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# **Statistics**

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**The Sage Development Team**

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## BASIC STATISTICS

This file contains basic descriptive functions. Included are the mean, median, mode, moving average, standard deviation, and the variance. When calling a function on data, there are checks for functions already defined for that data type.

The `mean()` function returns the arithmetic mean (the sum of all the members of a list, divided by the number of members). Further revisions may include the geometric and harmonic mean. The `median()` function returns the number separating the higher half of a sample from the lower half. The `mode()` returns the most common occurring member of a sample, plus the number of times it occurs. If entries occur equally common, the smallest of a list of the most common entries is returned. The `moving_average()` is a finite impulse response filter, creating a series of averages using a user-defined number of subsets of the full data set. The `std()` and the `variance()` return a measurement of how far data points tend to be from the arithmetic mean.

Functions are available in the namespace `stats`, i.e. you can use them by typing `stats.mean`, `stats.median`, etc.

REMARK: If all the data you are working with are floating point numbers, you may find `stats.TimeSeries` helpful, since it is extremely fast and offers many of the same descriptive statistics as in the module.

AUTHOR:

- Andrew Hou (11/06/2009)

`sage.stats.basic_stats.mean(v)`

Return the mean of the elements of `v`.

We define the mean of the empty list to be the (symbolic) NaN, following the convention of MATLAB, Scipy, and R.

This function is deprecated. Use `numpy.mean()` or `numpy.nanmean()` instead.

INPUT:

- `v` – list of numbers

OUTPUT: a number

EXAMPLES:

```
sage: mean([pi, e]) #_
↳needs sage.symbolic
doctest:warning...
DeprecationWarning: sage.stats.basic_stats.mean is deprecated;
use numpy.mean or numpy.nanmean instead
See https://github.com/sagemath/sage/issues/29662 for details.
1/2*pi + 1/2*e
sage: mean([]) #_
↳needs sage.symbolic
NaN
```

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```

sage: mean([I, sqrt(2), 3/5]) #_
↳needs sage.symbolic
1/3*sqrt(2) + 1/3*I + 1/5
sage: mean([RIF(1.0103,1.0103), RIF(2)]) #_
↳needs sage.rings.real_interval_field
1.5051500000000000?
sage: mean(range(4))
3/2
sage: v = stats.TimeSeries([1..100]) #_
↳needs numpy
sage: mean(v) #_
↳needs numpy
50.5

```

`sage.stats.basic_stats.median(v)`

Return the median (middle value) of the elements of *v*.

If *v* is empty, we define the median to be NaN, which is consistent with NumPy (note that R returns NULL). If *v* is comprised of strings, `TypeError` occurs. For elements other than numbers, the median is a result of `sorted()`.

This function is deprecated. Use `numpy.median()` or `numpy.nanmedian()` instead.

INPUT:

- *v* – list

OUTPUT: median element of *v*

EXAMPLES:

```

sage: median([1,2,3,4,5])
doctest:warning...
DeprecationWarning: sage.stats.basic_stats.median is deprecated;
use numpy.median or numpy.nanmedian instead
See https://github.com/sagemath/sage/issues/29662 for details.
3
sage: median([e, pi]) #_
↳needs sage.symbolic
1/2*pi + 1/2*e
sage: median(['sage', 'linux', 'python'])
'python'
sage: median([]) #_
↳needs sage.symbolic
NaN
sage: class MyClass:
....:     def median(self):
....:         return 1
sage: stats.median(MyClass())
1

```

`sage.stats.basic_stats.mode(v)`

Return the mode of *v*.

The mode is the list of the most frequently occurring elements in *v*. If *n* is the most times that any element occurs in *v*, then the mode is the list of elements of *v* that occur *n* times. The list is sorted if possible.

This function is deprecated. Use `scipy.stats.mode()` or `statistics.mode()` instead.

**Note**

The elements of  $v$  must be hashable.

INPUT:

- $v$  – list

OUTPUT: list (sorted if possible)

EXAMPLES:

```
sage: v = [1, 2, 4, 1, 6, 2, 6, 7, 1]
sage: mode(v)
doctest:warning...
DeprecationWarning: sage.stats.basic_stats.mode is deprecated;
use scipy.stats.mode or statistics.mode instead
See https://github.com/sagemath/sage/issues/29662 for details.
[1]
sage: v.count(1)
3
sage: mode([])
[]

sage: mode([1, 2, 3, 4, 5])
[1, 2, 3, 4, 5]
sage: mode([3, 1, 2, 1, 2, 3])
[1, 2, 3]
sage: mode([0, 2, 7, 7, 13, 20, 2, 13])
[2, 7, 13]

sage: mode(['sage', 'four', 'I', 'three', 'sage', 'pi'])
['sage']

sage: class MyClass:
....:     def mode(self):
....:         return [1]
sage: stats.mode(MyClass())
[1]
```

`sage.stats.basic_stats.moving_average(v, n)`

Return the moving average of a list  $v$ .

The moving average of a list is often used to smooth out noisy data.

If  $v$  is empty, we define the entries of the moving average to be NaN.

This method is deprecated. Use `pandas.Series.rolling()` instead.

INPUT:

- $v$  – list
- $n$  – the number of values used in computing each average

OUTPUT: list of length  $\text{len}(v) - n + 1$ , since we do not fabric any values

EXAMPLES:

```

sage: moving_average([1..10], 1)
doctest:warning...
DeprecationWarning: sage.stats.basic_stats.moving_average is deprecated;
use pandas.Series.rolling instead
See https://github.com/sagemath/sage/issues/29662 for details.
[1, 2, 3, 4, 5, 6, 7, 8, 9, 10]
sage: moving_average([1..10], 4)
[5/2, 7/2, 9/2, 11/2, 13/2, 15/2, 17/2]
sage: moving_average([], 1)
[]
sage: moving_average([pi, e, I, sqrt(2), 3/5], 2) #_
↪needs sage.symbolic
[1/2*pi + 1/2*e, 1/2*e + 1/2*I, 1/2*sqrt(2) + 1/2*I,
 1/2*sqrt(2) + 3/10]

```

We check if the input is a time series, and if so use the optimized `simple_moving_average()` method, but with (slightly different) meaning as defined above (the point is that the `simple_moving_average()` on time series returns  $n$  values:

```

sage: a = stats.TimeSeries([1..10]) #_
↪needs numpy
sage: stats.moving_average(a, 3) #_
↪needs numpy
[2.0000, 3.0000, 4.0000, 5.0000, 6.0000, 7.0000, 8.0000, 9.0000]
sage: stats.moving_average(list(a), 3) #_
↪needs numpy
[2.0, 3.0, 4.0, 5.0, 6.0, 7.0, 8.0, 9.0]

```

`sage.stats.basic_stats.std(v, bias=False)`

Return the standard deviation of the elements of  $v$ .

We define the standard deviation of the empty list to be NaN, following the convention of MATLAB, Scipy, and R.

This function is deprecated. Use `numpy.std()` or `numpy.nanstd()` instead.

INPUT:

- $v$  – list of numbers
- `bias` – boolean (default: `False`); if `False`, divide by  $\text{len}(v) - 1$  instead of  $\text{len}(v)$  to give a less biased estimator (sample) for the standard deviation.

OUTPUT: a number

EXAMPLES:

```

sage: # needs sage.symbolic
sage: std([1..6], bias=True)
doctest:warning...
DeprecationWarning: sage.stats.basic_stats.std is deprecated;
use numpy.std or numpy.nanstd instead
See https://github.com/sagemath/sage/issues/29662 for details.
doctest:warning...
DeprecationWarning: sage.stats.basic_stats.variance is deprecated;
use numpy.var or numpy.nanvar instead
See https://github.com/sagemath/sage/issues/29662 for details.
doctest:warning...
DeprecationWarning: sage.stats.basic_stats.mean is deprecated;

```

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```

use numpy.mean or numpy.nanmean instead
See https://github.com/sagemath/sage/issues/29662 for details.
1/2*sqrt(35/3)
sage: std([1..6], bias=False)
sqrt(7/2)
sage: std([e, pi])
sqrt(1/2)*abs(pi - e)
sage: std([])
NaN
sage: std([I, sqrt(2), 3/5])
1/15*sqrt(1/2)*sqrt((10*sqrt(2) - 5*I - 3)^2
+ (5*sqrt(2) - 10*I + 3)^2 + (5*sqrt(2) + 5*I - 6)^2)
sage: std([RIF(1.0103, 1.0103), RIF(2)])
0.6998235813403261?

sage: # needs numpy
sage: import numpy
sage: if int(numpy.version.short_version[0]) > 1:
....:     numpy.set_printoptions(legacy="1.25")
sage: x = numpy.array([1,2,3,4,5])
sage: std(x, bias=False)
1.5811388300841898
sage: x = stats.TimeSeries([1..100])
sage: std(x)
29.011491975882016

```

sage.stats.basic\_stats.**variance**(*v*, *bias=False*)

Return the variance of the elements of *v*.

We define the variance of the empty list to be NaN, following the convention of MATLAB, Scipy, and R.

This function is deprecated. Use `numpy.var()` or `numpy.nanvar()` instead.

INPUT:

- *v* – list of numbers
- *bias* – boolean (default: `False`); if `False`, divide by  $\text{len}(v) - 1$  instead of  $\text{len}(v)$  to give a less biased estimator (sample) for the standard deviation.

