Homework - Categorization and Model Comparison Part A (70/110 points)

by *Todd Gureckis* and *Brenden Lake* Computational Cognitive Modeling

NYU class webpage: https://brendenlake.github.io/CCM-site/ (https://brendenlake.github.io/ (<a h

This homework is due before midnight on April 25 2022.

```
In [1]: import string as str
import os
import numpy as np
import seaborn as sns
import pandas as pd
import math
from random import random, randint, shuffle, uniform
from scipy.optimize import fmin, brute
```

Background and Theory

In this homework we explore the cognitive mechanisms that support unsupervised pattern categorization in humans. In addition, we use this as an example of testing and comparing between models.

A simple (classic) unsupervised categorization experiment

In Posner and Keele (1968) report a now classic categorization experiment with humans. In the task participants viewed visual stimuli that are clouds of points (known as dot patterns) similar to a scatter plot of data on a graph. An examples of the stimuli is shown here:



Experiment Design

The experiment was divided into a training and test phase. During the training phase, for each subject a single random dot pattern was generated and considered to be the underlying "prototype" structure. A prototype is like a common template or reference pattern. The key is that participants never get to see the "prototype" pattern directly during training. Instead they see what are known as "distortions" of the prototype. A distortion of a pattern is made by adding random spatial noise to each point in a pattern to kind of "wiggle" the points away from their original position.

For example, here is a random prototype (top) and a bunch of random distortions of the prototype made by adding or subtracting small random values from the $\langle x, y \rangle$ value of each point in the pattern.



Posner and Keele created distortions that added more or less random noise. For example, "high" distortions add a lot of randomness to the underlying template pattern whereas "low" distortions add only a little bit of noise.

Training Phase

In the training phase of the experiment subjects view 10 training examples one at a time which are "high" distortions of a randomly generated prototype. The instructions are that subjects should look at these patterns, and that they come from a single category similar to if you viewed a series of pictures of dogs they would all come from the category dog. Subjects were try to figure out the pattern that related the different images to one another. Try it for yourself by looking at each of the "distortions" patterns above one by one and trying to detected the common structure.

Test Phase

During the test phase, participants view a series of dot patterns one at a time and have to judge: **Does the given pattern come from the same general category or family you studied earlier or is it a new pattern that is different?** This is an unsupervised categorization task because the subject has to abstract what the common structure is from the given patterns and then use that information to make classification decisions about new patterns.

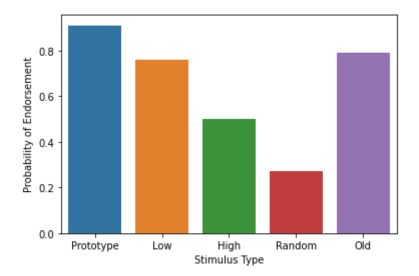
Unknown to participants the set of test items varied in a specific way with respect the training patterns. In particular, there were five particular types of patterns presented during test.

- The first type were "old" patterns which were identical to those presented during the training phase.
- The second type were "random" patterns which were from a complete new randomly generated prototype (thus had nothing to do with the items presented during training).
- The third type were new "high distortions" of the underlying prototype that was used to create the study set. These are thus similar to the "old" items but do not match exactly.
- The fourth type were "low distortions" of the underlying prototype that was used to crete the study set. These are more similar to the prototype pattern than the "high" distortions are.
- Finally the actual prototype used to generate the items during training was presented. This pattern is interesting because the prototype pattern was never seen exactly during training. However, people saw many high distortions of this item during training and given the instructions to detect what the common structure of the training patterns is, they may have learned something about this latent or hidden pattern.

Typical Results:

t.'>

This graph show example results that are typical for an experiment like this:



The height of the bars indicates the probability of endorsing a pattern as a member of the category during the test phase (high values mean that at test a subject is more likely to agree "yes this pattern fits with the one I learned"). Notice that the "old" items (the exact patterns studied during learning are endorsed at a relatively high rate. In contrast the "random" patterns (those coming from a completely different underlying pattern) are endorsed at a very low rate. The low and high distortions are endorsed at intermediate rates (with the low distoritions preferred). Interestingly, the prototype pattern is endorsed most strongly even though it was never presented during the study period! It is like during learning people figured out the underlying pattern that generated the stimuli!

Categorization: Memory for examples or abstractions?

The dot pattern experiments are interesting and have been replicated in various ways perhaps hundreds of time. How do people make these judgments? What information to people store during the study phase that would predict their performance in the test phase? What rule do they use for combining information from memory in order to make these kind of judgements? We are going to build up a model of categorization in a few simple steps.

Step 1: How are dot patterns represented in the mind?

Our first question concerns how people represent the dot patterns and the similarity between different dot patterns. A variety of work has suggested that the psychological similarity between pairs of dot patterns follows roughly a logarithmic transform of the average euclidean distance between the pairs of points (plus one). This was established by having people view pairs of dot patterns and rate how similar they seem. For example here is a plot from a paper by Smith and Minda showing a strong contgruence between dissimiliarty ratings between pairs of stimuli and log(distance)



In light of this lets define the similarity between two dot patterns i and j as s_{ij} and let it equal the following equation:

$$s_{ij} = log(1 + \frac{\sqrt{\sum_{d}[(i_{dx} - j_{dx})^{2} + (i_{dy} - j_{dy})^{2}]}}{9})$$

where i_{d_x} is the x position of the d-th dot in pattern i and j_{d_x} is the x position for pattern j (likewise for i_{d_y}). There are 9 dots total, so we divide by 9 to get the average. Because it can sometimes be ambiguous which dot aligns which which one in a pattern we choose the dots which are closest in the two patterns to compute this score.

Step 2: What is stored in memory?

The next consideration is what people actually store in memory during the training phase of the experiment. There are of course many alternatives. People could store an "average" of the points seen so far, or they could store each individual pattern that they have seen, or they could store nothing and try to figure it out at test, or they could store some verbal description of what the shapes "look like", or the shape of the outer edge of the dot-cloud (the "convex null"), etc...

