hw3p2

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0.1 Homework 3, Problem 2 on homogeneous Poisson processes

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0.2 Background

The goal of this notebook is to model a neuron as a homogeneous Poisson processes and evaluate its properties. We will consider a simulated neuron that has a cosine tuning curve described in equation (1.15) in TN (TN refers to Theoretical Neuroscience by Dayan and Abbott.)

$$\lambda(s) = r_0 + (r_{\text{max}} - r_0)\cos(s - s_{\text{max}})$$

where λ is the firing rate (in spikes per second), s is the reaching angle of the arm, s_{max} is the reaching angle associated with the maximum response r_{max} , and r_0 is an offset that shifts the tuning curve up from the zero axis. This will be referred as tuning equation in the following questions.

Let
$$r_0 = 35$$
, $r_{\text{max}} = 60$, and $s_{\text{max}} = \pi/2$.

Note: If you are not as familiar with Python, be aware that if 1 is of type int, then 1 / a where a is any int greater than 1 will return 0, rather than a real number between 0 and 1. This is because Python will return an int if both inputs are ints. If instead you write 1.0 / a, you will get out the desired output, since 1.0 is of type float.

```
[1]:

"""

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"""

import numpy as np
import matplotlib.pyplot as plt
import nsp as nsp # these are helper functions that we provide.
import scipy.special
from scipy.stats import poisson

# Load matplotlib images inline
%matplotlib inline

# Reloading any code written in external .py files.
%load_ext autoreload
%autoreload 2
```

0.2.1 (a) (6 points) Spike trains

For each of the following reaching condition ($s = k \cdot \pi/4$, where k = 0, 1, ..., 7), generate 100 spike trains according to a homogeneous Poisson process. Each spike train should have a duration of 1 second. You can think of each of each spike train sequence as a trial. Therefore, we generate 100 trials of the neuron spiking according to a homogeneous Poisson Process for 8 reach directions.

Your code for this section should populate a 2D numpy array, $spike_times$ which has dimensions $num_cons \times num_trials$ (i.e., it is 8×100). Each element of this 2D numpy array is a numpy array containing the spike times for the neuron on a given condition and trial. Note that this array may have a different length for each trial.

e.g., spike_times.shape should return (8, 100) and spike_times[0,0] should return the spike times on the first trial for a reach to the target at 0 degrees. In one instantiation, our code returns that spike times[0,0] is:

```
array([
         0.
                        5.94436383,
                                      10.85691999,
                                                     26.07821145,
              67.417219 ,
50.02836141,
                             74.2948356 , 119.19210112,
                                                           139.41789878,
176.59511596, 244.40788916,
                             267.3643421 ,
                                                   288.42590046, 324.3770265,
                                    460.76250631, 471.23773964,
340.26911602, 407.75730065,
                                                                  489.41659607,
514.60180131,
                     548.71822693,
                                    565.6036432 ,
                                                   586.20557118,
                                                                  601.11595447,
710.37485206,
              751.60837895, 879.93536952, 931.26983289,
                                                                  944.1130483 ,
949.38455374,
              963.22509374,
                             964.67365483,
                                                   966.3865719 , 974.3657882 ,
987.25729081])
```

