Assignment 1

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Question 1: Chromosome structures

The code used to solve this question:

```
import numpy as np
import pandas as pd
file_list = ['./ce10.chrom.sizes',
             './dm6.chrom.sizes',
             './ecoli.chrom.sizes',
             './hg38.chrom.sizes',
             './TAIR10.chrom.sizes',
             './tomato.chrom.sizes',
             './wheat.chrom.sizes',
             './yeast.chrom.sizes']
col_list = ['Total','Number', 'max_size', 'max_name', 'min_size', 'min_name', 'me
row list = ['ce10','dm6','ecoli','hg38','TAIR10','tomato','wheat','yeast']
def information(file_list, col_list, row_list):
    this function is used to produce a excel table of the chromosome information
    Parameters
    file list:list
    the files path
    col list:list
    the table column name list of the excel table
    row list:list
    the table row name list of the excel table
    # open a df data frame
    df = pd.DataFrame(columns=row_list, index = col_list)
    value_total = []
    value num = []
```

```
max value = []
    max value name = []
    min_value = []
    min value name = []
    mean = []
    i=0
    for name in file list:
        context = {}
        chrom = open(name, 'r')
        line = chrom.readline().replace('\n', '')
        while line != '':
            line list = line.split()
            context[line_list[0]] = int(line_list[1])
            line = chrom.readline().replace('\n', '')
        value_list = []
        for value in context.values():
            value_list.append(value)
        # calculate the information of the chromosome of a species and put it int
o a list
        value total = np.sum(value list)
        value_num = len(value_list)
        max value = np.max(value list)
        max value name = max(context, key=context.get)
        min_value = np.min(value_list)
        min_value_name = min(context, key=context.get)
        mean = np.mean(value_list)
        arr_value = [value_total, value_num, max_value, max_value_name,
            min value, min value name, mean]
        # write the list into dataframes
        df[row_list[i]] = arr_value
        i += 1
    df.to excel('excel.xls')
if __name__ == "__main__":
   information(file list, col list, row list)
```

the result table is save into an excel.xls file, the screen shoot of that file context is:

	ce10	dm6	ecoli	hg38	TAIR10	tomato	wheat	yeast
Total	1E+08	1.38E+0	463921	3.09E+0	1.19E+0	7.83E+0	1.45E+1	1215710
		8	1	9	8	8	0	5

Number	7	7	1	24	5	13	22	17
max_size	2092414	3207933	463921	2.49E+0	3042767	9086368	8.31E+0	1531933
	9	1	1	8	1	2	8	
max_nam	chrV	chr3R	Ecoli	chr1	Chr1	ch01	3B	chrIV
е								
min_size	13794	1348131	463921	4670998	1858505	9643250	4.74E+0	85779
			1	3	6		8	
min_nam	chrM	chr4	Ecoli	chr21	Chr4	ch00	6D	chrM
е								
mean	1432658	1964970	463921	1.29E+0	2382927	6019384	6.61E+0	715123.
	1	9	1	8	0	9	8	8

Question 2: Sequence content

Question 2.1

The code used to solve this question is:

```
import sys
def fafile2dict():
    this function can be used to calculate the As, Cs, Gs, Ts in entire genome
   STDIN:
    the fasta file
    run as 'python3 ques1.py yeast.fa'
    Return:
    base_a:int
    the number of As
   base_c:int
    the number of Cs
    base_g:int
    the number of Gs
    base_t:int
    the number of Ts
    line = sys.stdin.readline().replace('\n','')
    seq = \{\}
   while line != '':
```

```
if line[0] == '>':
            name = line.replace('\n','')
            seq[name] = ''
        else:
            seq[name] += line.replace('\n','').strip()
        line = sys.stdin.readline()
    base a = 0
    base_t = 0
    base c = 0
    base_g = 0
    for bp in seq.values():
        bp list = list(bp)
        for bp_sort in bp_list:
            if bp sort == 'A':
                base a += 1
            elif bp_sort == 'T':
                base_t += 1
            elif bp_sort == 'C':
                base c += 1
            else:
                base_g += 1
    print('A:',base_a,'T:',base_t,'C:',base_c,'G:',base_g)
    return base_a, base_t, base_c, base_g
if __name__ == "__main__":
    fafile2dict()
```