There are, however, two leading theories which have attracted considerable debate in the cognitive science literature: the prototype and exemplar theory.

Exemplar models

Exemplar models are a general class of psychological models related to nearest neighbor algorithms. The most important feature of these models is the idea that people have what appears to be a nearly infitite memory for the past and as a result you can store all past experiences or examples in memory. This seems crazy as we are forgetting things all the time but actually psychology is unclear about if we actually forget things or if we simply lose the ability to retreive a memory (i.e., more like losing the pointer to the memory rather than decay).

As mentioned in lecture, nearset neighbor classifiers use a similarity function (similar to the ones described above) to retrieve from memory the nearest labeled example and to predict the category membership based on the label for this item. This nearest neighbor algorithm can be relaxed slightly to consider k-nearest neighbors. According to this algorithm you find the k neighbor examples (with k > 1) to the current pattern and response based on what the majority of these examples say.

Now we can go a bit further and say that you compute the similarity to all past examples but weight their vote according to their similarity. So instead of picking the label of the closest or k-closest examples we compare the current pattern using a global match to all examples in the memory and weight their predictions based on similarity. Pretty neat!

Ok, but how does this help us in the case of **unsupervised** categorization such as in the dot pattern case? Here what we will assume is that we compute this similarity of the to-be-categorized item (the test item) to all the examples stored in memory and compare it to some criterion value. If the sum of the similarity to all the examples falls below this criterion then we assume the pattern is new and doesn't match what we learned. If it is above the criterion we judge the item is a good example of the category.

In the example model we will consider the probability of endorsing an item is going to be determined by the following equation:

$$P(A|i) = \frac{\sum_{j} e^{-c \cdot s_{ij}}}{\sum_{i} e^{-c \cdot s_{ij}} + k}$$

</br>

where P(A|i) is the probability of endorsing pattern i as a member of the category seen during study. s_{ij} is the similarity between pattern i and pattern j which is an example stored in memory during the study phase. k is the criterion against which the summed similarity is being compared. If k is zero then you endorse the item as a member of the category all the time irrespecitive of the similiary and if k gets large you become more and more less likely to endorse the item (i.e., more likely to say no).



The sum is with respect to an exponential sum which has some deeper relation to research on categorization that we do not have time to discuss. However, it is basically the idea that very close matches ($s_{ij} = 0$) are especially strong and things that are less similar count less. You can think of it as the the particular weighted nearest neighbor algorithm we think the mind uses. c is a free parameter that controls that weighting function and is often fitted to data.

Prototype models

The prototype model is different than the exemplar model because it assumes that instead of storing each of the training patterns in memory exactly, instead people store a single summary representation. For example, people might store a mentally computed "average" pattern. When you think about how you would perform the task you might think that you kind of compare the training patterns to one another and then compute some summary.



In the case of the dot pattern stimuli one way to do this is to store a special trace in memory called the prototype which is the average of all the patterns seen so far (averaging the < x, y > position of each point to find an average dot location.

According to the prototype model the probability of endorsing a test item pattern as a member of the category that was studied during training is:

$$P(A|i) = \frac{e^{-c \cdot s_{ip}}}{e^{-c \cdot s_{ip}} + k}$$

</br>

Note that nearly everything about this equation is the same except there is no longer a sum! Instead we simply compute the similarity between the test pattern and this special "prototype" pattern (p) which has been averaged from the training examples.



Parameters

k and c are "free parameters" in both the exemplar and prototype model which are assumed to modulate or alter the core psychological processes. These parameters might vary between subjects and as a function of condition. Thus, in order to assess the ability of the model to account for the data we often "fit" these parameters to our data.

Model Comparison

With these idea in mind, in this homework you are going to compare the exemplar and prototype model to account for some data from an actual dot pattern categorization task collected with human subjects. The goal is that by doing the homework you would develop some useful code that would let you more or less plug in a model that you might come across in your research, fit it to data, and verify that the fits are good, etc...

Reading in some data

The data/ folder that comes with this homework contains data from 14 human subjects who participated in a dot pattern classification task. The data describing each subject is in a text file (.dat) indexed by subject number (e.g., 1.dat, 2.dat, etc...).

The organization of these files is as follows:

The first 44 lines of the file contain a description of the stimulus that the subject saw on a given trial including the x, y coordinate of each dot. The first columns of these 44 lines is the number of the pattern (1-44). The second column is the type of pattern using the following codes:

- 1 = prototype
- 2 = 10 "high distortions" of the prototype that were used as study patterns during learning
- 3 = 10 new "high distortions" of the prototype presented during test
- 4 = 4 "low distortions" of the prototype that were presented during test
- 5 = 20 random items that come from different prototypes that were presented at test

The next 18 values of each row are the coordinates of the dots (with the x, y coordinates in sequence). So [x1, y1, x2, y2, x3, y3, ...]

The following 40 lines of the file show the sequence of items presented during the study phase. This is not all that important for our purposes, but basically the last column is which pattern was displayed (indexed from the patterns just described. Each of 10 "high distortions" were presented four times each during study in a random order.

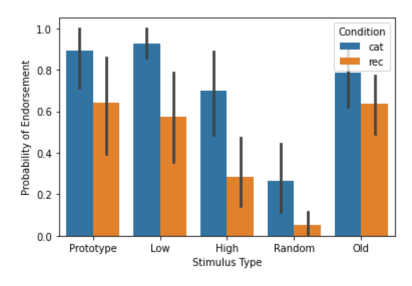
Finally the remaining lines of the file report the results of the test phase. The first column is the subject number, the second columns is the condition number, the next is the trial number in the experiment, the other columns worth mentioning are the last column (the pattern number from the beginning of the file), the second to last column (the type of stimulus it is according to the codes described above), and the reaction time in milliseconds.

In addition, participants in this experiment were assigned to one of two conditions: a recognition condition and a categorization condition. These conditions differed only in the instructions given to participants at the start of the test phase. In the recognition condition participants were told they would view a series of patterns and they should respond "yes" only if the patterns was **exactly** one they say in the previous study phase. In the categorization condition, participants were asked to respond "yes" only if the pattern belonged to the same general category or pattern that they observed in the training phase.

