Project 4 Group 2 final presentation

Table of contents ¶

- 1. Import data and libraries
- 2. Data adjustments
- 3. Basic k-means
- 4. Mini batch
- 5. <u>K++</u>
- 6. Clustering Function and plotting
- 7. Results

Import of Data and libraries

Libraries

```
In [1]:
```

```
import numpy as np
import matplotlib.pyplot as plt
import matplotlib.cm as cm
import scanpy as sc
from matplotlib import colors
from datetime import datetime
from sklearn.decomposition import PCA
from sklearn.cluster import KMeans, MiniBatchKMeans
from sklearn.ensemble import IsolationForest
from mpl_toolkits.mplot3d import Axes3D
```

Data

```
In [2]:
```

```
data = sc.read_10x_mtx('./data/filtered_gene_bc_matrices/hg19/', var_names='gene_symbol
s', cache=True)
```

Data adjustments

Filtering

```
In [3]:
```

```
sc.pp.filter_genes(data, min_cells=1)
filtered_data = np.array(data._X.todense())
```

All genes which are not expressed at least once are removed from the dataset. Conversion of data into a numpy array for easier handling.

Removal of outliers and PCA

In [4]:

```
def remove_outliers():
    global pca_data
   X_train = pca_data
   clf = IsolationForest(behaviour="new", contamination=.07, max_samples=0.25)
    clf.fit(X_train)
    y_pred_train = clf.predict(X_train)
    pca_data = X_train[np.where(y_pred_train == 1, True, False)]
def pca(d, rmo=False):
    global dim, pca_data, pbmcs, genes
    dim = d
    pca = PCA(n_components=dim)
    pca_data = pca.fit_transform(filtered_data)
   if rmo == True:
        remove outliers()
    print("Sum of explained variances: ""%.2f" % (sum(pca.explained_variance_ratio_)) +
"\n")
    pbmcs = pca_data.shape[0]
    genes = pca_data.shape[1]
```

PCA to reduce dimensionality of our data to d dimension. The variables **pbmcs** and **genes** describe the dimensionalty of our data. Removal of outliers by Isolation forest can be chosen as a parameter (**rmo**).

Basic k-means

Initial centroid generation

In [5]:

```
def random start centroids(starttype):
    global centroids_array, genes, pbmcs, genes
    pbmcs = pca_data.shape[0]
    genes = pca_data.shape[1]
    centroids_array = np.empty([0, genes])
    if starttype == "randcell":
        centroids_numbers = np.random.randint(pbmcs, size=k)
        i = 0
        while i < k:
            random_cell = centroids_numbers[i]
            centroids_array = np.append(centroids_array, [pca_data[random_cell, :]], ax
is=0)
            i += 1
    elif starttype == "randnum":
        centroids_array = (np.amax(pca_data) - np.amin(pca_data)) * np.random.random_sa
mple((k, genes)) + np.amin(
            pca_data)
    elif starttype == "k++":
        kppcentroids()
```

Creating an empty array of centroids (**centroids_array**) which can be extended by three differnet methods. **"randcell"** selects k random samples from the given data. **"randum"** creates k new centroids by creating random numbers inbetween the maximum and minimum of the data values. **"k++"** performs the k++ centroid function.

Assignment of datapoints to closest centroids

In [6]:

```
def dist(cell point, cluster number):
    return np.linalg.norm(pca_data[cell_point, :] - centroids_array[cluster_number - 1,
:])
def assign_centroids(data_array):
    global nearest_centroid
    i = 0
    nearest centroid = np.zeros([data array.shape[0], 1])
    # Loop over all datapoints
    while i < data_array.shape[0]:</pre>
        sml_distance = -1
        # Loop over every centroid
        j = 1
        while j <= k:
            if sml_distance == -1 or dist(i, j) < sml_distance:</pre>
                sml_distance = dist(i, j)
                nearest_centroid[i, 0] = j
            j += 1
        i += 1
```

dist function returns linear distance between a selected datapoint (i) and a selected centroid (j).

assign_centroids function creates nearest_centroid array in which the closest centroid for each data point is saved.

