```
In [1]: # Initialize Otter
import otter
grader = otter.Notebook("hw3-diatom.ipynb")
```

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```
In [2]: import numpy as np
import pandas as pd
import altair as alt
from sklearn.decomposition import PCA
```

Background: diatoms and paleoclimatology

Diatoms are a type of phytoplankton -- they are photosynthetic algae that function as primary producers in aquatic ecosystems. Diatoms are at the bottom of the food web: they are consumed by filter feeders, like clams, mussels, and many fish, which are in turn consumed by larger organisms like scavengers and predators and, well, us. As a result, changes in the composition of diatom species in marine ecosystems have ripple effects that can dramatically alter overall community structure in any environment of which marine life forms a part.

Diatoms have glass bodies. As a group of organisms, they display a great diversity of body shapes, and many are quite elaborate. The image below, taken from a Scientific American article, shows a small sample of their shapes and structures.



Because they are made of glass, diatoms preserve extraordinarily well over time. When they die, their bodies sink and form part of the sediment. Due to their abundance, there is a sort of steady rain of diatoms forming part of the sedimentation process, which produces sediment layers that are dense with diatoms.

Sedimentation is a long-term process spanning great stretches of time, and the deeper one looks in sediment, the older the material. Since diatoms are present in high density throughout sedimentation layers, and they preserve so well, it is possible to study their presence over longer time spans -- potentially hundreds of thousands of years.

A branch of paleoclimatology is dedicated to studying changes in biological productivity on geologic time scales, and much research in this area has involved studying the relative abundances of diatoms. In this assignment, you'll do just that on a small scale and work with data from sediment cores taken in the gulf of California at the location indicated on the map:



The data is publicly available:

Barron, J.A., et al. 2005. High Resolution Guaymas Basin Geochemical, Diatom, and Silicoflagellate Data. IGBP PAGES/World Data Center for Paleoclimatology Data Contribution Series # 2005-022. NOAA/NGDC Paleoclimatology Program, Boulder CO, USA.

Assignment objectives

In this assignment, you'll use the exploratory techniques we've been discussing in class to analyze the relative abundances of diatom taxa over a time span of 15,000 years. This will involve practicing the following skills.

Acquaint and tidy

- data import
- handling NaNs
- transforming values
- assessing time resolution

Exploratory analysis of individual variables

- visualizing summary statistics
- density histograms and kernel density estimates
- describing variation

Exploratory analysis of multiple variables

- examining correlation structure
- computing and selecting principal components
- interpreting principal component loadings
- using principal components to visualize multivariate data

Communication and critical thinking

- summarizing results in written form
- suggesting next steps

Have fun!

Collaboration

You are encouraged to collaborate with other students on the labs, but are expected to write up your own work for submission. Copying and pasting others' solutions is considered plaigarism and may result in penalties, depending on severity and extent.

If you choose to work with others, please list their names here.

Your name: Amy Nguyen

Collaborators:

O. Getting acquainted with the diatom data

In this assignment you'll focus less on tidying and more on exploration -- the data you'll work with are already tidy. So, in this initial part, you'll:

- · import the data;
- examine its structure to get acquainted; and
- perform some simple preprocessing transformations to facilitate exploratory analysis.

The data are diatom counts sampled from evenly-spaced depths in a sediment core from the gulf of California. In sediment cores, depth correlates with time before the present -- deeper layers are older -- and depths are typically chosen to obtain a desired temporal resolution. The counts were recorded by sampling material from sediment cores at each depth, and examining the sampled material for phytoplankton cells. For each sample, phytoplankton were identified at the taxon level and counts of diatom taxa were recorded along with the total number of phytoplankton cells identified. Thus:

- The observational units are sediment samples.
- The variables are depth (age), diatom abundance counts, and the total number of identified phytoplankton. Age is inferred from radiocarbon.
- One observation is made at each depth from 0cm (surface) to 13.71 cm.

The table below provides variable descriptions and units for each column in the dataframe.

Variable	Description	Units
Depth	Depth interval location of sampled material in sediment core	Centimeters (cm)
Age	Radiocarbon age	Thousands of years before present (KyrBP)
A_curv	Abundance of Actinocyclus curvatulus	Count (n)
A_octon	Abundance of Actinocyclus octonarius	Count (n)
ActinSpp	Abundance of Actinoptychus species	Count (n)
A_nodul	Abundance of Azpeitia nodulifer	Count (n)
CocsinSpp	Abundance of Coscinodiscus species	Count (n)
CyclotSpp	Abundance of Cyclotella species	Count (n)
Rop_tess	Abundance of Roperia tesselata	Count (n)
StephanSpp	Abundance of Stephanopyxis species	Count (n)
Num.counted	Number of diatoms counted in sample	Count (n)

The cell below imports the data.

```
In [3]: # import diatom data
    diatoms_raw = pd.read_csv('data/barron-diatoms.csv')
    diatoms_raw.head(5)
```

Out[3]:		Depth	Age	A_curv	A_octon	ActinSpp	A_nodul	CoscinSpp	CyclotSpp	Rop_tess	StephanSpp	Num.counted
	0	0.00	1.33	5.0	2.0	32	14.0	21	22.0	1.0	1.0	201
	1	0.05	1.37	8.0	2.0	31	16.0	20	16.0	7.0	2.0	200
	2	0.10	1.42	8.0	6.0	33	18.0	29	7.0	1.0	1.0	200
	3	0.15	1.46	11.0	1.0	21	1.0	12	28.0	25.0	3.0	200
	4	0.20	1.51	11.0	1.0	38	3.0	18	24.0	3.0	NaN	300

The data are already in tidy format, because each row is an observation (a set of measurements on one sample of sediment) and each column is a variable (one of age, depth, or counts). However, examine rows 3 and 4. These rows illustrate two noteworthy features of the raw data:

1. NaNs are present

2. The number of individuals counted in each sample varies by a lot from sample to sample.

Let's address those before conducting initial explorations.

'Missing' values

The NaNs are an artefact of the data recording -- if *no* diatoms in a particular taxa are observed, a – is entered in the table (you can verify this by checking the .csv file). In these cases the value isn't missing, but rather zero. These entries are parsed by pandas as NaNs, but they correspond to a value of 0 (no diatoms observed).

Q0 (a). Filling NaNs

Use .fill_na() to replace all NaNs by zeros, and store the result as diatoms_mod1. Store rows 4 and 5 (index, not integer location) of the resulting dataframe as diatoms_mod1_sample and print it out.

(*Hint*: check the documentation for fill_na().)