The following graph computes the probability of endorsement in the data set as a function of stimulus type and condition (Cat or Rec instructions).

```
# getcurve
        def getcurve(filename):
           prototypes = []
           low = []
           old = []
           high = []
           random = []
           mydata = readfile(filename)
           cond = mydata[-1][1]
           for line in mydata:
                if line[4] == 2 and len(line) == 9:
                   if line[7] == 1:
                       prototypes.append(line[5])
                   if line[7] == 2:
                       old.append(line[5])
                   if line[7] == 3:
                       high.append(line[5])
                   if line[7] == 4:
                       low.append(line[5])
                   if line[7] == 5:
                       random.append(line[5])
           #print([len(prototypes), len(low), len(high), len(random), len(ol
        d)])
           # print(prototypes)
           # print(low)
           # print(high)
           # print(random)
           # print(old)
           return [np.average(prototypes), np.average(low), np.average(high), n
        p.average(random), np.average(old), filename, cond]
        def readfile(filename):
           results = []
           fp = open(filename, 'r')
           for line in fp.readlines():
               myline = list(map(int, line.split(' ')[:-1]))
               results.append(myline[:])
           fp.close()
           return results
        def get all filenames(directoryname):
           files = filter(lambda x: x[-4:] == '.dat' and x[0] !=
                          '.', os.listdir(os.path.join(".", directoryname)))
           fn = map(lambda x: os.path.join(".", directoryname, x), files)
           # process each file and drop last 5 trials
           return list(fn)
        def create_df(subjnum, cond, pattern):
           nobs = len(pattern)
           df = pd.DataFrame({"Subject": [subjnum]*nobs, "Condition": [cond]*no
```

```
In [4]: sns.barplot(x="Stimulus Type", y="Probability of Endorsement", hue="Corition", data=get_human_results())
```



Problem 1 (20 points)

Using your own words explain the data pattern you see in the above figure. What is different between the conditions and stimulus type? Why do you suspect these patterns exist? Your answer will need to consider the nature of the experiments, what is manipulated, and even your intuitive psychological theory about what might be going on. Your response should take 3-4 sentences and appear in a cell below. Is any feature of this data surprising to you?

Answer: The two conditioning methods differ in how subjects decides to endorse the pattern as a certain stimulus type (therefore cause different P(endorsement) for each stimulus), but the actual relative probability of endorsing stimulus types follow a similar pattern for both categorization and recognition; nonetheless, Categorization has higher P(endorsement) than recognition's P(endorsement) in cases of observation. Also, there is a clear difference in P(endorsement) between the five stimulus types in that prototype, low, and old stimulus types have higher P(endorsement) than high and random stimulus types. I suspect these patterns exist because the nature of the experiment is based on human mind's natural instinct/memory to decide whether to endorse the pattern or not; therfore, it is easier to recognize general pattern (categorization) than finding the exact match (recognition) since human memory isn't likely to be infinite. As a result, protopyte, low, and old stimulus types are able to receive high endorsement, but finding the difference between them is very difficult for human congnition (shown in the graphs). Also, what is heavily manipulated in the experiment are high and random stimulus types compared to the original train patterns shown to the subjects, which makes sense the lower endorsement in those two stimulus types. One thing that surprised me was that over half of the subjects endorsed high stimulus type, which was surprising since I expected it would be much more difficult to endorse pattern when heavy noise is introduced.

Predictions for the exemplar model

The following cells set up the exemplar model using the equations described above.

```
# unitdist:
       # computes the euclidean distance between
       # two dots
       def unitdist(x, y):
          x1 = np.array(x)
          y1 = np.array(y)
          return math.sqrt(sum(pow(x-y, 2.0)))
       ##################################
       # computeresponse
       # computes the "activation" of each
       # trace in memory
       def computeresponse(target, memory, c, k):
          res = []
          for mem in memory:
             res.append(
                 math.log(1.0+np.average(list(map(lambda x, y: unitdist(x, y
       ), target, mem()))))
          resp = [math.exp(-c*x) for x in res]
          pofr = sum(resp)/(sum(resp)+k)
          return pofr
```

```
# exemplar model
        # stores all 10 study items in memory
        # and computes the probability of endorsement
        # for each item type
        def exemplarmodel(filename, c, k):
           data = readfile(filename)
           cond = data[-1][1]
           memory = []
           for line in data:
                if len(line) == 20 and line[1] == 2:
                   memory.append(np.resize(line[2:], (9, 2)))
           # print(memory)
           # prototype items
           proto = []
           for line in data:
               if len(line) == 20 and line[1] == 1:
                   item = np.resize(line[2:], (9, 2))
                   pofr = computeresponse(item, memory, c, k)
                   proto.append(pofr)
           # print(np.average(proto))
           # old items
           old = []
           for line in data:
               if len(line) == 20 and line[1] == 2:
                   item = np.resize(line[2:], (9, 2))
                   pofr = computeresponse(item, memory, c, k)
                   old.append(pofr)
            # print "p of r", old
           # print(np.average(old))
           # new high items
           newhigh = []
           for line in data:
               if len(line) == 20 and line[1] == 3:
                   item = np.resize(line[2:], (9, 2))
                   pofr = computeresponse(item, memory, c, k)
                   newhigh.append(pofr)
           # print(np.average(newhigh))
           # new low items
           newlow = []
           for line in data:
               if len(line) == 20 and line[1] == 4:
                   item = np.resize(line[2:], (9, 2))
                   pofr = computeresponse(item, memory, c, k)
                   newlow.append(pofr)
           # print(np.average(newlow))
           # random items
           random = []
```

