
CMCpy Documentation

Release 0.1

David H. Ardell, Peter Becich and Brian Stark

December 21, 2012

CONTENTS

1	cmcpy Package	3
1.1	cmcpy Package	3
1.2	__main__ Module	3
1.3	amino_acid_spaces Module	3
1.4	codon_spaces Module	4
1.5	evolvers Module	5
1.6	genetic_codes Module	7
1.7	misreading Module	8
1.8	observables Module	9
1.9	site_type_spaces Module	9
2	CMCpy – Code-Message Coevolution Models in Python	11
3	Dependencies	13
4	Installation	15
5	Usage	17
6	Documentation	19
7	Licensing and Attribution	21
8	Release Notes	23
9	References	25
10	Indices and tables	27
	Bibliography	29
	Python Module Index	31
	Index	33

Contents:

CMCPY PACKAGE

1.1 cmcpy Package

1.2 __main__ Module

1.3 amino_acid_spaces Module

class amino_acid_spaces.**RegionAminoAcidSpace** (*num_aas=None, coords=None, num_dims=1, seed=42, labels=None*)

Bases: amino_acid_spaces._AminoAcidSpace

Region amino acid spaces model amino acid (dis)similarities in bounded regions of a finite number of dimensions.

```
>>> aa = RegionAminoAcidSpace(num_aas = 5,num_dims = 2)
>>> map(lambda x:x.round(2),aa.coords)
[array([[ 0.16,  0.71]]), array([[ 0.37,  0.16]]), array([[ 0.6,  0.6]]), array([[ 0.73,  0.87]])]
>>> dm = aa.get_distance_matrix()
>>> dm.round(3)
array([[ 0.    ,  0.594,  0.455,  0.597,  1.027],
       [ 0.594,  0.    ,  0.498,  0.795,  0.584],
       [ 0.455,  0.498,  0.    ,  0.297,  0.647],
       [ 0.597,  0.795,  0.297,  0.    ,  0.837],
       [ 1.027,  0.584,  0.647,  0.837,  0.    ]])
```

class amino_acid_spaces.**RingAminoAcidSpace** (*num_aas=None, seed=42, coords=None, labels=None*)

Bases: amino_acid_spaces._AminoAcidSpace

Ring amino acid spaces model amino acid (dis)similarities in a one-dimensional circular physicochemical amino acid space

```
>>> aa = RingAminoAcidSpace(num_aas = 5)
>>> map(lambda x: x.round(3),aa.coords)
[array([[ 0.156]]), array([[ 0.375]]), array([[ 0.599]]), array([[ 0.732]]), array([[ 0.951]])]
>>> dm = aa.get_distance_matrix()
>>> dm.round(3)
array([[ 0.    ,  0.219,  0.443,  0.424,  0.205],
       [ 0.219,  0.    ,  0.224,  0.357,  0.424],
       [ 0.443,  0.224,  0.    ,  0.133,  0.352],
       [ 0.424,  0.357,  0.133,  0.    ,  0.219],
       [ 0.205,  0.424,  0.352,  0.219,  0.    ]])
```

1.4 codon_spaces Module

class `codon_spaces.RingCodonSpace` (*num_codons, mu*)

Bases: `codon_spaces._CodonSpace`

Ring codon spaces are wrapped linear mutation spaces where codons mutate only to their two immediate neighbors. For ring codon models, *mu* defines the probability of change to one of two codon neighbors. The probability of no change is $[1 - (2*\mu)]$

```
>>> codons = RingCodonSpace(num_codons = 5, mu = 0.1)
>>> mm = codons.get_mutation_matrix()
>>> mm.round(3)
array([[ 0.8,  0.1,  0. ,  0. ,  0.1],
       [ 0.1,  0.8,  0.1,  0. ,  0. ],
       [ 0. ,  0.1,  0.8,  0.1,  0. ],
       [ 0. ,  0. ,  0.1,  0.8,  0.1],
       [ 0.1,  0. ,  0. ,  0.1,  0.8]])
```

get_derivative_matrix()

get_mutation_eigensystem()

post_process_perturbative_solution (*lhtopy, vhtopy*)

class `codon_spaces.WordCodonSpace` (*num_bases, num_positions, mu, kappa=1.0*)

Bases: `codon_spaces._CodonSpace`

Word codon spaces model natural codons with a finite number of bases and a finite word-length.