```
In [4]: diatoms_mod1 = diatoms_raw.fillna(0)
        # print rows 4 and 5
        diatoms_mod1_sample = diatoms_mod1.iloc[[4,5]]
        print(diatoms_mod1_sample)
          Depth Age A_curv A_octon ActinSpp A_nodul CoscinSpp CyclotSpp
           0.20 1.51
                         11.0
                                1.0
                                        38
                                                   3.0
                                                                18
                                                                         24.0
           0.25 1.55
                          4.0
                                  9.0
                                             30
                                                    10.0
                                                                16
                                                                         14.0
          Rop tess StephanSpp Num.counted
                                       300
                          0.0
        5
              16.0
                           0.0
                                       203
In [5]:
        grader.check("q0_a")
```

Out [5]: **q0_a** passed!

Varying total counts

Since the total number of phytoplankton counted in each sample varies, the raw counts are not directly comparable -- e.g., a count of 18 is actually a *different* abundance in a sample with 200 individuals counted than in a sample with 300 individuals counted.

For exploratory analysis, you'll want the values to be comparable across rows. This can be achieved by a simple transformation so that the values are *relative* abundances: *proportions* of phytoplankton observed from each taxon.

Q0 (b). Counts to proportions

Convert the counts to proportions by dividing by the relevant entry in the Num. counted column. There are a few ways to do this, but here's one approach:

- 1. Set Depth and Age to row indices using _set_index(...) and store the result as diatoms_mod2.
- 2. Store the Num.counted column from diatoms_mod2 as sampsize.
- 3. Use .div(...) to divide entrywise every column in diatoms_mod2 by sampsize and store the result as diatoms_mod3.
- 4. Drop the Num.counted column from diatoms_mod3 and reset the index; store the result as diatoms.

Carry out these steps and print the first four rows of diatoms.

(*Hint*: careful with the axis = ... argument in .div(...); you may want to look at the documentation.)

```
In [6]: # set depth, age to indices
    diatoms_mod2 = diatoms_mod1.set_index(['Depth', 'Age'])

# store sample sizes
    sampsize = diatoms_mod2['Num.counted']

# divide
    diatoms_mod3 = diatoms_mod2.div(sampsize, axis = 0)

# drop num.counted and reset index
    diatoms = diatoms_mod3.drop(columns='Num.counted').reset_index()

# print
    diatoms.head()
```

A_curv

0.00 1.33 0.024876 0.009950

A_octon ActinSpp

0.159204 0.069652

	1	0.05	1.37	0.040000	0.010000	0.155000	0.080000	0.100000	0.080000	0.035000	0.010000
	2	0.10	1.42	0.040000	0.030000	0.165000	0.090000	0.145000	0.035000	0.005000	0.005000
	3	0.15	1.46	0.055000	0.005000	0.105000	0.005000	0.060000	0.140000	0.125000	0.015000
	4	0.20	1.51	0.036667	0.003333	0.126667	0.010000	0.060000	0.080000	0.010000	0.000000
In [7]:	grader.check("q0_b")										

A_nodul CoscinSpp CyclotSpp

0.104478

0.109453

Rop_tess StephanSpp

0.004975

0.004975

Now that the data are ready for exploratory analysis, take a moment to think about what the data represent. They are relative abundances over time; essentially, snapshots of the community composition of diatoms over time, and thus information about how ecological

Before diving in, it will be helpful to resolve two matters:

1. How far back in time do the data go?

community composition changes.

2. What is the time resolution of the data?

Q0 (c). Time span

Out[6]:

Out[7]:

Depth Age

q0_b passed!

What is the geological time span covered by the data? Compute the minimum and maximum age using <code>.aggregate(...)</code> and store as <code>min_max_age</code>.

Note: This may be a new function for you, but it's simple: it takes as an argument a list of functions that will be applied to the dataframe (columnwise by default). So for example, to get the mean and variance of each column in df, one would use df.aggregate(['mean', 'var']). See the documentation for further examples.

(Remember: age is reported as thousands of years before present, so Age = 2 means 2000 years ago.)

q0_c passed!

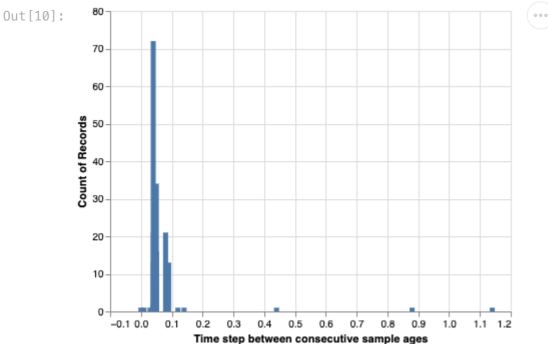
Q0 (d). Time resolution

How are the observations spaced in time?

(i) Make a histogram of the time steps between consecutive sample ages.

Follow these steps:

- 1. Extract the Age column from diatoms, sort the values in ascending order, compute the differences between consecutive rows, and store the result as diffs.
 - *Hint*: use <code>.sort_values()</code> and <code>.diff()</code>.
 - Notice: that the first difference is NaN, because there is no previous value to compare the first row with. Drop this entry when you store diffs.
- 1. Make a simple count histogram (no need to manually bin or convert to density scale) with bins of width 0.02 (20 years).
 - Label the x axis 'Time step between consecutive sample ages'



