## Code Sample S1: Template parameters file

```
# template_params.py
import numpy as np
import pandas as pd
import os
# set up the landscape
b4 = np.vstack([np.linspace(1, 0, 50) for _ in range(50)])
af = np.vstack([np.linspace(1, 0.5, 50) for _ in range(50)])
stable = np.vstack([np.linspace(1, 0, 50) for _ in range(50)])
K = np.ones((50, 50))
# NOTE: time_steps.CSV USED TO SET CLIMATE CHANGE
       TO START AT TIME STEP 2500
#
#
       AND FINISH AT TIME STEP 2750
if os.getcwd().split('/')[1] == 'home':
    steps = pd.read_csv(('/home/deth/Desktop/CAL/research/projects/sim/'
                         ch2/climate_change_adaptation_and_genomic_arch/sim/'
                        'time_steps.CSV'))
   steps = pd.read_csv(('/global/scratch/users/drewhart/'
                        'ch2/climate_change_adaptation_and_genomic_arch/sim/'
                        'time_steps.CSV'))
# set time when environmental change begins
change_T = int(steps[steps['name']=='start']['num'].values[0])
# set time when environmental change ends
T = int(steps[steps['name']=='end']['num'].values[0])
# show the landscape, for debugging, if requested
debug_landscape = False
if debug_landscape:
   import matplotlib pyplot as plt
   fig = plt.figure()
   ax1 = fig.add_subplot(221)
   im1 = ax1.imshow(b4, cmap='spring', vmin=0, vmax=1)
   plt.colorbar(im1)
   ax3 = fig.add_subplot(223)
   im3 = ax3.imshow(af, cmap='spring', vmin=0, vmax=1)
   plt.colorbar(im3)
   ax2 = fig.add_subplot(222)
   im2 = ax2.imshow(stable, cmap='winter', vmin=0, vmax=1)
   plt.colorbar(im2)
   ax4 = fig.add_subplot(224)
   im4 = ax4.imshow(K, cmap='autumn', vmin=0, vmax=1)
   plt.colorbar(im4)
   plt.show()
# This is a parameters file generated by Geonomics
# (by the gnx.make_parameters_file() function).
                                          :: :::::::#
                  #
            #:::::
                                            :: :: :::::::::::::::#
                                      #:::::::
                                      #::::::::
                                          MM MM IIIII CCCCC SSSSS #
    # GGGGG :EEEE: 00000
                                    00000
                         NN
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         EE OO OO NNN NN OO OO MM MM II CC
   # GG
                                                                 SS #
                                      OO MMM MMM
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            EE
                00
                       OO NN N NN OO
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                                                                 SSSSSS #
                                                      ΙI
   # GG GGG EEEE 00
                                         OO MM M MM
                                                                    SS #
                       OO NN NNN OO
                                                      ΙI
                                                          CC
                              NN OO OO MM MM II CC
   # GG G EE 00 00 NN
                          NN
    # GGGGG :EEEE: 00000
                               NN 00000 MM
                                                MM IIIIII CCCCC SSSSS #
     #
                                                            :: : #
       #:
                                                            :::::: #
         #
                                                            ::::: #
                                      ::::
                                                               #
                                                          #
                     # :: ::
```

```
#####################
#### LANDSCAPE ####
'landscape': {
   ###############
   #### main ####
   ################
        'main': {
           \#x,y (a.k.a. j,i) dimensions of the Landscape
            'dim':
                                        (50,50),
            #x,y resolution of the Landscape
                                        (1,1),
           #x,y coords of upper-left corner of the Landscape 'ulc': (0,0),
            #projection of the Landscape
            'prj':
                                        None,
           }, # <END> 'main'
   #################
   #### layers ####
   #################
        'layers': {
            #layer name (LAYER NAMES MUST BE UNIQUE!)
