinent given the increasing use of centralised HPCs or cloud computing, which often use per-hour cost calculations.

### **Latest aDNA practices**

nf-core/eager follows a similar structural foundation with the original EAGER. Given Illumina short-read FASTQ and/or BAM files, and a reference FASTA file, this can be split into four main stages:

- 1. Preprocessing
  - Sequencing quality control: FastQC (37/)
  - Sequencing artefact clean-up: AdapterRemoval2 (38)
  - Preprocessing statistics generation
- 2. Mapping and post-processing

- Alignment against reference genome: BWA (30, 39, 40), CircularMapper (29)
- Mapping quality filtering: Samtools (41)
- PCR duplicate removal: DeDup (29), Picard MarkDuplicates(42)
- Mapping statistics generation: PreSeq (43), Qualimap2(44)
- 3. aDNA Evaluation and Modification
  - Damage profiling: DamageProfiler (45)
  - aDNA reads selection: PMDTools (46)
  - Damage removal: Bamutils(47)
  - (Human) contamination estimation: ANGSD (48)
- 4. Genotyping and Consensus Sequencing: GATK (42), VCF2Genome (29)

In nf-core/eager, all tools also originally used in EAGER have been updated to latest versions, as available on Bioconda (49) and conda-forge (50) to ensure widespread accessibility and stability of utilized tools. The MapDamage2 (51) and Schmutzi (52) methods have not been carried over to nf-core/eager, the first because a more performant successor method (DamageProfiler) exists, and the latter because a stable release of the method could not be migrated to Bioconda. We anticipate that there will be an updated version of Schmutzi in the near future that will allow us to integrate the method again in nf-core/eager, once a version is released on Bioconda. Support for Bowtie2 (53) will be added in the near future, after consultation with the palaeogenetics community. New tools to the basic workflow include fastp (54) for the removal of poly-G sequencing artefacts that are common in 2-colour Illumina sequencing machines (such as the increasingly popular NextSeq and NovaSeq platforms). We have also included the FreeBayes genotyper as an alternative to the human-focused GATK tools. We have also maintained the possibility of using the now unsupported GATK UnifiedGenotyper, as the GATK HaplotypeCaller performs *de novo* assembly around possible variants, which may not be suitable for low-coverage aDNA data.

We have further extended functionality of the pipeline, primarily focusing on ancient metagenomic analysis to be run alongside the standard genomic analysis against a single reference genome. We have added the ability to screen all off-target reads (from mapping to the supplied reference FASTA) with two metagenomic profilers: MALT (55, 56) and Kraken2 (57). Characterisation of properties of authentic ancient DNA (58) from MALT alignments is carried out with the HOPS pipeline (59). Ancient metagenomic studies sometimes may include comparative samples from living individuals (60). To support open data, whilst respecting data privacy, nf-core/eager includes a 'strip\_fastq' script which creates raw FASTQ files but with reference-genome mapped reads removed. This then allows safe upload of sequencing data to public repositories with identifiable human data removed.

Additional functionality tailored for ancient bacterial genomics includes integration of a SNP alignment generation tool, MultiVCFAnalyzer (§), which allows assessment of levels of cross-mapping from different related taxa to a reference genome - a common challenge in ancient bacteria genome reconstruction (61). Simple coverage statistics of particular annotations (e.g. genes) of an input reference is offered by bedtools (62). When using a human reference genome, nf-core/eager also now can give estimates of the biological sex of a given individual with Sex.DetEERRmine (63). A dedicated 'endogenous DNA' calculator (endorS.py) is also included to provide an approximate level of on-target DNA yield within a sample.

A major upgrade in contrast to the previous EAGER version is that the new pipeline supports processing of complex sequencing strategies for many samples. Given the large amount of sequencing often required to yield sufficient genome coverage from ancient DNA data, palaeogeneticists tend to use multiple (differently treated) libraries or sequencing runs. As an alternative to direct paths to FASTQ or BAM files, the pipeline can also accept a TSV file which includes file paths and additional metadata such as sample name, library name, sequencing lane, colour chemistry, and UDG treatment. This allows simultaneous processing and appropriate merging of heterogeneous data from multiple sequencing runs and/or libraries types.

Finally, the original EAGER tabular report format has been replaced with a much more extensive MultiQC (64) report. The original EAGER pipeline required users to look through many independent output directories and files to make full assessment of their sequencing data. Aggregation of all log files into a single interactive report will assist users in making fuller assessment of their sequencing and analysis runs. Most tools within nf-core/eager have a corresponding MultiQC module to ensure as complete an evaluation as possible.

### Accessibility

Alongside the portable new pipeline report, we have written extensive documentation on all parts of running and interpreting the output of the pipeline. Given that a large fraction of aDNA researchers come from fields outside computational biology, and thus may have limited computational training, we have written documentation that also gives guidance on how to interpret each section of the report, specifically in the context of NGS and aDNA. This includes schematic images of best practices or expected output that published under CC-BY licenses to allow for use in other training material. We hope this open-access resource will make the aDNA discipline more accessible to researchers new to the field, by providing practical and 'applied' knowledge as to how aDNA characteristics translate to downstream analyses.