Empty check

```
In [7]:
```

```
def empty_check():
    i = 0
    while i < k:
        if list(nearest_centroid).count(i + 1) == 0:
             print("Empty cluster! Correcting centroids.")
             random_start_centroids("randnum")
             assign_centroids(pca_data)
             empty_check()
        i += 1</pre>
```

Ensuring, if "randnum" method is chosen, no empty clusters were generated.

Generation of new centroids

```
In [8]:
```

```
def new_centroids():
    global centroids_array, centroids_oldarray, nearest_centroid_squeeze
    centroids_oldarray = centroids_array # create copy of old array for threshold funci
on
    nearest_centroid_squeeze = np.squeeze(nearest_centroid.astype(int))
    centroids_array = np.empty([0, genes])

i = 1
    while i <= k:
        calc_means = np.mean(pca_data[nearest_centroid_squeeze == i], axis = 0)
        centroids_array = np.append(centroids_array, np.expand_dims(calc_means, axis =
0), axis = 0)
    i += 1</pre>
```

Centroids of last generation are saved as **centroids_oldarray** to calculate distance to the subsequent generation for the threshold function. Creation of second centroid generation by calculation of the mean of associated data points.

Threshholding

```
In [9]:
```

```
def improv():
    distances = []
    i = 0
    while i < k:
        d = np.linalg.norm(centroids_array[i, :] - centroids_oldarray[i,:])
        distances.append(d)
        i += 1
        c_str = np.array2string(np.array(distances), precision=2)
        print("Distances of clusters as compared to last generation: \n" + str(c_str))</pre>
```

Calculation of distance inbetween each cluster after n iterations

Runtime function

```
In [10]:
```

```
def runtime_start():
    global t1
    t1 = datetime.now().time()

def runtime_end():
    t2 = datetime.now().time()
    fmt = '%H:%M:%S.%f'
    elapsed = str(datetime.strptime(str(t2), fmt) - datetime.strptime(str(t1), fmt))
    return str("\truntime: " + elapsed)
```

To evaluate efficiency.

Complete basic k-means

In [11]:

```
def kmeans(start, k1, n_iterations, t):
    global k
    k = k1
    i = 0
    runtime_start()
    random_start_centroids(start)
    assign centroids(pca data)
    if start == "randnum":
        empty_check()
    if t == None:
        while i < n_iterations:</pre>
            new_centroids()
            assign_centroids(pca_data)
            i += 1
        improv()
    else:
        count = 0
        d = t
        while d >= t:
            new_centroids()
            assign_centroids(pca_data)
            d = np.linalg.norm(centroids_oldarray-centroids_array)
            count+=1
        print("%s iterations were performed" %count)
    print("\nKMEANS:")
    print("\ngroup 4_2 algorithm:")
    print(runtime_end())
    print("\twss: " + str(wss('self')))
```

Function performing a basic k-means with k(k1) clusters until change of clusters is below the threshold t. If no threshold is chosen n iterations are performed.

Mini Batch

In [12]:

```
def minibatch(k1, n iterations, b):
    global k, pca_data, nearest_centroid_squeeze, pca_data, bg, n_iterationsg, centroid
s_array, cnnew
    k = k1
    bg = b
    n_iterationsg = n_iterations
    runtime_start()
    v = np.zeros((k, 1))
    j = 1
    random start centroids("randcell")
    cnnew = centroids_array
    while (j <= n iterations):</pre>
        # Reduce data to batch
        pca_batch = pca_data[np.random.randint(pca_data.shape[0], size=b), :]
        # Start centroids
        assign_centroids(pca_batch)
        i = 0
        # Centroid movement along gradient
        while (i < b):
            c = cnnew[int(nearest_centroid[i, 0])-1, :]
            v[int((nearest\_centroid[i, 0]-1)), 0] = int(v[int((nearest\_centroid[i, 0]-1)), 0])
1)), 0]) + 1
            n = 1/v[int((nearest_centroid[i, 0]-1)), 0]
            cnnew[int(nearest\_centroid[i, 0])-1, :] = c * (1-n) + pca\_data[i, :] * n
            i+=1
        j+=1
    centroids_array = cnnew
    assign_centroids(pca_data)
    nearest_centroid_squeeze = np.squeeze(nearest_centroid.astype(int))
    print("\nMINI-BATCH:")
    print("\ngroup 4_2 algorithm:")
    print(runtime_end())
    print("\twss: " + str(wss('self')))
```

