

# graph\_energy

March 12, 2024

## 1 Graph Based Spin Lattice

### 1.0.1 Find lowest ground state of simple Ising Hamiltonian

For a graph,  $G = (E, V)$ , defined by a set of edges,  $E$ , and vertices,  $V$ , we want to represent an Ising model, where the edge weights,  $w_{ij}$  are given by the spin interactions, i.e.,  $w_{ij} = J_{ij}$ .

Given a configuration of spins (e.g.,  $\uparrow\downarrow\downarrow\uparrow\downarrow$ ) we can define the energy using what is referred to as an Ising Hamiltonian:

$$\hat{H} = \sum_{(i,j) \in E} J_{ij} s_i s_j$$

where,  $s_i = 1$  if the  $i^{th}$  spin is up and  $s_i = -1$  if it is down, and the summation runs over all edges in the graph.

```
[ ]: # Load relevant libraries. If you have errors you probably need to install them
      ↪ into your conda env

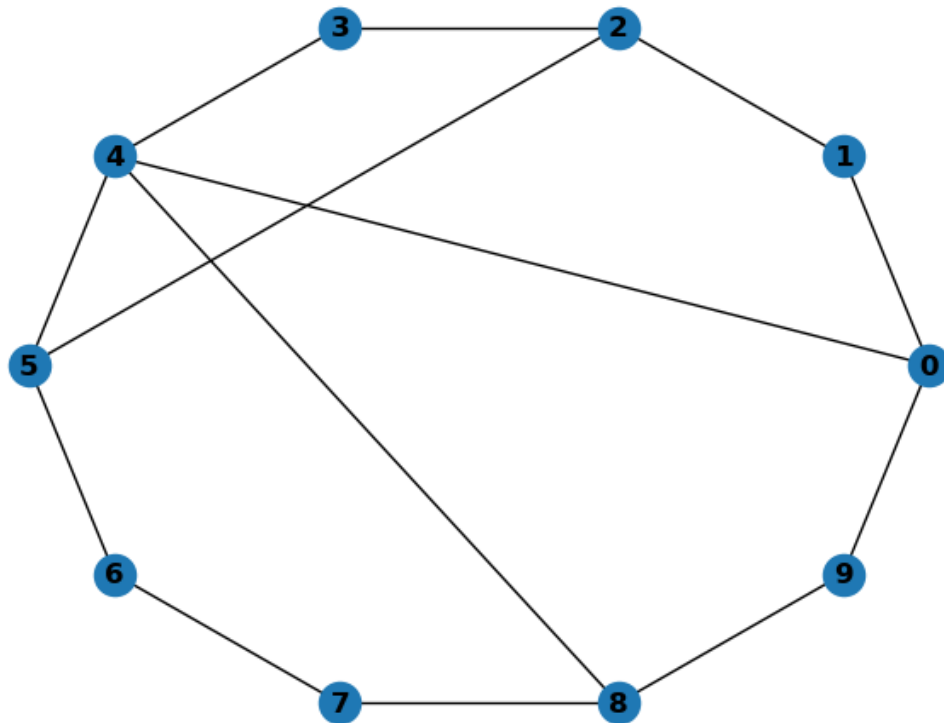
import numpy as np
import networkx as nx
import matplotlib.pyplot as plt
import random
# import scipy
random.seed(2)
```

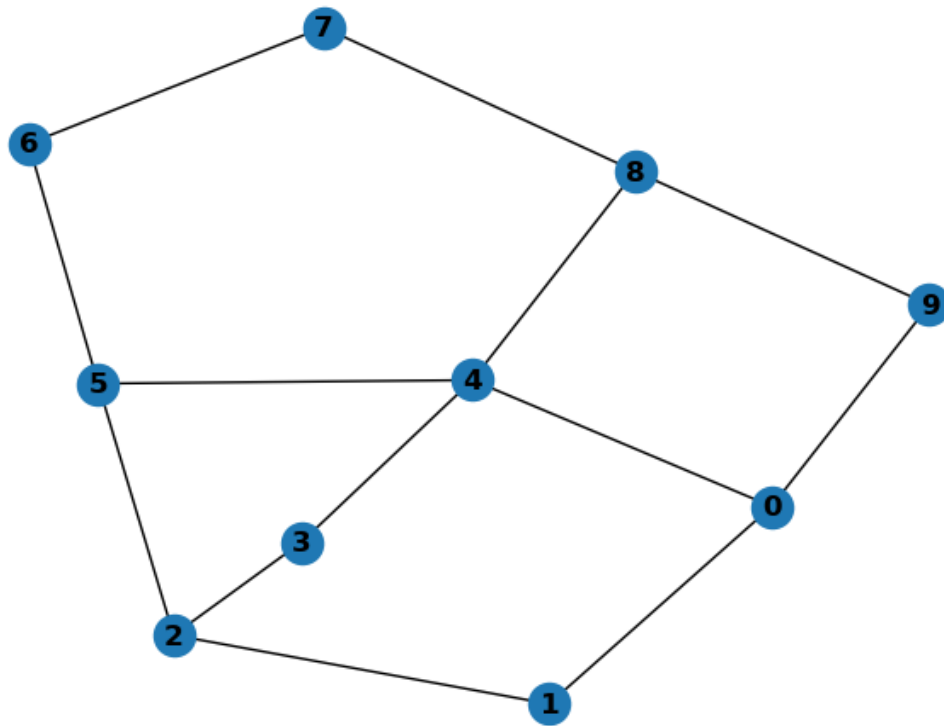
### 1.0.2 Create a graph that defines the Ising interactions

```
[ ]: G = nx.Graph()
G.add_nodes_from([i for i in range(10)])
G.add_edges_from([(i, (i+1)% G.number_of_nodes() ) for i in range(10)])
G.add_edge(2,5)
G.add_edge(4,8)
G.add_edge(4,0)
for e in G.edges:
    G.edges[e]['weight'] = 1.0

# Now Draw the graph. First we will draw it with the nodes arranged on the
  ↪ circle, then we will draw the same graph
# with the position of the nodes optimized for easier visualization
```