The development of nf-core/eager in Nextflow and the nf-core initiative will also improve open-source community contributions to the pipeline. While Nextflow is written primarily in Groovy, the Nextflow DSL simplifies a number of concepts to an intermediate level that bioinformaticians without Java/Groovy experience can easily access (regardless of own programming language experience). Furthermore, Nextflow places ubiquitous and more widely known command-line interfaces, such as bash, in a prominent position within the code, rather than custom java code and classes. We hope this will motivate further bug fixes and feature contributions from the community to keep the pipeline updated with standard practises during a longer life-cycle. This will also be supported by the active and welcoming nf-core community who provide general guidance and advice on developing Nextflow and nf-core pipelines.

#### **Conclusion**

nf-core/eager is an efficient, portable, and accessible pipeline for processing ancient DNA genomic data. This re-implementation of EAGER into Nextflow and nf-core will improve reproducibility and inclusion of rapidly increasing ancient DNA datasets, for both large and small laboratories. Extensive documentation also enables newcomers to the field get a practical understanding on how to interpret ancient DNA in the context of NGS data processing. Ultimately, nf-core/eager provides easier access to the latest tools and routine screening analyses commonly used in the field, and sets up the pipeline for staying at the forefront of palaeogenetic analysis.

#### Methods

#### **Installation**

nf-core/eager requires Java, Nextflow and either a functional conda installation *or* Docker/Singularity container installation. A quick installation guide to follow to get started can be found in the *Quickstart* section of the nf-core/eager <u>documentation</u>.

#### Running

After the installation, users can run the pipeline using standard test data by utilizing some of the test profiles we provide (e.g. using Docker):

```
nextflow run nf-core/eager -r 2.1.0 -profile test,docker
```

This will fetch test data automatically, run the pipeline locally with all software tools containerized in a Docker container and store the output of that run in the ./results folder of your current directory.

#### **Profiles**

We utilize a central configuration profile repository to enable users from various institutions to use pipelines on their particular infrastructure <u>more easily</u>. There are multiple resources listed in this repository with information on how to add your own configuration profile with help from the nf-core community.

Users can customize this infrastructure profile by themselves, with the nf-core community, or with their local system administrator to make sure that the pipeline runs successfully, and can then rely on the Nextflow and nf-core framework to ensure compatibility upon further infrastructure changes. For example, in order to run the nf-core/eager pipeline at the Max Planck Institute for the Science of Human History in Jena, users only have to run:

```
nextflow run nf-core/eager -r 2.1.0 -profile shh_cdag,test
```

This runs the testing profile of the nf-core/eager pipeline with parameters specifically adapted to the HPC system at the MPI-SHH. In some cases, similar institutional configs for other institutions may already exist (originally utilised for different nf-core pipelines), so users need not write their own.

#### **Inputs**

The pipeline can be started using (raw) FASTQ files from sequencing or pre-mapped BAM files. Additionally, the pipeline requires a FASTA reference genome. If users have complex setups, e.g. multiple sequencing lanes that require merging of files for example, the pipeline can be supplied with a tabular separated value (TSV) file to enable such complex data handling.

#### Monitoring

Users can either monitor their pipeline execution with the messages Nextflow prints to the console while running, or utilize projects such as <a href="Nextflow Tower">Nextflow Tower</a> to monitor their analysis pipeline during runtime.

#### **Output**

The pipeline produces several dozen output files in various file formats, with a more detailed listing available in the user documentation. This includes metrics, statistical analysis data and standardized output files (BAM, VCF) for close inspection and further downstream analysis, as well as a MultiQC report. If an emailing daemon is set up on the server, the latter can even be emailed to users automatically, when starting the pipeline with a dedicated option ( --email you@yourdomain.org ).

### Data and software availability

All code is available on github at <a href="https://github.com/nf-core/eager">https://github.com/nf-core/eager</a> and archived with Zenodo under the DOI <a href="https://github.com/nf-core/eager">10.5281/zenodo.1465061</a>. All test data is from the ENA public repository available under ENA IDs: **FIXME** 

### **Competing Interests**

No competing interests are declared.

### **Acknowledgements**

We thank the nf-core community for general support and suggestions during the writing of the pipeline. We also thank Arielle Munters, Hester van Schalkwyk, Irina Velsko, Katerine Eaton, Luc Venturini, Marcel Keller, Pierre Lindenbaum, Pontus Skoglund, Raphael Eisenhofer, Torsten Günter, and Kevin Lord for bug reports and feature suggestions. We are grateful to the members of the Department of Archaeogenetics at the Max Planck Institute for the Science of Human History who performed beta testing of the pipeline. We thank the aDNA twitter community for responding to polls regarding design decisions of the pipeline.

We also acknowledge Christina Warinner, Stephan Schiffels and the Max Planck Society who provided funds for travel to nf-core events.

