1 The Wang-Landau algorithm (density of states)

We determine thermodynamic quantities from the partition function by obtaining the density of states from a simulation.

```
import numpy as np
import matplotlib.pyplot as plt
from scipy import interpolate
```

The test system is the 2d Ising model.

```
class Ising:
        def __init__(self, n):
            self.n = n
            self.spins = np.sign(np.random.rand(n, n) - 0.5)
            self.E = self.energy()
            self.Ev = self.E
        def neighbors(self, i, j):
            return np.hstack([self.spins[:,j].take([i-1,i+1], mode='wrap'),
                              self.spins[i,:].take([j-1,j+1], mode='wrap')])
        def energy(self):
10
            return -0.5 * sum(np.sum(s * self.neighbors(i, j))
11
                             for (i, j), s in np.ndenumerate(self.spins))
12
        def propose(self):
13
            i, j = np.random.randint(self.n), np.random.randint(self.n)
            self.i, self.j = i, j
15
            dE = 2 * np.sum(self.spins[i, j] * self.neighbors(i, j))
            self.dE = dE
17
            self.Ev = self.E + dE
        def accept(self):
19
            self.spins[self.i, self.j] *= -1
            self.E = self.Ev
```

Note that this class-based approach adds some overhead. For speed, instances of Ising should be inlined into the simulation method.

A Wang-Landau algorithm, with quantities as logarithms and with montecarlo steps proportional to $f^{-1/2}$ (a "Zhou-Bhat schedule").

```
def flat(H, tol = 0.1):
    """Determines if an evenly-spaced histogram is approximately flat."""
Hμ = np.mean(H)
Hf = np.max(H)
Hθ = np.min(H)
return Hf / (1 + tol) < Hμ < Hθ / (1 - tol)</pre>
```

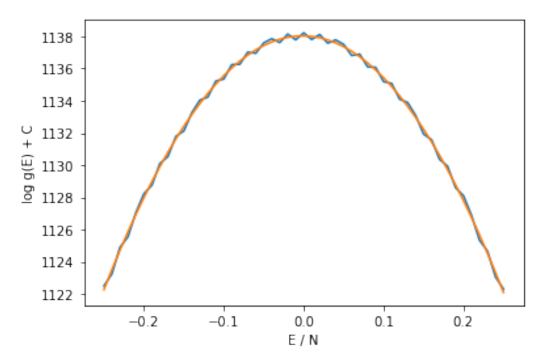
```
# Note: some parameters are hardcoded for testing
    def density_sim(system):
         randint = np.random.randint
         rand = np.random.rand
4
         exp = np.exp
         # Parameters
         M = 1_000_000 # Monte carlo step scale
         \varepsilon = 1e-6
         logftol = np.log(1 + \epsilon)
10
         logf0 = 1
11
         N = int(32**2 / 20) \# Energy bins
12
         E0 = -32**2 / 4
         Ef = 32**2 / 4
14
15
         \Delta E = (Ef - E\theta) / (N - 1)
16
         fiters = int(np.ceil(np.log2(logf0) - np.log2(logftol)))
         fiter = 0
18
         mciters = 0
19
         Es = np.linspace(E0, Ef, N)
20
         S = np.zeros(N) # Set all initial g's to 1
21
         H = np.zeros(N, dtype=int)
22
         logf = logf0
23
         # Linearly bin the energy
         i = max(0, min(N - 1, int(round((N - 1) * (system.E - E0) / (Ef - E0)))))
25
         print("ΔE = {}".format(ΔE))
26
         while logftol < logf:</pre>
27
             H[:] = 0
28
             logf /= 2
29
             iters = 0
30
             niters = int((M + 1) * exp(-logf / 2))
31
             fiter += 1
32
             while not flat(H[:-1]) and iters < niters:</pre>
33
                 system.propose()
34
                 Ev = system.Ev
35
                 j = max(0, min(N - 1, int(round((N - 1) * (Ev - E0) / (Ef - E0)))))
36
                 if E0 - \Delta E/2 \le Ev \le Ef + \Delta E/2 and (S[j] < S[i] \text{ or } rand() < exp(S[i] - S[j])):
37
                      system.accept()
38
                      i = j
                 H[i] += 1
40
                 S[i] += logf
41
                 iters += 1
42
             mciters += iters
             print("f: {} / {}\t({} / {} )\".format(fiter, fiters, iters, niters))
44
```

```
print("Done: {} total MC iterations.".format(mciters))
       return Es, S, H
47
   isingn = 32
   sys = Ising(isingn)
   Es, S, H = density_sim(sys);
   \Delta E = 10.24
   f: 1 / 20
                (46498 / 778801)
   f: 2 / 20
                (51746 / 882497)
   f: 3 / 20
                (78519 / 939414)
   f: 4 / 20
                (51944 / 969234)
   f: 5 / 20
                (62813 / 984497)
   f: 6 / 20
                (171583 / 992218)
   f: 7 / 20
                (168143 / 996102)
   f: 8 / 20
                (237575 / 998049)
   f: 9 / 20
                (303706 / 999024)
   f: 10 / 20 (280809 / 999512)
   f: 11 / 20 (577765 / 999756)
   f: 12 / 20 (999878 / 999878)
   f: 13 / 20 (927226 / 999939)
    f: 14 / 20 (999970 / 999970)
   f: 15 / 20 (999985 / 999985)
    f: 16 / 20 (999993 / 999993)
   f: 17 / 20 (867712 / 999997)
    f: 18 / 20 (999999 / 999999)
   f: 19 / 20 (1000000 / 1000000)
    f: 20 / 20 (1000000 / 1000000)
    Done: 10825864 total MC iterations.
```

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Calculating canonical ensemble averages 1.1

```
gspl = interpolate.splrep(Es, S, s=2*np.sqrt(2))
   gs = np.exp(interpolate.splev(Es, gspl) - min(S))
   plt.plot(Es / isingn**2, S)
plt.plot(Es / isingn**2, interpolate.splev(Es, gspl))
g plt.xlabel("E / N")
   plt.ylabel("log g(E) + C");
```



Translate energies to have minimum zero so that Z is representable.

```
nEs = Es - min(Es)

Z = lambda β: np.sum(gs * np.exp(-β * nEs))

Ensemble averages

βs = [np.exp(k) for k in np.linspace(-5, 0, 200)]

Eμ = lambda β: np.sum(nEs * gs * np.exp(-β * nEs)) / Z(β)

E2 = lambda β: np.sum(nEs**2 * gs * np.exp(-β * nEs)) / Z(β)

CV = lambda β: (E2(β) - Εμ(β)**2) * β**2

F = lambda β: -np.log(Z(β)) / β

Sc = lambda β: β*Εμ(β) + np.log(Z(β))

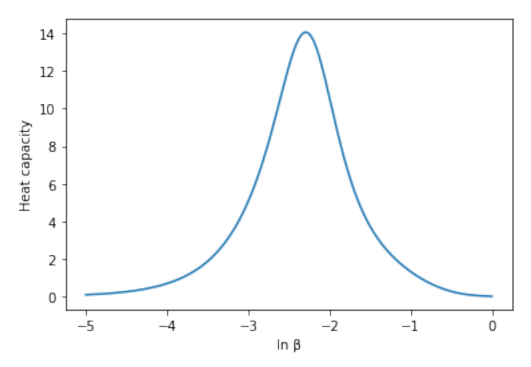
Heat capacity

plt.plot(np.log(βs), [CV(β) for β in βs])

plt.xlabel("ln β")

plt.ylabel("Heat capacity")

plt.show()
```



Entropy

```
plt.plot(np.log(\betas), [Sc(\beta) for \beta in \betas]) plt.xlabel("ln \beta")
```

- plt.ylabel(" $S(\beta) + C$ ")
- plt.show()