Of course, this varies based off of random seed. Also note that time at 0 is not a spike.

```
[2]: ## 2a
     bin_width = 20
                                                 # (ms)
                                                 # (radians)
     s = np.arange(8)*np.pi/4
     num_cons = np.size(s)
                                                  # num_cons = 8 in this case, number_
     →of directions
     r 0 = 35 \# (spikes/s)
     r_max = 60 \# (spikes/s)
     s_max = np.pi/2 # (radians)
     T = 1000 \#trial \ length \ (ms)
     num trials = 100 # number of spike trains to generate
     tuning = r_0 + (r_max-r_0)*np.cos(s-s_max) # tuning curve
     spike_times = np.empty((num_cons, num_trials), dtype=list)
     for con in range(num_cons):
         for rep in range(num_trials):
             # YOUR CODE HERE:
                 Generate homogeneous Poisson process spike trains.
                 You should populate the np.ndarray 'spike_times' according
                 to the above description.
```

```
Lambda=r_0+(r_max-r_0)*np.cos(s[con]-s_max)
spike_times[con, rep] = nsp.GeneratePoissonSpikeTrain(T,Lambda)
#=======#
# END YOUR CODE
#=======#
```

```
[3]:

s_labels = ['0', '$\pi$/4', '$\pi$/2', '3$\pi$/4', '$\pi$', '5$\pi$/4', '3$\pi$/

-2', '7$\pi$/4']

num_plot_rows = 5

num_plot_cols = 3

subplot_indx = [9, 6, 2, 4, 7, 10, 14, 12]

num_rasters_to_plot = 5 # per condition

# Generate and plot homogeneous Poisson process spike trains

plt.figure(figsize=(10,8))

for con in range(num_cons):

# Plot spike rasters

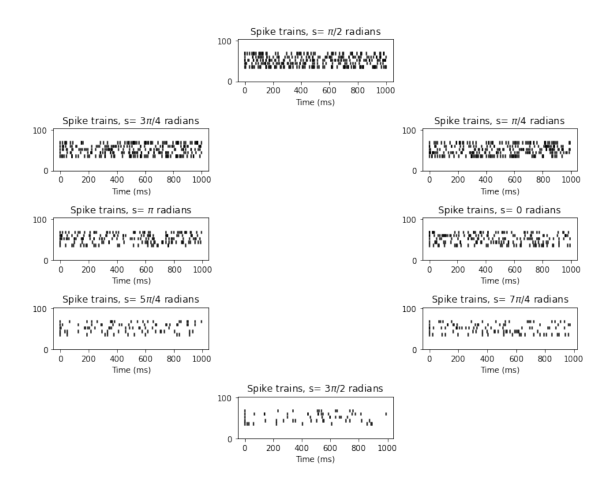
plt.subplot(num_plot_rows, num_plot_cols, subplot_indx[con])

nsp.PlotSpikeRaster(spike_times[con, 0:num_rasters_to_plot])

plt.title('Spike trains, s= '+s_labels[con]+' radians')

plt.tight_layout()

plt.savefig("test.png")
```



0.2.2 Plotting the spike rasters.

The following code plot 5 spike trains for each reaching angle in the same format as shown in Figure 1.6(A) in TN. You should take a look at this code to understand what it's doing. You may also want to look at the PlotSpikeRaster function from nsp.

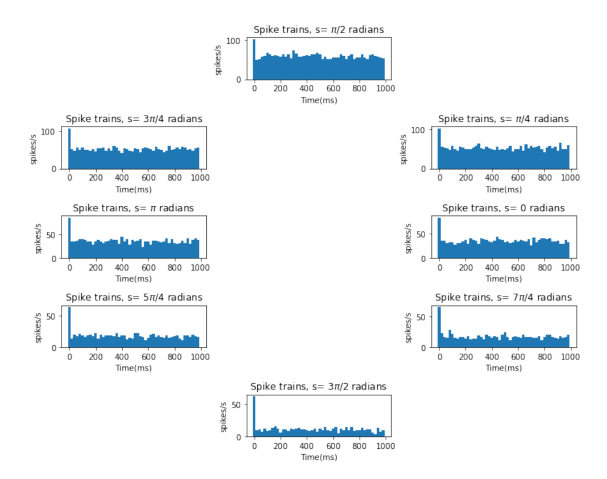
The plots should make intuitive sense given the tuning parameters.

0.2.3 (b) (5 points) Plot spike histograms

For each reaching angle, find the spike histogram by taking spike counts in non-overlapping 20 ms bins, then averaging across the 100 trials. Plot the 8 resulting spike histograms around a circle, as in part (a). This time, as we'll allow you to represent the data as you like, you will have to also plot each histogram on your own. The spike histograms should have firing rate (in spikes / second) as the vertical axis and time (in msec, not time bin index) as the horizontal axis.