```
In [11]: grader.check("q1_d_i")
```

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Out [11]: **q1_d_i** passed!

(ii) What is the typical time step in years?

The typical time step is 40 years.

1. Exploring diatom taxon abundances

Recall that the first type of exploratory analysis question has to do with exploring variation in each variable; to begin, you'll examine the variation in relative abundance over time for the eight individual taxa.

Here are some initial questions in this spirit that will help you to hone in and develop more focuesed exploratory questions:

- Which taxa are most and least abundant on average over time?
- Which taxa vary the most over time?

These can be answered by computing simple summary statistics for each column in the diatom data.

Q1 (a). Summary statistics

Use .aggregate(...) to find the mean and standard deviation of relative abundances for each taxon. Follow these steps:

- 1. See Q0 (c) for an explanation of .aggregate(...).
- 2. Drop the depth and age variables before performing the aggregation.
- 3. Use transpose() to ensure that the table is rendered in long form (8 rows by 2 columns rather than 2 columns by 8 rows).
- 4. Store the result as diatom_summary and print the dataframe.

```
In [12]: diatom_summary = diatoms.drop(columns=['Depth', 'Age']).aggregate(['mean', 'std']).transpose()
# print the dataframe
diatom_summary
```

```
Out[12]: mean std

A_curv 0.028989 0.018602
A_octon 0.018257 0.016465
ActinSpp 0.135900 0.053797
A_nodul 0.072940 0.092677
CoscinSpp 0.085925 0.031795
CyclotSpp 0.070366 0.042423
Rop_tess 0.060448 0.076098
StephanSpp 0.002447 0.007721
```

```
In [13]: grader.check("q1_a")
```

q1_a passed!

It will be easier to determine which taxa are most/least abundant and most variable by displaying this information visually.

Out[13]:

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Q1 (b). Visualizing summary statistics

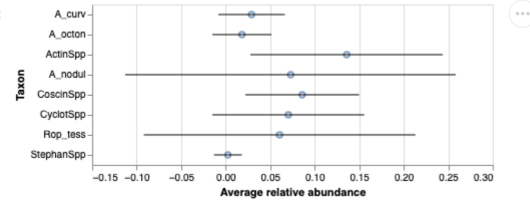
Create a plot of the average relative abundances and their variation over time by following these steps:

- 1. Reset the index of diatom_summary so that the taxon names are stored as a column and not an index. Store the result as plot_df.
- 2. Create an Altair chart based on plot_df with no marks -- just alt.Chart(...).encode(...) -- and pass the column of taxon names to the Y encoding channel with the title 'Taxon' and sorted in descending order of mean relative abundance. Store the result as base.
 - *Hint*: alt.Y(..., sort = {'field': 'column', 'order': 'descending'}) will sort the Y channel by 'column' in descending order.
- 1. Modify base to create a point plot of the average relative abundances for each taxon; store the result as means.
 - Average relative abundance (the mean you calculated in Q1 (a)) should appear on the x axis, and taxon on the y axis.
 - Since the Y encoding was already specified in base, you do not need to add a Y encoding at this stage.
 - Give the x axis the title 'Average relative abundance'.
- 1. Modify base to create a plot with bars spanning two standard deviations in either direction from the mean. Store the result as bars.
 - First use base.transform_calculate(...) to compute lwr and upr for the positions of the bar endpoints:
 - $lwr = mean 2 \times std$
 - lacksquare upr = mean $+2 imes exttt{std}$.
 - Then append .mark_errorbar().encode(...) to the chain:
 - pass lwr:Q to the X encoding channel with the title 'Average relative abundance' (to match the point plot)
 - pass upr:Q to the X2 encoding channel (no specific title needed).
- 1. Layer the plots: means + bars.

It may help to have a look at this example. Once you make the plot, answer questions (i) - (iii) below.

```
In [14]: # reset index
         plot df = diatom summary.reset index()
         # create base chart
         base = alt.Chart(plot df).encode(
             y = alt.Y('index',
                       title = 'Taxon',
                       sort = {'field': 'column', 'order': 'descending'})
         # create point plot
         means = base.mark_point().encode(
             x = alt.X('mean', title = 'Average relative abundance')
         # create bar plot
         bars = base.transform_calculate(
             lwr = 'datum.mean - 2 * datum.std',
             upr = 'datum.mean + 2 * datum.std'
         ).mark_errorbar().encode(
             x = alt.X('lwr:Q', title = 'Average relative abundance'),
             x2 = 'upr:Q
         # layer
         means + bars
```

Out[14]:



(i) Which taxon is most abundant on average over time?

Actinoptychus species is the most abundant taxon on average over time.

(ii) Which taxon is most rare on average over time?

Stephanopyxis species is the most rare taxon on average over time.

(iii) Which taxon varies most in relative abundance over time?

Azpeitia nodulifer taxon varies most in relative abundance over time.

Now that you have a sense of the typical abundances for each taxon (measured by means) and the variations in abundance (measured by standard deviations), you'll dig in a bit further and examine the variation in abundance of the most variable taxon.

For the next few questions, it may help you to follow code examples from lab 4.

Q1 (c). Distribution of Azpeitia nodulifer abundance over time

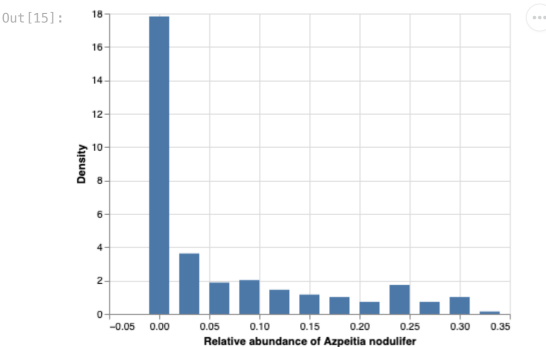
Here you'll construct a few plots that will help you answer the following key exploratory questions:

- Which values are common?
- Which values are rare?
- How spread out are the values?
- Are values spread evenly or irregularly?

(i) Construct a density scale histogram of the relative abundances of Azpeitia nodulifer.

Use the diatoms dataframe and a bin width of 0.03 and store the histogram as hist.

Hint: It may help to look at Q0 in Lab4 on Smoothing



(ii) Construct a kernel density estimate of the distribution.

Create and store the KDE curve as smooth and layer it on top of the density histogram from part i.

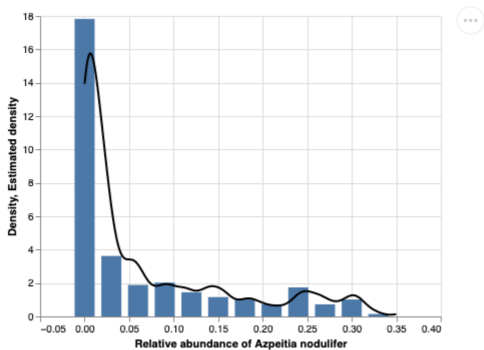
(Remember: experiment with the bandwidth parameter, and find a value that you feel captures the shape best.)