For word codon models, *mu* defines the total probability of change of a base to any neighbor. The probability of no change of a single base is defined as $(1 - \mu)$.

If *kappa* is not equal to 1.0, then *num_bases* must be even.

```
>>> codons = WordCodonSpace(num_bases = 2, num_positions = 2, mu = 0.1)
>>> mm = codons.get_mutation_matrix()
>>> mm.round(3)
array([[ 0.81,  0.09,  0.09,  0.01],
       [ 0.09,  0.81,  0.01,  0.09],
       [ 0.09,  0.01,  0.81,  0.09],
       [ 0.01,  0.09,  0.09,  0.81]])

>>> codons = WordCodonSpace(num_bases = 4, num_positions = 2, mu = 0.2, kappa = 2)
>>> mm = codons.get_mutation_matrix()
>>> mm.round(3)
array([[ 0.64,  0.08,  0.04,  0.04,  0.08,  0.01,  0.005,  0.005,
         0.04,  0.005,  0.003,  0.003,  0.04,  0.005,  0.003,  0.003],
       [ 0.08,  0.64,  0.04,  0.04,  0.01,  0.08,  0.005,  0.005,
         0.005,  0.04,  0.003,  0.003,  0.005,  0.04,  0.003,  0.003],
       [ 0.04,  0.04,  0.64,  0.08,  0.005,  0.005,  0.08,  0.01,
         0.003,  0.003,  0.04,  0.005,  0.003,  0.003,  0.04,  0.005],
       [ 0.04,  0.04,  0.08,  0.64,  0.005,  0.005,  0.01,  0.08,
         0.003,  0.003,  0.005,  0.04,  0.003,  0.003,  0.005,  0.04 ],
       [ 0.08,  0.01,  0.005,  0.005,  0.64,  0.08,  0.04,  0.04,
         0.04,  0.005,  0.003,  0.003,  0.04,  0.005,  0.003,  0.003],
       [ 0.01,  0.08,  0.005,  0.005,  0.08,  0.64,  0.04,  0.04,
         0.005,  0.04,  0.003,  0.003,  0.005,  0.04,  0.003,  0.003],
       [ 0.005,  0.005,  0.08,  0.01,  0.04,  0.04,  0.64,  0.08,
         0.003,  0.003,  0.04,  0.005,  0.003,  0.003,  0.04,  0.005],
       [ 0.005,  0.005,  0.01,  0.08,  0.04,  0.04,  0.08,  0.64,
         0.003,  0.003,  0.005,  0.04,  0.003,  0.003,  0.005,  0.04 ]],
```

```
[ 0.04 , 0.005, 0.003, 0.003, 0.04 , 0.005, 0.003, 0.003,
  0.64 , 0.08 , 0.04 , 0.04 , 0.08 , 0.01 , 0.005, 0.005],
[ 0.005, 0.04 , 0.003, 0.003, 0.005, 0.04 , 0.003, 0.003,
  0.08 , 0.64 , 0.04 , 0.04 , 0.01 , 0.08 , 0.005, 0.005],
[ 0.003, 0.003, 0.04 , 0.005, 0.003, 0.003, 0.04 , 0.005,
  0.04 , 0.04 , 0.64 , 0.08 , 0.005, 0.005, 0.08 , 0.01 ],
[ 0.003, 0.003, 0.005, 0.04 , 0.003, 0.003, 0.005, 0.04 ,
  0.04 , 0.04 , 0.08 , 0.64 , 0.005, 0.005, 0.01 , 0.08 ],
[ 0.04 , 0.005, 0.003, 0.003, 0.04 , 0.005, 0.003, 0.003,
  0.08 , 0.01 , 0.005, 0.005, 0.64 , 0.08 , 0.04 , 0.04 ],
[ 0.005, 0.04 , 0.003, 0.003, 0.005, 0.04 , 0.003, 0.003,
  0.01 , 0.08 , 0.005, 0.005, 0.08 , 0.64 , 0.04 , 0.04 ],
[ 0.003, 0.003, 0.04 , 0.005, 0.003, 0.003, 0.04 , 0.005,
  0.005, 0.005, 0.08 , 0.01 , 0.04 , 0.04 , 0.64 , 0.08 ],
[ 0.003, 0.003, 0.005, 0.04 , 0.003, 0.003, 0.005, 0.04 ,
  0.005, 0.005, 0.01 , 0.08 , 0.04 , 0.04 , 0.08 , 0.64 ]]
```