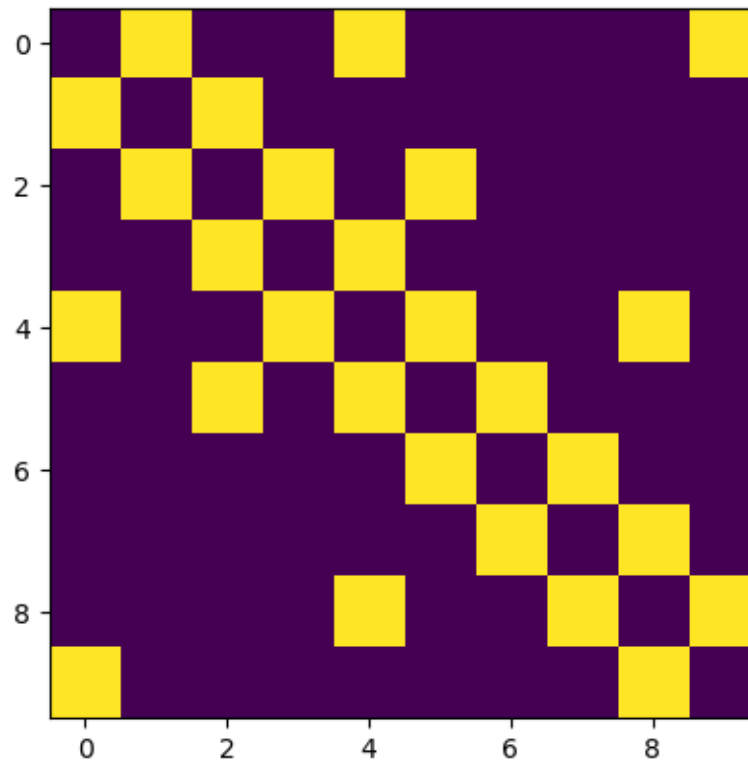
```
plt.figure(1)
nx.draw(G, with_labels=True, font_weight='bold', pos=nx.circular_layout(G))
plt.figure(2)
nx.draw(G, with_labels=True, font_weight='bold')
plt.show()
```





```
[ ]: A = nx.adjacency_matrix(G).todense()
      display(A)
      plt.imshow(A);
```

```
array([[0., 1., 0., 0., 1., 0., 0., 0., 0., 1.],
       [1., 0., 1., 0., 0., 0., 0., 0., 0., 0.],
       [0., 1., 0., 1., 0., 1., 0., 0., 0., 0.],
       [0., 0., 1., 0., 1., 0., 0., 0., 0., 0.],
       [1., 0., 0., 1., 0., 1., 0., 0., 1., 0.],
       [0., 0., 1., 0., 1., 0., 1., 0., 0., 0.],
       [0., 0., 0., 0., 0., 1., 0., 1., 0., 0.],
       [0., 0., 0., 0., 0., 0., 1., 0., 1., 0.],
       [0., 0., 0., 0., 1., 0., 0., 1., 0., 1.],
       [1., 0., 0., 0., 0., 0., 0., 0., 0., 1.]])
```



### 1.0.3 Add your BitString class below

```
[ ]: import numpy as np
import math

class BitString:
    """
    Simple class to implement a config of bits
    """
    def __init__(self, N):
        self.N = N
        self.config = np.zeros(N, dtype=int)

    def __repr__(self):
        selfString = ''
        for bit in self.config:
            selfString += str(bit)
        return selfString

    def __eq__(self, other):
        return (self.config == other.config).all()
```

```

def __len__(self):
    return self.N

def on(self):
    num_on = 0
    for bit in self.config:
        if bit == 1:
            num_on += 1
    return num_on

def off(self):
    num_off = 0
    for bit in self.config:
        if bit == 0:
            num_off += 1
    return num_off

def flip_site(self,i):
    self.config[i] ^= 1

def int(self):
    return int(str(self), 2)

def set_config(self, s:list[int]):
    self.config = s

def set_int_config(self, dec:int):
    self.config = np.zeros(self.N, dtype=int)

    i = 1
    while dec != 0:
        self.config[-i] = dec % 2
        dec = dec // 2
        i += 1

```

```

[ ]: # translating 0 and 1 to -1 and 1
def z1_to_n1(bit1, bit2):
    if bit1 == bit2:
        return 1
    else:
        return -1

def energy(bs: BitString, G: nx.Graph):
    """Compute energy of configuration, `bs`

    .. math::

```

```


$$E = \langle \hat{H} \rangle$$


Parameters
-----
bs    : Bitstring
        input configuration
G      : Graph
        input graph defining the Hamiltonian
Returns
-----
energy : float
        Energy of the input configuration
"""
# energy = sum of J_ij * S_i * S_j = J_ij * translation_thiny(S_i, S_j)
# J represents whether or not the qubits are adjacent

array_J = nx.adjacency_matrix(G).todense()
nrg = 0
# for i in range(bs.N):
#     for j in range(i, bs.N):
#         nrg += array_J[i,j] * z1_to_n1(bs.config[i], bs.config[j])
for (i,j) in G.edges:
    nrg += z1_to_n1(bs.config[i], bs.config[j])

return nrg

```

#### 1.0.4 Naive minimization

Loop over all configurations and compute energies to find the lowest

```

[ ]: x = [] # Store list of indices
     y = [] # Store list of energies
     xmin = None # configuration of minimum energy configuration
     emin = 0 # minimum of energy
     my_bs = BitString(10)

     for i in range(2 ** 10):
         my_bs.set_int_config(i)
         y.append(energy(my_bs, G))
         x.append(i)
     emin = min(y)
     xmin = y.index(emin)
     print(emin)

     # Print out all the energies
     plt.plot(x,y);

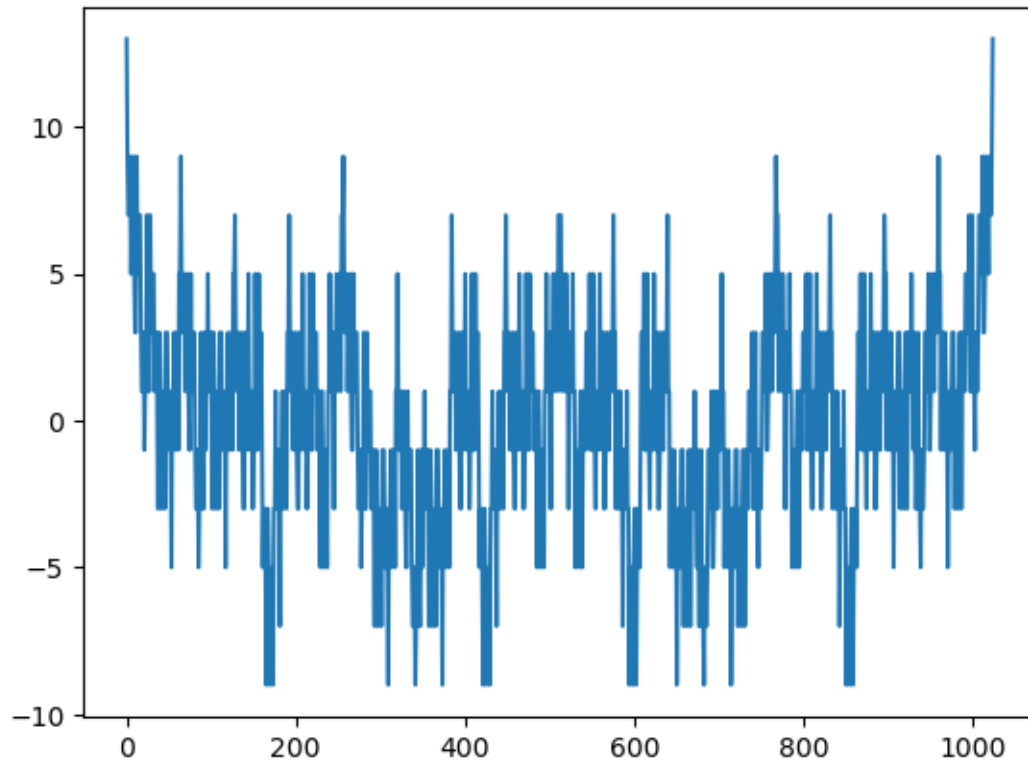
```

```
# Print out the lowest energy configuration
my_bs.set_int_config(xmin)
print("Lowest energy %12.8f: %s" %(emin, my_bs), ' (' , my_bs.int(), ') ',
      ↪sep='')

assert(abs(energy(my_bs, G) - -9) < 1e-12)
```

-9

Lowest energy -9.00000000: 0010100101 (165)



### 1.0.5 Visualize ground state

Print out the graph again, this time coloring each node according to it's state (0 or 1)

```
[ ]: print("Configuration: %s" %my_bs)
      nx.draw(G, node_color=my_bs.config)
```

Configuration: 0010100101