            'shift': {
       #--- layer num. 0: init parameters ---#
                #initiating parameters for this layer
                'init': {
                    #parameters for a 'defined'-type Layer
                    'defined': {
                        #raster to use for the Laver
                        'rast':
                        #point coordinates
                        'pts':
                                                  None,
                        #point values
                        'vals':
                                                  None,
                        #interpolation method {None, 'linear', 'cubic',
                        #'nearest'}
                        'interp_method':
                                                  None,
                        }, # <END> 'defined'
                   }, # <END> 'init'
           #--- layer num. 0: change parameters ---#
                #landscape-change events for this Layer
                'change': {
                    0: {
                        #array or file for final raster of event, or directory
                        #of files for each stepwise change in event
                        'change_rast':
                                                  af,
                        #starting timestep of event
                        'start_t':
                                           change_T,
                        #ending timestep of event
                        'end_t': T,
                        #number of stepwise changes in event
                        'n_steps':
                                           T-change_T,
                        }, # <END> event 0
                    }, # <END> 'change'
                }, # <END> layer num. 0
```

```
#layer name (LAYER NAMES MUST BE UNIQUE!)
    'stable': {
#--- layer num. 1: init parameters ---#
       #initiating parameters for this layer
       'init': {
            #parameters for a 'defined'-type Layer
            'defined': {
               #raster to use for the Layer
                'rast':
                                         stable,
               #point coordinates
                'pts':
                                         None,
               #point values
                'vals':
                                         None,
               #interpolation method {None, 'linear', 'cubic',
               #'nearest'}
               'interp_method':
                                         None,
               }, # <END> 'defined'
           }, # <END> 'init'
        }, # <END> layer num. 1
    #layer name (LAYER NAMES MUST BE UNIQUE!)
#--- layer num. 2: init parameters ---#
       #initiating parameters for this layer
        'init': {
            #parameters for a 'defined'-type Layer
            'defined': {
               #raster to use for the Layer
                'rast':
               #point coordinates
               'pts':
                                         None,
               #point values
                'vals':
                                         None,
               #interpolation method {None, 'linear', 'cubic',
               #'nearest'}
                'interp_method':
                                         None,
               }, # <END> 'defined'
           }, # <END> 'init'
        }, # <END> layer num. 2
#layer name (LAYER NAMES MUST BE UNIQUE!)
'move': {
#----#
#--- layer num. 2: init parameters ---#
        #initiating parameters for this layer
        'init': {
            #parameters for a 'defined'-type Layer
            'defined': {
               #raster to use for the Layer
                'rast':
                                         np.ones((50,50)),
```

```
#point coordinates
                       'pts':
                                               None,
                      #point values
                       'vals':
                                               None,
                      #interpolation method {None, 'linear', 'cubic',
                      #'nearest'}
                       'interp_method':
                                               None,
                      }, # <END> 'defined'
                   }, # <END> 'init'
               }
   #### NOTE: Individual Layers' sections can be copy-and-pasted (and
   #### assigned distinct keys and names), to create additional Layers.
           } # <END> 'layers'
       }, # <END> 'landscape'
#####################
#### COMMUNITY ####
##############################
    'comm': {
       'species': {
           #species name (SPECIES NAMES MUST BE UNIQUE!)
           'spp_0': {
           #-----#
           #--- spp num. 0: init parameters ---#
               'init': {
                   #starting number of individs
                                    1000,
                   #carrying-capacity Layer name
'K_layer': 'K',
                   'K_layer': 'K',
#multiplicative factor for carrying-capacity layer
                   'K_factor':
                                     2.5,
                   }, # <END> 'init'
           #----#
           #--- spp num. 0: mating parameters ---#
               'mating'
                        : {
                   #age(s) at sexual maturity (if tuple, female first)
                   'repro_age':
                   #whether to assign sexes
                   'sex':