### 1. Complete Genomes Reveal Signatures of Demographic and Genetic Declines in the Woolly Mammoth

Eleftheria Palkopoulou, Swapan Mallick, Pontus Skoglund, Jacob Enk, Nadin Rohland, Heng Li, Ayça Omrak, Sergey Vartanyan, Hendrik Poinar, Anders Götherström, ... Love Dalén *Current Biology* (2015-05) https://doi.org/34d

DOI: <u>10.1016/j.cub.2015.04.007</u> · PMID: <u>25913407</u> · PMCID: <u>PMC4439331</u>

### 2. Recalibrating Equus evolution using the genome sequence of an early Middle Pleistocene horse

Ludovic Orlando, Aurélien Ginolhac, Guojie Zhang, Duane Froese, Anders Albrechtsen, Mathias Stiller, Mikkel Schubert, Enrico Cappellini, Bent Petersen, Ida Moltke, ... Eske Willerslev *Nature* (2013-06-26) https://doi.org/q7n

DOI: 10.1038/nature12323 · PMID: 23803765

#### 3. Ancient pigs reveal a near-complete genomic turnover following their introduction to Europe

Laurent A. F. Frantz, James Haile, Audrey T. Lin, Amelie Scheu, Christina Geörg, Norbert Benecke, Michelle Alexander, Anna Linderholm, Victoria E. Mullin, Kevin G. Daly, ... Greger Larson *Proceedings of the National Academy of Sciences* (2019-08-27) <a href="https://doi.org/gf9hnf">https://doi.org/gf9hnf</a>
DOI: <a href="https://doi.org/gf9hnf">10.1073/pnas.1901169116</a> · PMID: <a href="https://doi.org/gf9hnf">31405970</a> · PMCID: <a href="https://doi.org/gf9hnf">PMC6717267</a>

#### 4. Ancient DNA reveals the Arctic origin of Viking Age cod from Haithabu, Germany

Bastiaan Star, Sanne Boessenkool, Agata T. Gondek, Elena A. Nikulina, Anne Karin Hufthammer, Christophe Pampoulie, Halvor Knutsen, Carl André, Heidi M. Nistelberger, Jan Dierking, ... James H. Barrett

*Proceedings of the National Academy of Sciences* (2017-08-22) <a href="https://doi.org/gbt8b2">https://doi.org/gbt8b2</a>
DOI: <a href="https://doi.org/gbt8b2">10.1073/pnas.1710186114</a> · PMID: <a href="https://doi.org/gbt8b2">28784790</a> · PMCID: <a href="https://doi.org/gbt8b2">PMCID: PMC5576834</a>

#### 5. 137 ancient human genomes from across the Eurasian steppes

Peter de Barros Damgaard, Nina Marchi, Simon Rasmussen, Michaël Peyrot, Gabriel Renaud, Thorfinn Korneliussen, J. Víctor Moreno-Mayar, Mikkel Winther Pedersen, Amy Goldberg, Emma Usmanova, ... Eske Willerslev

Nature (2018-05-09) https://doi.org/gd8hs5

DOI: <u>10.1038/s41586-018-0094-2</u> · PMID: <u>29743675</u>

#### 6. A High-Coverage Genome Sequence from an Archaic Denisovan Individual

M. Meyer, M. Kircher, M.-T. Gansauge, H. Li, F. Racimo, S. Mallick, J. G. Schraiber, F. Jay, K. Prufer, C. de Filippo, ... S. Paabo

Science (2012-08-30) https://doi.org/q8p

DOI: <u>10.1126/science.1224344</u> · PMID: <u>22936568</u> · PMCID: <u>PMC3617501</u>

#### 7. The genome of the offspring of a Neanderthal mother and a Denisovan father

Viviane Slon, Fabrizio Mafessoni, Benjamin Vernot, Cesare de Filippo, Steffi Grote, Bence Viola, Mateja Hajdinjak, Stéphane Peyrégne, Sarah Nagel, Samantha Brown, ... Svante Pääbo *Nature* (2018-08-22) <a href="https://doi.org/cs64">https://doi.org/cs64</a>

DOI: <u>10.1038/s41586-018-0455-x</u> · PMID: <u>30135579</u> · PMCID: <u>PMC6130845</u>

# 8. Pre-Columbian mycobacterial genomes reveal seals as a source of New World human tuberculosis

Kirsten I. Bos, Kelly M. Harkins, Alexander Herbig, Mireia Coscolla, Nico Weber, Iñaki Comas, Stephen A. Forrest, Josephine M. Bryant, Simon R. Harris, Verena J. Schuenemann, ... Johannes Krause

Nature (2014-08-20) https://doi.org/f6nk4g

DOI: 10.1038/nature13591 · PMID: 25141181 · PMCID: PMC4550673

# 9. Integrative approach using *Yersinia pestis* genomes to revisit the historical landscape of plague during the Medieval Period

Amine Namouchi, Meriam Guellil, Oliver Kersten, Stephanie Hänsch, Claudio Ottoni, Boris V. Schmid, Elsa Pacciani, Luisa Quaglia, Marco Vermunt, Egil L. Bauer, ... Barbara Bramanti *Proceedings of the National Academy of Sciences* (2018-12-11) <a href="https://doi.org/ggfn3h">https://doi.org/ggfn3h</a>
DOI: <a href="https://doi.org/ggfn3h">10.1073/pnas.1812865115</a> · PMID: <a href="https://doi.org/ggfn3h">30478041</a> · PMCID: <a href="https://doi.org/ggfn3h">PMC6294933</a>