Suggestion: you can use plt.bar to plot the histogram, it is important to set the width for this function, e.g. width = 12.

```
[4]: ## 2b
   plt.figure(figsize=(10,8))
   for con in range(num_cons):
      plt.subplot(num_plot_rows,num_plot_cols,subplot_indx[con])
       # YOUR CODE HERE:
          Generate and plot spike histogram for this condition
       #========#
      bins=np.arange(0,50)*20
      counts=np.zeros(50)
      for rep in range(num_trials):
         rbins,rcounts=np.unique(spike_times[con][rep][0:]//
    →20,return_counts=True)
         for i,b in enumerate(rbins):
             counts[int(b)]+=rcounts[i]
      plt.bar(bins,(counts/num_trials)/(20*10**-3),width=20)
      plt.ylabel('spikes/s')
      plt.xlabel('Time(ms)')
       # END YOUR CODE
      #========#
      plt.title('Spike trains, s= '+s_labels[con]+' radians')
      plt.tight_layout()
```



0.2.4 (c) (4 points) Tuning curve

For each trial, count the number of spikes across the entire trial. Plots these points on the axes like shown in Figure 1.6(B) in TN, where the x-axis is reach angle and the y-axis is firing rate. There should be 800 points in the plot (but some points may be on top of each other due to the discrete nature of spike counts). For each reaching angle, find the mean firing rate across the 100 trials, and plot the mean firing rate using a red point on the same plot. Now, plot the tuning curve of this neuron in green on the same plot.

```
[5]: ## 2c

spike_counts = np.zeros((num_cons, num_trials)) # each element in spike_counts_

is the total spike count for this reach direction and trial

#========#

# YOUR CODE HERE:

# Plot the single trial spike counts and the tuning curve

# on top of each other.

#========#

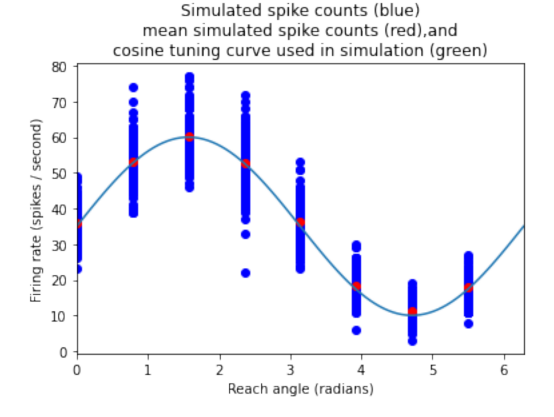
MeanSquaredDeviation=np.empty(num_cons)

for con in range(num_cons):
```

```
for trial in range(num_trials):
        spike_counts[con,trial]=spike_times[con,trial].shape[0]
        plt.plot(s[con],spike_counts[con,trial],"o",color="blue")
    plt.plot(s[con],np.mean(spike_counts[con]),"o",color="red")
    MeanSquaredDeviation[con]=(tuning[con]-np.mean(spike_counts[con]))**2
#plot tuning curve
tuning_curve=lambda s: r_0 + (r_max-r_0)*np.cos(s-s_max)
s_plot=np.linspace(0,2*np.pi,100)
plt.plot(s_plot,tuning_curve(s_plot))
print("RMS deviation is", np.sqrt(np.mean(MeanSquaredDeviation)))
plt.xlabel('Reach angle (radians)')
plt.ylabel('Firing rate (spikes / second)')
plt.title('Simulated spike counts (blue)\n'+
           'mean simulated spike counts (red),and\n'+
           'cosine tuning curve used in simulation (green)')
plt.xlim(0, 2*np.pi)
```

RMS deviation is 0.859082934619929

[5]: (0.0, 6.283185307179586)



Question: Do the mean firing rates lie near the tuning curve?