get_base_mutation_matrix()

Returns the mutation matrix of bases in one site.

get_derivative_matrix()

This function exists to serve the perturbative solution draft in evolvers.py

get_mutation_eigensystem()

post_process_perturbative_solution (*lpert*, *vpert*)

1.5 evolvers Module

1.5.1 evolvers – cmcpy module for abstract base class of Ardell Sella Evolvers

class evolvers.ArdellSellaEvolverAbstractBase (*initial_code*, *site_types*, *delta*, *epsilon*, *observables*)

Bases: object

Abstract Base Class for ArdellSellaEvolvers for Code-Message Coevolution corresponding to models published in Ardell and Sella (2001, 2002) and Sella and Ardell (2002, 2006). Concrete Implementations subclass from this for different implementations of Eigenvalue solutions.

compute_code_fitness_given_messages (*equilibrated_messages*, *effective_code_matrix*)

Implement eg eqns. 2-7 from Sella and Ardell(2006)

compute_max_fitness_code_mutation ()

equilibrate_messages ()

Compute eigensystems in site-types for an established genetic code.

This finds eigensystems (codon frequencies and growth rates) for different site-types given the genetic code.

Abstract method: subclasses must:

1) store their results by setting `self.eigenvalues` and `self.eigenmatrix`

2.set `self.equilibrated` to True

3) call `super(<<SubClass>>, self).equilibrate_messages()` to print observables at end of subclass method where `<<SubClass>>` is the subclass name to print observables.

evolve_code_unless_frozen()

Mutate genetic code.

Unless genetic code is frozen, mutate it according to the Ardell and Sella models, and update code to most fit mutant if it exists. If no more fit mutant code exists, set the “frozen” attribute to True.

evolve_one_step()

Mutate genetic code and equilibrate messages.

Unless genetic code is frozen, mutate it according to the Ardell and Sella models, update code to most fit mutant if it exists, and equilibrate messages to the new mutant genetic code. If no more fit mutant code exists, set the “frozen” attribute to True.

evolve_until_frozen()

Iteratively evolve genetic code and messages.

Until genetic code is frozen, mutate it according to the Ardell and Sella models, update code to most fit mutant if it exists, and equilibrate messages to the new mutant genetic code. Once no more fit mutant code exists, set the “frozen” attribute to True.

get_eigenvalue (*msm*, *eigenvec*)

get_mutation_selection_matrix (*alpha*)

get_selection_matrix (*alpha*)

growth_rate ()

growth_rate_from_lambda ()

initial_equilibrate_messages ()

Compute eigensystems in site-types for an established genetic code.