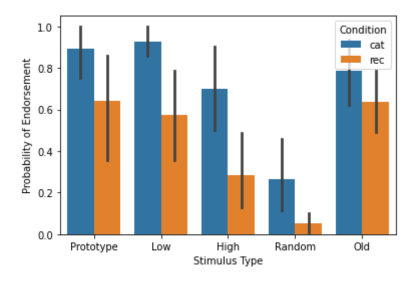
```
for line in data:
    if len(line) == 20 and line[1] == 5:
        item = np.resize(line[2:], (9, 2))
        pofr = computeresponse(item, memory, c, k)
        random.append(pofr)
# print(np.average(random))

return [np.average(proto), np.average(newlow), np.average(newhigh),
np.average(random), np.average(old), filename, cond]
```

```
In [7]:
    def get_exemplar_results(c_cat, k_cat, c_rec, k_rec):
        allres = {fn: readfile(fn) for fn in get_all_filenames('data')}
        cat = []
        rec = []
        for filename in allres.keys():
            if allres[filename][-1][1] == 0:
                res = exemplarmodel(filename, c_cat, k_cat)
                cat.append(create_df(filename, 'cat', res[:-2]))
        else:
            res = exemplarmodel(filename, c_rec, k_rec)
                rec.append(create_df(filename, 'rec', res[:-2]))
        cat, rec = pd.concat(cat), pd.concat(rec)
        return pd.concat([cat, rec])
```

First let's replot the human results:

```
In [8]: sns.barplot(x="Stimulus Type", y="Probability of Endorsement", hue="Corition", data=get_human_results())
```

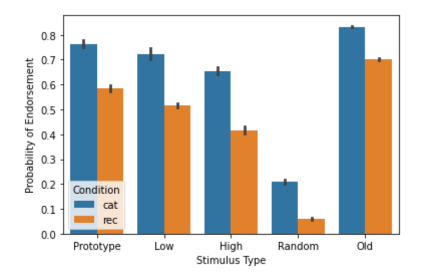


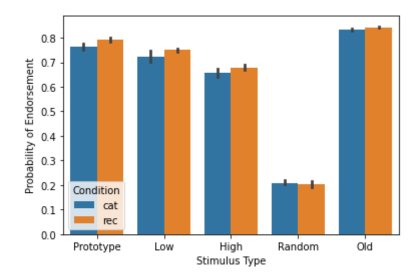
Problem 2 (10 points)

By hand adjust the setting of the model parameters to roughly fit the human data pattern shown above. How close can you get? What parameters did you find (report them) and you assessment of how well they fit. Was it a good fit or are there systematic problems with the fit? In addition, what are the parameter values and do they make sense in light of the equations described above? When the parameters are the same for recognition and categorization instructions why do the bars look a little different?

Answer: I can get the model to roughly a similar fit to the human data pattern in terms of stimulus types, but it has lower P(endorsement) for prototype, low, and high stimulus types and Old stimulus type has much higher probability than rest of the stimulus which is different from human pattern. The parameters are {c_cat, k_cat, c_rec, k_rec = 1.7, 0.3, 1.95, 0.6}, and overall the shape of the pattern follows that of human data, but I believe the order of endorsement is different and high distortion stimulus seems too highly endorsed; therefore, I also believe there exists systematic problems with the fit. Considering the equations from above, it makes sense k value is higher for recognition condition because it is much harder to find exact match of pattern than just a general matching pattern. As mentioned above, k is the criterion that summed similarity is being compared to, so higher the k value is the less likely it is to endorse the pattern. The reason why bars are different when c_cat == c_rec and k_cat == k_rec is because of introduced random noise in the data.

The following cell lets you plot the model predictions for the exemplar model fitted to the stimuli that participants in this experiment actually viewed. There is a k and a c parameter for both categorization and recogniton.





Predictions for the prototype model

The following cells set up the prototype model using the equations described above.

```
# prototype model
         # stores an averate of the study items in memory
         # and computes the probability of endorsement
         # for each item type
         def prototypemodel(filename, c, k):
             data = readfile(filename)
             cond = data[-1][1]
             # average all the old items in memory
             memory = []
             for line in data:
                 if len(line) == 20 and line[1] == 2:
                    memory.append(line[2:])
            memory = [
                np.resize(list(map(np.average, np.transpose(np.array(memory)))),
         (9, 2))]
             # prototype items
             proto = []
             for line in data:
                 if len(line) == 20 and line[1] == 1:
                    item = np.resize(line[2:], (9, 2))
                    pofr = computeresponse(item, memory, c, k)
                    proto.append(pofr)
             # print(np.average(proto))
             # old items
             old = []
             for line in data:
                 if len(line) == 20 and line[1] == 2:
                    item = np.resize(line[2:], (9, 2))
                    pofr = computeresponse(item, memory, c, k)
                    old.append(pofr)
             # print "p of r", old
             # print(np.average(old))
             # new high items
             newhigh = []
             for line in data:
                 if len(line) == 20 and line[1] == 3:
                    item = np.resize(line[2:], (9, 2))
                    pofr = computeresponse(item, memory, c, k)
                    newhigh.append(pofr)
             # print(np.average(newhigh))
             # new low items
             newlow = []
             for line in data:
                if len(line) == 20 and line[1] == 4:
                    item = np.resize(line[2:], (9, 2))
                    pofr = computeresponse(item, memory, c, k)
                    newlow.append(pofr)
             # print(np.average(newlow))
             # random items
```

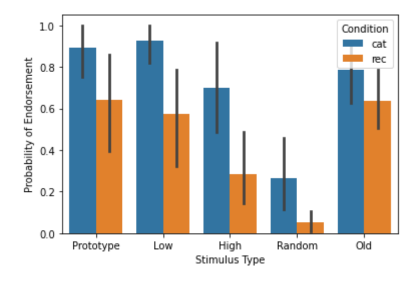
```
random = []
for line in data:
    if len(line) == 20 and line[1] == 5:
        item = np.resize(line[2:], (9, 2))
        pofr = computeresponse(item, memory, c, k)
        random.append(pofr)
# print(np.average(random))

return [np.average(proto), np.average(newlow), np.average(newhigh),
np.average(random), np.average(old), filename, cond]
```

```
In [17]: def get_prototype_results(c_cat, k_cat, c_rec, k_rec):
    allres = {fn: readfile(fn) for fn in get_all_filenames('data')}
    cat = []
    rec = []
    for filename in allres.keys():
        if allres[filename][-1][1] == 0:
            res = prototypemodel(filename, c_cat, k_cat)
            cat.append(create_df(filename, 'cat', res[:-2]))
    else:
        res = prototypemodel(filename, c_rec, k_rec)
        rec.append(create_df(filename, 'rec', res[:-2]))
    cat, rec = pd.concat(cat), pd.concat(rec)
    return pd.concat([cat, rec])
```