#### 10. Ancient genomes reveal a high diversity of Mycobacterium leprae in medieval Europe

Verena J. Schuenemann, Charlotte Avanzi, Ben Krause-Kyora, Alexander Seitz, Alexander Herbig, Sarah Inskip, Marion Bonazzi, Ella Reiter, Christian Urban, Dorthe Dangvard Pedersen, ... Johannes Krause

PLOS Pathogens (2018-05-10) <a href="https://doi.org/gdrj4v">https://doi.org/gdrj4v</a>

DOI: <u>10.1371/journal.ppat.1006997</u> · PMID: <u>29746563</u> · PMCID: <u>PMC5944922</u>

#### 11. Ancient hepatitis B viruses from the Bronze Age to the Medieval period

Barbara Mühlemann, Terry C. Jones, Peter de Barros Damgaard, Morten E. Allentoft, Irina Shevnina, Andrey Logvin, Emma Usmanova, Irina P. Panyushkina, Bazartseren Boldgiv, Tsevel Bazartseren, ... Eske Willerslev

Nature (2018-05-09) <a href="https://doi.org/gddxvj">https://doi.org/gddxvj</a>

DOI: <u>10.1038/s41586-018-0097-z</u> · PMID: <u>29743673</u>

#### 12. Neolithic and medieval virus genomes reveal complex evolution of hepatitis B

Ben Krause-Kyora, Julian Susat, Felix M Key, Denise Kühnert, Esther Bosse, Alexander Immel, Christoph Rinne, Sabin-Christin Kornell, Diego Yepes, Sören Franzenburg, ... Johannes Krause *eLife* (2018-05-10) <a href="https://doi.org/gdhck2">https://doi.org/gdhck2</a>

DOI: 10.7554/elife.36666 · PMID: 29745896 · PMCID: PMC6008052

# 13. Ancient reveals the timing and persistence of organellar genetic bottlenecks over 3,000 years of sunflower domestication and improvement

Nathan Wales, Melis Akman, Ray H. B. Watson, Fátima Sánchez Barreiro, Bruce D. Smith, Kristen J. Gremillion, M. Thomas P. Gilbert, Benjamin K. Blackman

Evolutionary Applications (2018-02-13) https://doi.org/gf568v

DOI: <u>10.1111/eva.12594</u> · PMID: <u>30622634</u> · PMCID: <u>PMC6304678</u>

#### 14. The origins and adaptation of European potatoes reconstructed from historical genomes

Rafal M. Gutaker, Clemens L. Weiß, David Ellis, Noelle L. Anglin, Sandra Knapp, José Luis Fernández-Alonso, Salomé Prat, Hernán A. Burbano

Nature Ecology & Evolution (2019-06-24) https://doi.org/ggxkk8

DOI: <u>10.1038/s41559-019-0921-3</u> · PMID: <u>31235927</u>

### 15. The Prevotella copri Complex Comprises Four Distinct Clades Underrepresented in Westernized Populations

Adrian Tett, Kun D. Huang, Francesco Asnicar, Hannah Fehlner-Peach, Edoardo Pasolli, Nicolai Karcher, Federica Armanini, Paolo Manghi, Kevin Bonham, Moreno Zolfo, ... Nicola Segata *Cell Host & Microbe* (2019-11) https://doi.org/ggc9dc

DOI: 10.1016/j.chom.2019.08.018 · PMID: 31607556 · PMCID: PMC6854460

# 16. CoproID predicts the source of coprolites and paleofeces using microbiome composition and host DNA content

Maxime Borry, Bryan Cordova, Angela Perri, Marsha Wibowo, Tanvi Prasad Honap, Jada Ko, Jie Yu,

Kate Britton, Linus Girdland-Flink, Robert C. Power, ... Christina Warinner

PeerJ (2020-04-17) https://doi.org/dr8x

DOI: 10.7717/peerj.9001 · PMID: 32337106 · PMCID: PMC7169968

#### 17. Pathogens and host immunity in the ancient human oral cavity

Christina Warinner, João F Matias Rodrigues, Rounak Vyas, Christian Trachsel, Natallia Shved, Jonas Grossmann, Anita Radini, Y Hancock, Raul Y Tito, Sarah Fiddyment, ... Enrico Cappellini *Nature Genetics* (2014-02-23) <a href="https://doi.org/r4n">https://doi.org/r4n</a>

DOI: <u>10.1038/ng.2906</u> · PMID: <u>24562188</u> · PMCID: <u>PMC3969750</u>

#### 18. Neanderthal behaviour, diet, and disease inferred from ancient DNA in dental calculus

Laura S. Weyrich, Sebastian Duchene, Julien Soubrier, Luis Arriola, Bastien Llamas, James Breen, Alan G. Morris, Kurt W. Alt, David Caramelli, Veit Dresely, ... Alan Cooper