Your answer: Yes, it does. The RMS deviation between the two is 1.3hz

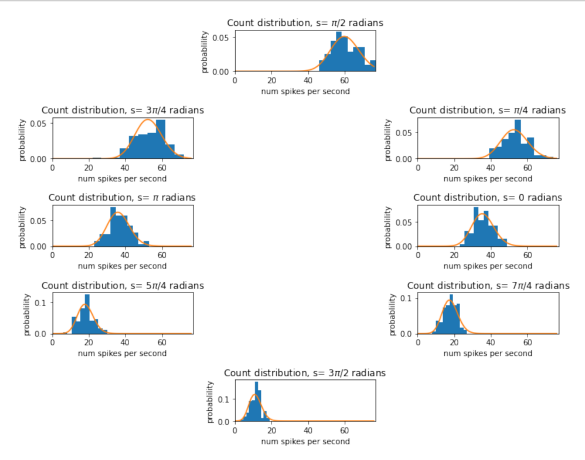
0.2.5 (d) (6 points) Count distribution

For each reaching angle, plot the *normalized* distribution (i.e., normalized so that the area under the distribution equals one) of spike counts (using the same counts from part (c)). Plot the 8 distributions around a circle, as in part (a). Fit a Poisson distribution to each empirical distribution and plot it on top of the corresponding empirical distribution.

Please plot the empirical distribution as well as the fit

```
[6]: ##2d
   plt.figure(figsize=(10,8))
   max count = np.max(spike counts)
   spike_count_bin_centers = np.arange(0,max_count,1)
   for con in range(num_cons):
      plt.subplot(num_plot_rows,num_plot_cols,subplot_indx[con])
      #----#
      # YOUR CODE HERE:
         Calculate the empirical mean for the Poisson spike
         counts, and then generate a curve reflecting the probability
         mass function of the Poisson distribution as a function
         of spike counts.
      mean=np.mean(spike_counts[con])
      #=======#
      # END YOUR CODE
      #-----#
      # YOUR CODE HERE:
         Plot the empirical count distribution, and on top of it
         plot your fit Poisson distribution.
      #----#
      plt.hist(spike_counts[con],density=True)
      plt.plot(np.arange(0,max_count),poisson.pmf(np.arange(0,max_count), mean))
      plt.ylabel("probablility")
      plt.xlabel("num spikes per second")
      # END YOUR CODE
      #----#
```

```
plt.xlim([0, max_count])
plt.title('Count distribution, s= '+ s_labels[con]+' radians')
plt.tight_layout()
```



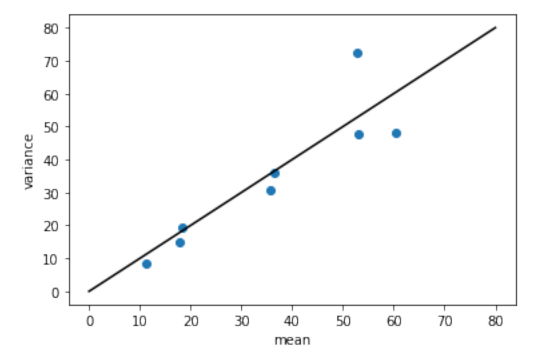
Question: Are the empirical distributions well-fit by Poisson distributions?

Your answer: Yes

0.2.6 (e)(4 points) Fano factor

For each reaching angle, find the mean and variance of the spike counts across the 100 trials (using the same spike counts from part (c)). Plot the obtained mean and variance on the axes shown in Figure 1.14(A) in TN. There should be 8 points in this plot – one per reaching angle.

```
[7]: ## 2e
#========#
# YOUR CODE HERE:
# Calculate and plot the mean and variance for each of
# the 8 reaching conditions. Mean should be on the
# x-axis and variance on the y-axis.
```



Question: Do these points lie near the 45 deg diagonal, as would be expected of a Poisson distribution?