Again lets replot the human results for easy reference.

```
In [18]: sns.barplot(x="Stimulus Type", y="Probability of Endorsement", hue="Corition", data=get_human_results())
```

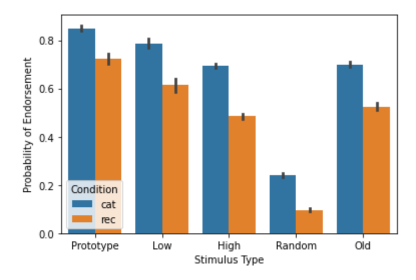


Problem 3 (10 points)

By hand adjust the setting of the model parameters in the next cell to roughly fit the human data pattern shown above. How close can you get? What parameters did you find (report them) and you assessment of how well they fit. Was it a good fit or are there systematic problems with the fit? In addition, what are the parameter values and do they make sense in light of the equations described above?

Answer: My parameters are {c_cat, k_cat, c_rec, k_rec = 1.4, 0.06, 1.5, 0.13}. It is hard to say the model fit well and I think there are systematic problems with the fit. The endorsement probabilities made sense for prototype,low, and random stimulus, but since high and old stimulus endorsement probabilities were very similar, I didn't think it fit the human pattern precisely. Also, human data had highest endorsement for low stimulus type, different than the model below. Considering the equation above, k parameter makes sense it is higher for recognition condition because it is much harder to find exact match of pattern than finding a general matching pattern; in this note, higher k value means less endorsement.

```
In [65]: c_cat, k_cat, c_rec, k_rec = 1.4, 0.06, 1.5, 0.13
    res=get_prototype_results(c_cat, k_cat, c_rec, k_rec)
    sns.barplot(x="Stimulus Type", y="Probability of Endorsement", hue="Condition", data=res)
```



Fitting the models using RMSE

Next we would like to come up with a more quantitative way to assess the quality of the model fits. The first technique we will use is the "goodness of fit" measures that were discussed in lecture. One of the most common measures of goodness of fit is the Root Mean Squared Error (RMSE). This measure compares the value of each data point x to each prediction y using the following equation:

$$RMSE = \sqrt{\frac{\sum_{i}(x_i - y_i)^2}{N}}$$

</br>

Often the RMSE is computed between the AVERAGE prediction of the model and the AVERAGE estimates of the behavior to all the subjects in an experiment. Using the code we developed above we can find the average endorsement curves for humans and both models like this:

```
In [66]: human_results=get_human_results()
    exemplar_predictions = get_exemplar_results(c_cat, k_cat, c_rec, k_rec)
    prototype_predictions = get_prototype_results(c_cat, k_cat, c_rec, k_rec
)
```

Out[67]:

	Condition	Stimulus Type	Probability of Endorsement
0	cat	High	0.700000
1	cat	Low	0.928571
2	cat	Old	0.785714
3	cat	Prototype	0.892857
4	cat	Random	0.264286
5	rec	High	0.285714
6	rec	Low	0.571429
7	rec	Old	0.635714
8	rec	Prototype	0.642857
9	rec	Random	0.050000

Out[68]:

	Condition	Stimulus Type	Probability of Endorsement
0	cat	High	0.939436
1	cat	Low	0.952741
2	cat	Old	0.967979
3	cat	Prototype	0.960152
4	cat	Random	0.751648
5	rec	High	0.869711
6	rec	Low	0.901237
7	rec	Old	0.933689
8	rec	Prototype	0.918836
9	rec	Random	0.505024

Out[69]:

	Condition	Stimulus Type	Probability of Endorsement
0	cat	High	0.694033
1	cat	Low	0.786957
2	cat	Old	0.699643
3	cat	Prototype	0.849233
4	cat	Random	0.240838
5	rec	High	0.484085
6	rec	Low	0.615297
7	rec	Old	0.524251
8	rec	Prototype	0.721810
9	rec	Random	0.096520

Problem 4 (20 points)

First, write a function below called `rmse` that computes the RMSE between two `numpy` vectors.

```
In [70]: def rmse(human, model):
    return np.sqrt(np.mean((human-model)**2))
```

Write your code above. This code will then by used in the provided functions below to evaluate the fit of the prototype and exemplar models. The parameters to the model is provided as a list with [c_cat, k_cat, c_rec, k_rec] the implied order.

```
In [71]: def fit exemplar model rmse(params, human results):
             [c_cat, k_cat, c_rec, k_rec] = params
             predictions = get_exemplar_results(c_cat, k_cat, c_rec, k_rec)
             avgpredict=predictions.groupby(['Condition', 'Stimulus Type'],as_ind
         ex=False).mean()
             model_results = avgpredict['Probability of Endorsement'].values
             return rmse(human_results, model_results)
         def fit_prototype_model_rmse(params, human_results):
             [c_cat, k_cat, c_rec, k_rec] = params
             predictions = get_prototype_results(c_cat, k_cat, c_rec, k_rec)
             avgpredict=predictions.groupby(['Condition', 'Stimulus Type'],as_ind
         ex=False).mean()
             model_results = avgpredict['Probability of Endorsement'].values
             return rmse(human results, model results)
In [72]: human results=get human results()
         avghuman=human results.groupby(['Condition', 'Stimulus Type'],as index=F
         alse).mean()
         human results = avghuman['Probability of Endorsement'].values
         print(fit exemplar model rmse([0.1, 2.0, 0.1, 2.0], human results))
         print(fit prototype model rmse([0.1, 2.0, 0.1, 2.0], human results))
```

0.35330178202197526 0.3792712020204823

Next adjust the parameters by hand for both the exemplar and prototype models to find values that appear to minimize the RMSE. Copy the code above for plotting the predictions of the models given your best fit parameters. Which model do you think fits better according to this fit statistic?