```
In [16]:
    smooth = base.transform_density(
        density = 'A_nodul',
        as_ = ['Relative abundance of Azpeitia nodulifer', 'Estimated density'],
        bandwidth = 0.01,
        extent = [0,0.35],
        steps = 500
    ).mark_line(color='black').encode(
        x = 'Relative abundance of Azpeitia nodulifer:Q',
        y = 'Estimated density:Q'
    )
    hist + smooth
```

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Out[16]:



(iii) Which values are common?

0.0 is the most common value for the relative abundance of Azpeitia nodulifer.

(iv) Which values are rare?

Values greater than 0.10 are rare.

(v) How spread out are the values and how are they spread out?

The values range from 0 to 0.35, where the most common value of the Relative abundance of Azpeitia nodulifer is 0 and the density estimates drop drastically for values greater than zero. Additionally, as relative abundance increases, the estimated density decreases.

(vi) How would you describe the shape?

The shape of the distribution for the relative abundance of Azpeitia nodulifer is right-skewed.

Comment: 'zero inflation'

There are a disproportionately large number of zeroes, because in many samples no *Azpeitia nodulifer* diatoms were observed. This is a common phenomenon in ecological data, and even has a name: it results in a 'zero inflated' distribution of values. The statistician to identify and name the phenomenon was Diane Lambert, whose highly influential work on the subject (>4k citations) was published in 1992.

Zero inflation can present a variety of challenges. You may have noticed, for example, that there was no bandwidth parameter for the KDE curve that *both* captured the shape of the histogram near zero *and* away from zero -- it either got the height near zero right but was too wiggly, or got the shape away from zero right but was too low near zero.

Conditioning on a climate event

There was a major climate event during the time span covered by the diatom data. The oldest data points in the diatom data correspond to the end of the Pleistocene epoch (ice age), at which time there was a pronounced warming (Late Glacial Interstadial, 14.7 - 12.9 KyrBP) followed by a return to glacial conditions (Younger Dryas, 12.9 - 11.7 KyrBP).

This fluctuation can be seen from temperature reconstructions. Below is a plot of sea surface temperature reconstructions off the coast of Northern California. Data come from the following source:

Barron *et al.*, 2003. Northern Coastal California High Resolution Holocene/Late Pleistocene Oceanographic Data. IGBP PAGES/World Data Center for Paleoclimatology. Data Contribution Series # 2003-014. NOAA/NGDC Paleoclimatology Program, Boulder CO, USA.

The shaded region indicates the time window with unusually large flucutations in sea surface temperature; this window roughly corresponds to the dates of the climate event.

```
)
).mark_area(opacity = 0.2, color = 'orange').encode(
    y = 'SST',
    x = alt.X('upr', title = 'Thousands of years before present'),
    x2 = 'lwr'
)

# add smooth trend
smooth = line.transform_loess(
    on = 'Age',
    loess = 'SST',
    bandwidth = 0.2
).mark_line(color = 'black')