Nature (2017-03-08) https://doi.org/f9szrm

DOI: <u>10.1038/nature21674</u> · PMID: <u>28273061</u>

#### 19. Fifty thousand years of Arctic vegetation and megafaunal diet

Eske Willerslev, John Davison, Mari Moora, Martin Zobel, Eric Coissac, Mary E. Edwards, Eline D. Lorenzen, Mette Vestergård, Galina Gussarova, James Haile, ... Pierre Taberlet *Nature* (2014-02-05) <a href="https://doi.org/f2zr4s">https://doi.org/f2zr4s</a>

DOI: 10.1038/nature12921 · PMID: 24499916

#### 20. Neandertal and Denisovan DNA from Pleistocene sediments

Viviane Slon, Charlotte Hopfe, Clemens L. Weiß, Fabrizio Mafessoni, Marco de la Rasilla, Carles Lalueza-Fox, Antonio Rosas, Marie Soressi, Monika V. Knul, Rebecca Miller, ... Matthias Meyer *Science* (2017-05-12) https://doi.org/b6jd

DOI: 10.1126/science.aam9695 · PMID: 28450384

#### 21. Plasmodium vivax Malaria Viewed through the Lens of an Eradicated European Strain

Lucy van Dorp, Pere Gelabert, Adrien Rieux, Marc de Manuel, Toni de-Dios, Shyam Gopalakrishnan, Christian Carøe, Marcela Sandoval-Velasco, Rosa Fregel, Iñigo Olalde, ... Carles Lalueza-Fox *Molecular Biology and Evolution* (2020-03) <a href="https://doi.org/ggqzq2">https://doi.org/ggqzq2</a>

DOI: <u>10.1093/molbev/msz264</u> · PMID: <u>31697387</u> · PMCID: <u>PMC7038659</u>

### 22. Paging through history: parchment as a reservoir of ancient DNA for next generation sequencing

M. D. Teasdale, N. L. van Doorn, S. Fiddyment, C. C. Webb, T. O'Connor, M. Hofreiter, M. J. Collins, D. G. Bradley

*Philosophical Transactions of the Royal Society B: Biological Sciences* (2015-01-19) <a href="https://doi.org/gggzq3">https://doi.org/gggzq3</a>

DOI: 10.1098/rstb.2013.0379 · PMID: 25487331 · PMCID: PMC4275887

#### 23. A 5700 year-old human genome and oral microbiome from chewed birch pitch

Theis Z. T. Jensen, Jonas Niemann, Katrine Højholt Iversen, Anna K. Fotakis, Shyam Gopalakrishnan, Åshild J. Vågene, Mikkel Winther Pedersen, Mikkel-Holger S. Sinding, Martin R. Ellegaard, Morten E. Allentoft, ... Hannes Schroeder

Nature Communications (2019-12-17) <a href="https://doi.org/ggfm6x">https://doi.org/ggfm6x</a>

DOI: 10.1038/s41467-019-13549-9 · PMID: 31848342 · PMCID: PMC6917805

# 24. Ancient DNA from mastics solidifies connection between material culture and genetics of mesolithic hunter-gatherers in Scandinavia

Natalija Kashuba, Emrah Kırdök, Hege Damlien, Mikael A. Manninen, Bengt Nordqvist, Per Persson, Anders Götherström

Communications Biology (2019-05-15) https://doi.org/gggzgz

DOI: 10.1038/s42003-019-0399-1 · PMID: 31123709 · PMCID: PMC6520363

#### 25. The Beaker phenomenon and the genomic transformation of northwest Europe

Iñigo Olalde, Selina Brace, Morten E. Allentoft, Ian Armit, Kristian Kristiansen, Thomas Booth, Nadin Rohland, Swapan Mallick, Anna Szécsényi-Nagy, Alissa Mittnik, ... David Reich

Nature (2018-02-21) https://doi.org/gcx74m

DOI: <u>10.1038/nature25738</u> · PMID: <u>29466337</u> · PMCID: <u>PMC5973796</u>

#### 26. The genomic history of southeastern Europe

Iain Mathieson, Songül Alpaslan-Roodenberg, Cosimo Posth, Anna Szécsényi-Nagy, Nadin Rohland, Swapan Mallick, Iñigo Olalde, Nasreen Broomandkhoshbacht, Francesca Candilio, Olivia Cheronet, ... David Reich

*Nature* (2018-02-21) <a href="https://doi.org/gc2n9h">https://doi.org/gc2n9h</a>

DOI: <u>10.1038/nature25778</u> · PMID: <u>29466330</u> · PMCID: <u>PMC6091220</u>

#### 27. A draft genome of Yersinia pestis from victims of the Black Death

Kirsten I. Bos, Verena J. Schuenemann, G. Brian Golding, Hernán A. Burbano, Nicholas Waglechner, Brian K. Coombes, Joseph B. McPhee, Sharon N. DeWitte, Matthias Meyer, Sarah Schmedes, ... Johannes Krause