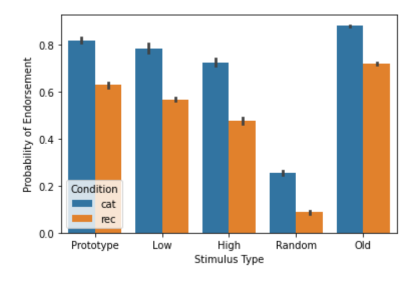
```
In [80]: human_results=get_human_results()
    avghuman=human_results.groupby(['Condition', 'Stimulus Type'],as_index=F
    alse).mean()
    human_results = avghuman['Probability of Endorsement'].values

    print(fit_exemplar_model_rmse([1.75, 0.2, 1.8, 0.6], human_results))
    print(fit_prototype_model_rmse([1.6, 0.05, 1.5, 0.2], human_results))
```

0.08953586057610792 0.10145102738580922 Answer: With hand adjustment, I believe examplar model is a better fit to this fit statistic because RMSE=0.08953586057610792 for exemplar model which is lower than RMSE=0.10145102738580922 of prototype model. The overall shape of expamplar doesn't fit the human model but it does fit better than prototype when it comes to endorsement of Random and Old Stimulus types.

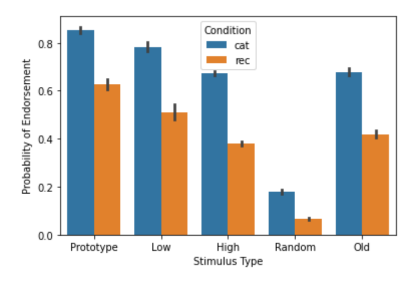
Examplar by Hand

```
In [81]: c_cat, k_cat, c_rec, k_rec = 1.75, 0.2, 1.8, 0.6
    exemplar_results = get_exemplar_results(c_cat, k_cat, c_rec, k_rec)
    sns.barplot(x="Stimulus Type", y="Probability of Endorsement", hue="Condition", data=exemplar_results)
```



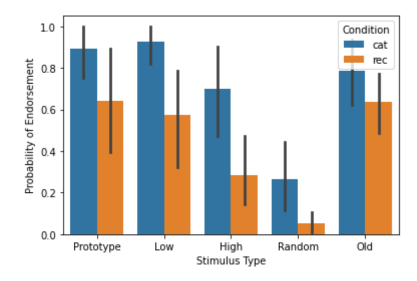
Prototype by Hand

```
In [82]: c_cat, k_cat, c_rec, k_rec = 1.6, 0.05, 1.5, 0.2
    prototype_results = get_prototype_results(c_cat, k_cat, c_rec, k_rec)
    sns.barplot(x="Stimulus Type", y="Probability of Endorsement", hue="Condition", data=prototype_results)
```



Human

```
In [83]: human_results = get_human_results()
    sns.barplot(x="Stimulus Type", y="Probability of Endorsement", hue="Cond
    ition", data=human_results)
```



Problem 5 (10 points)

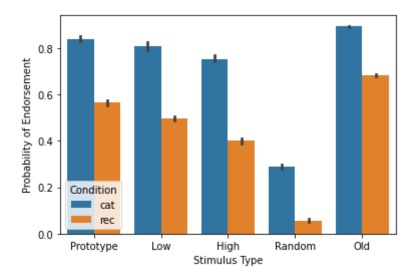
Read about the scipy 'fmin' function. Use fmin to algorithmically search for the best parameters for each model using the RMSE evaluation function described above.

```
In [73]: initial x0 = [0.1, 2.0, 0.1, 2.0]
         best examplar param = fmin(fit exemplar model rmse, x0=initial x0, args=
         (human results,), full output=True)
         best prototype param = fmin(fit prototype model rmse, x0=initial x0, arg
         s=(human_results,), full_output=True)
         print('best examplar param',best examplar param[0])
         print('best_prototype param',best_prototype param[0])
         Optimization terminated successfully.
                  Current function value: 0.076819
                  Iterations: 260
                  Function evaluations: 435
         Optimization terminated successfully.
                  Current function value: 0.077796
                  Iterations: 442
                  Function evaluations: 743
         best examplar param [1.73818548 0.17362409 1.92580663 0.66919728]
         best prototype param [1.62604888 0.02939519 1.62430878 0.14084599]
In [85]: human results=get human results()
         avghuman=human results.groupby(['Condition', 'Stimulus Type'], as index=F
         alse).mean()
         human results = avghuman['Probability of Endorsement'].values
         print(fit exemplar model rmse([1.73818548, 0.17362409, 1.92580663, 0.669
         19728], human results))
         print(fit prototype model rmse([1.62604888, 0.02939519, 1.62430878, 0.14
         084599], human results))
         0.07681866353753997
         0.07779553421390219
```

Here, exemplar model (RMSE=0.07681866353753997) also does slightly better than prototype model (RMSE=0.07779553421390219) in terms of RMSE.

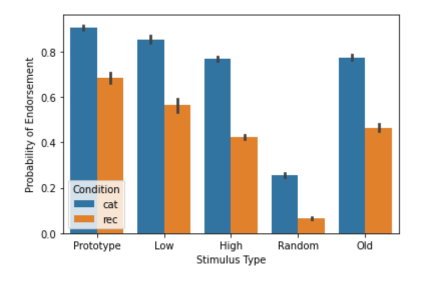
Examplar

```
In [86]: c_cat, k_cat, c_rec, k_rec = 1.73818548, 0.17362409, 1.92580663, 0.669:
    728
    exemplar_results = get_exemplar_results(c_cat, k_cat, c_rec, k_rec)
    sns.barplot(x="Stimulus Type", y="Probability of Endorsement", hue="Condition", data=exemplar_results)
```



Prototype

```
In [87]: c_cat, k_cat, c_rec, k_rec = 1.62604888, 0.02939519, 1.62430878, 0.1408
599
prototype_results = get_prototype_results(c_cat, k_cat, c_rec, k_rec)
sns.barplot(x="Stimulus Type", y="Probability of Endorsement", hue="Condition", data=prototype_results)
```



In []:	

Homework - Categorization and Model Comparison Part B (40/110 points)

by *Todd Gureckis* and *Brenden Lake* Computational Cognitive Modeling

NYU class webpage: https://brendenlake.github.io/CCM-site/ (https://brendenlake.github.io/ (<a h

This homework is due before midnight on April 25 2022.

```
In [1]: from IPython.display import display
    import string as str
    import os
    import numpy as np
    import seaborn as sns
    import pandas as pd
    import math
    from random import random, randint, shuffle, uniform
    from scipy.optimize import fmin, brute
    from scipy.special import comb # gets the combinations function
    from exemplarproto import * # this grabs much of the code from Part A o
    f the homework
```

Fitting the models using maximum likelihood

As mentioned in the lecture, RMSE is not always an ideal mechanism for fitting models. One reason is that it is insensitive to the number of observations that define each data point. For example, remember in our experiment that participants saw the prototype item four times at test. In contrast, there were 20 different "new" patterns. This means there are five times as many trials contributing to the "new" bar in this graph as for the prototype patterns. Since RMSE measures the raw deviation of the average model predictions from those of the model it doesn't take into account these issues. Thus, we would like to also evaluate these two models using maximum likelihood.

The key to this is going to be the provided function below which computes the likelihood of a particular set of data under a binomial probability model.:

```
# computeLogLikelihood
       \# N = number of observations
       # S = number of "successes" (i.e., endorsements)
       # p = predicted probability of successes by the model
       def computeLogLikelihood(N, S, p):
           p = p if p > 0.0 else 0.0+1e-10
           p = p if p < 1.0 else 1.0-1e-10
           try:
              result = math.log(comb(N, S)) + (S*math.log(p) + (N-S)*math.log(
       1.0-p))
           except:
              print(N, S, p) # this shouldn't happen but just in case
              result = 0
           return result
       def pandas ll(row):
           return computeLogLikelihood(row['Total'], row['N Yes'], row['Probabi
       lity of Endorsement'])
```

A short explanation may be in order: the models predictions take the form of probabilities of endorsement for each of the prototype, low, high, random, and old items. If you find out that people endorse the prototype on 2 out of 2 trials how likely is this outcome given that the model (for a particular set of parameters) predicts an endorsement of p=0.8? Three numbers are required to do this for each data point N, the number of trials/presentations within the stimulus class, S the number of successes observed (S<=N), and p the predicted probability. Then you can turn the crank on the above computelogLikelihood() function which returns the probability that you would get S successes in N trials if the true probability was p (make sure you understand what is happening in computelogLikelihood). You can sum these log likelihoods for each stimulus class (prototype, low, high, random, old) to compute a total log(likelihood) of the data for any given model with any set of parameters. For this homework will we focus on fitting the group data rather than to individuals.

To get the data formatted into an appropriate shape for fitting likelihoods we provide a function get_human_results_ll() which returns a Pandas data frame containing the number of times a pattern of a particular type was endorsed and the number of times it was presented for each subject.

```
In [3]: human_res=get_human_results_ll()
human_res
```

Out[3]:

	Subject	Condition	Stimulus Type	N_Yes	Total
0	./data/8.dat	cat	Prototype	4	4
1	./data/8.dat	cat	Low	4	4
2	./data/8.dat	cat	High	10	10
3	./data/8.dat	cat	Random	15	20
4	./data/8.dat	cat	Old	20	20
0	./data/7.dat	rec	Prototype	3	4
1	./data/7.dat	rec	Low	4	4
2	./data/7.dat	rec	High	8	10
3	./data/7.dat	rec	Random	0	20
4	./data/7.dat	rec	Old	19	20

70 rows × 5 columns

This reorganizes the data per condition.

```
In [4]: human_data=human_res.groupby(['Condition','Stimulus Type']).sum()
human_data
```

Out[4]:

		N_Yes	Total
Condition	Stimulus Type		
cat	High	49	70
	Low	26	28
	Old	110	140
	Prototype	25	28
	Random	37	140
rec	High	20	70
	Low	16	28
	Old	89	140
	Prototype	18	28
	Random	7	140

Finally these function allow us to compute the negative log likelihood of the data given the model.

```
In [5]: def fit exemplar model_nll(params, human_results):
            [c cat, k_cat, c_rec, k_rec] = params
            k_cat = k_cat if k_cat > 0.0 else 0.0
            k_rec = k_rec if k_rec > 0.0 else 0.0
            predictions = get exemplar results(c cat, k cat, c rec, k rec)
            model = predictions.groupby(
                ['Condition', 'Stimulus Type'], as_index=False).mean()
            fitted data = pd.merge(model, human results)
            return -1.0*fitted_data.apply(pandas_ll, axis=1).sum()
        def fit prototype model nll(params, human results):
            [c_cat, k_cat, c_rec, k_rec] = params
            k_cat = k_cat if k_cat > 0.0 else 0.0
            k_rec = k_rec if k_rec > 0.0 else 0.0
            predictions = get prototype results(c_cat, k_cat, c_rec, k_rec)
            model = predictions.groupby(
                ['Condition', 'Stimulus Type'], as_index=False).mean()
            fitted_data = pd.merge(model, human_results)
            return -1.0*fitted data.apply(pandas 11, axis=1).sum()
```

Problem 6 (20 points)

The cell blocks below allow you to fit the exemplare model and the prototype model to the dataset we considered in Part A of the homework. Make sure you understand and follow the code provided above and in the provided library (exemplarproto.py). Next, try altering the parameters to minimize the negative log likelihood score. When you think you have found the best fit parameters for both the exemplar and prototype models report your final parameter values along with the plot of the resulting model predictions. In a markdown cell describe which model you believe fits better. Is this conclusion the same or different from what you considered in Part 4 of the homework? If the fit looks different, why?