This finds eigensystems (codon frequencies and growth rates) for different site-types given the genetic code.

messages ()

print_initial_observables ()

print_observables ()

print_observables_header ()

class `evolvers.ArdellSellaEvolverHomotopy` (*initial_code*, *site_types*, *num_processes*, *observables*, *delta=1e-32*, *epsilon=1e-12*, *num_iterations=1000*)

Bases: `evolvers.ArdellSellaEvolverAbstractBase`

compute_eigensystem (*alpha*, *max_time=60*, *numpy_type=<type 'numpy.float64'>*)

equilibrate_messages ()

class `evolvers.ArdellSellaEvolverNumpy` (*initial_code*, *site_types*, *num_processes*, *observables*, *delta=1e-32*, *epsilon=1e-12*, *num_iterations=1000*)

Bases: `evolvers.ArdellSellaEvolverAbstractBase`

compute_eigensystem (*alpha*, *max_time=60*, *numpy_type=<type 'numpy.float64'>*)

equilibrate_messages ()

class `evolvers.ArdellSellaEvolverNumpyMulticore` (*initial_code*, *site_types*, *num_processes*, *observables*, *delta=1e-32*, *epsilon=1e-12*, *num_iterations=1000*)

Bases: `evolvers.ArdellSellaEvolverAbstractBase`

equilibrate_messages ()

```

class evolvers.ArdellSellaEvolverNumpyProcessChild(in_queue, out_queue)
    Bases: multiprocessing.process.Process

    Finds the eigensystem (message equilibrium and growth rate) for an established genetic code

    run()

class evolvers.ArdellSellaEvolverPowerCUDA(initial_code, site_types, delta, num_processes, epsilon,
                                              observables, num_iterations=1000)
    Bases: evolvers.ArdellSellaEvolverAbstractBase

    equilibrate_messages()

class evolvers.ArdellSellaEvolverPowerMethod(initial_code, site_types, num_processes,
                                              observables, delta=1e-32, epsilon=1e-12,
                                              max_order=5)
    Bases: evolvers.ArdellSellaEvolverAbstractBase

    compute_eigensystem(alpha, max_time=60, numpy_type=<type 'numpy.float64'>)
    equilibrate_messages()

class evolvers.ArdellSellaEvolverPowerMethodProcessChild(in_queue, out_queue,
                                                           num_codons, delta,
                                                           max_time=60)
    Bases: multiprocessing.process.Process

    Finds the eigensystem (message equilibrium and growth rate) for an established genetic code

    run()

class evolvers.ArdellSellaEvolverPowerMulticore(initial_code, site_types, num_processes,
                                                  observables, delta=1e-32, epsilon=1e-12,
                                                  num_iterations=1000)
    Bases: evolvers.ArdellSellaEvolverAbstractBase

    equilibrate_messages()

    in_queue = <multiprocessing.queues.JoinableQueue object at 0x101c420d0>
    out_queue = <multiprocessing.queues.JoinableQueue object at 0x101c56350>

class evolvers.eigensystem_CUDA_implementation(parent, max_time=60, delta=1e-32)

    calculate()
    done()
    error_check()
    get_eigenmatrix()
    get_eigenvalue(alpha)
    get_eigenvalues()

```

1.6 genetic_codes Module

```

class genetic_codes.GeneticCodeMutation(code, codon, aa)

    get_effective_code_matrix()

```

```
class genetic_codes.InitiallyAmbiguousGeneticCode(codons, amino_acids, misreading=None)
```

```
    Bases: genetic_codes._GeneticCode
```

```
class genetic_codes.UserInitializedGeneticCode(codons, amino_acids, code_matrix=None,
                                                code_dict=None, misreading=None)
```

```
    Bases: genetic_codes._GeneticCode
```

User-Initialized Genetic Codes are initialized with a numpy.ndarray code matrix or a dict of codons mapping to indices (not labels) of amino acids

```
>>> codons = codon_spaces.WordCodonSpace(num_bases = 4, num_positions = 2, mu = 0.2, kappa = 2)
>>> aas = amino_acid_spaces.RegionAminoAcidSpace(num_aas = 20, seed = 40)
>>> cm = numpy.eye(16)
>>> cm = numpy.hstack((cm, numpy.zeros((16, 4))))
>>> cm.shape
(16, 20)
>>> cm[0][1] = 1
>>> cm /= cm.sum(axis = 1).reshape(16, 1)
>>> gc = UserInitializedGeneticCode(codons, aas, code_matrix = cm)
>>> gc.num_codons
16
>>> gc.num_amino_acids
20
>>> gc.ambiguous_codons()
set([0])
>>> gc.encoded_aas
set([1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15])
>>> print gc
|* b c d|
|e f g h|
|i j k l|
|m n o p|
>>> gc.as_labelled_dict()
{0: '*', 1: 'b', 2: 'c', 3: 'd', 4: 'e', 5: 'f', 6: 'g', 7: 'h', 8: 'i', 9: 'j', 10: 'k', 11: 'l', 12: 'm', 13: 'n', 14: 'o', 15: 'p'}
>>> gc.as_dict()
{0: '*', 1: 1, 2: 2, 3: 3, 4: 4, 5: 5, 6: 6, 7: 7, 8: 8, 9: 9, 10: 10, 11: 11, 12: 12, 13: 13, 14: 14, 15: 15}
>>> gc2 = UserInitializedGeneticCode(codons, aas, code_dict = {0: '*', 1: 1, 2: 2, 3: 3, 4: 4, 5: 5})
>>> print gc2
|* b c d|
|e f g h|
|i j k l|
|m n o p|
>>> print gc2.code_matrix[0]
[ 0.05  0.05  0.05  0.05  0.05  0.05  0.05  0.05  0.05  0.05  0.05  0.05
  0.05  0.05  0.05  0.05  0.05  0.05  0.05  0.05]
```

1.7 misreading Module

```
class misreading.PositionalMisreading(codons, misreading)
```

```
    Bases: misreading._Misreading
```

Positional misreading models misreading on word codon spaces which model natural codons with a finite number of bases and a finite word-length.

For positional misreading, the misreading parameter is a list of positional misreading parameters mr_i which define the total misreading probability of a base at position i to any neighbor. The probability of no misreading of a single base at position i is defined as $(1 - mr_i)$.

```
>>> codons = codon_spaces.WordCodonSpace(num_bases = 2, num_positions = 2, mu = 0.1)
>>> misreading = PositionalMisreading(codons, [0.1, 0.01])
>>> mr = misreading.get_misreading_matrix()
>>> mr.round(3)
array([[ 0.891,  0.099,  0.009,  0.001],
       [ 0.099,  0.891,  0.001,  0.009],
       [ 0.009,  0.001,  0.891,  0.099],
       [ 0.001,  0.009,  0.099,  0.891]])
```

class misreading.**RingMisreading**(codons, misreading)
 Bases: misreading._Misreading

Ring misreading is one-dimensional misreading uniform over all other codons. For ring misreading, (mr/(nc - 1)) is the probability of misreading as a specific codon. The probability of no misreading is (1 - (mr)).

The misreading parameter is a list with one element, mr.

```
>>> codons = codon_spaces.RingCodonSpace(num_codons = 5, mu = 0.1)
>>> misreading = RingMisreading(codons, [0.1])
>>> mr = misreading.get_misreading_matrix()
>>> mr.round(3)
array([[ 0.9 ,  0.025,  0.025,  0.025,  0.025],
       [ 0.025,  0.9 ,  0.025,  0.025,  0.025],
       [ 0.025,  0.025,  0.9 ,  0.025,  0.025],
       [ 0.025,  0.025,  0.025,  0.9 ,  0.025],
       [ 0.025,  0.025,  0.025,  0.025,  0.9 ]])
```

1.8 observables Module

Control and select output from CMCpy simulations

class observables.**Observables**(show_codes=True, show_messages=False, show_initial_parameters=True, show_matrix_parameters=False, show_fitness_statistics=False, show_code_evolution_statistics=False, show_frozen_results_only=False, print_precision=6, show_all=False)

1.9 site_type_spaces Module

Site-type fitness matrices are intended as site-types over rows and amino acids over columns

class site_type_spaces.**MirroringSiteTypeSpace**(amino_acids, phi, weights=None)

