Topological data analysis of SNP array data exposes the genetic differentiation between lberians and Canary Islanders









José M. Lorenzo-Salazar¹, Ana Díaz-de Usera¹, **Adrián Muñoz-Barrera¹**, Luis A. Rubio-Rodríguez¹, Beatriz Guillen-Guio², Almudena Corrales^{2,3}, Itahisa Marcelino-Rodríguez², David Comas⁴, Rafaela González-Montelongo², Santos Alonso⁵, Carlos Flores^{1,2,3}

¹Genomics Division, Instituto Tecnológico y de Energías Renovables (ITER), Santa Cruz de Tenerife, Spain; ²Research Unit, Hospital Universitario N.S. de Candelaria, Universidad de La Laguna, Santa Cruz de Tenerife, Spain; ³CIBER de Enfermedades Respiratorias, Instituto de Salud Carlos III, Madrid, Spain; ⁴Department of Experimental and Health Sciences, Institut de Biologia Evolutiva (CSIC-UPF), Universitat Pompeau Fabra, Barcelona, Spain; ⁵Department of Genetics, Phsyical Anthropology and Animal Physiology, University of the Basque Country UPV/EHU, Leioa, Bizkaia, Spain.

Introduction

Unraveling global patterns of human genetic variation is of main interest for the scientific community. The 1000 Genomes Project (1KGP) highlighted that a typical genome differs roughly at 4.1-5.0 million positions from the reference human genome. The population from the Canary Islands (CAN), a Spanish archipelago situated in the Atlantic Ocean 100 Km off the NW African coast, has a unique genetic pool due to isolation, local adaptation and recent admixture of Europeans (EUR), North-Africans (NAF) and sub-Saharan Africans (SSA)².

Assessing the genetic structure of populations often requires a multidimensionality reduction approach, typically assessed by Principal Component (PC) Analysis (PCA)³. However, such procedure most commonly focuses on few main dimensions limiting the possibilities to excavate fine-grained strata. Here we used Topological Data Analysis (TDA)⁴ to explore the genetic dissimilarity of Iberians and Canary Islanders by embedding high-dimensionality SNP array and whole-genome sequencing (WGS) data to explore the genetic differentiation between populations into a low-dimensional space. New WGS data from NAF were also included for comparative purposes.

Materials and Methods

<u>Sample data</u>: WGS data from 46 Canary Islanders (CAN) and 23 North Africans (NAF) obtained with a HiSeq 4000 (Illumina) to an average of 30x, together with 740 subjects genotyped for the Spain Biobank Array (SBA, Thermo-Fisher Scientific). Additionally, data from 478 individuals from 1KGP were included as reference of EUR and SSA populations. All individuals were unrelated.

Quality control: See the procedures shown on the **Workflow** diagram below.

Statistical analyses: PCA and TDA were assessed on WGS and SBA data using PLINK⁵ v1.9 and umap⁴ v0.2.0 library for R.

Workflow

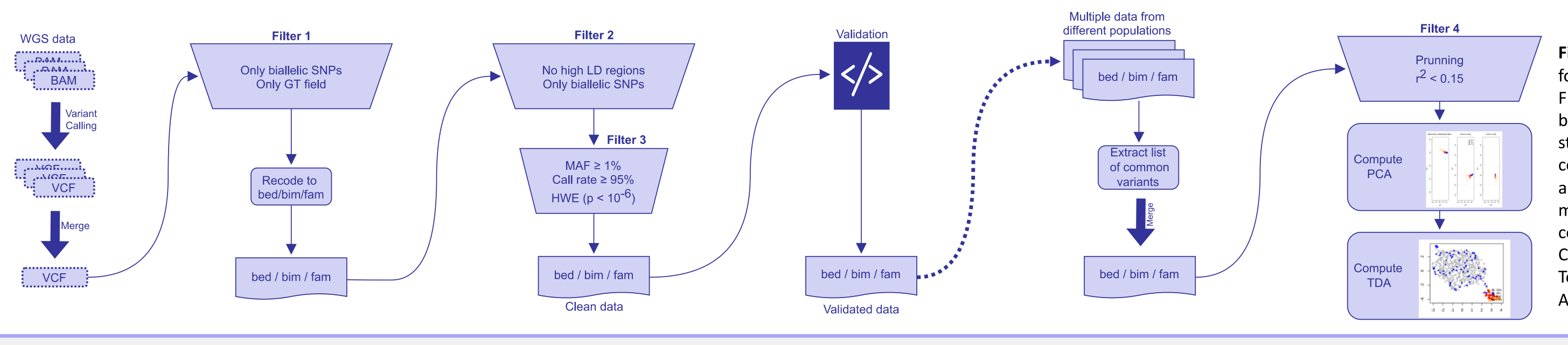
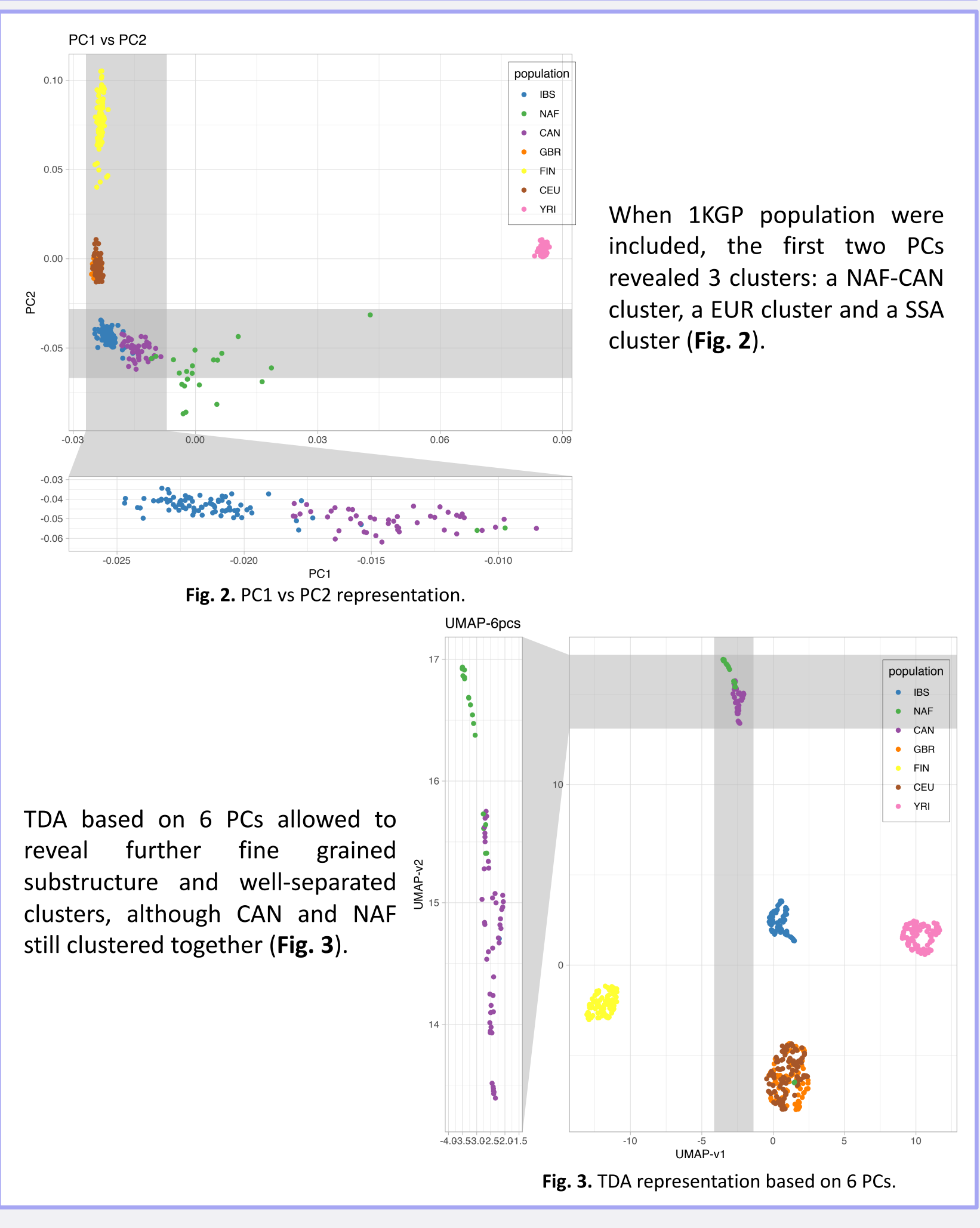


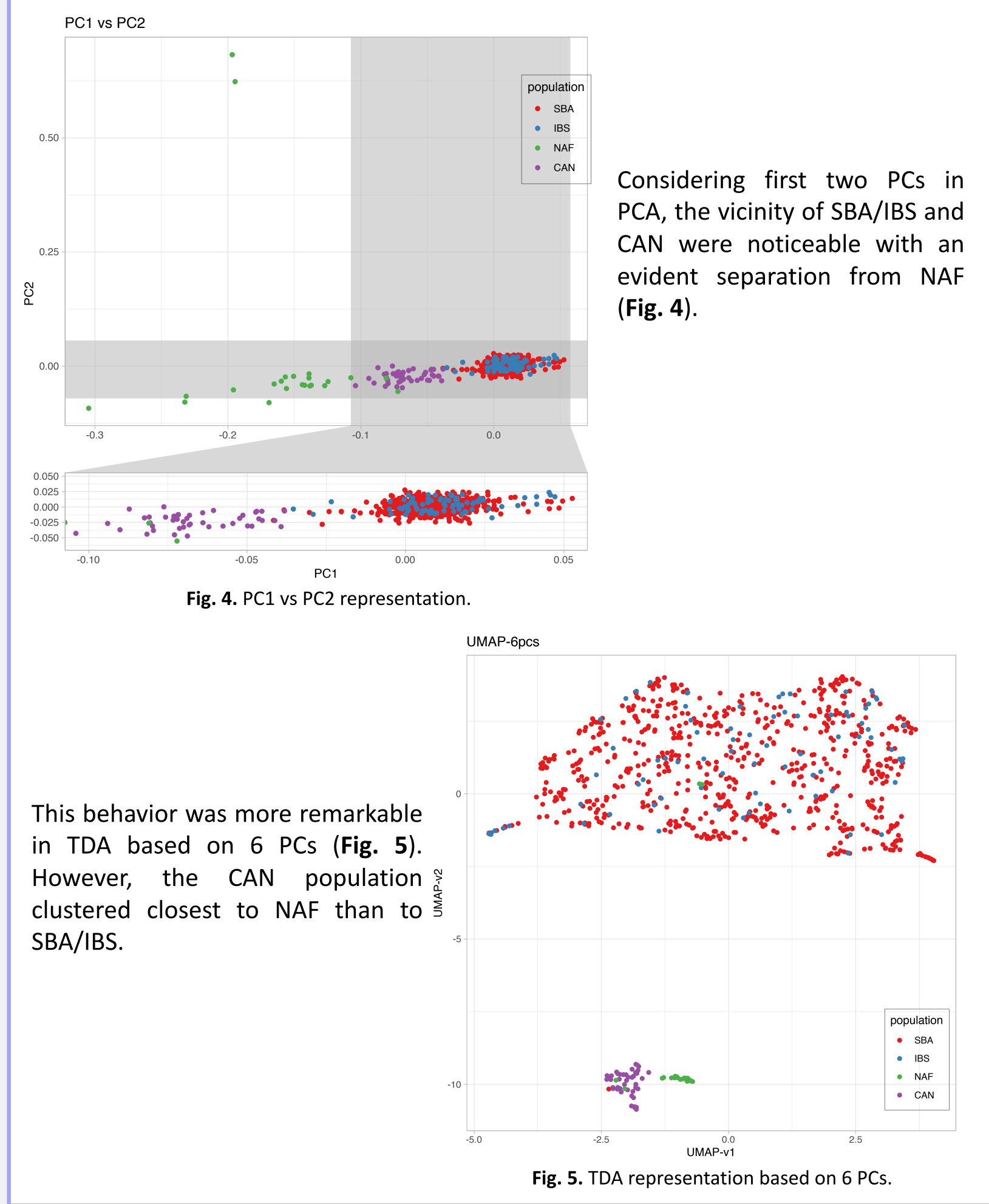
Fig. 1. Workflow stages data preparation: Filter and recode data to bed/bim/fam format, allele strand validation⁶ data from and merge multiple population to compute Principal Components and Topological Data Analysis.

Results

Comparing CAN, NAF and 1KGP populations



Comparing CAN, NAF and SBA/IBS populations



Conclusions

TDA provides an optimal alternative to reveal previously unrecognized fine structure separating IBS individuals from CAN, a result compatible with genetic drift and African admixture in the latter. Co-clustering of CAN both with NAF and IBS supports wide interindividual variation in ancestries.

In addition, the observed structure of present CAN-IBS and the genetic distance with the rest of EUR populations highlights the unique genetic features of current Canary Islanders.

Contact

Funding

Ministerio de Ciencia, Innovación y Universidades (RTC-2017-6471-1; MINECO/AEI/FEDER, UE), agreement OA17/008 with ITER. Fellowship by Spanish Ministry of Education, Culture, and Sports to ADU (FPU16/01435) and ACIISI co-funded by European Social Fund to B.G.G (TESIS2015010057).

The authors declare no conflict of interest.







References

3026.

1. The 1000 Genomes Project Consortium. *Nature* 2015; 526: 68-74.

2. Guillen-Guio et al. Mol Biol Evol 2018; 35: 3010-

4. McInnes et al. *arXiv 2018*; 1802.03426v2.5. Chang et al. *GigaScience 2015*; 4: 7.

6. W. Rayner, URL:

https://www.well.ox.ac.uk/~wrayner/tools/i