# layer
line + highlight + smooth
```

Out [17]:

12

10

8

4

2

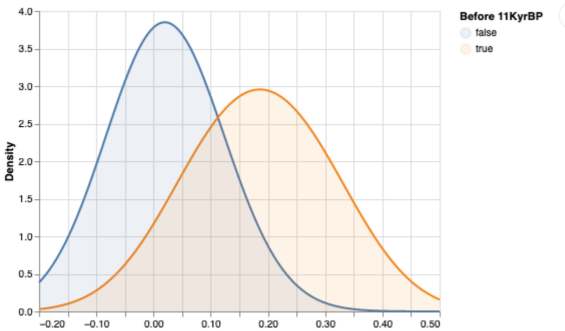
Q1 (d). Conditional distributions of relative abundance

Thousands of years before present

Does the distribution of relative abundance of Azpeitia nodulifer differ when variation in sea temperatures was higher (before 11KyrBP)?

- (i) Plot kernel density estimates to show the distribution of relative abundances before and after 11KyrBP.
- 1. Use .transform_caluculate(...) to calculate an indicator variable, pre_dryas, that indicates whether Age exceeds 11.
- 2. Use transform_density(...) to compute KDEs separately for observations of relative abundance before and after 11KyrBP.
 - *Hint*: group by pre_dryas
- 1. Plot the KDEs distinguished by color; give the color legend the title 'Before 11KyrBP' and store the plot as kdes.
- 2. Add a shaded area beneath the KDE curves. Adjust the opacity of the area to your liking.

Out[18]:



Relative abundance of Azpeitia nodulifer

(ii) Describe the variation in relative abundance of Azpeitia nodulifer between now and 11,000 years ago.

Between now and 11,000 years ago, there were little to no Azpeitia nodulifer diatoms observed which is characterized by the sharp peak about 0. The relative abundance of Azpeitia nodulifer is rare for values greater than 0.05 after 11KyrBP.

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(iii) Describe the variation in relative abundance of Azpeitia nodulifer between 11,000 and 14,000 years ago.

Between 11,000 and 14,000 years ago, Azpeitia nodulifer diatoms were relatively more abundant reaching the most abundant a little below 0.15 and a little above 0.25. Relative abundance below 0.05 and above 0.35 were rare.

2. Visualizing community composition with PCA

So far you've seen that the abundances of one taxon -- *Azpeitia nodulifer* -- change markedly before and after a shift in climate conditions. In this part you'll use PCA to explore variation in community composition *among* all eight taxa.

Throughout this part, it may help you to refer to code examples from lab 5.

Q2 (a). Pairwise correlations in relative abundances

Before carrying out PCA it is a good idea to inspect the correlations between relative abundances directly. Here you'll compute and then visualize the correlation matrix.

(i) Compute the pairwise correlations between relative abundances.

Be sure to remove or set to indices the Depth and Age variables before computing the correlation matrix. Save the matrix as corr_mx and print the result.

Hint: See the pandas documentation for .corr().

In [19]: corr mx = diatoms.drop(columns=['Depth', 'Age']).corr() corr_mx Out[19]: A_octon ActinSpp A_nodul CoscinSpp CyclotSpp Rop_tess StephanSpp A_curv 1.000000 0.390898 -0.446778 A_curv 0.111480 0.091222 0.219439 -0.062690 0.151909 A_octon 0.111480 1.000000 -0.005009 -0.217992 0.049589 0.065249 -0.023047 -0.041017 1.000000 -0.363475 -0.055732 **ActinSpp** 0.390898 -0.005009 0.306021 -0.343410 0.058494 **A_nodul** -0.446778 -0.217992 -0.363475 1.000000 -0.010920 -0.407338 -0.471941 -0.151409 CoscinSpp 0.091222 0.049589 0.306021 -0.010920 1.000000 -0.266157 -0.016332 -0.341755 CyclotSpp 1.000000 0.018149 0.070684 0.219439 0.065249 -0.055732 -0.407338 -0.266157 -0.062690 -0.023047 0.032607 Rop_tess -0.343410 -0.471941 -0.341755 0.018149 1.000000

-0.016332

0.070684

0.032607

1.000000

In [20]: grader.check("q2_a_i")

StephanSpp

0.151909

Out [20]: **q2_a_i** passed!

(ii) Visualize the correlation matrix as a heatmap

-0.041017

0.058494

-0.151409

Have a look at either lab 5 or this example (or both!). Notice that to make a heatmap of a matrix, you'll need to melt it into long format.

1. Melt corr_mx to obtain a dataframe with three columns:

- row , which contains the values of the index of corr_mx (taxon names);
- column, which contains the names of the columns of corr_mx (also taxon names); and
- Correlation , which contains the values of corr_mx .

Store the result as corr_mx_long.

- 1. Create an Altair chart based on corr_mx_long and construct the heatmap by following the examples indicated above.
 - Adjust the color scheme to **blueorange** over the extent (-1, 1) to obtain a diverging color gradient where a correlation of zero is blank (white).
 - Adjust the color legend to indicate the color values corresponding to correlations of 1, 0.5, 0, -0.5, and -1.
 - Sort the rows and columns in ascending order of correlation.

```
In [21]: # melt corr_mx
         corr_mx_long = corr_mx.reset_index().rename(
             columns = {'index':'row'}
         ) • melt(
             id_vars = 'row',
             var name = 'col',
             value_name = 'Correlation'
         # construct plot
         heatmap = alt.Chart(corr_mx_long).mark_rect().encode(
             x = alt.X('col', title = '', sort = {'field': 'Correlation', 'order': 'ascending'}),
             y = alt.Y('row', title = '', sort = {'field': 'Correlation', 'order': 'ascending'}),
             color = alt.Color('Correlation',
                                scale = alt.Scale(scheme = 'blueorange',
                                                  domain = (-1, 1),
                                                  type = 'sqrt'),
                               legend = alt.Legend(tickCount = 5))
         ).properties(width = 200, height = 200)
         heatmap
```

Correlation Out[21]: A_nodul 1.0 Rop_tess 0.5 A_curv CyclotSpp ActinSpp 0.0 CoscinSpp A_octon StephanSpp -0.5A_curv CyclotSpp ActinSpp CoscinSpp

```
In [22]: grader.check("q2_a_ii")
```

Out [22]: **q2_a_ii** passed!

(iii) How does the relative abundance of Azpeitia nodulifer seem to covary with the other taxa?

Based off the correlation heatmap, the relative abundance of Azpeitia nodulifer appears to be negatively correlated with the abundance of all other taxons. The relative abundance of Azpeitia nodulifer has the weakest correlation with the Coscinodiscus species, indicating that this pair has nearly no correspondence with one another in comparison to the other taxons.

Q2 (b). Computing and selecting principal components

Here you'll perform all of the calculations involved in PCA and check the variance ratios to select an appropriate number of principal components. The parts of this question correspond to the individual steps in this process.

(i) Center and scale the data columns.

For PCA it is usually recommended to center and scale the data; set Depth and Age as indices and center and scale the relative abundances. Store the normalized result as pcdata.

```
In [23]: # helper variable pcdata_raw; set Depth and Age as indices
    pcdata_raw = diatoms.set_index(['Depth', 'Age'])

# center and scale the relative abundances
    pcdata = (pcdata_raw - pcdata_raw.mean())/pcdata_raw.std()
In [24]: grader.check("q2_b_i")
```

Out [24]: **q2_b_i** passed!

5/13/22, 4:55 PM

(ii) Compute the principal components.

Compute all 8 principal components. (For this part you do not need to show any specific output.)

q2_b_ii passed!

(iii) Examine the variance ratios.

Create a dataframe called pcvars with the variance information by following these steps:

- 1. Store the proportion of variance explained (called _explained_variance_ratio_ in the PCA output) as a dataframe named pcvars with just one column named Proportion of variance explained.
- 2. Add a column named Component to pcvars with the integers 1 through 8 as values (indicating the component number).
- 3. Add a column named Cumulative variance explained to powers that is the cumulative sum of Proportion of variance explained.
 - Hint: slice the Proportion of variance explained column and use .cumsum(axis = ...).

Print the dataframe pcvars.