Minibatch algorithm was implemented to the specifications of "Web-Scale K-Means Clustering" by D. Sculley.

K++

In [13]:

```
def kppcentroids():
    global centroids_array, dist_array, prob_array
    first_centroid = np.random.randint(pbmcs, size=1)
    i = 0
    centroids_array = np.append(centroids_array, pca_data[first_centroid, :], axis=0)
    dist_array = np.empty ([0,pbmcs])
    prob_array = np.empty ([0,pbmcs])
    j = 0
    while i < k - 1:
        z = centroids_array.shape[0] + 1
        while j < pbmcs:</pre>
            sml_distance = -1
            1 = 1
            while 1 < z:
                if sml_distance == -1 or dist(j, 1) < sml_distance:</pre>
                    sml_distance = dist(j, 1)
            dist_array = np.append(dist_array,sml_distance **2)
        prob_array = dist_array / np.sum(dist_array)
        s = np.random.choice(pbmcs,p = prob_array )
        centroids_array = np.append(centroids_array, np.expand_dims(pca data[s, :], axi
s=0), axis=0)
        i += 1
```

Creation of emtpy centroids array similar to random method. First centroid is chosen randomly. For every conecutive centroid an array of the squared distances between each data point to its closest centroids is created (dist_array). According probability array is created with identical dimension (prob_array). Centroid array is appeneded by selecting a random datapoint with according probability until k centroids are generated. Afterwards clustering follows basic k-means algorythm.

Clustering Function and plotting

Sklearn k-means

In [14]:

```
def sklearn kmeans function(var, k, start):
    global y_sklearnkmeans, sklearn_kmeans, pca_data
    runtime start()
    if start == "randcell" or start == "randnum":
        if var == "kmeans":
            sklearn_kmeans = KMeans(init='random', n_clusters=k).fit(pca_data)
        if var == "mini":
            sklearn_kmeans = MiniBatchKMeans(n_clusters=k, init = 'random', max_iter=n_
iterationsg, batch_size=bg).fit(pca_data)
    if start == "k++":
        if var == "kmeans":
            sklearn kmeans = KMeans(n clusters=k).fit(pca data)
        if var == "mini":
            sklearn_kmeans = MiniBatchKMeans(n_clusters=k, max_iter=n_iterationsg, batc
h_size=bg).fit(pca_data)
    y_sklearnkmeans = sklearn_kmeans.predict(pca_data)
    print("\nsklearn kmeans:")
    print(runtime_end())
    print("\twss: " + str(wss('sklearn')))
```