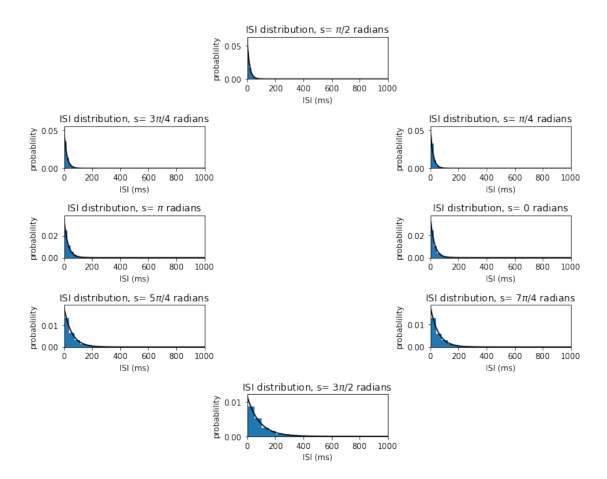
Your answer: yes

0.2.7 (f) (5 points) Interspike interval (ISI) distribution

For each reaching angle, plot the normalized distribution of ISIs. Plot the 8 distributions around a circle, as in part (a). Fit an exponential distribution to each empirical distribution and plot it on top of the corresponding empirical distribution.

Please plot the empirical distribution as well as the fit

```
[8]: ## 2f
   plt.figure(figsize=(10,8))
   ISIs=[]
   for con in range(num_cons) :
      plt.subplot(num_plot_rows,num_plot_cols,subplot_indx[con])
      #----#
      # YOUR CODE HERE:
       Calculate the interspike interval (ISI) distribution
        by finding the empirical mean of the ISI's, which
       is the inverse of the rate of the distribution.
      ISI=[]
      for trial in range(num_trials):
        ISI+=list(spike_times[con,trial][1:]-spike_times[con,trial][:-1])
      l=1/np.mean(ISI)
      plt.xlim(0,T)
      # END YOUR CODE
      #========#
      #----#
      # YOUR CODE HERE:
      # Plot Interspike interval (ISI) distribution
      #========#
      plt.hist(ISI,density=True)
      plt.plot(np.arange(0,T),l*np.exp(-l*np.arange(0,T)),"black")
      ISIs.append(ISI)
      plt.ylabel("probablility")
      plt.xlabel("ISI (ms)")
      # END YOUR CODE
      plt.title('ISI distribution, s= '+ s_labels[con]+' radians')
      plt.tight_layout()
```



Question: Are the empirical distributions well-fit by exponential distributions?

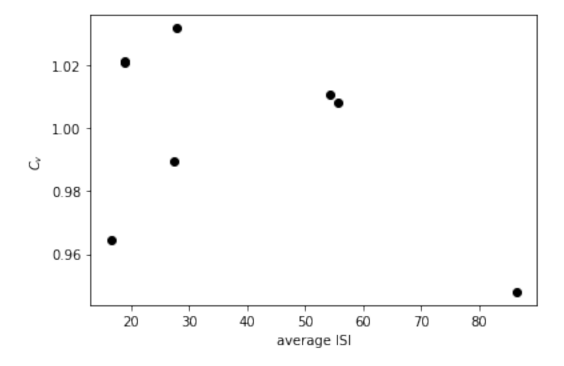
Your answer: yes

0.2.8 (g) (5 points) Coefficient of variation (C_V)

For each reaching angle, find the average ISI and C_V of the ISIs. Plot the resulting values on the axes shown in Figure 1.16 in TN. There should be 8 points in this plot.

```
[9]: #2g
#========#
# YOUR CODE HERE:
# Calculate and plot coeffcient of variation
#=========#
for ISI in ISIs:
    plt.plot(np.mean(ISI),np.std(ISI)/np.mean(ISI),"o",color="black")
    plt.xlabel("average ISI")
    plt.ylabel("$C_v$")
#=========#
```





Question: Do the C_V values lie near unity, as would be expected of a Poisson process?

Your answer: yes