Nature (2011-10-12) https://doi.org/fk87wk

DOI: <u>10.1038/nature10549</u> · PMID: <u>21993626</u> · PMCID: <u>PMC3690193</u>

# 28. Characterization of ancient and modern genomes by SNP detection and phylogenomic and metagenomic analysis using PALEOMIX

Mikkel Schubert, Luca Ermini, Clio Der Sarkissian, Hákon Jónsson, Aurélien Ginolhac, Robert Schaefer, Michael D Martin, Ruth Fernández, Martin Kircher, Molly McCue, ... Ludovic Orlando *Nature Protocols* (2014-04-10) <a href="https://doi.org/f5x3qm">https://doi.org/f5x3qm</a>

DOI: 10.1038/nprot.2014.063 · PMID: 24722405

#### 29. EAGER: efficient ancient genome reconstruction

Alexander Peltzer, Günter Jäger, Alexander Herbig, Alexander Seitz, Christian Kniep, Johannes Krause, Kay Nieselt

Genome Biology (2016-03-31) https://doi.org/gggzpk

DOI: <u>10.1186/s13059-016-0918-z</u> · PMID: <u>27036623</u> · PMCID: <u>PMC4815194</u>

#### 30. Fast and accurate short read alignment with Burrows-Wheeler transform

H. Li, R. Durbin

Bioinformatics (2009-05-18) https://doi.org/dqt59j

DOI: <u>10.1093/bioinformatics/btp324</u> · PMID: <u>19451168</u> · PMCID: <u>PMC2705234</u>

#### 31. mapDamage: testing for damage patterns in ancient DNA sequences

Aurelien Ginolhac, Morten Rasmussen, M. Thomas P. Gilbert, Eske Willerslev, Ludovic Orlando *Bioinformatics* (2011-08-01) https://doi.org/cn45v7

DOI: <u>10.1093/bioinformatics/btr347</u> · PMID: <u>21659319</u>

#### 32. Early Divergent Strains of Yersinia pestis in Eurasia 5,000 Years Ago

Simon Rasmussen, Morten Erik Allentoft, Kasper Nielsen, Ludovic Orlando, Martin Sikora, Karl-Göran Sjögren, Anders Gorm Pedersen, Mikkel Schubert, Alex Van Dam, Christian Moliin Outzen Kapel, ... Eske Willerslev

Cell (2015-10) https://doi.org/f3mxqd

DOI: 10.1016/j.cell.2015.10.009 · PMID: 26496604 · PMCID: PMC4644222

#### 33. The Stone Age Plague and Its Persistence in Eurasia

Aida Andrades Valtueña, Alissa Mittnik, Felix M. Key, Wolfgang Haak, Raili Allmäe, Andrej Belinskij, Mantas Daubaras, Michal Feldman, Rimantas Jankauskas, Ivor Janković, ... Johannes Krause *Current Biology* (2017-12) https://doi.org/cgmv

DOI: 10.1016/j.cub.2017.10.025 · PMID: 29174893

#### 34. Nextflow enables reproducible computational workflows

Paolo Di Tommaso, Maria Chatzou, Evan W Floden, Pablo Prieto Barja, Emilio Palumbo, Cedric Notredame

Nature Biotechnology (2017-04-11) https://doi.org/gfj52z

DOI: 10.1038/nbt.3820 · PMID: 28398311

#### 35. The nf-core framework for community-curated bioinformatics pipelines

Philip A. Ewels, Alexander Peltzer, Sven Fillinger, Harshil Patel, Johannes Alneberg, Andreas Wilm, Maxime Ulysse Garcia, Paolo Di Tommaso, Sven Nahnsen

Nature Biotechnology (2020-02-13) https://doi.org/ggk3qh

DOI: 10.1038/s41587-020-0439-x · PMID: 32055031

#### 36. Improving ancient DNA read mapping against modern reference genomes

Mikkel Schubert, Aurelien Ginolhac, Stinus Lindgreen, John F Thompson, Khaled AS AL-Rasheid, Eske Willerslev, Anders Krogh, Ludovic Orlando

BMC Genomics (2012) https://doi.org/gb3ff7

DOI: <u>10.1186/1471-2164-13-178</u> · PMID: <u>22574660</u> · PMCID: <u>PMC3468387</u>

# 37. Babraham Bioinformatics - FastQC A Quality Control tool for High Throughput Sequence Data <a href="https://www.bioinformatics.babraham.ac.uk/projects/fastgc/">https://www.bioinformatics.babraham.ac.uk/projects/fastgc/</a>

#### 38. AdapterRemoval v2: rapid adapter trimming, identification, and read merging

Mikkel Schubert, Stinus Lindgreen, Ludovic Orlando

BMC Research Notes (2016-02-12) https://doi.org/gfzqhb

DOI: <u>10.1186/s13104-016-1900-2</u> · PMID: <u>26868221</u> · PMCID: <u>PMC4751634</u>