This class models site-types in one-to-one correspondence with amino acids as according to the published models of Ardell and Sella.

```
>>> aa = amino_acid_spaces.RingAminoAcidSpace(num_aas = 5)
>>> dm = aa.get_distance_matrix()
>>> dm.round(3)
array([[ 0. ,  0.219,  0.443,  0.424,  0.205],
       [ 0.219,  0. ,  0.224,  0.357,  0.424],
       [ 0.443,  0.224,  0. ,  0.133,  0.352],
       [ 0.424,  0.357,  0.133,  0. ,  0.219],
       [ 0.205,  0.424,  0.352,  0.219,  0. ]])
>>> st = MirroringSiteTypeSpace(aa, phi = 0.96)
>>> fm = st.get_fitness_matrix()
```

```
>>> fm.round(3)
array([[ 1.    ,  0.991,  0.982,  0.983,  0.992],
       [ 0.991,  1.    ,  0.991,  0.986,  0.983],
       [ 0.982,  0.991,  1.    ,  0.995,  0.986],
       [ 0.983,  0.986,  0.995,  1.    ,  0.991],
       [ 0.992,  0.983,  0.986,  0.991,  1.    ]])
```

```
get_fitness_matrix()
```

```
get_site_type_weights()
```

CMCPY – CODE-MESSAGE COEVOLUTION MODELS IN PYTHON

CMCpy provides an object-oriented python API, together with command-line interface executables, that implement “Code-Message Coevolution” models. These published evolutionary models pertain to the evolution by natural selection of a genetic code in coevolution with a population of protein-coding genes.

Formally, CMC models are sets of quasispecies coupled together for their fitness through a genetic code. The system alternates between quasispecies equilibration and adaptive hill-climbing through codon assignments and reassignment by code mutation.

CMCpy can reproduce the statistics and results of [\[Ardell_and_Sella_2001\]](#), [\[Sella_and_Ardell_2002\]](#), [\[Ardell_and_Sella_2002\]](#) and [\[Sella_and_Ardell_2006\]](#). CMCpy additionally implements additional extensions that have not yet been studied in published work. It is easily feasible to extend the present code-base to implement the model studied by [\[Vetsigian_et_al_2006\]](#).

CMC evolutionary trajectories are partly a sequence of eigensystem solutions. Qualitative differences in results on different platforms can originate from differences in convergence criteria when power method-based eigensystem solvers are used, or from differences in floating point representations. Python defers to the platform C library for float representation. The default eigensystem solver is the `eig()` function in Numpy.

DEPENDENCIES

CMCpy relies heavily on, and absolutely requires, `numpy` as a prerequisite. You should install `numpy` with the `easy_install` framework to be detected as installed when installing this package.

If you wish to play with an experimental CUDA-based power-method eigensystem solver, you must install `pycuda`. This implementation is not faster than the NumPy default solver for many systems.

INSTALLATION

This installer requires `setuptools`, the most recent python packaging framework. If you do not already have this installed, this package will install it for you, so long as you have network access. Otherwise preinstall the correct version of `setuptools` using the EasyInstall installation instructions at <http://peak.telecommunity.com/DevCenter/EasyInstall#installation-instructions>

If you need to install this package somewhere other than the main `site-packages` directory, install `setuptools` using the instructions for Custom Installation Locations before installing this package. The instructions are here: <http://peak.telecommunity.com/DevCenter/EasyInstall#custom-installation-locations>

If you have downloaded the source-code package, the easiest way to install the package is to execute (from within the source root directory):

```
easy_install .
```

Mac users may need to run this command with “`sudo`” prepended.

You may also try simply executing:

```
easy_install CMCpy
```


USAGE

CMCpy comes with an executable inside the bin subdirectory to the installation source package, a UNIX-compatible script called “cmc”.

Additionally, a platform-specific executable may be automatically generated on installation.

Published results with CMC models may be (at least qualitatively) reproduced through the `–demo` option to the executables.

Also try running the `–help` option to the executables after installation and for a command-line example.

Programmers may use the executable in bin as a guide and template for how to program against the cmcpy API.

DOCUMENTATION

Some documentation of the cmcpcy API is available within the “doc” subdirectory of the source distribution. HTML, pdf and texinfo alternative formats are provided.

LICENSING AND ATTRIBUTION

The CMCpy project is distributed under the terms of the Apache License 2.0 as described in the file LICENSE.txt
Please cite Becich et al. (2012) in all scientific works that use this code.

RELEASE NOTES

The most recent version is 0.1 released October 2012.

See `CHANGES.txt` for version-related changes to the CMCpy code-base.

REFERENCES

INDICES AND TABLES

- *genindex*
- *modindex*
- *search*

BIBLIOGRAPHY

- [Ardell_andSella₂001] D.H. Ardell and G. Sella (2001). On the evolution of redundancy in genetic codes. *Journal of Molecular Evolution* 53(4/5):269-281.
- [Ardell_andSella₂002] D.H. Ardell and G. Sella (2002). No accident: genetic codes freeze in error-correcting patterns of the standard genetic code. *Philosophical Transactions of the Royal Society of London B* 357:1625-1642.
- [Sella_andArdell₂002] 7. Sella and D.H. Ardell (2002). The impact of message mutation on the fitness of a genetic code. *Journal of Molecular Evolution* 54(5):638-651.
- [Sella_andArdell₂006] 7. Sella and D.H. Ardell (2006). The coevolution of genes and genetic codes: Crick's frozen accident revisited. *J. Mol. Evol.* 63(3):297-313.
- [Vetsigian_et_al₂006] Vetsigian K., Woese C. R., Goldenfeld N. (2006). Collective evolution and the genetic code. *Proc. Natl. Acad. Sci. U.S.A.* 103, 10696-10701.

PYTHON MODULE INDEX

`__init__`, 3
`__main__`, 3

a

`amino_acid_spaces`, 3

c

`codon_spaces`, 4

e

`evolvers`, 5

g

`genetic_codes`, 7

m

`misreading`, 8

o

`observables`, 9

s

`site_type_spaces`, 9

INDEX

Symbols

`__init__` (module), 3
`__main__` (module), 3

A

`amino_acid_spaces` (module), 3
`ArdellSellaEvolverAbstractBase` (class in `evolvers`), 4
`ArdellSellaEvolverNumpy` (class in `evolvers`), 6
`ArdellSellaEvolverNumpyMulticore` (class in `evolvers`), 6
`ArdellSellaEvolverNumpyProcessChild` (class in `evolvers`), 6
`ArdellSellaEvolverPerturbative` (class in `evolvers`), 6
`ArdellSellaEvolverPowerCUDA` (class in `evolvers`), 6
`ArdellSellaEvolverPowerMethod` (class in `evolvers`), 6
`ArdellSellaEvolverPowerMethodProcessChild` (class in `evolvers`), 6
`ArdellSellaEvolverPowerMulticore` (class in `evolvers`), 6

C

`calculate()` (`evolvers.eigensystem_CUDA_implementation` method), 7
`choose()` (`evolvers.ArdellSellaEvolverPerturbative` method), 6
`codon_spaces` (module), 4
`compute_code_fitness_given_messages()` (`evolvers.ArdellSellaEvolverAbstractBase` method), 5
`compute_eigensystem()` (`evolvers.ArdellSellaEvolverNumpy` method), 6
`compute_eigensystem()` (`evolvers.ArdellSellaEvolverPerturbative` method), 6
`compute_eigensystem()` (`evolvers.ArdellSellaEvolverPowerMethod` method), 6
`compute_max_fitness_code_mutation()` (`evolvers.ArdellSellaEvolverAbstractBase` method), 5

D

`done()` (`evolvers.eigensystem_CUDA_implementation` method), 7

E

`eigensystem_CUDA_implementation` (class in `evolvers`), 7
`equilibrate_messages()` (`evolvers.ArdellSellaEvolverAbstractBase` method), 5
`equilibrate_messages()` (`evolvers.ArdellSellaEvolverNumpy` method), 6
`equilibrate_messages()` (`evolvers.ArdellSellaEvolverNumpyMulticore` method), 6
`equilibrate_messages()` (`evolvers.ArdellSellaEvolverPerturbative` method), 6
`equilibrate_messages()` (`evolvers.ArdellSellaEvolverPowerCUDA` method), 6
`equilibrate_messages()` (`evolvers.ArdellSellaEvolverPowerMethod` method), 6
`equilibrate_messages()` (`evolvers.ArdellSellaEvolverPowerMulticore` method), 6
`error_check()` (`evolvers.eigensystem_CUDA_implementation` method), 7
`evolve_code_unless_frozen()` (`evolvers.ArdellSellaEvolverAbstractBase` method), 5
`evolve_one_step()` (`evolvers.ArdellSellaEvolverAbstractBase` method), 5
`evolve_until_frozen()` (`evolvers.ArdellSellaEvolverAbstractBase` method), 5
`evolvers` (module), 4

G

`genetic_codes` (module), 7
`GeneticCodeMutation` (class in `genetic_codes`), 7
`get_derivative_matrix()` (`codon_spaces.RingCodonSpace` method), 4
`get_derivative_matrix()` (`codon_spaces.WordCodonSpace` method), 4
`get_effective_code_matrix()` (`genetic_codes.GeneticCodeMutation` method), 7
`get_eigenmatrix()` (`evolvers.eigensystem_CUDA_implementation` method), 7
`get_eigenvalue()` (`evolvers.ArdellSellaEvolverAbstractBase` method), 7

method), 5
get_eigenvalue() (evolvers.eigensystem_CUDA_implementation
method), 7
get_eigenvalues() (evolvers.eigensystem_CUDA_implementation
method), 7
get_fitness_matrix() (site_type_spaces.MirroringSiteTypeSpace
method), 9
get_mutation_selection_matrix()
(evolvers.ArdellSellaEvolverAbstractBase
method), 5
get_selection_matrix() (evolvers.ArdellSellaEvolverAbstractBase
method), 5
get_site_type_weights() (site_type_spaces.MirroringSiteTypeSpace
method), 9
growth_rate() (evolvers.ArdellSellaEvolverAbstractBase
method), 5
growth_rate_from_lambda()
(evolvers.ArdellSellaEvolverAbstractBase
method), 5

I

in_queue (evolvers.ArdellSellaEvolverPowerMulticore
attribute), 6
initial_equilibrate_messages()
(evolvers.ArdellSellaEvolverAbstractBase
method), 5
InitiallyAmbiguousGeneticCode (class in genetic_codes),
7

M

messages() (evolvers.ArdellSellaEvolverAbstractBase
method), 5
MirroringSiteTypeSpace (class in site_type_spaces), 9
misreading (module), 8

O

Observables (class in observables), 8
observables (module), 8
out_queue (evolvers.ArdellSellaEvolverPowerMulticore
attribute), 6

P

PositionalMisreading (class in misreading), 8
post_process_perturbative_solution()
(codon_spaces.RingCodonSpace method),
4
post_process_perturbative_solution()
(codon_spaces.WordCodonSpace method),
4
print_initial_observables()
(evolvers.ArdellSellaEvolverAbstractBase
method), 5
print_observables() (evolvers.ArdellSellaEvolverAbstractBase
method), 5

print_observables_header()
(evolvers.ArdellSellaEvolverAbstractBase
method), 5

R

RegionAminoAcidSpace (class in amino_acid_spaces), 3
RingAminoAcidSpace (class in amino_acid_spaces), 3
RingCodonSpace (class in codon_spaces), 4
RingMisreading (class in misreading), 8
run() (evolvers.ArdellSellaEvolverNumpyProcessChild
method), 6
run() (evolvers.ArdellSellaEvolverPowerMethodProcessChild
method), 6

S

site_type_spaces (module), 9

U

UserInitializedGeneticCode (class in genetic_codes), 7

W

WordCodonSpace (class in codon_spaces), 4