Plotting

```
In [15]:
```

```
def plots(add = ""):
    global fig1, fig2
    # 2D plots:
    additional = ""
    if add == "mini":
        additional = " (mini-batch)"
    # Kmeans
    fig1 = plt.figure(1, figsize=[10, 5], dpi=200)
    plt1, plt2 = fig1.subplots(1, 2)
    plt1.scatter(pca data[:, 0], pca data[:, 1], c=nearest centroid squeeze, s=0.5, cma
p='viridis')
    # plt1.plot(centroids_array[:, 0], centroids_array[:, 1], markersize=5, marker="s",
linestyle='None', c='k')
    plt1.set_title('kmeans' + additional)
    # Sklearnkmeans
    plt2.scatter(pca_data[:, 0], pca_data[:, 1], c=y_sklearnkmeans, s=0.5, cmap='viridi
    # plt2.plot(sklearn_kmeans.cluster_centers_[:, 0], sklearn_kmeans.cluster_centers_
[:, 1], markersize=5, marker="s", linestyle='None', c='k')
    plt2.set_title('sklearn kmeans' + additional)
    # 3D plots
    if dim >= 3:
        fig2 = plt.figure(2, figsize=[15,10], dpi=200)
        plt21 = fig2.add_subplot(221, projection = '3d')
        plt21.scatter(pca_data[:, 1], pca_data[:, 2], pca_data[:, 0], s=2, c = nearest_
centroid_squeeze, cmap='viridis')
        # plt21.plot(centroids_array[:, 0], centroids_array[:, 1], centroids_array[:,
2], markersize=5, marker="s", linestyle='None', c='k')
        plt21.set_title('3d kmeans' + additional)
        # Sklearnkmeans
        plt22 = fig2.add subplot(222, projection = '3d')
        plt22.scatter(pca_data[:, 1], pca_data[:, 2], pca_data[:, 0], s=2, c = y_sklear
nkmeans, cmap='viridis')
        # plt22.plot(sklearn_kmeans.cluster_centers_[:, 0], sklearn_kmeans.cluster_cent
ers_[:, 1], sklearn_kmeans.cluster_centers_[:, 2], markersize=5, marker="s", linestyle
        plt22.set title('3D kmeans by sklearn' + additional)
```

Quality control

In [16]:

```
def wss(where):
    i = 0
    wsssum = 0
    while (i < len(pca_data)):
        if where == "self":
            assigned_centroid = int(nearest_centroid[i,0])
            centr_val = centroids_array[assigned_centroid-1]
        if where == "sklearn":
            assigned_centroid = int(y_sklearnkmeans[i])
            centr_val = sklearn_kmeans.cluster_centers_[assigned_centroid]
        point_val = pca_data[i]
        i+=1
        sqdist = np.linalg.norm(centr_val - point_val)**2
        wsssum += np.trunc(sqdist)
    return(wsssum)</pre>
```

Calculation of within-cluster sum of squares.

Final Clustering Function

In [17]:

```
def cluster(pcas = 5, rmo=True, variant = 'kmeans', start='randcell', k = 3, max_iterat
ions = 10, threshold = 0.00001, batch_size = 2000):
    pca(pcas, rmo)

if variant == "kmeans":
        kmeans(start, k, max_iterations, threshold)
        sklearn_kmeans_function("kmeans", k, start)
        plots()

if variant == "mini":
        minibatch(k, max_iterations, batch_size)
        sklearn_kmeans_function("mini", k, start)
        plots("mini")
```

The depicted cluster funtion allows us to perform kmeans clustering while giving us the ability to select following parameters:

- 1. Amount of principal components
- 2. Removal of outliers (Y/N)
- 3. Performance of k-means or mini batch k-means algorythm
- 4. Method for creating the initial cluster Generation (random cells, random numbers, k++)
- 5. Cluster-amount
- 6. Maximum iterations (not utilizied if a threshold is defined)
- 7. Minimum threshhold for change of cluster centers
- 8. Batch size for mini batch k-means

By combining these features we are able to utilize the same function for all three subprojects.

