Library selection demo

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```
In [1]: from os import listdir
       from os.path import join
       from fnmatch import fnmatch
       import pickle
       import pandas as pd
       import numpy as np
       from scipy.stats import gaussian_kde
       from sklearn.decomposition import IncrementalPCA
       from sklearn.manifold import TSNE
       from rdkit import DataStructs
       from rdkit import Chem
       from rdkit.Chem import AllChem
       from rdkit.Chem import Descriptors
        from rdkit.Chem import Draw
       from rdkit.Chem.Fingerprints import FingerprintMols
       from matplotlib import pyplot as plt
       import py3Dmol
        from tqdm import tqdm_notebook as tqdm
```

```
import warnings
        warnings.filterwarnings('ignore')
In [2]: def draw3D(m,p=None,confId=-1):
            draw chemical structures using 3D stick depictions
            adapted from http://rdkit.bloqspot.com/2016/07/a-recent-post-on-in-pipeline-talked.html
            mb = Chem.MolToMolBlock(m,confId=confId)
            if p is None:
                p = py3Dmol.view(width=400,height=400)
            p.removeAllModels()
            p.addModel(mb,'sdf')
            p.setStyle({'stick':{}})
            p.setBackgroundColor('0xeeeeee')
            p.zoomTo()
            return p.show()
        def gaussianKDE(x, y):
            kernel density estimation for coloring scatter plots
            see https://docs.scipy.org/doc/scipy/reference/generated/scipy.stats.gaussian_kde.html
            xy = np.vstack([x,y])
            z = gaussian_kde(xy)(xy)
            return z
        def load_library(library):
            read structure files with .sdf and .smi extensions
            if library.split('.')[-1] == 'sdf':
                mols = Chem.SDMolSupplier(join('data', library.split('/')[-1]))
            elif library.split('.')[-1] == 'smi':
```

```
mols = Chem.SDMolSupplier(join('data', library.split('/')[-1]))
else:
    print('ERROR: please provide a path to a molecule data file in valid .smi or .sdf format')
return mols
```

1 Simplified Molecular Input Line Entry System (SMILES)

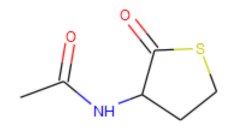
1.1 Advantages:

• Memory efficient: relatively small data files

1.2 Disadvantages:

- Only capable of encoding 2D structure information
- Can only include one identifier for each compound (i.e. no other molecule attributes)

```
In [3]: mols = Chem.SmilesMolSupplier(join('data', 'Maybridge_Ro3_Diversity_Set1.smi')) # read SMILES file
    m = mols[1] # second molecule
    print(Chem.MolToSmiles(m)) # SMILES pattern for second molecule
    Draw.MolToImage(m) # draw 2D structure
CC(=0)NC1CCSC1=0
Out[3]:
```



Structure Data File (SDF)

2.1 Advantages:

- Capable of encoding 3D coordinatesCan include as many properties for each molecule as desired

2.2 Disadvantages:

• Larger files (require $10 - 100 \times \text{more memory}$) In [4]: mols = Chem.SDMolSupplier(join('data', 'Maybridge_Ro3_Diversity_Set1.sdf')) # read SD file (coordinates are 2D) m = mols[1] # second molecule print(Chem.MolToMolBlock(m)) # 2D block for second molecule RDKit 2D 10 10 0 0 0 0 0 0 0 0999 V2000 0 0 4.6250 -5.5000 0.0000 C 0 0 0 0 0 0 5.3417 -4.2542 0.0000 N 0 0 0 4.6292 -4.6750 0.0000 C 0 0 0 0 0 0 3.9167 -5.9167 0.0000 S 0 0 0 0 0 0 0 0 0 0 6.0542 -4.6625 0.0000 C 0 0 0 0 0 0 0 5.3417 -5.9125 0.0000 0 0 0 0 0 0 0 0 0 0 0 6.0667 -5.4875 0.0000 🛭 0 0 0 0 0 0 0 0 0 3.2042 -4.6667 0.0000 C 0 0 0 0 0 0 0 0 0 3.2042 -5.4917 0.0000 C $\ \, 0 \ \, 0$ 6.7667 0.0000 C 0 0 0 0 0 0 0 0 0 0 0 -4.2417 2 3 1 0 3 1 1 0 4 1 1 0 5 2 1 0 6 1 2 0 7 5 2 0 8 3 1 0 9 4 1 0 10 5 1 0 8 9 1 0 M END

```
In [5]: draw3D(m) # show 3D depiction
```

2.3 2D molecules can be converted to 3D using RDKit

- 3D structure coordinates are needed to calculate 3D molecular descriptors (see normalized principle moment ratio and plane of best fit later on)
- After 3D coordinates are generated, they can be saved in SDF format

RDKit ЗD 19 19 0 0 0 0 0 0 0 0999 V2000 -1.3912 -0.7729 1.3471 C 0 0 0 0 0 0 0.8149 -0.2635 0.4168 N 0 0 0 -0.5829 0.4393 C 0.1418 0.7518 S -3.0360 -0.9679 1.7695 0.5369 -0.1621 C -0.9751 2.3644 0 -1.3091 0 0 0 -0.7817 O 1.5013 1.5607 0 0 -1.2598 0.0308 -0.9249 C 0 0 0 0 -0.6829 C -2.7563 0.1070 0 0 0 0 0 3.1829 0.0334 C 0.0599 1.1099 -0.9218 1.1331 H 0 0 0 0 -0.6496 0.8313 H 1.1641 0 -0.9370 0.8272 -1.6038 H 0 0 0 0 0 0 0 0 0 -1.0022 -0.9206 -1.4107 H 0 0 0 0 0 0 0 -0.4334 H -3.0643 1.1273 0 0 0 0 0 0 0 0 -3.3289 -0.2370 -1.5482 H 0 0 0 0 0 0 0 0 3.6838 0.7108 0.7545 H 0 0 0 0 0 0 0 0 0

```
3.2128 -0.9696
                  0.4012 H 0 0 0 0 0 0 0 0
                 -0.9251 H 0 0 0 0 0 0 0 0 0 0 0
   3.7081
          0.0959
 2 3 1 0
 3 1 1 0
 4 1 1 0
 5 2 1 0
 6 1 2 0
 7 5 2 0
 8 3 1 0
 9 4 1 0
 10 5 1 0
 8 9 1 0
 2 11 1 0
 3 12 1 0
 8 13 1 0
 8 14 1 0
 9 15 1 0
 9 16 1 0
 10 17 1 0
10 18 1 0
10 19 1 0
M END
```

In [7]: draw3D(m2) # show 3D representation

2.4 Additional molecular properties are stored for each molecule after the end of the structure coordinates (M END) and before the compound delimiter (\$\$\$\$)

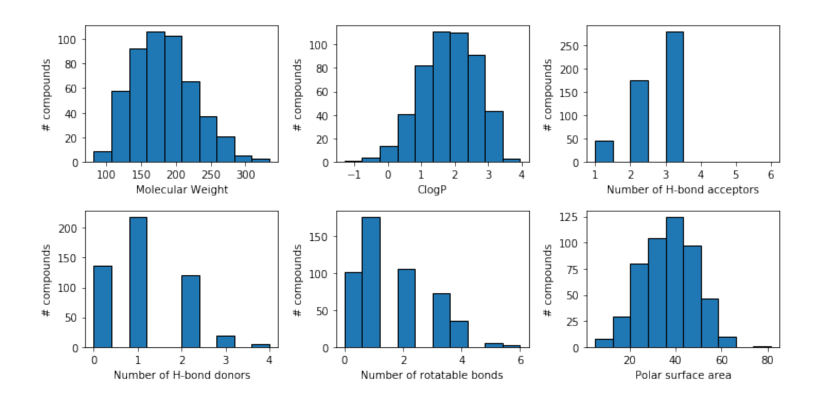
```
'Appearance',
'casno',
'product_name',
'c_log_p',
'acd_code',
'Flexibility',
'PSA',
'h_bond_donors',
'h_bond_acceptors',
'parent_mw',
'Heavy_Atom_Count',
'PAINS_Free',
'Set']
```

3 2D descriptors:

- 3.1 A popular school of thought holds that fragment libraries should contain molecules with the following attributes:
 - Molecular Weight < 300 g/mol
 - $ClogP \le 3$
 - Number of hydrogen bond donors ≤ 3
 - Number of hydrogen bond acceptors ≤ 3
 - Number of rotatable bonds ≤ 3
 - Polar surface area < 60 Å