OUTPUT: a number

EXAMPLES:

```

sage: variance([1..6])
doctest:warning...
DeprecationWarning: sage.stats.basic_stats.variance is deprecated;
use numpy.var or numpy.nanvar instead
See https://github.com/sagemath/sage/issues/29662 for details.
7/2
sage: variance([1..6], bias=True)
35/12
sage: variance([e, pi])
↪needs sage.symbolic #_
1/2*(pi - e)^2
sage: variance([])
NaN
sage: variance([I, sqrt(2), 3/5])
↪needs sage.symbolic #_

```

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```

1/450*(10*sqrt(2) - 5*I - 3)^2 + 1/450*(5*sqrt(2) - 10*I + 3)^2
+ 1/450*(5*sqrt(2) + 5*I - 6)^2
sage: variance([RIF(1.0103, 1.0103), RIF(2)])
0.4897530450000000?
sage: import numpy #_
↳needs numpy
sage: if int(numpy.version.short_version[0]) > 1: #_
↳needs numpy
.....: numpy.set_printoptions(legacy="1.25") #_
↳needs numpy
sage: x = numpy.array([1,2,3,4,5]) #_
↳needs numpy
sage: variance(x, bias=False) #_
↳needs numpy
2.5
sage: x = stats.TimeSeries([1..100])
sage: variance(x)
841.6666666666666
sage: variance(x, bias=True)
833.25
sage: class MyClass:
.....:     def variance(self, bias=False):
.....:         return 1
sage: stats.variance(MyClass())
1
sage: class SillyPythonList:
.....:     def __init__(self):
.....:         self.__list = [2, 4]
.....:     def __len__(self):
.....:         return len(self.__list)
.....:     def __iter__(self):
.....:         return self.__list.__iter__()
.....:     def mean(self):
.....:         return 3
sage: R = SillyPythonList()
sage: variance(R)
2
sage: variance(R, bias=True)
1

```

## C INT LISTS

This is a class for fast basic operations with lists of C ints. It is similar to the double precision TimeSeries class. It has all the standard C int semantics, of course, including overflow. It is also similar to the Python list class, except all elements are C ints, which makes some operations much, much faster. For example, concatenating two IntLists can be over 10 times faster than concatenating the corresponding Python lists of ints, and taking slices is also much faster.

AUTHOR:

- William Stein, 2010-03

**class** sage.stats.intlist.IntList

Bases: object

A list of C int's.

**list** ()

Return Python list version of self with Python ints as entries.

EXAMPLES:

```
sage: a = stats.IntList([1..15]); a
[1, 2, 3, 4, 5 ... 11, 12, 13, 14, 15]
sage: a.list()
[1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15]
sage: list(a) == a.list()
True
sage: type(a.list()[0])
<... 'int'>
```

**max** (*index=False*)

Return the largest value in this time series. If this series has length 0 we raise a `ValueError`

INPUT:

- *index* – boolean (default: `False`); if `True`, also return index of maximum entry

OUTPUT:

- integer – largest value
- integer – index of largest value; only returned if *index=True*

EXAMPLES:

```
sage: v = stats.IntList([1,-4,3,-2,-4,3])
sage: v.max()
3
```

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```
sage: v.max(index=True)
(3, 2)
```

**min** (*index=False*)

Return the smallest value in this integer list. If this series has length 0 we raise a `ValueError`.

INPUT:

- `index` – boolean (default: `False`); if `True`, also return index of minimal entry

OUTPUT:

- float – smallest value
- integer – index of smallest value; only returned if `index=True`

EXAMPLES:

```
sage: v = stats.IntList([1,-4,3,-2,-4])
sage: v.min()
-4
sage: v.min(index=True)
(-4, 1)
```

**plot** (*\*args, \*\*kws*)

Return a plot of this `IntList`.

This just constructs the corresponding double-precision floating point `TimeSeries` object, passing on all arguments.

EXAMPLES:

```
sage: stats.IntList([3,7,19,-2]).plot() #_
↪needs sage.plot
Graphics object consisting of 1 graphics primitive
sage: stats.IntList([3,7,19,-2]).plot(color='red', #_
↪needs sage.plot
....:                                     pointsize=50, points=True)
Graphics object consisting of 1 graphics primitive
```

**plot\_histogram** (*\*args, \*\*kws*)

Return a histogram plot of this `IntList`.

This just constructs the corresponding double-precision floating point `TimeSeries` object, and plots it, passing on all arguments.

EXAMPLES:

```
sage: stats.IntList([1..15]).plot_histogram() #_
↪needs sage.plot
Graphics object consisting of 50 graphics primitives
```

**prod** ()

Return the product of the entries of `self`.

EXAMPLES:

```
sage: a = stats.IntList([1..10]); a
[1, 2, 3, 4, 5, 6, 7, 8, 9, 10]
sage: a.prod()
3628800
sage: factorial(10)
3628800
```

Note that there can be overflow:

```
sage: a = stats.IntList([2^30, 2]); a
[1073741824, 2]
sage: a.prod()
-2147483648
```

### **sum()**

Return the sum of the entries of `self`.

EXAMPLES:

```
sage: stats.IntList([1..100]).sum()
5050
```

Note that there can be overflow, since the entries are C ints:

```
sage: a = stats.IntList([2^30, 2^30]); a
[1073741824, 1073741824]
sage: a.sum()
-2147483648
```

### **time\_series()**

Return `TimeSeries` version of `self`, which involves changing each entry to a double.

EXAMPLES:

```
sage: T = stats.IntList([-2, 3, 5]).time_series(); T
[-2.0000, 3.0000, 5.0000]
sage: type(T)
<... 'sage.stats.time_series.TimeSeries'>
```

`sage.stats.intlist.unpickle_intlist_v1(v, n)`

Version 1 unpickle method.

INPUT:

- `v` – a raw char buffer

EXAMPLES:

```
sage: v = stats.IntList([1, 2, 3])
sage: s = v.__reduce__()[1][0]
sage: type(s) == type(b'')
True
sage: sage.stats.intlist.unpickle_intlist_v1(s, 3)
[1, 2, 3]
sage: sage.stats.intlist.unpickle_intlist_v1(s+s, 6)
[1, 2, 3, 1, 2, 3]
sage: sage.stats.intlist.unpickle_intlist_v1(b'', 0)
[]
```



## HIDDEN MARKOV MODELS

This is a complete pure-Cython optimized implementation of Hidden Markov Models. It fully supports Discrete, Gaussian, and Mixed Gaussian emissions.

The best references for the basic HMM algorithms implemented here are:

- Tapas Kanungo’s “Hidden Markov Models”
- **Jackson’s HMM tutorial:**  
<http://personal.ee.surrey.ac.uk/Personal/P.Jackson/tutorial/>

LICENSE: Some of the code in this file is based on reading Kanungo’s GPLv2+ implementation of discrete HMM’s, hence the present code must be licensed with a GPLv2+ compatible license.

AUTHOR:

- William Stein, 2010-03

**class** sage.stats.hmm.hmm.**DiscreteHiddenMarkovModel**

Bases: *HiddenMarkovModel*

A discrete Hidden Markov model implemented using double precision floating point arithmetic.

INPUT:

- *A* – list of lists or a square  $N \times N$  matrix, whose  $(i, j)$  entry gives the probability of transitioning from state  $i$  to state  $j$ .
- *B* – list of  $N$  lists or a matrix with  $N$  rows, such that  $B[i, k]$  gives the probability of emitting symbol  $k$  while in state  $i$ .
- *pi* – the probabilities of starting in each initial state, i.e., *pi*[*i*] is the probability of starting in state  $i$ .
- *emission\_symbols* – None or list (default: None); if None, the emission\_symbols are the ints  $[0 \dots N-1]$ , where  $N$  is the number of states. Otherwise, they are the entries of the list *emissions\_symbols*, which must all be hashable.
- *normalize* – boolean (default: True); if given, input is normalized to define valid probability distributions, e.g., the entries of *A* are made nonnegative and the rows sum to 1, and the probabilities in *pi* are normalized.

EXAMPLES:

```
sage: m = hmm.DiscreteHiddenMarkovModel([[0.4, 0.6], [0.1, 0.9]],
....:                                     [[0.1, 0.9], [0.5, 0.5]],
....:                                     [.5, .5]); m
Discrete Hidden Markov Model with 2 States and 2 Emissions
Transition matrix:
[0.4 0.6]
[0.1 0.9]
```

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```
Emission matrix:
[0.1 0.9]
[0.5 0.5]
Initial probabilities: [0.5000, 0.5000]
sage: m.log_likelihood([0,1,0,1,0,1])
-4.66693474691329...
sage: m.viterbi([0,1,0,1,0,1])
([1, 1, 1, 1, 1, 1], -5.378832842208748)
sage: m.baum_welch([0,1,0,1,0,1])
(0.0, 22)
sage: m # rel tol 1e-10
Discrete Hidden Markov Model with 2 States and 2 Emissions
Transition matrix:
[1.0134345614745788e-70          1.0]
[          1.0 3.9974352713558623e-19]
Emission matrix:
[ 7.380221566254936e-54          1.0]
[          1.0 3.9974352626002193e-19]
Initial probabilities: [0.0000, 1.0000]
sage: m.sample(10)
[0, 1, 0, 1, 0, 1, 0, 1, 0, 1]
sage: m.graph().plot() #_
↪needs sage.plot
Graphics object consisting of 6 graphics primitives
```

A 3-state model that happens to always outputs 'b':

```
sage: m = hmm.DiscreteHiddenMarkovModel([[1/3]*3]*3, [[0,1,0]]*3, [1/3]*3, ['a','b',
↪','c'])
sage: m.sample(10)
['b', 'b', 'b', 'b', 'b', 'b', 'b', 'b', 'b', 'b']
```

**baum\_welch** (*obs*, *max\_iter=100*, *log\_likelihood\_cutoff=0.0001*, *fix\_emissions=False*)

Given an observation sequence *obs*, improve this HMM using the Baum-Welch algorithm to increase the probability of observing *obs*.

INPUT:

- *obs* – list of emissions
- *max\_iter* – integer (default: 100); maximum number of Baum-Welch steps to take
- *log\_likelihood\_cutoff* – positive float (default: 1e-4); the minimal improvement in likelihood with respect to the last iteration required to continue. Relative value to log likelihood.
- *fix\_emissions* – boolean (default: False); if True, do not change emissions when updating

OUTPUT:

changes the model in place, and returns the log likelihood and number of iterations.

EXAMPLES:

```
sage: m = hmm.DiscreteHiddenMarkovModel([[0.1,0.9],[0.9,0.1]],
.....:                                     [[.5,.5],[0,1]],
.....:                                     [.2,.8])
sage: m.baum_welch([1,0]*20, log_likelihood_cutoff=0)
(0.0, 4)
sage: m # rel tol 1e-14
```

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```

Discrete Hidden Markov Model with 2 States and 2 Emissions
Transition matrix:
[1.3515269707707603e-51          1.0]
[          1.0          0.0]
Emission matrix:
[          1.0 6.462537138850569e-52]
[          0.0          1.0]
Initial probabilities: [0.0000, 1.0000]

```

The following illustrates how Baum-Welch is only a local optimizer, i.e., the above model is far more likely to produce the sequence  $[1,0]^*20$  than the one we get below:

```

sage: m = hmm.DiscreteHiddenMarkovModel([[0.5,0.5],[0.5,0.5]],
.....:                                  [[.5,.5],[.5,.5]],
.....:                                  [.5,.5])
sage: m.baum_welch([1,0]*20, log_likelihood_cutoff=0)
(-27.725887222397784, 1)
sage: m
Discrete Hidden Markov Model with 2 States and 2 Emissions
Transition matrix:
[0.5 0.5]
[0.5 0.5]
Emission matrix:
[0.5 0.5]
[0.5 0.5]
Initial probabilities: [0.5000, 0.5000]

```

We illustrate fixing emissions:

```

sage: m = hmm.DiscreteHiddenMarkovModel([[0.1,0.9],[0.9,0.1]],
.....:                                  [[.5,.5],[.2,.8]],
.....:                                  [.2,.8])
sage: set_random_seed(0); v = m.sample(100)
sage: m.baum_welch(v, fix_emissions=True)
(-66.98630856918774, 100)
sage: m.emission_matrix()
[0.5 0.5]
[0.2 0.8]
sage: m = hmm.DiscreteHiddenMarkovModel([[0.1,0.9],[0.9,0.1]],
.....:                                  [[.5,.5],[.2,.8]],
.....:                                  [.2,.8])
sage: m.baum_welch(v)
(-66.782360659293..., 100)
sage: m.emission_matrix() # rel tol 1e-14
[ 0.5303085748626447 0.46969142513735535]
[ 0.2909775550173978 0.7090224449826023]

```

### `emission_matrix()`

Return the matrix whose  $i$ -th row specifies the emission probability distribution for the  $i$ -th state.

More precisely, the  $i, j$  entry of the matrix is the probability of the Markov model outputting the  $j$ -th symbol when it is in the  $i$ -th state.

OUTPUT: a Sage matrix with real double precision (RDF) entries.

EXAMPLES:

```
sage: m = hmm.DiscreteHiddenMarkovModel([[0.4,0.6],[0.1,0.9]],
.....:                                  [[0.1,0.9],[0.5,0.5]],
.....:                                  [.5,.5])
sage: E = m.emission_matrix(); E
[0.1 0.9]
[0.5 0.5]
```

The returned matrix is mutable, but changing it does not change the transition matrix for the model:

```
sage: E[0,0] = 0; E[0,1] = 1
sage: m.emission_matrix()
[0.1 0.9]
[0.5 0.5]
```

**generate\_sequence** (*length*, *starting\_state=None*)

Return a sample of the given length from this HMM.

INPUT:

- *length* – positive integer
- *starting\_state* – integer (or None); if specified, generate a sequence using this model starting with the given state instead of the initial probabilities to determine the starting state

OUTPUT:

- an *IntList* or list of emission symbols
- *IntList* of the actual states the model was in when emitting the corresponding symbols

EXAMPLES:

In this example, the emission symbols are not set:

```
sage: set_random_seed(0)
sage: a = hmm.DiscreteHiddenMarkovModel([[0.1,0.9],[0.1,0.9]],
.....:                                  [[1,0],[0,1]],
.....:                                  [0,1])
sage: a.generate_sequence(5)
([1, 0, 1, 1, 1], [1, 0, 1, 1, 1])
sage: list(a.generate_sequence(1000)[0]).count(0)
90
```

Here the emission symbols are set:

```
sage: set_random_seed(0)
sage: a = hmm.DiscreteHiddenMarkovModel([[0.5,0.5],[0.1,0.9]],
.....:                                  [[1,0],[0,1]],
.....:                                  [0,1], ['up', 'down'])
sage: a.generate_sequence(5)
(['down', 'up', 'down', 'down', 'down'], [1, 0, 1, 1, 1])
```

Specify the starting state:

```
sage: set_random_seed(0); a.generate_sequence(5, starting_state=0)
(['up', 'up', 'down', 'down', 'down'], [0, 0, 1, 1, 1])
```

**log\_likelihood** (*obs*, *scale=True*)

Return the logarithm of the probability that this model produced the given observation sequence. Thus the output is a nonpositive number.

## INPUT:

- `obs` – sequence of observations
- `scale` – boolean (default: `True`); if `True`, use rescaling to overoid loss of precision due to the very limited dynamic range of floats. You should leave this as `True` unless the `obs` sequence is very small.

## EXAMPLES:

```
sage: m = hmm.DiscreteHiddenMarkovModel([[0.4,0.6],[0.1,0.9]],
....:                                  [[0.1,0.9],[0.5,0.5]],
....:                                  [.2,.8])
sage: m.log_likelihood([0, 1, 0, 1, 1, 0, 1, 0, 0, 0])
-7.3301308009370825
sage: m.log_likelihood([0, 1, 0, 1, 1, 0, 1, 0, 0, 0], scale=False)
-7.330130800937082
sage: m.log_likelihood([])
0.0

sage: m = hmm.DiscreteHiddenMarkovModel([[0.4,0.6],[0.1,0.9]],
....:                                  [[0.1,0.9],[0.5,0.5]],
....:                                  [.2,.8], ['happy','sad'])
sage: m.log_likelihood(['happy','happy'])
-1.6565295199679506
sage: m.log_likelihood(['happy','sad'])
-1.4731602941415523
```

## Overflow from not using the scale option:

```
sage: m = hmm.DiscreteHiddenMarkovModel([[0.4,0.6],[0.1,0.9]],
....:                                  [[0.1,0.9],[0.5,0.5]],
....:                                  [.2,.8])
sage: m.log_likelihood([0,1]*1000, scale=True)
-1433.820666652728
sage: m.log_likelihood([0,1]*1000, scale=False)
-inf
```

**viterbi** (*obs*, *log\_scale=True*)

Determine “the” hidden sequence of states that is most likely to produce the given sequence `seq` of observations, along with the probability that this hidden sequence actually produced the observation.

## INPUT:

- `seq` – sequence of emitted ints or symbols
- `log_scale` – boolean (default: `True`); whether to scale the sequence in order to avoid numerical overflow

## OUTPUT:

- `list` – “the” most probable sequence of hidden states, i.e., the Viterbi path
- `float` – log of probability that the observed sequence was produced by the Viterbi sequence of states

## EXAMPLES:

```
sage: a = hmm.DiscreteHiddenMarkovModel([[0.1,0.9],[0.1,0.9]],
....:                                  [[0.9,0.1],[0.1,0.9]],
....:                                  [0.5,0.5])
sage: a.viterbi([1,0,0,1,0,0,1,1])
([1, 0, 0, 1, ..., 0, 1, 1], -11.06245322477221...)
```

We predict the state sequence when the emissions are 3/4 and 'abc':

```
sage: a = hmm.DiscreteHiddenMarkovModel([[0.1, 0.9], [0.1, 0.9]],
....:                                     [[0.9, 0.1], [0.1, 0.9]],
....:                                     [0.5, 0.5], [3/4, 'abc'])
```

Note that state 0 is common below, despite the model trying hard to switch to state 1:

```
sage: a.viterbi([3/4, 'abc', 'abc'] + [3/4]*10)
[0, 1, 1, 0, 0 ... 0, 0, 0, 0, 0], -25.299405845367794
```

**class** sage.stats.hmm.hmm.**HiddenMarkovModel**

Bases: object

Abstract base class for all Hidden Markov Models.

**graph** (*eps=0.001*)

Create a weighted directed graph from the transition matrix, not including any edge with a probability less than *eps*.

INPUT:

- *eps* – nonnegative real number

OUTPUT: a `DiGraph`

EXAMPLES:

```
sage: m = hmm.DiscreteHiddenMarkovModel([[.3, 0, .7], [0, 0, 1], [.5, .5, 0]],
....:                                     [[.5, .5, .2]]*3,
....:                                     [1/3]*3)
sage: G = m.graph(); G #_
↪needs sage.graphs
Looped digraph on 3 vertices
sage: G.edges(sort=True) #_
↪needs sage.graphs
[(0, 0, 0.3), (0, 2, 0.7), (1, 2, 1.0), (2, 0, 0.5), (2, 1, 0.5)]
sage: G.plot() #_
↪needs sage.graphs sage.plot
Graphics object consisting of 11 graphics primitives
```

**initial\_probabilities** ()

Return the initial probabilities as a `TimeSeries` of length *N*, where *N* is the number of states of the Markov model.

EXAMPLES:

```
sage: m = hmm.DiscreteHiddenMarkovModel([[0.4, 0.6], [0.1, 0.9]],
....:                                     [[0.1, 0.9], [0.5, 0.5]],
....:                                     [.2, .8])
sage: pi = m.initial_probabilities(); pi
[0.2000, 0.8000]
sage: type(pi)
<... 'sage.stats.time_series.TimeSeries'>
```

The returned time series is a copy, so changing it does not change the model:

```
sage: pi[0] = .1; pi[1] = .9
sage: m.initial_probabilities()
[0.2000, 0.8000]
```

Some other models:

```
sage: m = hmm.GaussianHiddenMarkovModel([[.1,.9],[.5,.5]],
....:                                  [(1,1), (-1,1)],
....:                                  [.1,.9])
sage: m.initial_probabilities()
[0.1000, 0.9000]
sage: m = hmm.GaussianMixtureHiddenMarkovModel(
....:   [[.9,.1],[.4,.6]],
....:   [[(.4,(0,1)), (.6,(1,0.1))], [(1,(0,1))]],
....:   [.7,.3])
sage: m.initial_probabilities()
[0.7000, 0.3000]
```

**sample** (*length*, *number=None*, *starting\_state=None*)

Return number samples from this HMM of given length.

INPUT:

- *length* – positive integer
- *number* – (default: None) if given, compute list of this many sample sequences
- *starting\_state* – integer (or None); if specified, generate a sequence using this model starting with the given state instead of the initial probabilities to determine the starting state

OUTPUT:

- if *number* is not given, return a single TimeSeries.
- if *number* is given, return a list of TimeSeries.

EXAMPLES:

```
sage: set_random_seed(0)
sage: a = hmm.DiscreteHiddenMarkovModel([[0.1,0.9],[0.1,0.9]],
....:                                  [[1,0],[0,1]],
....:                                  [0,1])
sage: print(a.sample(10, 3))
[[1, 0, 1, 1, 1, 1, 0, 1, 1, 1],
 [1, 1, 0, 0, 1, 1, 1, 1, 1, 1],
 [1, 1, 1, 1, 0, 1, 0, 1, 1, 1]]
sage: a.sample(15)
[1, 1, 1, 1, 0 ... 1, 1, 1, 1, 1]
sage: a.sample(3, 1)
[[1, 1, 1]]
sage: list(a.sample(1000)).count(0)
88
```

If the emission symbols are set:

```
sage: set_random_seed(0)
sage: a = hmm.DiscreteHiddenMarkovModel([[0.5,0.5],[0.1,0.9]],
....:                                  [[1,0],[0,1]], [0,1],
....:                                  ['up', 'down'])
sage: a.sample(10)
['down', 'up', 'down', 'down', 'down', 'down', 'up', 'up', 'up', 'up']
```

Force a starting state:

```
sage: set_random_seed(0); a.sample(10, starting_state=0)
['up', 'up', 'down', 'down', 'down', 'down', 'up', 'up', 'up', 'up']
```

**transition\_matrix()**

Return the state transition matrix.

OUTPUT: a Sage matrix with real double precision (RDF) entries.