Results

Basic k-means

In [18]:

```
cluster(variant = 'kmeans', start = "randcell", k=2)
```

Sum of explained variances: 0.60

8 iterations were performed

KMEANS:

group 4_2 algorithm:

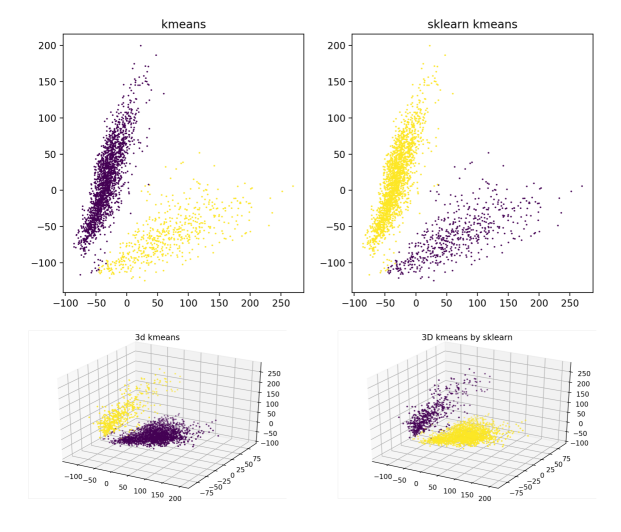
runtime: 0:00:00.419982

wss: 11497775.0

sklearn kmeans:

runtime: 0:00:00.100032

wss: 11497736.0



Mini batch k-means

In [21]:

```
cluster(variant = 'mini', start = "randcell", k=2)
```

Sum of explained variances: 0.60

MINI-BATCH:

group 4_2 algorithm:

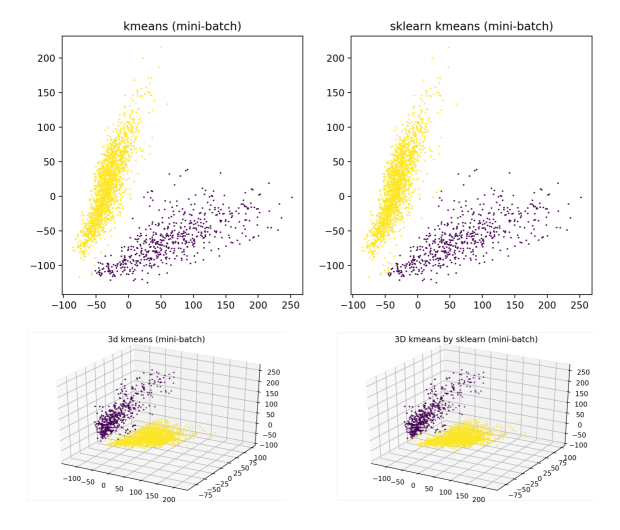
runtime: 0:00:00.696185

wss: 11665541.0

sklearn kmeans:

runtime: 0:00:00.044014

wss: 11645561.0



K++

In [20]:

```
cluster(variant = 'kmeans', start = "k++", k=2)
```

Sum of explained variances: 0.60

24 iterations were performed

KMEANS:

group 4_2 algorithm:

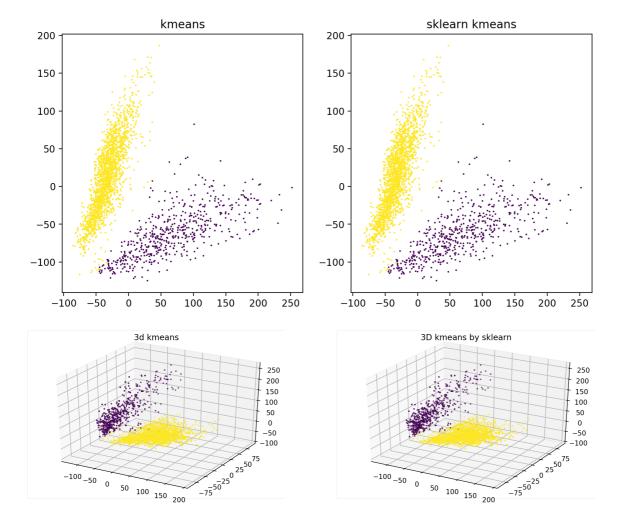
runtime: 0:00:01.255997

wss: 11504310.0

sklearn kmeans:

runtime: 0:00:00.087998

wss: 11504313.0



In []: