BME-230B Spring 2019 HW 2 Question 1

James Casaletto, Andrew Davidson, Yuanqing Xue, Jim Zheng

Question 1.a, 1.b see euclid knn.py (euclid knn.py)

ref:

- Single-Cell Analysis in Python (https://scanpy.readthedocs.io/en/stable/api/index.html#tools-tl)
- · data exploration (exploreData.ipynb)
- scanpy.tl.umap (https://icb-scanpy.readthedocs-hosted.com/en/stable/api/scanpy.tl.umap.html)
- · scanpy.api.pl no longer exists
- scanpy,pl.umap_(https://icb-scanpy,readthedocs-hosted.com/en/stable/api/scanpy,pl.umap.html#scanpy,pl.umap

```
In [1]: from euclid_knn import KnnG
  import matplotlib.pyplot as plt
  import numpy as np
  import scanpy.api as sc
  import scanpy
```

In [2]: anndata = sc.read("PBMC.merged.h5ad")

```
In [3]: %%time
# run PCA, compute pairwise distance, k-nearest-neighbors
# trick to replacing scanpy implementation with our own is to
# update the anndata object with our intermediate values
#
# we replace the scanpy version of PCA by updating
# .adata.obsm['X_pca'] = our PCA() output
#
# we replace the scanpy version of k-nearest-neighbors by updating
# self.adata.uns['neighbors']['connectivities'] = our knn() output
# self.adata.uns['neighbors']['distances'] = out knn() output

knng = KnnG(anndata, n_neighbors=12, runPCA=True, nPC=50)
```

CPU times: user 3min 41s, sys: 1.06 s, total: 3min 42s Wall time: 3min 36s

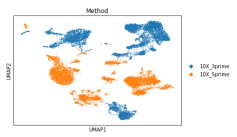
```
In [4]: 
# umap() reduce results to 2 deminsions so that we can plot the data
sc.tl.umap(anndata)
```

CPU times: user 16.3 s, sys: 147 ms, total: 16.4 s Wall time: 13.9 s

1.c. [5 pts] Turn in a UMAP plot of your 12-NN graph calculated from the combined chemistry PBMC dataset colored by batch (the chemistry used)

```
In [5]: # color can be any thing in obs
plt.figure(figsize=(10,10))
scanpy.pl.umap(anndata, color=['Method'])
```

<Figure size 720x720 with 0 Axes>



1.d. [5 pts] Turn in another UMAP plot of your 12-NN graph calculated from the combined chemistry PBMC dataset but colored by cell type