EXAMPLES:

```
sage: M = hmm.DiscreteHiddenMarkovModel([[0.7,0.3],[0.9,0.1]],
....:                                  [[0.5,.5],[.1,.9]],
....:                                  [0.3,0.7])
sage: T = M.transition_matrix(); T
[0.7 0.3]
[0.9 0.1]
```

The returned matrix is mutable, but changing it does not change the transition matrix for the model:

```
sage: T[0,0] = .1; T[0,1] = .9
sage: M.transition_matrix()
[0.7 0.3]
[0.9 0.1]
```

Transition matrices for other types of models:

```
sage: M = hmm.GaussianHiddenMarkovModel([[.1,.9],[.5,.5]],
....:                                  [(1,1), (-1,1)],
....:                                  [.5,.5])
sage: M.transition_matrix()
[0.1 0.9]
[0.5 0.5]
sage: M = hmm.GaussianMixtureHiddenMarkovModel(
....:   [[.9,.1],[.4,.6]],
....:   [[(.4,(0,1)), (.6,(1,0.1))],[ (1,(0,1))]],
....:   [.7,.3])
sage: M.transition_matrix()
[0.9 0.1]
[0.4 0.6]
```

`sage.stats.hmm.hmm.unpickle_discrete_hmm_v0(A, B, pi, emission_symbols, name)`

`sage.stats.hmm.hmm.unpickle_discrete_hmm_v1(A, B, pi, n_out, emission_symbols, emission_symbols_dict)`

Return a *DiscreteHiddenMarkovModel*, restored from the arguments.

This function is used internally for unpickling.

## CONTINUOUS EMISSION HIDDEN MARKOV MODELS

AUTHOR:

- William Stein, 2010-03

**class** sage.stats.hmm.chmm.**GaussianHiddenMarkovModel**

Bases: *HiddenMarkovModel*

Gaussian emissions Hidden Markov Model.

INPUT:

- A – matrix; the  $N \times N$  transition matrix
- B – list of pairs (*mu*, *sigma*) that define the distributions
- pi – initial state probabilities
- normalize – boolean (default: True)

EXAMPLES:

We illustrate the primary functions with an example 2-state Gaussian HMM:

```
sage: m = hmm.GaussianHiddenMarkovModel([[.1, .9], [.5, .5]],
.....:                                     [(1,1), (-1,1)],
.....:                                     [.5, .5]); m
Gaussian Hidden Markov Model with 2 States
Transition matrix:
[0.1 0.9]
[0.5 0.5]
Emission parameters:
[(1.0, 1.0), (-1.0, 1.0)]
Initial probabilities: [0.5000, 0.5000]
```

We query the defining transition matrix, emission parameters, and initial state probabilities:

```
sage: m.transition_matrix()
[0.1 0.9]
[0.5 0.5]
sage: m.emission_parameters()
[(1.0, 1.0), (-1.0, 1.0)]
sage: m.initial_probabilities()
[0.5000, 0.5000]
```

We obtain a sample sequence with 10 entries in it, and compute the logarithm of the probability of obtaining this sequence, given the model:

```

sage: obs = m.sample(5); obs # random
[-1.6835, 0.0635, -2.1688, 0.3043, -0.3188]
sage: log_likelihood = m.log_likelihood(obs)
sage: counter = 0
sage: n = 0
sage: def add_samples(i):
.....:     global counter, n
.....:     for _ in range(i):
.....:         n += 1
.....:         obs2 = m.sample(5)
.....:         if all(abs(obs2[i] - obs[i]) < 0.25 for i in range(5)):
.....:             counter += 1

sage: add_samples(10000)
sage: while abs(log_likelihood - log(counter*1.0/n/0.5^5)) < 0.1:
.....:     add_samples(10000)

```

We compute the Viterbi path, and probability that the given path of states produced obs:

```

sage: m.viterbi(obs) # random
([1, 0, 1, 0, 1], -8.714092684611794)

```

We use the Baum-Welch iterative algorithm to find another model for which our observation sequence is more likely:

```

sage: try:
.....:     p, s = m.baum_welch(obs)
.....:     assert p > log_likelihood
.....:     assert (1 <= s <= 500)
.....: except RuntimeError:
.....:     pass

```

Notice that running Baum-Welch changed our model:

```

sage: m # random
Gaussian Hidden Markov Model with 2 States
Transition matrix:
[ 0.4154981366185841 0.584501863381416]
[ 0.9999993174253741 6.825746258991804e-07]
Emission parameters:
[(0.4178882427119503, 0.5173109664360919),
 (-1.5025208631331122, 0.5085512836055119)]
Initial probabilities: [0.0000, 1.0000]

```

**baum\_welch** (*obs*, *max\_iter*=500, *log\_likelihood\_cutoff*=0.0001, *min\_sd*=0.01, *fix\_emissions*=False, *v*=False)

Given an observation sequence *obs*, improve this HMM using the Baum-Welch algorithm to increase the probability of observing *obs*.

INPUT:

- *obs* – a time series of emissions
- *max\_iter* – integer (default: 500); maximum number of Baum-Welch steps to take
- *log\_likelihood\_cutoff* – positive float (default: 1e-4); the minimal improvement in likelihood with respect to the last iteration required to continue. Relative value to log likelihood.
- *min\_sd* – positive float (default: 0.01); when reestimating, the standard deviation of emissions is not allowed to be less than *min\_sd*.



- `fix_emissions` – boolean (default: `False`); if `True`, do not change emissions when updating

OUTPUT:

changes the model in place, and returns the log likelihood and number of iterations.

EXAMPLES:

```
sage: m = hmm.GaussianHiddenMarkovModel([[.1,.9],[.5,.5]],
....:                                  [(1,.5), (-1,3)],
....:                                  [.1,.9])
sage: m.log_likelihood([-2,-1,.1,0.1])
-8.858282215986275
sage: m.baum_welch([-2,-1,.1,0.1])
(4.534646052182..., 7)
sage: m.log_likelihood([-2,-1,.1,0.1])
4.534646052182...
sage: m # rel tol 3e-14
Gaussian Hidden Markov Model with 2 States
Transition matrix:
[  0.9999999992430161  7.569839394440382e-10]
[  0.49998462791192644    0.5000153720880736]
Emission parameters:
[(0.09999999999999999, 0.01), (-1.4999508147591902, 0.5000710504895474)]
Initial probabilities: [0.0000, 1.0000]
```

We illustrate bounding the standard deviation below. Note that above we had different emission parameters when the `min_sd` was the default of 0.01:

```
sage: m = hmm.GaussianHiddenMarkovModel([[.1,.9],[.5,.5]],
....:                                  [(1,.5), (-1,3)],
....:                                  [.1,.9])
sage: m.baum_welch([-2,-1,.1,0.1], min_sd=1)
(-4.07939572755..., 32)
sage: m.emission_parameters()
[(-0.2663018798..., 1.0), (-1.99850979..., 1.0)]
```

We watch the log likelihoods of the model converge, step by step:

```
sage: m = hmm.GaussianHiddenMarkovModel([[.1,.9],[.5,.5]],
....:                                  [(1,.5), (-1,3)],
....:                                  [.1,.9])
sage: v = m.sample(10)
sage: l = stats.TimeSeries([m.baum_welch(v, max_iter=1)[0]
....:                       for _ in range(len(v))])
sage: all(l[i] <= l[i+1] + 0.0001 for i in range(9))
True
sage: l # random
[-20.1167, -17.7611, -16.9814, -16.9364, -16.9314,
-16.9309, -16.9309, -16.9309, -16.9309, -16.9309]
```

We illustrate fixing emissions:

```
sage: m = hmm.GaussianHiddenMarkovModel([[.1,.9],[.9,.1]],
....:                                  [(1,2), (-1,.5)],
....:                                  [.3,.7])
sage: set_random_seed(0); v = m.sample(100)
sage: m.baum_welch(v, fix_emissions=True)
(-164.72944548204..., 23)
```

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```
sage: m.emission_parameters()
[(1.0, 2.0), (-1.0, 0.5)]
sage: m = hmm.GaussianHiddenMarkovModel([[.1, .9], [.9, .1]],
....:                                     [(1, 2), (-1, .5)],
....:                                     [.3, .7])
sage: m.baum_welch(v)
(-162.854370397998..., 49)
sage: m.emission_parameters() # rel tol 3e-14
[(1.2722419172602375, 2.371368751761901),
 (-0.9486174675179113, 0.5762360385123765)]
```

**emission\_parameters()**

Return the parameters that define the normal distributions associated to all of the states.

OUTPUT:

a list B of pairs  $B[i] = (\mu, \text{std})$ , such that the distribution associated to state  $i$  is normal with mean  $\mu$  and standard deviation  $\text{std}$ .

EXAMPLES:

```
sage: M = hmm.GaussianHiddenMarkovModel([[.1, .9], [.5, .5]],
....:                                     [(1, .5), (-1, 3)],
....:                                     [.1, .9])
sage: M.emission_parameters()
[(1.0, 0.5), (-1.0, 3.0)]
```

**generate\_sequence** (*length, starting\_state=None*)

Return a sample of the given length from this HMM.

INPUT:

- length – positive integer
- starting\_state – integer (or None); if specified then generate a sequence using this model starting with the given state instead of the initial probabilities to determine the starting state.

OUTPUT:

- an *IntList* or list of emission symbols
- TimeSeries of emissions

EXAMPLES:

```
sage: m = hmm.GaussianHiddenMarkovModel([[.1, .9], [.5, .5]],
....:                                     [(1, .5), (-1, 3)],
....:                                     [.1, .9])
sage: m.generate_sequence(5) # random
[[-3.0505, 0.5317, -4.5065, 0.6521, 1.0435], [1, 0, 1, 0, 1]]
sage: m.generate_sequence(0)
([], [])
sage: m.generate_sequence(-1)
Traceback (most recent call last):
...
ValueError: length must be nonnegative
```

Verify numerically that the starting state is 0 with probability about 0.1:

```

sage: counter = 0
sage: n = 0
sage: def add_samples(i):
.....:     global counter, n
.....:     for i in range(i):
.....:         n += 1
.....:         if m.generate_sequence(1)[1][0] == 0:
.....:             counter += 1

sage: add_samples(10^5)
sage: while abs(counter*1.0 / n - 0.1) > 0.01: add_samples(10^5)

```

Example in which the starting state is 0 (see [Issue #11452](#)):

```

sage: set_random_seed(23); m.generate_sequence(2)
([0.6501, -2.0151], [0, 1])

```

Force a starting state of 1 even though as we saw above it would be 0:

```

sage: set_random_seed(23); m.generate_sequence(2, starting_state=1)
([-3.1491, -1.0244], [1, 1])

```

### **log\_likelihood** (*obs*)

Return the logarithm of a continuous analogue of the probability that this model produced the given observation sequence.

Note that the “continuous analogue of the probability” above can be bigger than 1, hence the logarithm can be positive.

INPUT:

- *obs* – sequence of observations

OUTPUT: float

EXAMPLES:

```

sage: m = hmm.GaussianHiddenMarkovModel([[.1, .9], [.5, .5]],
.....:                                  [(1, .5), (-1, 3)],
.....:                                  [.1, .9])
sage: m.log_likelihood([1, 1, 1])
-4.297880766072486
sage: s = m.sample(20)
sage: -80 < m.log_likelihood(s) < -20
True

```

### **viterbi** (*obs*)

Determine “the” hidden sequence of states that is most likely to produce the given sequence *obs* of observations, along with the probability that this hidden sequence actually produced the observation.

INPUT:

- *obs* – sequence of emitted ints or symbols

OUTPUT:

- list – “the” most probable sequence of hidden states, i.e., the Viterbi path
- float – log of probability that the observed sequence was produced by the Viterbi sequence of states

EXAMPLES:

We find the optimal state sequence for a given model:

```
sage: m = hmm.GaussianHiddenMarkovModel([[0.5,0.5],[0.5,0.5]],
....:                                  [(0,1),(10,1)],
....:                                  [0.5,0.5])
sage: m.viterbi([0,1,10,10,1])
([0, 0, 1, 1, 0], -9.0604285688230...)
```

Another example in which the most likely states change based on the last observation:

```
sage: m = hmm.GaussianHiddenMarkovModel([[.1,.9],[.5,.5]],
....:                                  [(1,.5), (-1,3)],
....:                                  [.1,.9])
sage: m.viterbi([-2,-1,.1,0.1])
([1, 1, 0, 1], -9.61823698847639...)
sage: m.viterbi([-2,-1,.1,0.3])
([1, 1, 1, 0], -9.566023653378513)
```

**class** sage.stats.hmm.chmm.**GaussianMixtureHiddenMarkovModel**

Bases: *GaussianHiddenMarkovModel*

Gaussian mixture Hidden Markov Model.

INPUT:

- *A* – matrix; the  $N \times N$  transition matrix
- *B* – list of mixture definitions for each state. Each state may have a varying number of gaussians with selection probabilities that sum to 1 and encoded as  $(p, (\mu, \sigma))$
- *pi* – initial state probabilities
- *normalize* – boolean (default: True); if given, input is normalized to define valid probability distributions, e.g., the entries of *A* are made nonnegative and the rows sum to 1, and the probabilities in *pi* are normalized.

EXAMPLES:

```
sage: A = [[0.5,0.5],[0.5,0.5]]
sage: B = [[(0.9,(0.0,1.0)), (0.1,(1,10000))],[(1,(1,1)), (0,(0,0.1))]]
sage: hmm.GaussianMixtureHiddenMarkovModel(A, B, [1,0])
Gaussian Mixture Hidden Markov Model with 2 States
Transition matrix:
[0.5 0.5]
[0.5 0.5]
Emission parameters:
[0.9*N(0.0,1.0) + 0.1*N(1.0,10000.0), 1.0*N(1.0,1.0) + 0.0*N(0.0,0.1)]
Initial probabilities: [1.0000, 0.0000]
```

**baum\_welch** (*obs*, *max\_iter*=1000, *log\_likelihood\_cutoff*=1e-12, *min\_sd*=0.01, *fix\_emissions*=False)

Given an observation sequence *obs*, improve this HMM using the Baum-Welch algorithm to increase the probability of observing *obs*.

INPUT:

- *obs* – a time series of emissions
- *max\_iter* – integer (default: 1000); maximum number of Baum-Welch steps to take
- *log\_likelihood\_cutoff* – positive float (default: 1e-12); the minimal improvement in likelihood with respect to the last iteration required to continue. Relative value to log likelihood.

- `min_sd` – positive float (default: 0.01); when reestimating, the standard deviation of emissions is not allowed to be less than `min_sd`.
- `fix_emissions` – boolean (default: False); if True, do not change emissions when updating

OUTPUT:

changes the model in place, and returns the log likelihood and number of iterations.

EXAMPLES:

```
sage: m = hmm.GaussianMixtureHiddenMarkovModel(
.....:     [[.9, .1], [.4, .6]],
.....:     [[(.4, (0,1)), (.6, (1,0.1))], [(1, (0,1))]],
.....:     [.7, .3])
sage: set_random_seed(0); v = m.sample(10); v
[0.3576, -0.9365, 0.9449, -0.6957, 1.0217,
 0.9644, 0.9987, -0.5950, -1.0219, 0.6477]
sage: m.log_likelihood(v)
-8.31408655939536...
sage: m.baum_welch(v)
(2.18905068682..., 15)
sage: m.log_likelihood(v)
2.18905068682...
sage: m # rel tol 6e-12
Gaussian Mixture Hidden Markov Model with 2 States
Transition matrix:
[ 0.8746363339773399  0.12536366602266016]
[                    1.0  1.451685202290174e-40]
Emission parameters:
[0.500161629343*N(-0.812298726239,0.173329026744)
 + 0.499838370657*N(0.982433690378,0.029719932009),
 1.0*N(0.503260056832,0.145881515324)]
Initial probabilities: [0.0000, 1.0000]
```

We illustrate bounding the standard deviation below. Note that above we had different emission parameters when the `min_sd` was the default of 0.01:

```
sage: m = hmm.GaussianMixtureHiddenMarkovModel(
.....:     [[.9, .1], [.4, .6]],
.....:     [[(.4, (0,1)), (.6, (1,0.1))], [(1, (0,1))]],
.....:     [.7, .3])
sage: m.baum_welch(v, min_sd=1)
(-12.617885761692..., 1000)
sage: m.emission_parameters() # rel tol 6e-12
[0.503545634447*N(0.200166509595,1.0) + 0.496454365553*N(0.200166509595,1.0),
 1.0*N(0.0543433426535,1.0)]
```

We illustrate fixing all emissions:

```
sage: m = hmm.GaussianMixtureHiddenMarkovModel(
.....:     [[.9, .1], [.4, .6]],
.....:     [[(.4, (0,1)), (.6, (1,0.1))], [(1, (0,1))]],
.....:     [.7, .3])
sage: set_random_seed(0); v = m.sample(10)
sage: m.baum_welch(v, fix_emissions=True)
(-7.58656858997..., 36)
sage: m.emission_parameters()
[0.4*N(0.0,1.0) + 0.6*N(1.0,0.1),
 1.0*N(0.0,1.0)]
```

**emission\_parameters()**

Return a list of all the emission distributions.

OUTPUT: list of Gaussian mixtures

EXAMPLES:

```
sage: m = hmm.GaussianMixtureHiddenMarkovModel([[.9, .1], [.4, .6]],
.....:                                     [[(.4, (0, 1)), (.6, (1, 0.1))],
↪ [(1, (0, 1))]],
.....:                                     [.7, .3])
sage: m.emission_parameters()
[0.4*N(0.0, 1.0) + 0.6*N(1.0, 0.1), 1.0*N(0.0, 1.0)]
```

sage.stats.hmm.chmm.**unpickle\_gaussian\_hmm\_v0**(A, B, pi, name)

EXAMPLES:

```
sage: m = hmm.GaussianHiddenMarkovModel([[1]], [(0, 1)], [1])
sage: sage.stats.hmm.chmm.unpickle_gaussian_hmm_v0(m.transition_matrix(), m.
↪emission_parameters(), m.initial_probabilities(), 'test')
Gaussian Hidden Markov Model with 1 States
Transition matrix:
[1.0]
Emission parameters:
[(0.0, 1.0)]
Initial probabilities: [1.0000]
```

sage.stats.hmm.chmm.**unpickle\_gaussian\_hmm\_v1**(A, B, pi, prob, n\_out)

EXAMPLES:

```
sage: m = hmm.GaussianHiddenMarkovModel([[1]], [(0, 1)], [1])
sage: loads(dumps(m)) == m # indirect test
True
```

sage.stats.hmm.chmm.**unpickle\_gaussian\_mixture\_hmm\_v1**(A, B, pi, mixture)

EXAMPLES:

```
sage: m = hmm.GaussianMixtureHiddenMarkovModel([[1]], [[(.4, (0, 1)), (.6, (1, 0.
↪1))]], [1])
sage: loads(dumps(m)) == m # indirect test
True
```

## DISTRIBUTIONS USED IN IMPLEMENTING HIDDEN MARKOV MODELS

These distribution classes are designed specifically for HMM's and not for general use in statistics. For example, they have fixed or non-fixed status, which only make sense relative to being used in a hidden Markov model.

AUTHOR:

- William Stein, 2010-03

**class** sage.stats.hmm.distributions.**DiscreteDistribution**

Bases: *Distribution*

**class** sage.stats.hmm.distributions.**Distribution**

Bases: object

A distribution.

**plot** (\*args, \*\*kws)

Return a plot of the probability density function.

INPUT:

- args and kws, passed to the Sage *plot()* function

OUTPUT: a *Graphics* object

EXAMPLES:

```
sage: P = hmm.GaussianMixtureDistribution([(0.2, -10, 0.5), (0.6, 1, 1), (0.2, 20, 0.5)])
sage: P.plot(-10, 30) #_
↔needs sage.plot
Graphics object consisting of 1 graphics primitive
```

**prob** (x)

The probability density function evaluated at *x*.

INPUT:

- *x* – object

OUTPUT: float

EXAMPLES:

This method must be defined in a derived class:

```
sage: import sage.stats.hmm.distributions
sage: sage.stats.hmm.distributions.Distribution().prob(0)
Traceback (most recent call last):
...
NotImplementedError
```

**sample** (*n=None*)

Return either a single sample (the default) or *n* samples from this probability distribution.

INPUT:

- *n* – None or a positive integer

OUTPUT: a single sample if *n* is 1; otherwise many samples

EXAMPLES:

This method must be defined in a derived class:

```
sage: import sage.stats.hmm.distributions
sage: sage.stats.hmm.distributions.Distribution().sample()
Traceback (most recent call last):
...
NotImplementedError
```

**class** sage.stats.hmm.distributions.**GaussianDistribution**

Bases: *Distribution*

**class** sage.stats.hmm.distributions.**GaussianMixtureDistribution**

Bases: *Distribution*

A probability distribution defined by taking a weighted linear combination of Gaussian distributions.

EXAMPLES:

```
sage: P = hmm.GaussianMixtureDistribution([(0.3,1,2), (.7,-1,1)]); P
0.3*N(1.0,2.0) + 0.7*N(-1.0,1.0)
sage: P[0]
(0.3, 1.0, 2.0)
sage: P.is_fixed()
False
sage: P.fix(1)
sage: P.is_fixed(0)
False
sage: P.is_fixed(1)
True
sage: P.unfix(1)
sage: P.is_fixed(1)
False
```

**fix** (*i=None*)

Set that this *GaussianMixtureDistribution* (or its *i*-th component) is fixed when using Baum-Welch to update the corresponding HMM.

INPUT:

- *i* – None (default) or integer; if given, only fix the *i*-th component

EXAMPLES:



```

sage: P = hmm.GaussianMixtureDistribution([(0.2, -10, .5), (.6, 1, 1), (.2, 20, .5)])
sage: P.fix(1); P.is_fixed()
False
sage: P.is_fixed(1)
True
sage: P.fix(); P.is_fixed()
True

```

**is\_fixed** (*i=None*)

Return whether or not this *GaussianMixtureDistribution* is fixed when using Baum-Welch to update the corresponding HMM.

INPUT:

- *i* – None (default) or integer; if given, only return whether the *i*-th component is fixed

EXAMPLES:

```

sage: P = hmm.GaussianMixtureDistribution([(0.2, -10, .5), (.6, 1, 1), (.2, 20, .5)])
sage: P.is_fixed()
False
sage: P.is_fixed(0)
False
sage: P.fix(0); P.is_fixed()
False
sage: P.is_fixed(0)
True
sage: P.fix(); P.is_fixed()
True

```

**prob** (*x*)

Return the probability of *x*.

Since this is a continuous distribution, this is defined to be the limit of the *p*'s such that the probability of [*x*,*x*+*h*] is *p*\**h*.

INPUT:

- *x* – float

OUTPUT: float

EXAMPLES:

```

sage: P = hmm.GaussianMixtureDistribution([(0.2, -10, .5), (.6, 1, 1), (.2, 20, .5)])
sage: P.prob(.5)
0.21123919605857971
sage: P.prob(-100)
0.0
sage: P.prob(20)
0.1595769121605731

```

**prob\_m** (*x, m*)

Return the probability of *x* using just the *m*-th summand.

INPUT:

- *x* – float
- *m* – integer

OUTPUT: float

EXAMPLES:

```
sage: P = hmm.GaussianMixtureDistribution([(0.2, -10, .5), (.6, 1, 1), (.2, 20, .5)])
sage: P.prob_m(.5, 0)
2.7608117680508...e-97
sage: P.prob_m(.5, 1)
0.21123919605857971
sage: P.prob_m(.5, 2)
0.0
```

**sample** (*n=None*)

Return a single sample from this distribution (by default), or if  $n > 1$ , return a `TimeSeries` of samples.

INPUT:

- $n$  – integer or `None` (default: `None`)

OUTPUT:

- float if  $n$  is `None` (default); otherwise a `TimeSeries`

EXAMPLES:

```
sage: P = hmm.GaussianMixtureDistribution([(0.2, -10, .5), (.6, 1, 1), (.2, 20, .5)])
sage: type(P.sample())
<class 'float'>
sage: l = P.sample(1)
sage: len(l)
1
sage: type(l)
<class 'sage.stats.time_series.TimeSeries'>
sage: l = P.sample(5)
sage: len(l)
5
sage: type(l)
<class 'sage.stats.time_series.TimeSeries'>
sage: l = P.sample(0)
sage: len(l)
0
sage: type(l)
<class 'sage.stats.time_series.TimeSeries'>
sage: P.sample(-3)
Traceback (most recent call last):
...
ValueError: n must be nonnegative
```

**unfix** (*i=None*)

Set that this *GaussianMixtureDistribution* (or its  $i$ -th component) is not fixed when using Baum-Welch to update the corresponding HMM.

INPUT:

- $i$  – `None` (default) or integer; if given, only fix the  $i$ -th component

EXAMPLES:

```
sage: P = hmm.GaussianMixtureDistribution([(0.2, -10, .5), (.6, 1, 1), (.2, 20, .5)])
sage: P.fix(1); P.is_fixed(1)
```

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```
True
sage: P.unfix(1); P.is_fixed(1)
False
sage: P.fix(); P.is_fixed()
True
sage: P.unfix(); P.is_fixed()
False
```

```
sage.stats.hmm.distributions.unpickle_gaussian_mixture_distribution_v1(c0, c1,
                                                                    param,
                                                                    fixed)
```

Used in unpickling *GaussianMixtureDistribution* objects.

#### EXAMPLES:

```
sage: P = hmm.GaussianMixtureDistribution([(0.2, -10, 0.5), (0.6, 1, 1), (0.2, 20, 0.5)])
sage: loads(dumps(P)) == P          # indirect doctest
True
```



## HIDDEN MARKOV MODELS – UTILITY FUNCTIONS

AUTHOR:

- William Stein, 2010-03

**class** sage.stats.hmm.util.HMM\_Util

Bases: object

A class used in order to share cdef's methods between different files.

**initial\_probs\_to\_TimeSeries** (*pi*, *normalize*)

This function is used internally by the `__init__` methods of various Hidden Markov Models.

INPUT:

- *pi* – vector, list, or TimeSeries
- *normalize* – if True, replace negative entries by 0 and rescale to ensure that the sum of the entries in each row is equal to 1. If the sum of the entries in a row is 0, replace them all by  $1/N$ .

OUTPUT: a TimeSeries of length *N*

EXAMPLES:

```
sage: import sage.stats.hmm.util
sage: u = sage.stats.hmm.util.HMM_Util()
sage: u.initial_probs_to_TimeSeries([0.1,0.2,0.9], True)
[0.0833, 0.1667, 0.7500]
sage: u.initial_probs_to_TimeSeries([0.1,0.2,0.9], False)
[0.1000, 0.2000, 0.9000]
```

**normalize\_probability\_TimeSeries** (*T*, *i*, *j*)

This function is used internally by the Hidden Markov Models code.

Replace entries of  $T[i:j]$  in place so that they are all nonnegative and sum to 1. Negative entries are replaced by 0 and  $T[i:j]$  is then rescaled to ensure that the sum of the entries in each row is equal to 1. If all entries are 0, replace them by  $1/(j-i)$ .

INPUT:

- *T* – a TimeSeries
- *i* – nonnegative integer
- *j* – nonnegative integer

OUTPUT: *T* is modified

EXAMPLES:

```

sage: import sage.stats.hmm.util
sage: T = stats.TimeSeries([.1, .3, .7, .5])
sage: u = sage.stats.hmm.util.HMM_Util()
sage: u.normalize_probability_TimeSeries(T,0,3)
sage: T
[0.0909, 0.2727, 0.6364, 0.5000]
sage: u.normalize_probability_TimeSeries(T,0,4)
sage: T
[0.0606, 0.1818, 0.4242, 0.3333]
sage: abs(T.sum()-1) < 1e-8      # might not exactly equal 1 due to rounding
True

```

**state\_matrix\_to\_TimeSeries** (*A*, *N*, *normalize*)

This function is used internally by the `__init__` methods of Hidden Markov Models to make a transition matrix from *A*.

INPUT:

- *A* – matrix, list, list of lists, or TimeSeries
- *N* – number of states
- *normalize* – if True, replace negative entries by 0 and rescale to ensure that the sum of the entries in each row is equal to 1. If the sum of the entries in a row is 0, replace them all by  $1/N$ .

OUTPUT: a TimeSeries

EXAMPLES:

```

sage: import sage.stats.hmm.util
sage: u = sage.stats.hmm.util.HMM_Util()
sage: u.state_matrix_to_TimeSeries([[.1, .7], [3/7, 4/7]], 2, True)
[0.1250, 0.8750, 0.4286, 0.5714]
sage: u.state_matrix_to_TimeSeries([[.1, .7], [3/7, 4/7]], 2, False)
[0.1000, 0.7000, 0.4286, 0.5714]

```

## DISCRETE GAUSSIAN SAMPLERS OVER THE INTEGERS

This class realizes oracles which returns integers proportionally to  $\exp(-(x-c)^2/(2\sigma^2))$ . All oracles are implemented using rejection sampling. See `DiscreteGaussianDistributionIntegerSampler.__init__()` for which algorithms are available.

AUTHORS:

- Martin Albrecht (2014-06-28): initial version

EXAMPLES:

We construct a sampler for the distribution  $D_{3,c}$  with width  $= 3$  and center  $c = 0$ :

```
sage: from sage.stats.distributions.discrete_gaussian_integer import _
      ↪ DiscreteGaussianDistributionIntegerSampler
sage: sigma = 3.0
sage: D = DiscreteGaussianDistributionIntegerSampler(sigma=sigma)
```

We ask for 100000 samples:

```
sage: from collections import defaultdict
sage: counter = defaultdict(Integer)
sage: n = 0
sage: def add_samples(i):
.....:     global counter, n
.....:     for _ in range(i):
.....:         counter[D()] += 1
.....:         n += 1
sage: add_samples(100000)
```

These are sampled with a probability proportional to  $\exp(-x^2/18)$ . More precisely we have to normalise by dividing by the overall probability over all integers. We use the fact that hitting anything more than 6 standard deviations away is very unlikely and compute:

```
sage: bound = (6*sigma).floor()
sage: norm_factor = sum([exp(-x^2/(2*sigma^2)) for x in range(-bound,bound+1)])
sage: norm_factor
7.519...
```

With this normalisation factor, we can now test if our samples follow the expected distribution:

```
sage: expected = lambda x : ZZ(round(n*exp(-x^2/(2*sigma^2))/norm_factor))
sage: observed = lambda x : counter[x]
sage: add_samples(10000)
```

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```
sage: while abs(observed(0)*1.0/expected(0) - 1.0) > 5e-2: add_samples(10000)
sage: while abs(observed(4)*1.0/expected(4) - 1.0) > 5e-2: add_samples(10000)
sage: while abs(observed(-10)*1.0/expected(-10) - 1.0) > 5e-2: add_samples(10000) #_
↪long time
```

We construct an instance with a larger width:

```
sage: from sage.stats.distributions.discrete_gaussian_integer import_
↪DiscreteGaussianDistributionIntegerSampler
sage: sigma = 127
sage: D = DiscreteGaussianDistributionIntegerSampler(sigma=sigma, algorithm=
↪'uniform+online')
```

ask for 100000 samples:

```
sage: from collections import defaultdict
sage: counter = defaultdict(Integer)
sage: n = 0
sage: def add_samples(i):
.....:     global counter, n
.....:     for _ in range(i):
.....:         counter[D()] += 1
.....:         n += 1
sage: add_samples(100000)
```

and check if the proportions fit:

```
sage: expected = lambda x, y: (
.....:     exp(-x^2/(2*sigma^2))/exp(-y^2/(2*sigma^2)).n())
sage: observed = lambda x, y: float(counter[x])/counter[y]

sage: while not all(v in counter for v in (0, 1, -100)): add_samples(10000)

sage: while abs(expected(0, 1) - observed(0, 1)) > 2e-1: add_samples(10000)
sage: while abs(expected(0, -100) - observed(0, -100)) > 2e-1: add_samples(10000)
```

We construct a sampler with  $c\%1! = 0$ :

```
sage: from sage.stats.distributions.discrete_gaussian_integer import_
↪DiscreteGaussianDistributionIntegerSampler
sage: sigma = 3
sage: D = DiscreteGaussianDistributionIntegerSampler(sigma=sigma, c=1/2)
sage: s = 0
sage: n = 0
sage: def add_samples(i):
.....:     global s, n
.....:     for _ in range(i):
.....:         s += D()
.....:         n += 1
.....:
sage: add_samples(100000)
sage: while abs(float(s)/n - 0.5) > 5e-2: add_samples(10000)
```

REFERENCES:

- [DDLL2013]



**class** sage.stats.distributions.discrete\_gaussian\_integer.  
**DiscreteGaussianDistributionIntegerSampler**

Bases: SageObject

A Discrete Gaussian Sampler using rejection sampling.

**\_\_init\_\_** (*sigma*, *c=0*, *tau=6*, *algorithm=None*, *precision='mp'*)

Construct a new sampler for a discrete Gaussian distribution.

INPUT:

- *sigma* – samples  $x$  are accepted with probability proportional to  $\exp(-(x - c)^2/(2^2))$
- *c* – the mean of the distribution. The value of  $c$  does not have to be an integer. However, some algorithms only support integer-valued  $c$  (default: 0)
- *tau* – samples outside the range  $(\lfloor c \rfloor - \tau, \dots, \lfloor c \rfloor + \tau)$  are considered to have probability zero. This bound applies to algorithms which sample from the uniform distribution (default: 6)
- **algorithm** – see list below (default: 'uniform+table' for  $t$  bounded by `DiscreteGaussianDistributionIntegerSampler.table_cutoff` and 'uniform+online' for bigger)
- *precision* – either 'mp' for multi-precision where the actual precision used is taken from *sigma* or 'dp' for double precision. In the latter case results are not reproducible. (default: 'mp')

ALGORITHMS:

- 'uniform+table' – classical rejection sampling, sampling from the uniform distribution and accepted with probability proportional to  $\exp(-(x - c)^2/(2^2))$  where  $\exp(-(x - c)^2/(2^2))$  is precomputed and stored in a table. Any real-valued  $c$  is supported.
- 'uniform+logtable' – samples are drawn from a uniform distribution and accepted with probability proportional to  $\exp(-(x - c)^2/(2^2))$  where  $\exp(-(x - c)^2/(2^2))$  is computed using logarithmically many calls to Bernoulli distributions. See [DDLL2013] for details. Only integer-valued  $c$  are supported.
- 'uniform+online' – samples are drawn from a uniform distribution and accepted with probability proportional to  $\exp(-(x - c)^2/(2^2))$  where  $\exp(-(x - c)^2/(2^2))$  is computed in each invocation. Typically this is very slow. See [DDLL2013] for details. Any real-valued  $c$  is accepted.
- 'sigma2+logtable' – samples are drawn from an easily samplable distribution with  $\sigma = k \cdot 2$  with  $2 = \sqrt{1/(2 \log 2)}$  and accepted with probability proportional to  $\exp(-(x - c)^2/(2^2))$  where  $\exp(-(x - c)^2/(2^2))$  is computed using logarithmically many calls to Bernoulli distributions (but no calls to `exp`). See [DDLL2013] for details. Note that this sampler adjusts  $\sigma$  to match  $k \cdot 2$  for some integer  $k$ . Only integer-valued  $c$  are supported.

EXAMPLES:

```
sage: from sage.stats.distributions.discrete_gaussian_integer import_
↳ DiscreteGaussianDistributionIntegerSampler
sage: DiscreteGaussianDistributionIntegerSampler(3.0, algorithm=
↳ 'uniform+online')
Discrete Gaussian sampler over the Integers with sigma = 3.000000 and c = 0.
↳ 000000
sage: DiscreteGaussianDistributionIntegerSampler(3.0, algorithm='uniform+table
↳ ')
Discrete Gaussian sampler over the Integers with sigma = 3.000000 and c = 0.
↳ 000000
sage: DiscreteGaussianDistributionIntegerSampler(3.0, algorithm=
↳ 'uniform+logtable')
```

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```
Discrete Gaussian sampler over the Integers with sigma = 3.000000 and c = 0.
↪000000
```

Note that 'sigma2+logtable' adjusts :

```
sage: DiscreteGaussianDistributionIntegerSampler(3.0, algorithm=
↪'sigma2+logtable')
Discrete Gaussian sampler over the Integers with sigma = 3.397287 and c = 0.
↪000000
```

**\_\_call\_\_()**

Return a new sample.

EXAMPLES:

```
sage: from sage.stats.distributions.discrete_gaussian_integer import _
↪DiscreteGaussianDistributionIntegerSampler
sage: DiscreteGaussianDistributionIntegerSampler(3.0, algorithm=
↪'uniform+online')() # random
-3
sage: DiscreteGaussianDistributionIntegerSampler(3.0, algorithm='uniform+table
↪')() # random
3
```

**algorithm**

**c**

**sigma**

**table\_cutoff = 1000000**

**tau**

## DISCRETE GAUSSIAN SAMPLERS FOR $\mathbb{Z}[X]$

This class realizes oracles which returns polynomials in  $\mathbb{Z}[x]$  where each coefficient is sampled independently with a probability proportional to  $\exp(-(x - c)^2/(2^2))$ .

AUTHORS:

- Martin Albrecht, Robert Fitzpatrick, Daniel Cabracas, Florian Göpfert, Michael Schneider: initial version

EXAMPLES:

```
sage: from sage.stats.distributions.discrete_gaussian_polynomial import_
↳DiscreteGaussianDistributionPolynomialSampler
sage: sigma = 3.0; n = 1000
sage: l = [DiscreteGaussianDistributionPolynomialSampler(ZZ['x'], 64, sigma)()
....:      for _ in range(n)]
sage: l = [vector(f).norm().n() for f in l] #_
↳needs sage.symbolic
sage: from numpy import mean #_
↳needs numpy
sage: mean(l), sqrt(64)*sigma # abs tol 5e-1 #_
↳needs numpy sage.symbolic
(24.0, 24.0)
```

```
class sage.stats.distributions.discrete_gaussian_polynomial.DiscreteGaussianDistributionPo
```

Bases: SageObject

Discrete Gaussian sampler for polynomials.

EXAMPLES:

```
sage: from sage.stats.distributions.discrete_gaussian_polynomial import_
↳DiscreteGaussianDistributionPolynomialSampler
sage: p = DiscreteGaussianDistributionPolynomialSampler(ZZ['x'], 8, 3.0)()
sage: p.parent()
Univariate Polynomial Ring in x over Integer Ring
sage: p.degree() < 8
True
sage: gs = DiscreteGaussianDistributionPolynomialSampler(ZZ['x'], 8, 3.0)
sage: [gs() for _ in range(3)] # random
[4*x^7 + 4*x^6 - 4*x^5 + 2*x^4 + x^3 - 4*x + 7, -5*x^6 + 4*x^5 - 3*x^3 + 4*x^2 +_
↳x, 2*x^7 + 2*x^6 + 2*x^5 - x^4 - 2*x^2 + 3*x + 1]
```

```
__init__(P, n, sigma)
```

Construct a sampler for univariate polynomials of degree  $n-1$  where coefficients are drawn independently with standard deviation `sigma`.

INPUT:

- `P` – a univariate polynomial ring over the Integers
- `n` – number of coefficients to be sampled
- `sigma` – coefficients  $x$  are accepted with probability proportional to  $\exp(-x^2/(2^2))$ . If an object of type `sage.stats.distributions.discrete_gaussian_integer.DiscreteGaussianDistributionIntegerSampler` is passed, then this sampler is used to sample coefficients.

EXAMPLES:

```
sage: from sage.stats.distributions.discrete_gaussian_polynomial import ↵
↵DiscreteGaussianDistributionPolynomialSampler
sage: p = DiscreteGaussianDistributionPolynomialSampler(ZZ['x'], 8, 3.0) ()
sage: p.parent()
Univariate Polynomial Ring in x over Integer Ring
sage: p.degree() < 8
True
sage: gs = DiscreteGaussianDistributionPolynomialSampler(ZZ['x'], 8, 3.0)
sage: [gs() for _ in range(3)] # random
[4*x^7 + 4*x^6 - 4*x^5 + 2*x^4 + x^3 - 4*x + 7, -5*x^6 + 4*x^5 - 3*x^3 + 4*x^
↵2 + x, 2*x^7 + 2*x^6 + 2*x^5 - x^4 - 2*x^2 + 3*x + 1]
```

`__call__()`

Return a new sample.

EXAMPLES:

```
sage: from sage.stats.distributions.discrete_gaussian_polynomial import ↵
↵DiscreteGaussianDistributionPolynomialSampler
sage: sampler = DiscreteGaussianDistributionPolynomialSampler(ZZ['x'], 8, 12.
↵0)
sage: sampler().parent()
Univariate Polynomial Ring in x over Integer Ring
sage: sampler().degree() <= 7
True
```

## DISCRETE GAUSSIAN SAMPLERS OVER LATTICES

This file implements oracles which return samples from a lattice following a discrete Gaussian distribution. That is, if  $\sigma$  is big enough relative to the provided basis, then vectors are returned with a probability proportional to  $\exp(-|x - c|_2^2/(2\sigma^2))$ . More precisely lattice vectors in  $x \in \Lambda$  are returned with probability:

$$\frac{\exp(-|x - c|_2^2/(2\sigma^2))}{\sum_{v \in \Lambda} \exp(-|v|_2^2/(2\sigma^2))}$$

### AUTHORS:

- Martin Albrecht (2014-06-28): initial version
- Gareth Ma (2023-09-22): implement non-spherical sampling

### EXAMPLES:

```
sage: D = distributions.DiscreteGaussianDistributionLatticeSampler(ZZ^10, 3.0)
sage: D(), D(), D() # random
((3, 0, -5, 0, -1, -3, 3, 3, -7, 2), (4, 0, 1, -2, -4, -4, 4, 0, 1, -4), (-3, 0, 4, 5,
↪ 0, 1, 3, 2, 0, -1))
sage: a = D()
sage: a.parent()
Ambient free module of rank 10 over the principal ideal domain Integer Ring
```

```
class sage.stats.distributions.discrete_gaussian_lattice.DiscreteGaussianDistributionLatticeSampler
```

Bases: SageObject

GPV sampler for Discrete Gaussians over Lattices.