                                             False,
                   #ratio of males to females
                   'sex_ratio':
                   #whether P(birth) should be weighted by parental dist
                   'dist_weighted_birth':
                                              False,
                   #intrinsic growth rate
                                              0.5,
                   'R':
                   #intrinsic birth rate (MUST BE 0<=b<=1)</pre>
                   'b':
                                              0.5,
                   #expectation of distr of n offspring per mating pair
                   'n_births_distr_lambda':
                                             1,
                   #whether n births should be fixed at n_births_dist_lambda
                   'n_births_fixed':
                                             True,
                   #radius of mate-search area
                   'mating_radius':
                       #whether individs should choose nearest neighs as mates
                   'choose_nearest_mate':
                                             False,
```

```
#whether mate-choice should be inverse distance-weighted
       'inverse_dist_mating': False,
       }, # <END> 'mating'
#-----#
#--- spp num. 0: mortality parameters ---#
#----#
    'mortality' : {
       #maximum age
       'max age':
       #min P(death) (MUST BE 0 \le d_min \le 1)
       'd min':
       #max P(death) (MUST BE 0<=d_max<=1)</pre>
       #width of window used to estimate local pop density
       'density_grid_window_width': None,
       }, # <END> 'mortality'
#--- spp num. 0: movement parameters ---#
    'movement': {
       #whether or not the species is mobile
       'move':
                         True,
       #mode of distr of movement direction
       'direction_distr_mu': 0,
       #concentration of distr of movement direction
       'direction_distr_kappa': 0,
       #mean of distr of movement distance
       'movement_distance_distr_param1':
                                              0.25,
       #variance of distr of movement distance
       'movement_distance_distr_param2': 0.5,
           #movement distance distr to use ('lognormal','levy','wald')
       'movement_distance_distr':
                                            'wald',
       #mean of distr of dispersal distance
       'dispersal_distance_distr_param1':
       #variance of distr of dispersal distance
       'dispersal_distance_distr_param2': 0.5,
          #dispersal distance distr to use ('lognormal','levy','wald')
                                             'wald',
       'dispersal_distance_distr':
       }, # <END> 'movement'
#--- spp num. 0: genomic architecture parameters ---#
       #file defining custom genomic arch
       'gen_arch_file': None,
       #num of loci
       'L':
                                1000,
       #value to use for fixed starting allele freqs (None to draw)
       'start_p_fixed': 0.5,
       #whether to start neutral locus freqs at 0
       'start_neut_zero': True,
       #genome-wide per-base neutral mut rate (0 to disable)
       'mu neut':
                     0.
       #genome-wide per-base deleterious mut rate (0 to disable)
                                 Ο,
       'mu delet':
       #shape of distr of deleterious effect sizes
       'delet_alpha_distr_shape': 0.2,
       #scale of distr of deleterious effect sizes
       'delet_alpha_distr_scale': 0.2,
       #NOTE: MAIN SCRIPT OVERRIDES THE FOLLOWING TWO PARAMS
       #TO SET RECOMBINATION RATES TO A FIXED VALUE OF
       #0.5, 0.05, OR 0.005 FOR INDEPENDENT,
       #WEAK, OR STRONG LINKAGE VALUES
       #alpha of distr of recomb rates
       'r_distr_alpha':
                                 1000
       #beta of distr of recomb rates
       'r_distr_beta':
```

```
#whether loci should be dominant (for allele '1')
                           False,
'dom':
#whether to allow pleiotropy
'pleiotropy':
                           False,
#custom fn for drawing recomb rates
'recomb_rate_custom_fn': None,
#number of recomb paths to hold in memory
'n_recomb_paths_mem':
                          int(1e4),
#total number of recomb paths to simulate
'n_recomb_paths_tot':
                        int(1e5),
   #num of crossing-over events (i.e. recombs) to simulate
'n_recomb_sims':
                           100_000,
#whether to generate recombination paths at each timestep
'allow_ad_hoc_recomb':
                           False,
#whether to save mutation logs
'mut_log':
                           False.
    #whether to jitter recomb bps, to correctly track num_trees
'jitter_breakpoints': False,
  #whether to use tskit (to record full spatial pedigree)
'use_tskit': True,
     #time step interval for simplication of tskit tables
'tskit_simp_interval': 100,
'traits': {
   #----#
   #---trait 0 parameters ---#
   #----#
   #trait name (TRAIT NAMES MUST BE UNIQUE!)
    'trait_0': {
       #trait-selection Layer name
        'laver':
                               'shift',
       #polygenic selection coefficient
        'phi':
                               1,
       #NOTE: MAIN SCRIPT CHANGES NEXT PARAM TO 4, 20, OR 100
       #FOR LOW-REDUNDANCY SCENARIOS OF DIFF. POLYGENICITY,
       #OR 8, 40, OR 200 FOR HIGH-REDUNDANCY SCENARIOS #number of loci underlying trait
        'n loci':
                               50.