#### 39. Fast and accurate long-read alignment with Burrows-Wheeler transform

Heng Li, Richard Durbin

Bioinformatics (2010-03-01) https://doi.org/cm27kg

DOI: 10.1093/bioinformatics/btp698 · PMID: 20080505 · PMCID: PMC2828108

#### 40. Aligning sequence reads, clone sequences and assembly contigs with BWA-MEM

Heng Li

arXiv (2013-05-28) https://arxiv.org/abs/1303.3997

#### 41. The Sequence Alignment/Map format and SAMtools

H. Li, B. Handsaker, A. Wysoker, T. Fennell, J. Ruan, N. Homer, G. Marth, G. Abecasis, R. Durbin, 1000 Genome Project Data Processing Subgroup

Bioinformatics (2009-06-08) https://doi.org/ff6426

DOI: <u>10.1093/bioinformatics/btp352</u> · PMID: <u>19505943</u> · PMCID: <u>PMC2723002</u>

### 42. The Genome Analysis Toolkit: A MapReduce framework for analyzing next-generation DNA sequencing data

A. McKenna, M. Hanna, E. Banks, A. Sivachenko, K. Cibulskis, A. Kernytsky, K. Garimella, D. Altshuler, S. Gabriel, M. Daly, M. A. DePristo

Genome Research (2010-07-19) https://doi.org/bnzbn6

DOI: 10.1101/gr.107524.110 · PMID: 20644199 · PMCID: PMC2928508

#### 43. Predicting the molecular complexity of sequencing libraries

Timothy Daley, Andrew D Smith

Nature Methods (2013-02-24) https://doi.org/gfx6f5

DOI: 10.1038/nmeth.2375 · PMID: 23435259 · PMCID: PMC3612374

#### 44. Qualimap 2: advanced multi-sample quality control for high-throughput sequencing data

Konstantin Okonechnikov, Ana Conesa, Fernando García-Alcalde

Bioinformatics (2015-10-01) https://doi.org/ggxrmx

DOI: <u>10.1093/bioinformatics/btv566</u> · PMID: <u>26428292</u> · PMCID: <u>PMC4708105</u>

#### 45. Integrative-Transcriptomics/DamageProfiler: DamageProfiler v0.4.9

Judith Neukamm, Alexander Peltzer

Zenodo (2019-11-29) https://doi.org/ggxrmz

DOI: 10.5281/zenodo.3557708

### 46. Separating endogenous ancient DNA from modern day contamination in a Siberian Neandertal

Pontus Skoglund, Bernd H. Northoff, Michael V. Shunkov, Anatoli P. Derevianko, Svante Pääbo, Johannes Krause, Mattias Jakobsson

Proceedings of the National Academy of Sciences (2014-02-11) https://doi.org/f2z5sw

DOI: <u>10.1073/pnas.1318934111</u> · PMID: <u>24469802</u> · PMCID: <u>PMC3926038</u>

### 47. An efficient and scalable analysis framework for variant extraction and refinement from population-scale DNA sequence data

Goo Jun, Mary Kate Wing, Gonçalo R. Abecasis, Hyun Min Kang

Genome Research (2015-06) https://doi.org/f7dz2d

DOI: <u>10.1101/gr.176552.114</u> · PMID: <u>25883319</u> · PMCID: <u>PMC4448687</u>

#### 48. ANGSD: Analysis of Next Generation Sequencing Data

Thorfinn Sand Korneliussen, Anders Albrechtsen, Rasmus Nielsen

BMC Bioinformatics (2014-11-25) https://doi.org/gb8wpz

DOI: 10.1186/s12859-014-0356-4 · PMID: 25420514 · PMCID: PMC4248462

#### 49. Bioconda: sustainable and comprehensive software distribution for the life sciences

Björn Grüning, Ryan Dale, Andreas Sjödin, Brad A. Chapman, Jillian Rowe, Christopher H. Tomkins-Tinch, Renan Valieris, Johannes Köster, The Bioconda Team

Nature Methods (2018-07-02) https://doi.org/gd2xzp

DOI: 10.1038/s41592-018-0046-7 · PMID: 29967506

#### 50. conda-forge | community driven packaging for conda https://conda-forge.org/

#### 51. mapDamage2.0: fast approximate Bayesian estimates of ancient DNA damage parameters

Hákon Jónsson, Aurélien Ginolhac, Mikkel Schubert, Philip L. F. Johnson, Ludovic Orlando *Bioinformatics* (2013-07) <a href="https://doi.org/gb5g2t">https://doi.org/gb5g2t</a>

DOI: 10.1093/bioinformatics/btt193 · PMID: 23613487 · PMCID: PMC3694634

# 52. Schmutzi: estimation of contamination and endogenous mitochondrial consensus calling for ancient DNA

Gabriel Renaud, Viviane Slon, Ana T. Duggan, Janet Kelso

Genome Biology (2015-10-12) https://doi.org/f72mvg

DOI: 10.1186/s13059-015-0776-0 · PMID: 26458810 · PMCID: PMC4601135

#### 53. Fast gapped-read alignment with Bowtie 2

Ben Langmead, Steven L Salzberg

Nature Methods (2012-03-04) https://doi.org/gd2xzn

DOI: 10.1038/nmeth.1923 · PMID: 22388286 · PMCID: PMC3322381

#### 54. fastp: an ultra-fast all-in-one FASTQ preprocessor

Shifu Chen, Yanqing Zhou, Yaru Chen, Jia Gu

Bioinformatics (2018-09-01) https://doi.org/gd9mrb

DOI: 10.1093/bioinformatics/bty560 · PMID: 30423086 · PMCID: PMC6129281

### 55. MALT: Fast alignment and analysis of metagenomic DNA sequence data applied to the Tyrolean Iceman

Alexander Herbig, Frank Maixner, Kirsten I. Bos, Albert Zink, Johannes Krause, Daniel H. Huson *bioRxiv* (2016-04-27) <a href="https://doi.org/ggxkk9">https://doi.org/ggxkk9</a>