### EXAMPLES:

```
sage: D = distributions.DiscreteGaussianDistributionLatticeSampler(ZZ^10, 3.0); D
Discrete Gaussian sampler with Gaussian parameter sigma = 3.000000000000000, c=(0, 0, 0,
↪ 0, 0, 0, 0, 0, 0, 0) over lattice with basis

[1 0 0 0 0 0 0 0 0 0]
[0 1 0 0 0 0 0 0 0 0]
```

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```
[0 0 1 0 0 0 0 0 0 0]
[0 0 0 1 0 0 0 0 0 0]
[0 0 0 0 1 0 0 0 0 0]
[0 0 0 0 0 1 0 0 0 0]
[0 0 0 0 0 0 1 0 0 0]
[0 0 0 0 0 0 0 1 0 0]
[0 0 0 0 0 0 0 0 1 0]
[0 0 0 0 0 0 0 0 0 1]
```

We plot a histogram:

```
sage: import warnings
sage: warnings.simplefilter('ignore', UserWarning)
sage: D = distributions.DiscreteGaussianDistributionLatticeSampler(identity_
↳matrix(2), 3.0)
sage: S = [D() for _ in range(2^12)]
sage: l = [vector(v.list() + [S.count(v)]) for v in set(S)]
sage: list_plot3d(l, point_list=True, interpolation='nn') #_
↳needs sage.plot
Graphics3d Object
```

REFERENCES:

- [GPV2008]

`__init__` (*B*, *sigma*=1, *c*=0, *r*=None, *precision*=None, *sigma\_basis*=False)

Construct a discrete Gaussian sampler over the lattice  $\Lambda(B)$  with parameter *sigma* and center *c*.

INPUT:

- *B* – a (row) basis for the lattice, one of the following:
  - an integer matrix,
  - an object with a `.matrix()` method, e.g.  $\mathbb{Z}\mathbb{Z}^n$ , or
  - an object where `matrix(B)` succeeds, e.g. a list of vectors
- *sigma* – Gaussian parameter, one of the following:
  - a real number  $\sigma > 0$  (spherical),
  - a positive definite matrix  $\Sigma$  (non-spherical), or
  - any matrix-like *S*, equivalent to  $\Sigma = SS^T$ , when `sigma_basis` is set
- *c* – (default: 0) center *c*, any vector in  $\mathbb{Z}^n$  is supported, but  $c \in \Lambda(B)$  is faster
- *r* – (default: None) rounding parameter *r* as defined in [Pei2010]; ignored for spherical Gaussian parameter; if not provided, set to be the maximal possible such that  $\Sigma - rBB^T$  is positive definite
- *precision* – bit precision  $\geq 53$
- **sigma\_basis** – boolean (default: False); when set, **sigma** is treated as a (row) basis, i.e. the covariance matrix is computed by  $\Sigma = SS^T$

**Todo**  
 Rename class methods like `.f` and hide most of them (at least behind something like `.data`).

EXAMPLES:

```

sage: n = 2; sigma = 3.0
sage: D = distributions.DiscreteGaussianDistributionLatticeSampler(ZZ^n,
↳sigma)
sage: f = D.f
sage: nf = D._normalisation_factor_zz(); nf #_
↳needs sage.symbolic
56.5486677646...

sage: from collections import defaultdict
sage: counter = defaultdict(Integer); m = 0
sage: def add_samples(i):
....:     global counter, m
....:     for _ in range(i):
....:         counter[D()] += 1
....:         m += 1

sage: v = vector(ZZ, n, (-3, -3))
sage: v.set_immutable()
sage: while v not in counter: add_samples(1000)
sage: while abs(m*f(v)*1.0/nf/counter[v] - 1.0) >= 0.1: #_
↳needs sage.symbolic
....:     add_samples(1000)

sage: counter = defaultdict(Integer); m = 0
sage: v = vector(ZZ, n, (0, 0))
sage: v.set_immutable()
sage: while v not in counter:
....:     add_samples(1000)
sage: while abs(m*f(v)*1.0/nf/counter[v] - 1.0) >= 0.1: #_
↳needs sage.symbolic
....:     add_samples(1000)

```

Spherical covariance are automatically handled:

```

sage: distributions.DiscreteGaussianDistributionLatticeSampler(ZZ^3,
↳sigma=Matrix(3, 3, 2))
Discrete Gaussian sampler with Gaussian parameter  $\sigma = 2.000000000000000$ ,  $c=(0,$ 
↳ $0, 0)$  over lattice with basis

[1 0 0]
[0 1 0]
[0 0 1]

```

The sampler supports non-spherical covariance in the form of a Gram matrix:

```

sage: n = 3
sage: Sigma = Matrix(ZZ, [[5, -2, 4], [-2, 10, -5], [4, -5, 5]])
sage: c = vector(ZZ, [7, 2, 5])
sage: D = distributions.DiscreteGaussianDistributionLatticeSampler(ZZ^n,
↳Sigma, c)
sage: f = D.f
sage: nf = D._normalisation_factor_zz(); nf # This has not been properly_
↳implemented
78.6804...

```

We can compute the expected number of samples before sampling a vector:

```
sage: v = vector(ZZ, n, (11, 4, 8))
sage: v.set_immutable()
sage: 1 / (f(v) / nf)
2553.9461...

sage: counter = defaultdict(Integer); m = 0
sage: while v not in counter:
....:     add_samples(1000)
sage: sum(counter.values()) # random
3000
sage: while abs(m*f(v)*1.0/nf/counter[v] - 1.0) >= 0.1: #_
↳needs sage.symbolic
....:     add_samples(1000)
```

If the covariance provided is not positive definite, an error is thrown:

```
sage: Sigma = Matrix(ZZ, [[0, 1], [1, 0]])
sage: distributions.DiscreteGaussianDistributionLatticeSampler(ZZ^2, Sigma)
Traceback (most recent call last):
...
RuntimeError: Sigma(=[0.0000000000000000  1.0000000000000000]
[ 1.0000000000000000  0.0000000000000000]) is not positive definite
```

The sampler supports passing a basis for the covariance:

```
sage: n = 3
sage: S = Matrix(ZZ, [[2, 0, 0], [-1, 3, 0], [2, -1, 1]])
sage: D = distributions.DiscreteGaussianDistributionLatticeSampler(ZZ^n, S,
↳sigma_basis=True)
sage: D.sigma()
[ 4.0000000000000000 -2.0000000000000000  4.0000000000000000]
[-2.0000000000000000  10.0000000000000000 -5.0000000000000000]
[ 4.0000000000000000 -5.0000000000000000  6.0000000000000000]
```

The non-spherical sampler supports offline computation to speed up sampling. This will be useful when changing the center  $c$  is supported. The difference is more significant for larger matrices. For 128x128 we observe a 4x speedup (86s -> 20s):

```
sage: D.offline_samples = []
sage: T = 2**12
sage: L = [D() for _ in range(T)] # 560ms
sage: D.add_offline_samples(T) # 150ms
sage: L = [D() for _ in range(T)] # 370ms
```

We can also initialise with matrix-like objects:

```
sage: qf = matrix(3, [2, 1, 1, 1, 2, 1, 1, 1, 2])
sage: D = distributions.DiscreteGaussianDistributionLatticeSampler(qf, 3.0); D
Discrete Gaussian sampler with Gaussian parameter  $\sigma = 3.0000000000000000$ ,  $c=(0, \_$ 
↳0, 0) over lattice with basis

[2 1 1]
[1 2 1]
[1 1 2]
sage: D().parent() is D.c().parent()
True
```



**\_\_call\_\_()**

Return a new sample.

EXAMPLES:

```

sage: D = distributions.DiscreteGaussianDistributionLatticeSampler(ZZ^3, 3.0,
↳c=(1,0,0))
sage: L = [D() for _ in range(2^12)]
sage: mean_L = sum(L) / len(L)
sage: norm(mean_L.n() - D.c()) < 0.25
True

sage: D = distributions.DiscreteGaussianDistributionLatticeSampler(ZZ^3, 3.0,
↳c=(1/2,0,0))
sage: L = [D() for _ in range(2^12)] # long time
sage: mean_L = sum(L) / len(L) # long time
sage: norm(mean_L.n() - D.c()) < 0.25 # long time
True

sage: import numpy
sage: M = matrix(ZZ, [[1,2],[0,1]])
sage: D = distributions.DiscreteGaussianDistributionLatticeSampler(M, 20.0)
sage: L = [D() for _ in range(2^12)]
↳ # long time
sage: div = numpy.mean([abs(x) for x,y in L]) / numpy.mean([abs(y) for x,y,
↳in L]) # long time
sage: 0.9 < div < 1.1
↳ # long time
True
    
```

**add\_offline\_samples (cnt=1)**

Precompute samples from  $B^{-1}D_1$  to be used in `_call_non_spherical()`.

EXAMPLES:

```

sage: Sigma = Matrix([[5, -2, 4], [-2, 10, -5], [4, -5, 5]])
sage: D = distributions.DiscreteGaussianDistributionLatticeSampler(ZZ^3,
↳Sigma)
↳Sigma)
sage: assert not D.is_spherical
sage: D.add_offline_samples(2^12)
sage: L = [D() for _ in range(2^12)] # Takes less time
    
```

**c()**

Center  $c$ .

Samples from this sampler will be centered at  $c$ .

EXAMPLES:

```

sage: D = distributions.DiscreteGaussianDistributionLatticeSampler(ZZ^3, 3.0,
↳c=(1,0,0)); D
Discrete Gaussian sampler with Gaussian parameter  $\sigma = 3.000000000000000$ ,  $c=(1,$ 
↳0, 0) over lattice with basis

[1 0 0]
[0 1 0]
[0 0 1]
    
```

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```
sage: D.c()
(1, 0, 0)
```

**static compute\_precision** (*precision, sigma*)

Compute precision to use.

INPUT:

- *precision* – integer  $\geq 53$  nor None
- *sigma* – if *precision* is None then the precision of *sigma* is used

EXAMPLES:

```
sage: DGL = distributions.DiscreteGaussianDistributionLatticeSampler
sage: DGL.compute_precision(100, RR(3))
100
sage: DGL.compute_precision(100, RealField(200)(3))
100
sage: DGL.compute_precision(100, 3)
100
sage: DGL.compute_precision(None, RR(3))
53
sage: DGL.compute_precision(None, RealField(200)(3))
200
sage: DGL.compute_precision(None, 3)
53
```

**f** (*x*)

Compute the Gaussian  $\rho_{\Lambda, c, \Sigma}$ .

EXAMPLES:

```
sage: Sigma = Matrix(ZZ, [[5, -2, 4], [-2, 10, -5], [4, -5, 5]])
sage: D = distributions.DiscreteGaussianDistributionLatticeSampler(ZZ^3,
↪Sigma)
sage: D.f([1, 0, 1])
0.802518797962478
sage: D.f([1, 0, 3])
0.00562800641440405
```

**set\_c** (*c*)

Modifies center *c*.

EXAMPLES:

```
sage: D = distributions.DiscreteGaussianDistributionLatticeSampler(ZZ^3, 3.0,
↪c=(1, 0, 0))
sage: D.set_c((2, 0, 0))
sage: D
Discrete Gaussian sampler with Gaussian parameter  $\sigma = 3.000000000000000$ ,  $c=(2,
↪0, 0)$  over lattice with basis

[1 0 0]
[0 1 0]
[0 0 1]
```

**sigma()**

Gaussian parameter  $\sigma$ .

If  $\sigma$  is a real number, samples from this sampler will have expected norm  $\sqrt{n}\sigma$  where  $n$  is the dimension of the lattice.

EXAMPLES:

```
sage: D = distributions.DiscreteGaussianDistributionLatticeSampler(ZZ^3, 3.0,   
↪C=(1, 0, 0))  
sage: D.sigma()  
3.0000000000000000
```



## T-TEST USING R

```
sage.stats.r.ttest(x, y, conf_level=0.95, **kw)
```

T-Test using R.

INPUT:

- `x, y` – vectors of same length
- `conf_level` – confidence level of the interval, [0,1) in percent

OUTPUT:

Tuple: (p-value, R return object)

EXAMPLES:

```
sage: a, b = ttest([1,2,3,4,5],[1,2,3,3.5,5.121]); a # abs tol 1e-12 # optional_
↪- rpy2
0.9410263720274274
```



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