```
In [27]: # store proportion of variance explained as a dataframe
    pcvars = pd.DataFrame({'Proportion of variance explained': pca.explained_variance_ratio_})

# add component number as a new column
    pcvars['Component'] = np.arange(1,9)

# add cumulative variance explained as a new column
    pcvars['Cumulative variance explained'] = pcvars.iloc[:, 0].cumsum(axis = 0)

# print
    pcvars
```

Proportion of variance explained Component Cumulative variance explained 0 0.255513 1 0.255513 0.478867 1 0.223354 2 2 0.132145 3 0.611012 3 0.122549 0.733560 4 0.110833 0.844394 5 0.922382 5 0.077988 6 6 0.067303 7 0.989684 0.010316 1.000000

```
In [28]: grader.check("q2_b_iii")
```

Out [28]: **q2_b_iii** passed!

Out[27]:

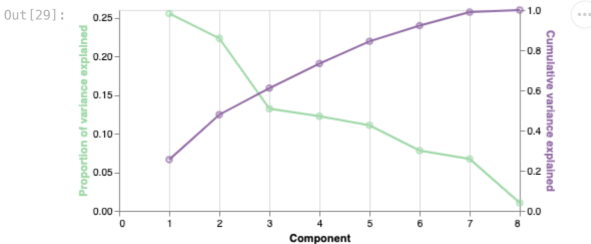
(iv) Plot the variance explained by each PC.

Use pcvars to construct a dual-axis plot showing the proportion of variance explained (left y axis) and cumulative variance explained (right y axis) as a function of component number (x axis), with points indicating the variance ratios and lines connecting the points.

Follow these steps:

- 1. Construct a base chart that encodes only Component on the X channel. Store this as base.
- 2. Make a base layer for the proportion of variance explained that modifies base by encoding Proportion of variance explained on the Y channel. Store the result as prop_var_base.
 - Give the Y axis title a distinct color of your choosing via alt.Y(..., axis = alt.Axis(titleColor = ...)).
- 1. Make a base layer for the cumulative variance explained that modifies base by endocing Cumulative variance explained on the Y channel. Store the result as cum_var_base.
 - Give the Y axis title another distinct color of your choosing via alt.Y(..., axis = alt.Axis(titleColor = ...)).
- 1. Create a plot layer for the proportion of variance explained by combining points (prop_var_base.mark_point()) with lines (prop_var_base.mark_line()). Store the result as cum_var.
 - Apply the color you chose for the axis title to the points and lines.
- 1. Repeat the previous step for the cumulative variance explained.
 - Apply the color you chose for the axis title to the points and lines.

1. Layer the plots together using alt.layer(l1, l2).resolve_scale(y = 'independent').



(v) How many PCs?

How many principal components capture a high proportion of covariation in relative abundances? How much total variation do these explain together?

The first two principal components each capture, respectively, roughly 25% and 22% of the variation in relative abundancies. Together, the two PCs explain nearly half of the total variation.

Q2 (c). Interpreting component loadings

Now that you've performed the calculations for PCA, you can move on to the fun part: figuring out what they say about the data!

The first step in this process is to examine the loadings. Each principal component is a linear combination of the relative abundances by taxon, and the loadings tell you *how* that combination is formed; the loadings are the linear combination coefficients, and thus correspond to the weight of each taxon in the corresponding principal component. Some useful points to keep in mind:

- a high loading value (negative or positive) indicates that a variable strongly influences the principal component;
- a negative loading value indicates that
 - increases in the value of a variable decrease the value of the principal component
 - and decreases in the value of a variable increase the value of the principal component;
- a positive loading value indicates that
 - increases in the value of a variable increase the value of the principal component
 - and decreases in the value of a variable decrease the value of the principal component;
- similar loadings between two or more variables indicate that the principal component reflects their average;
- divergent loadings between two sets of variables indicates that the principal component reflects their difference.

(i) Extract the loadings from pca.

Store the loadings for the first two principal components (called **.components_** in the PCA output) in a dataframe named **loading_df**. Name the columns PC1 and PC2, and append a column Taxon with the corresponding variable names, and print the resulting dataframe.

```
PC1
                             PC2
Out[30]:
                                        Taxon
           0 -0.521378
                       -0.157880
                                       A_curv
           1 -0.194520
                         0.001639
                                      A_octon
             -0.373815
                       -0.477144
                                      ActinSpp
               0.611563
                        -0.181503
                                      A_nodul
              -0.041199 -0.548427
                                    CoscinSpp
             -0.345726
                        0.285330
                                    CyclotSpp
             -0.116786
                         0.575665
                                     Rop_tess
             -0.204250
                         0.029760 StephanSpp
```

```
In [31]: grader.check("q2_c_i")
```

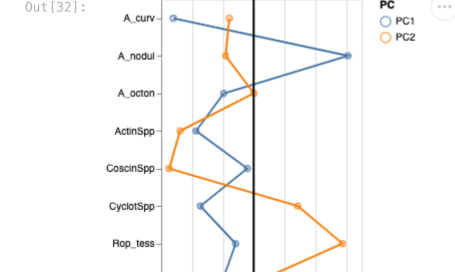
Out [31]: **q2_c_i** passed!

(ii) Loading plots

Construct a line-and-point plot connecting the loadings of the first two principal components. Display the value of the loading on the y axis and the taxa names on the x axis, and show points indicating the loading values. Distinguish the PC's by color, and add lines connecting the loading values for each principal component.

Hint: you will need to first melt loading_df to long form with three columns -- the taxon name, the principal component (1 or 2), and the value of the loading.