DOI: <u>10.1101/050559</u>

#### 56. Salmonella enterica genomes from victims of a major sixteenth-century epidemic in Mexico

Åshild J. Vågene, Alexander Herbig, Michael G. Campana, Nelly M. Robles García, Christina Warinner, Susanna Sabin, Maria A. Spyrou, Aida Andrades Valtueña, Daniel Huson, Noreen Tuross, ... Johannes Krause

Nature Ecology & Evolution (2018-01-15) https://doi.org/ggxkk7

DOI: 10.1038/s41559-017-0446-6 · PMID: 29335577

#### 57. Improved metagenomic analysis with Kraken 2

Derrick E. Wood, Jennifer Lu, Ben Langmead

Genome Biology (2019-11-28) https://doi.org/ggfk55

DOI: <u>10.1186/s13059-019-1891-0</u> · PMID: <u>31779668</u> · PMCID: <u>PMC6883579</u>

#### 58. Patterns of damage in genomic DNA sequences from a Neandertal

A. W. Briggs, U. Stenzel, P. L. F. Johnson, R. E. Green, J. Kelso, K. Prufer, M. Meyer, J. Krause, M. T. Ronan, M. Lachmann, S. Paabo

Proceedings of the National Academy of Sciences (2007-08-21) https://doi.org/bs4w7h

DOI: 10.1073/pnas.0704665104 · PMID: 17715061 · PMCID: PMC1976210

#### 59. HOPS: automated detection and authentication of pathogen DNA in archaeological remains

Ron Hübler, Felix M. Key, Christina Warinner, Kirsten I. Bos, Johannes Krause, Alexander Herbig *Genome Biology* (2019-12-16) https://doi.org/ggxkmb

DOI: 10.1186/s13059-019-1903-0 · PMID: 31842945 · PMCID: PMC6913047

### 60. Microbial differences between dental plaque and historic dental calculus are related to oral biofilm maturation stage

Irina M. Velsko, James A. Fellows Yates, Franziska Aron, Richard W. Hagan, Laurent A. F. Frantz, Louise Loe, Juan Bautista Rodriguez Martinez, Eros Chaves, Chris Gosden, Greger Larson, Christina Warinner

Microbiome (2019-07-06) https://doi.org/ggxkmc

DOI: <u>10.1186/s40168-019-0717-3</u> · PMID: <u>31279340</u> · PMCID: <u>PMC6612086</u>

#### 61. A Robust Framework for Microbial Archaeology

Christina Warinner, Alexander Herbig, Allison Mann, James A. Fellows Yates, Clemens L. Weiß, Hernán A. Burbano, Ludovic Orlando, Johannes Krause

Annual Review of Genomics and Human Genetics (2017-08-31) <a href="https://doi.org/gf5wqv">https://doi.org/gf5wqv</a>

DOI: <u>10.1146/annurev-genom-091416-035526</u> · PMID: <u>28460196</u> · PMCID: <u>PMC5581243</u>

#### 62. BEDTools: a flexible suite of utilities for comparing genomic features

Aaron R. Quinlan, Ira M. Hall

Bioinformatics (2010-03-15) <a href="https://doi.org/cmrms3">https://doi.org/cmrms3</a>

DOI: 10.1093/bioinformatics/btg033 · PMID: 20110278 · PMCID: PMC2832824

#### 63. Ancient Fennoscandian genomes reveal origin and spread of Siberian ancestry in Europe

Thiseas C. Lamnidis, Kerttu Majander, Choongwon Jeong, Elina Salmela, Anna Wessman, Vyacheslav Moiseyev, Valery Khartanovich, Oleg Balanovsky, Matthias Ongyerth, Antje Weihmann, ... Stephan Schiffels

Nature Communications (2018-11-27) <a href="https://doi.org/ggxkk6">https://doi.org/ggxkk6</a>

DOI: <u>10.1038/s41467-018-07483-5</u> · PMID: <u>30479341</u> · PMCID: <u>PMC6258758</u>

#### 64. MultiQC: summarize analysis results for multiple tools and samples in a single report

Philip Ewels, Måns Magnusson, Sverker Lundin, Max Käller *Bioinformatics* (2016-10-01) <a href="https://doi.org/f3s996">https://doi.org/f3s996</a>

DOI: 10.1093/bioinformatics/btw354 · PMID: 27312411 · PMCID: PMC5039924