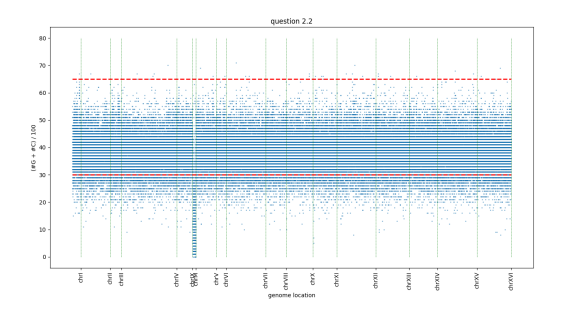
the result of this code is:

A: 3766349 T: 3753080 C: 2320576 G: 2317100

Question 2.2

```
line = sys.stdin.readline().replace('\n','')
    seq = \{\}
    num = []
    bp list = []
    n = 0
    chrom_name = []
    while line != '':
        if line[0] == '>':
            name = line.replace('\n','')
            seq[name] = ''
        else:
            seq[name] += line.replace('\n','').strip()
        line = sys.stdin.readline()
    for chrom, bp in seq.items():
        bp_list += bp
        n = n + len(bp)/100
        num.append(n)
        chrom_name.append(chrom.replace('>',''))
    count_list = []
    bp list = list(bp list)
    for i in range(int(len(bp_list)/100)):
        bp_frag = bp_list[100*i:(100*i+100)]
        base\_gc = 0
        for bp_sort in bp_frag:
            if bp sort == 'G' or bp sort == 'C':
                base_gc += 1
        count_list.append(base_gc)
    # draw the figure
    plt.xlabel('genome location')
    plt.ylabel('(#G + #C) / 100')
    plt.title('question 2.2')
    plt.vlines(num, 0, 80, 'g', 'dashed', linewidths=0.5)
    plt.hlines([30,65], 0, len(count_list),'r', 'dashed', linewidths=2)
    plt.scatter(range(len(count_list)), count_list, s=1, alpha=0.5)
    plt.xticks(num, chrom_name, rotation=90)
    plt.show()
if __name__ == "__main__":
    fafile2dict()
```

the result of the question is:

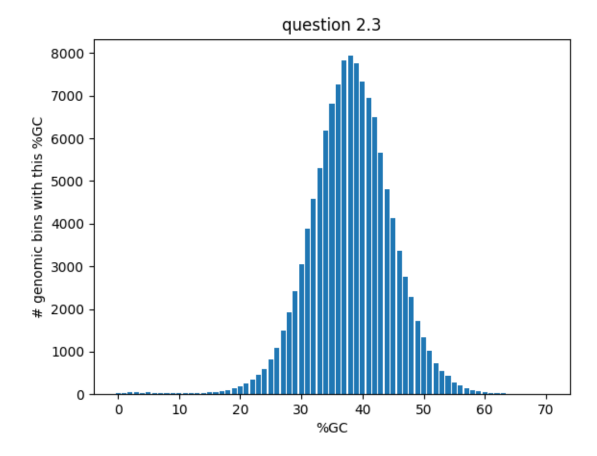


Question 2.3:

```
import sys
import matplotlib.pyplot as plt
import numpy as np
def fafile2dict():
    Make a histogram of the number of genomic bins of a given %GC
    STDIN:
    the fasta file
    run as 'python3 ques3.py < yeast.fa'</pre>
    line = sys.stdin.readline().replace('\n','')
    seq = \{\}
    while line != '':
        if line[0] == '>':
            name = line.replace('\n','')
            seq[name] = ''
        else:
            seq[name] += line.replace('\n','').strip()
```

```
line = sys.stdin.readline()
    count_list = []
    for bp in seq.values():
        bp list = list(bp)
        for i in range(int(len(bp_list)/100)):
            bp_frag = bp_list[(100*i-100):100*i]
            base gc = 0
            for bp_sort in bp_frag:
                if bp_sort == 'G' or bp_sort == 'C':
                    base_gc += 1
            count_list.append(base_gc)
    count_set = set(count_list)
    percentage = []
    num_gc = []
    for item in count_set:
        percentage.append(item)
        num_gc.append(count_list.count(item))
    num_gc = list(map(int, num_gc))
    percentage = list(map(int, percentage))
    plt.bar(percentage, num_gc)
    plt.xlabel("%GC")
    plt.ylabel("# genomic bins with this %GC")
    plt.title('question 2.3')
    plt.show()
if __name__ == "__main__":
   fafile2dict()
```

the result is:



Question2.4:

From the result of question 2.2, we can know that chrM will sequence poorly, because there is lots of GC $\leq 30\%$.