```
In [32]: # melt from wide to long
         loading_plot_df = loading_df.melt(
             id_vars = 'Taxon',
             var_name = 'PC',
             value_name = 'Loading'
         # create base layer with encoding
         base = alt.Chart(loading_plot_df).encode(
             y = alt.X('Taxon', title = ''),
             x = 'Loading',
             color = 'PC'
         # store horizontal line at zero
         rule = alt.Chart(pd.DataFrame({'Loading': 0}, index = [0])).mark_rule().encode(
             x = 'Loading', size = alt.value(2)
         # layer points + lines + rule to construct loading plot
         loading_plot = base.mark_point() + base.mark_line() + rule
         # show
         loading_plot.properties(height = 300, width = 200)
```



-0.2

(iii) Interpret the first principal component.

0.0

Loading

0.2

Indicate the following:

-0.6 - 0.4

StephanSpp

- which taxa are up-weighted and which are down-weighted;
- how you would describe the principal component in context (e.g., average abundance among a group, differences in abundances, etc.);
- and how you would interpret a positive value of the PC versus a negative value of the PC in terms of diatom community composition.

First looking at the first principal component, Azpeitia nodulifer taxon has the largest and only positive loading while all other taxa are negatively loaded for PC1, but more specifically Actinocyclus curvatulus has the largest negative loading. This suggests that sediment samples with a high value for PC1 will have a larger average relative of abundance for Azpeitia nodulifer, and a lower than average relative abundance of all other taxa, especially Actinocyclus curvatulus. In contrast, a lower than average relative abundance of Azpeitia nodulifer and higher than average relative abundance of all other taxa, especially Actinocyclus curvatulus, will yield a negative PC1 value. Thus, a negative PC1 value would indicate that Azpeitia nodulifer was absent while all other taxa were abundant in the the community composition and a positive PC1 value would mean Azpeitia nodulifer was highly present in the community composition while all other taxa were rarely found.

(iv) Interpret the second principal component.

Indicate the following:

- which taxa are up-weighted and which are down-weighted;
- how you would describe the principal component in context (e.g., average abundance among a group, differences in abundances, etc.);
- and how you would interpret a positive value of the PC versus a negative value of the PC in terms of diatom community composition.

For PC2, Roperia tesselata taxon has the largest positive loading while Coscinodiscus species and Actinoptychus species have the largest negative loadings. Stephanopyxis species and Actinocyclus octonarius have a loading value that is near zero meaning these taxa are not correlated with PC2. The second principal component increases when Roperia tesselata and Cyclotella have a higher than average relative abundance in the sample, and Coscinodiscus and Actinoptychus have a low relative abundance. Similarly, PC2 will have a negative value when Roperia tesselata and Cyclotella have a lower than average relative abundance in the sample, and Coscinodiscus and Actinoptychus have a higher average relative abundance. This implies that in the diatom community composition, samples where Roperia tesselata and Cyclotella species are most present in turn means Coscinodiscus and Actinoptychus were absent, and the reverse true for a negative PC2 value

Q2 (d). Visualizing community composition

Take a moment to recall that there was a shift in the *A. nodulifer* abundance before and after around 11,000 years ago, which roughly corresponded to a major transition in the earth's climate.

Well, you can now use PCA to investigate whether not just individual abundances but *community composition* may have shifted around that time. To that end, let's think of the principal components as 'community composition indices':

- consider PC1 a nodulifer/non-nodulifer community composition index; and
- consider PC2 a complex community composition index.

A pattern of variation or covariation in the principal components can be thought of as reflecting a particular ecological community composition dynamic -- a way that community composition varies throughout time. Here you'll look for distinct patterns of variation/covariation before and after 11,000 years ago via an exploratory plot of the principal components.

(i) Project the centered and scaled data onto the first two component directions.

This sounds a little more complicated than it is -- all that means is compute the values of the principal components for each data point.

Create a dataframe called projected_data containing just the first two principal components as two columns named PC1 and PC2, and two additional columns with the Age and Depth variables.

Print the first four rows of projected_data .

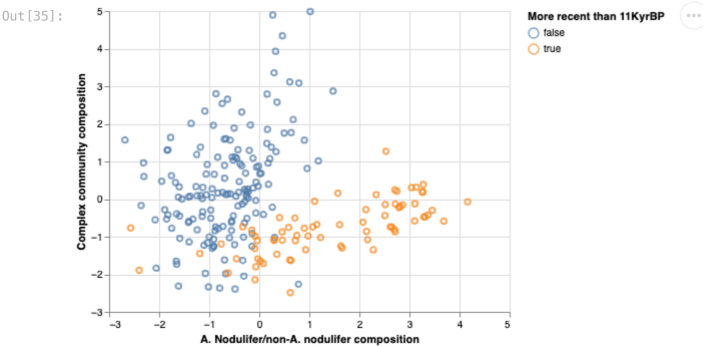
```
# project pcdata onto first two components; store as data frame
         projected_data = pd.DataFrame(pca.fit_transform(pcdata)).iloc[:, 0:2].rename(columns = {0: 'PC1', 1: 'PC2'})
          # add index and reset
         projected_data.index = pcdata.index
         projected_data = projected_data.reset_index()
          # print first four rows
         projected_data.head()
            Depth Age
                                      PC2
Out[33]:
                            PC1
             0.00 1.33 -0.294522 -0.633176
             0.05 1.37 -0.554702 -0.618875
              0.10 1.42 -0.307745 -2.050236
              0.15 1.46 -1.771066
                                 1.637274
             0.20 1.51 -0.292806
                                 0.259430
In [34]:
         grader.check("q2_d_i")
Out[34]:
```

q2_d_i passed!

Follow these steps to construct a scatterplot of the principal components.

- 1. Create an Altair chart based on projected_data and use .transform_calculate(...) to define a variable since_11KyrBP that indicates whether Age is older than 11,000 years. Store the result as base .
- 2. Modify base to add points with the following encodings.
 - Pass PC1 to the X encoding channel and title the axis 'A. Nodulifer/non-A. nodulifer composition'.
 - Pass PC2 to the Y encoding channel and title the axis 'Complex community composition'.
 - Pass the variable you created in step 1. to the color encoding channel and title it 'More recent than 11KyrBP'. Store the result as

Show the scatterplot once you complete these steps.



(iii) Add top and side panels with KDE curves.

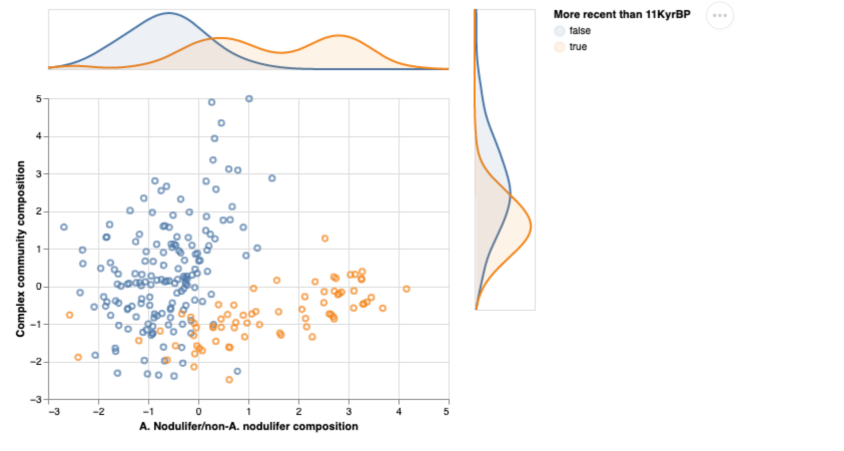
Construct plots of kernel density estimates for each principal component conditional on age being older than 11,000 years:

- modify base to create a top_panel plot with the KDE curves for PC1, with color corresponding to the age indicator from the .transform_calculate(...) step in making the base layer;
- modify base again to create a side_panel plot with the KDE curves for PC2, rotated 90 degrees relative to the usual orientation (flip the typical axes), and with color corresponding to the age indicator from the base layer.

Then, resize these panels appropriately (top should be thin, side should be narrow), and use Altair's faceting operators & (vertical concatenation) and | (horizontal concatenation) to combine them with your scatterplot.