       #mutation rate at loci underlying trait
        'mu':
                               0.
       #mean of distr of effect sizes
       'alpha_distr_mu' :
                               Θ,
       #variance of distr of effect size
        'alpha_distr_sigma':
                               Θ,
       #max allowed magnitude for an alpha value
       'max_alpha_mag':
                               None,
       #curvature of fitness function
        'gamma':
                               1,
       #whether the trait is universally advantageous
        'univ_adv':
                               False
       }, # <END> trait 0
   #----#
   #---trait 1 parameters ---#
   #----#
   #trait name (TRAIT NAMES MUST BE UNIQUE!)
    'trait_1': {
       #trait-selection Layer name
       'layer':
                               'stable',
       #polygenic selection coefficient
        'phi':
                               1,
       #NOTE: MAIN SCRIPT CHANGES NEXT PARAM TO 4, 20, OR 100
       #FOR LOW-REDUNDANCY SCENARIOS OF DIFF. POLYGENICITY,
       #OR 8, 40, OR 200 FOR HIGH-REDUNDANCY SCENARIOS #number of loci underlying trait
        'n loci':
                               50.
       #mutation rate at loci underlying trait
        'mu':
                               Θ,
       #mean of distr of effect sizes
        'alpha_distr_mu' :
                               Θ,
       #variance of distr of effect size
        'alpha_distr_sigma':
                               Θ,
       #max allowed magnitude for an alpha value
```

```
#curvature of fitness function
                          'gamma':
                         #whether the trait is universally advantageous
                         'univ_adv':
                         }, # <END> trait 1
   #### NOTE: Individual Traits' sections can be copy-and-pasted (and
   #### assigned distinct keys and names), to create additional Traits.
                      }, # <END> 'traits'
                  }, # <END> 'gen_arch'
              }, # <END> spp num. 0
   #### NOTE: individual Species' sections can be copy-and-pasted (and
   #### assigned distinct keys and names), to create additional Species.
          }, # <END> 'species'
       }, # <END> 'comm'
#################
#### MODEL ####
################
    'model': {
       # NOTE: NEXT PARAM OVERRIDDEN BY MAIN SCRIPT
       #total Model runtime (in timesteps)
       'T':
                     100000.
       #min burn-in runtime (in timesteps)
       'burn_T':
                      30,
       #seed number
       'num':
                      None,
       #####################################
       #### iterations parameters ####
       'its': {
          #num iterations
           'n_its':
           #whether to randomize Landscape each iteration
           'rand_landscape':
                            False,
           #whether to randomize Community each iteration
           'rand_comm':
                             False,
             #whether to randomize GenomicArchitectures each iteration
           'rand_genarch':
                             True,
           #whether to burn in each iteration
           'repeat_burn':
                             False,
           }, # <END> 'iterations'
       #### data-collection parameters ####
       'data': {
           'sampling': {
              #sampling scheme {'all', 'random', 'point', 'transect'}
'scheme': 'all',
              #sample size at each point, for point & transect sampling
              'n':
                                    1000,
              #coords of collection points, for point sampling
              'points':
                                    None,
              #coords of transect endpoints, for transect sampling
              'transect_endpoints':
                                    None,
```

'max\_alpha\_mag':

None,

```
#num points along transect, for transect sampling
             'n_transect_points':
                                        None,
             #collection radius around points, for point & transect sampling
             'radius':
                                         None,
             #when to collect data
              'when':
                                        [change_T-1,
                                        int((change_T-1+T-1)/2),
                                        T-1],
             #whether to save current Layers when data is collected
             'include_landscape':
                                        False,
             #whether to include fixed loci in VCF files
             'include_fixed_sites': True,
         'format': {
             #format for genetic data {'vcf', 'fasta'}
             'gen_format': ['vcf'],
#format for vector geodata {'csv', 'shapefile', 'geojson'}
'geo_vect_format': 'csv',
#format for rast geodata {'geotiff', 'txt'}
              'geo_rast_format': 'geotiff',
                #format for files containing non-neutral loci
             'nonneut_loc_format':
                                           'csv',
             },
         }, #<END> 'data'
    } # <END> 'model'
} # <END> params
```