1. Sequencing data

DrosEU & Dros-RTEC data:

Automated downloading and processing of 246 paired-end Pool-Seq libraries

DGN data:

Automated downloading and processing of single-individual libraries from 25 populations with ≥ 5 collected individuals and one *D. simulans* sample

Masked gSYNC file

BED file of masked pos.

2. Metadata

Sample metadata

Collection date Collector
Catching method Sample size
Inversion status SRA number
Sampling coordinates

Environmental metadata

WorldClimWeather-dataInterpolated
bioclimatic
trendsCurrent climatic
conditions from
weather stations

3. Mapping pipeline

3. Mapping pipeline		
Operation	Program/Script	Output file(s)
Sequence quality check	FASTQC	FASTQC output files
Adapter trimming & PE-read merging	cutdadapt bbmerge	
Mapping against hologenome reference	bwa mem GATK	Hologenome BAM
D. simulans decontamination	fix_bam.py	Decontaminated BAM
Conversion to pileup & gSYNC format	samtools mpileup mpileup2sync.py	Genome-wide pileup file Genome-wide gSYNC Indel position file
		Read-depth info
,		
Maakina		

MaskSYNC.py

4. SNP calling

PoolSNP

SNP calling based on heuristic parameters (common SNPs)

Merging of individual gSYNC files to joint SYNC files

Identifying polymorphic positions based on allele frequency (MAF) and allele counts (MAC)

Creating joint tbi-indexed SYNC/VCF file

Snape

probabilistic SNP calling (population-specific SNPs)

Probabilistic SNP calling based from individual pileup files

Filtering of SNAPE output and conversion to gSYNC file format

Merging individual gSYNC files to joint tbi-indexed SYNC/VCF file

5. Quality control

Basic summary statistics: Read depth; # of (private) SNPs; D. simulans contamination

p_N/p_s ratio and SFS at different MAF/MAC cutoffs

Comparison of AFs to Kapun *et al.* 2020 and Machado *et al.* 2020

Basic biological analyses: Principal component analyses

SNP calling parameters:

6. Resources generated

Extensible pipeline

We provide an highly automated and platform-independent mapping and SNP calling pipeline which is highly flexible and which allows to easily extend existing datasets.

Data availability

Using our novel pipeline we combined whole-genome sequencing data from 272 world-wide populations and provide high-confidence allele frequency data and corresponding meta information in multiple file formats

Genome browser

Population genetic estimates and allele frequency data can be visualized in and downloaded from an easy-to-use genome browser available at our website: http://dest.bio

Demography informative SNP markers

We generated a set of informative SNP markers, which allow to assign population samples to predefined demographic clusters

Part of Docker pipeline image

Masking

gSYNC file