

# FREGENE datasets: 'Population C'

## SUMMARY OF MODELLING ASSUMPTIONS USED TO SIMULATE WORLDWIDE HUMAN POPULATION.

Population C mimics the principle features of genetic variation in major worldwide human populations. This simulation required the following steps:

- **Founding population in Africa:** an homogeneous population (N=25K sequences) evolves for 125K generations.
- **Expansion in Africa:** the population expands from 25K to 48K sequences and evolves during 17K further generations.
- **Out of Africa (OoA) split and bottleneck:** among the 48K sequences, 8.5% (= 4,080) leave Africa. Simultaneously, the population in Africa encounters a bottleneck of size ratio 0.8%, leaving 380 sequences in that subpopulation.
- **African and OoA expansion:** African population expands back to N=48K. Similarly the OoA population expands to N=15.4K and evolves for 3.5K generations.
- **Asian and European split:** the OoA population encounters a bottleneck of size N=1,360, and splits with N=320 moving to Europe and N=1,040 to Asia.
- **Asian and European expansion:** Asian and European populations both expand to N=15.4K, and evolve for 2K generations. During this stage, migration occurs symmetrically, first between Asia and Africa (with rate  $0.8 \times 10^{-5}$  per chromosome), and between Europe and Africa (with rate  $3.2 \times 10^{-5}$  per chromosome).
- **Independent evolution of the three populations:** African, Asian and European populations evolve, without migration, during 200, 400 and 350 generations respectively, while each population expands to reach a final population size of N=50K sequences.

Each sequence is 1 *Mb* long. Two simulations are available: one neutral (see results in `Output_neutral`) and one with selection (see results in `Output_selected`). Scripts used to generate these simulations are also provided (`fregene_POPNC_neutral.sh` and `fregene_POPNC_sel.sh` respectively). The population at the end of each step is recorded in the `Input` directory (see `rin_*.xml`). Modelling assumptions are recorded in the input files (in `Input` directory) and summarized below. Finally one R script that generate the evolution of diversity along generations can be found in `R_Scripts` together with the corresponding figure (in `Figures`).

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<b>General parameters</b>	
Chromosome Length	1 <i>Mb</i>
Per-site mutation rate	$1.5 \times 10^{-8}$

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<b>Recombination model</b>	
Per site crossover rate:	$1.1 \times 10^{-8}$
Per site Gene conversion rate:	$4.5 \times 10^{-9}$
Proportion of recombination events occurring in hotspots	80%
Hotspot length:	2.0 <i>kb</i>
Gene conversion length:	0.5 <i>kb</i>
Mean distance between hotspots:	8.5 <i>kb</i>

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<b>Selection parameters (if applicable)</b>	
Prop. of sites under selection:	$5 \times 10^{-4}$
Proportion of selected sites locally under selection:	0.5
Mean # generations before selected sites are switched off	50,000
selection coefficient:	$s \sim 0.1 \times \mathcal{N}(0.005, 0.005^2)$ $+0.9 \times \mathcal{N}(-0.01, 0.005^2)$
dominance coefficient:	$h \sim 0.8 \times \mathcal{N}(0.5, 0.2)$ $+0.3 \times \mathcal{N}(1.2, 0.2^2)$

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