```
In [36]: # construct upper panel (kdes for pc1)
         top_panel = base.transform_density(
             density = 'PC1',
             as_ = ['A. Nodulifer/non-A. nodulifer composition', 'Estimated density'],
             groupby = ['since_11KyrBP'],
             bandwidth = 0.4,
             extent = [-3,5]
         ).mark line().encode(
             x = alt.X('A. Nodulifer/non-A. nodulifer composition:Q',
                       title = '',
                       axis = None
                      ),
             y = alt.Y('Estimated density:Q',
                       title = '',
                       axis = None
                      ),
             color = alt.Color('since_11KyrBP:N')
         ).properties(height = 60)
         # construct side panel (kdes for pc2)
         side panel = base.transform density(
             density = 'PC2',
             as = ['Complex community composition', 'Estimated density'],
```

Out[36]:



3. Communicating results

Take a moment to review and reflect on the results of your analysis in the previous parts. Think about how you would describe succinctly what you've learned from the diatom data.

Q3 (a). Summary

Write a brief paragraph (3-5 sentences) that addresses the following questions by referring to your final plot in Q2 (d).

- How would you characterize the typical ecological community composition of diatom taxa before and after 11,000 years ago?
 - Hint: focus on the side and top panels and the typical values of each index in the two time periods.
- Does the variation in ecological community composition over time seem to differ before and after 11,000 years ago?
 - Hint: focus on the shape of data scatter.

Before 11,000 years ago, the typical ecological community composition of diatom taxa was absent of Azpeitia nodulifer which is exemplified in the KDE curves. The density estimate for A. nodulifer is centered about 0 during this earlier time period. However, after 11,000 years ago, the density estimates shift indicating that A. nodulifer was more abundant after the major climate change, meanwhile all other diatom taxa became less abundant. Additionally, there appeared to be more variation in ecological community composition before 11,000 years ago which is characterized by the larger spread of the scatterplot while after 11,000 years ago, there is a roughly linear relationship between A. nodulifer and complex community composition.

Q3 (b). Further work

What more might you like to know, given what you've learned? Pose a question that your exploratory analysis raises for you.

Answer

I would be curious to take a closer look at whether there were any major outliers, and if there were if they 'make sense' in relation to both the time period and the sea temperature. I think it would be interesting to merge the sea temperature dataset with the diatoms dataset to further explore the distributions of taxa with respect to sea temp. So that leads me to pose the question: What is the approximate relationship between diatom community composition and sea temperature?

Submission Checklist

- 1. Save file to confirm all changes are on disk
- 2. Run Kernel > Restart & Run All to execute all code from top to bottom
- 3. Save file again to write any new output to disk
- 4. Select File > Download as > HTML.
- 5. Open in Google Chrome and print to PDF on A3 paper in portrait orientation.
- 6. Submit to Gradescope

To double-check your work, the cell below will rerun all of the autograder tests.

```
In [37]: grader.check_all()

Out[37]: q0_a results: All test cases passed!

q0_b results: All test cases passed!

q0_c results: All test cases passed!

q1_a results: All test cases passed!

q1_d_i results: All test cases passed!

q2_a_i results: All test cases passed!

q2_a_ii results: All test cases passed!

q2_b_i results: All test cases passed!

q2_b_ii results: All test cases passed!

q2_b_iii results: All test cases passed!

q2_b_iii results: All test cases passed!

q2_c_i results: All test cases passed!

q2_d_i results: All test cases passed!
```