Answer:

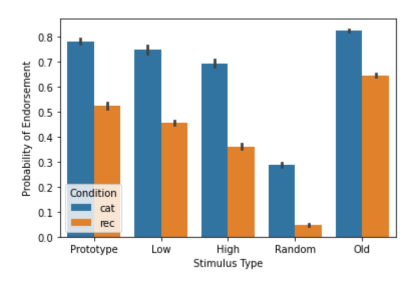
best_examplar_param [1.48769296 0.35961247 1.93041277 0.78398669] -> NL score 28.268922073692416 best_prototype_param [1.58816955 0.03164535 1.87904483 0.08402034] -> NL score 32.246756097276645

I believe exemplar model fits better than prototype model because its negative log (NL) score is lower than that of prototype model as shown above. Looking at the endorsement probability graph, exemplar model does much better in old and high stimulus, but it is worse in prototype and low stimulus types. The general trend was that exemplars tend to underestimate and prototypes tend to overestimate compared to human model. This conclusion is same as what I considered in Part 4 because exmplar did better with well performing old and high stimuli. The fit looks similar as well with both exemplar and prototype models.

```
In [8]: human = human_res.groupby(['Condition', 'Stimulus Type'], as_index=Fals
        ).sum()
        initial_x0 = [1.73818548, 0.17362409, 1.92580663, 0.66919728]
        best examplar param = fmin(fit exemplar model nll, x0=initial x0, args=(
        human,), full_output=True)
        best prototype param = fmin(fit prototype model_nll, x0=initial_x0, args
        =(human,), full output=True)
        print('best_examplar_param',best_examplar_param[0])
        print('best_prototype param',best_prototype param[0])
        Optimization terminated successfully.
                 Current function value: 28.268922
                 Iterations: 105
                 Function evaluations: 189
        Optimization terminated successfully.
                 Current function value: 32.246756
                 Iterations: 245
                 Function evaluations: 405
        best_examplar_param [1.48769296 0.35961247 1.93041277 0.78398669]
        best_prototype_param [1.58816955 0.03164535 1.87904483 0.08402034]
```

Exemplar model

The negative log score is 28.268922073692416



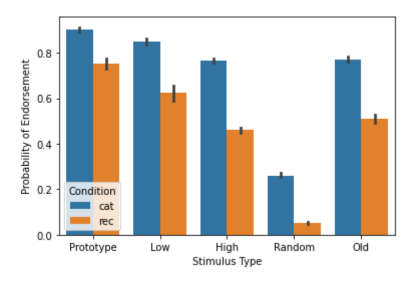
Prototype Model

```
In [10]: # human = human_res.groupby(['Condition', 'Stimulus Type'], as_index=Fi
se).sum()

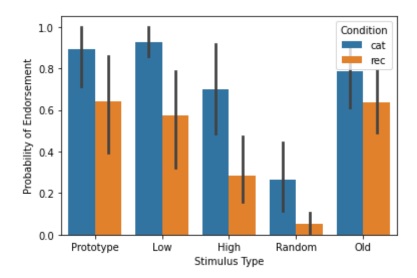
params = [1.58816955, 0.03164535, 1.87904483, 0.08402034]
nllfit = fit_prototype_model_nll(params, human)
print(f'The negative log score is {nllfit}')

# now plot the data
c_cat, k_cat, c_rec, k_rec = params
res = get_prototype_results(c_cat, k_cat, c_rec, k_rec)
sns.barplot(x="Stimulus Type", y="Probability of Endorsement",
hue="Condition", data=res)
```

The negative log score is 32.246756097276645



Human data again for reference



Problem 7 (10 points)

A famous saying is the "All models are wrong, but some are useful" (George Box). Do you think the exemplar or prototype model provides the best account of the data? Refer to particular patterns in the data that you believe the different models do a better job with.

With both RMSE and NLL scoring methods, examplar model performed better than prototype model. However, I believe this is difficult to say one method is much better than the other because each method has different upsides. Examplar model takes into account multiple patterns and finds the similarity which make it better for finding more distorted patterns and specific patterns from the original training data. This is shown in the graphs where examplar performs well with high distortion, random, and old stimuli. On the other hand, prototype model creates a representative prototype and finds similarity with the train data. This make this method stronger at finding averaged/less distorted data, proven by the graphs that show good performance on prototype, low, and random stimuli. As a result, in the five stimulus types, examplar model did slightly better on average, but not by a huge margin.

Problem 8 (5 points)

Thinking about how these models work explain why both the exemplar and prototype models have relatively high endorsement for the prototype item even though it was never presented during the training phase. In addition, explain in your own words why the models are able to explain the high endorsement rates for the old items.

Both exemplar and prototype models have relatively high endorsement for the prototype stimulus because both methods are able to find similarity in the given pattern through mulitple examples (in the case of exemplar) and through creating a single representative prototype (in the case of prototype). Since both methods are essentially doing the same comparision, it makes sense they perform well for endorsing prototype item. In addition, these models are also able to explain the old items with high endorsement rate because examplar keeps the memory of train data, and prototype can compute high similarity with old items to its representative data. Nonetheless, examplar has higher endorsement rate because it can recognize the exact patterns it has seen previously.

Problem 9 (5 points)

Are the exemplar model and the prototype model we considered nested? Would we compare them using AIC, BIC, or the G^2 statistic (or something else)?

Exemplar and prototype models are not nested. By definition, a nested model is simply a regression model that contains a subset of the predictor variables in another regression model. Exemplar model stores all training examples/data and use them for similarity calculation, but prototype model stores the representation of the training data. This means one model cannot be a subset of another model for both exemplar and prototype models. When we compare the two models, AIC and BIC may be valid since both methods combine a term reflecting how well the model fits the data with a term that penalizes the model in proportion to its number of parameters. However, G^2 statistic is widely used for comparing the fit of nested models, which may not be a good fit for comparing exemplar and prototype models.

In []:	
---------	--