```
'Number of H-bond donors' : zeros, 'Number of rotatable bonds' : zeros,
                               'Polar surface area' : zeros})
            df.name = library
            for x, tq in zip(xrange, tqdm(xrange, desc='calculating...')):
                try:
                    df['Molecular Weight'][x] = Descriptors.MolWt(mols[x])
                    df['ClogP'][x] = Descriptors.MolLogP(mols[x])
                    df['Number of H-bond acceptors'][x] = AllChem.CalcNumLipinskiHBA(mols[x])
                    df['Number of H-bond donors'][x] = AllChem.CalcNumLipinskiHBD(mols[x])
                    df['Number of rotatable bonds'][x] = Descriptors.NumRotatableBonds(mols[x])
                    df['Polar surface area'][x] = Descriptors.TPSA(mols[x])
                except:
                    pass
            return df
       def plot_descriptors(library):
           plot histograms for Ro3 descriptors from a collection of molecules
           df = calc_descriptors(library)
           fig, ax = plt.subplots(2, 3, figsize=(10, 5))
           c = 0
           for i in range(len(ax)):
               for j in range(len(ax[0])):
                    desc = list(df)[c]
                    ax[i][j].hist(df[desc], edgecolor='k', label=df.name)
                    ax[i][j].set_xlabel(desc)
                    ax[i][j].set_ylabel("# compounds")
                    c += 1
            plt.tight_layout()
           plt.show()
           plt.close()
In [10]: library = 'data/Maybridge_Ro3_Diversity_Set1.sdf' # Note: can change this to any other structure file
```

plot_descriptors(library)



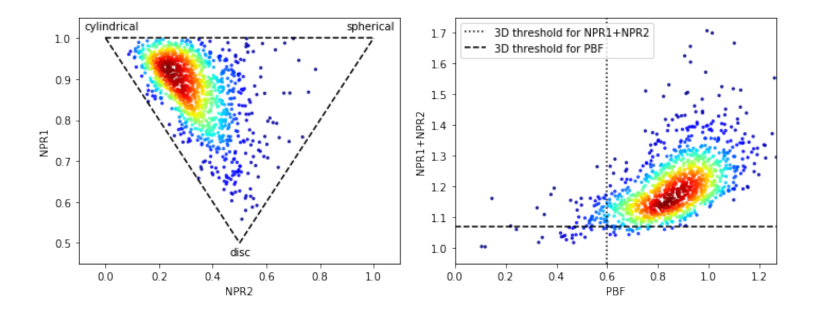
4 3-dimensionality (vs. flat structures) as a strategy to increase library diversity

4.1 Plots quantifying molecule 3D character depicted below

- 2-dimensional normalized ratios of principle moments of inertia
- Plane of best fit (PBF) score versus sum of NPRs

```
In [11]: def calc_3D_descriptors(library):
             Calculate 3D descriptors from collection of molecules
            mols = load_library(library)
            xrange = range(len(mols))
             zeros = np.zeros(len(mols))
             df = pd.DataFrame({'NPR1' : zeros, 'NPR2' : zeros, 'PBF' : zeros})
             df.name = library
            for x, tq in zip(xrange, tqdm(xrange, desc='calculating...')):
                 try:
                     m2 = Chem.AddHs(mols[x])
                     AllChem.EmbedMolecule(m2)
                    AllChem.MMFFOptimizeMolecule(m2)
                    df['NPR1'][x] = AllChem.CalcNPR1(m2)
                     df['NPR2'][x] = AllChem.CalcNPR2(m2)
                     df['PBF'][x] = AllChem.CalcPBF(m2)
                 except:
                     pass
             return df
        def plot_3D_descriptors(library):
             111
             Create plots to assess molecule 3D character
            df = calc_3D_descriptors(library)
            fig, ax = plt.subplots(1, 2, figsize=(10, 4))
```

```
ax[0].scatter(df.NPR1, df.NPR2, c=gaussianKDE(df.NPR1, df.NPR2), cmap='jet', s=5, alpha=0.8)
            ax[0].set_ylim([0.45, 1.05])
            ax[0].set_xlim([-.1, 1.1])
            ax[0].plot(np.linspace(0, 1, 100), np.repeat(1, 100), c='k', linestyle='--')
            ax[0].plot(np.linspace(0, 0.5, 100), np.linspace(0, 0.5, 100)*-1+1, c='k', linestyle='--')
            ax[0].plot(np.linspace(0.5, 1, 100), np.linspace(0.5, 1, 100)*1, c='k', linestyle='--')
            ax[0].text(-.08, 1.02, 'cylindrical')
            ax[0].text(0.90, 1.02, 'spherical')
            ax[0].text(0.465, 0.47, 'disc')
            ax[0].set_xlabel('NPR2')
            ax[0].set_ylabel('NPR1')
            ax[1].scatter(df.PBF, df.NPR1 + df.NPR2, c=gaussianKDE(df.PBF, df.NPR1 + df.NPR2),
                           cmap='jet', s=5, alpha=0.8)
            ax[1].set_ylim([0.95, 1.75])
            ax[1].set_xlim([0, np.max(df.PBF)])
            ax[1].plot(np.repeat(0.6, 100), np.linspace(0.95, 1.75, 100), c='k', linestyle=':',
                      label='3D threshold for NPR1+NPR2')
            ax[1].plot(np.linspace(0, np.max(df.PBF), 100), np.repeat(1.07, 100), c='k', linestyle='--',
                      label='3D threshold for PBF')
            ax[1].set_xlabel('PBF')
            ax[1].set_ylabel('NPR1+NPR2')
            ax[1].legend()
            plt.tight_layout()
            plt.show()
            plt.close()
In [12]: library = 'data/Enamine_3D_Shaped_Diverse_Fragments.sdf' # Note: can change this to any other structure file
         plot_3D_descriptors(library)
```



5 Fingerprinting and molecular similarity

5.1 click here for a more detailed explanation of the theory behind fingerprints

• In general, molecular substructures are converted into bitmaps that can be compared mathematically for similarity

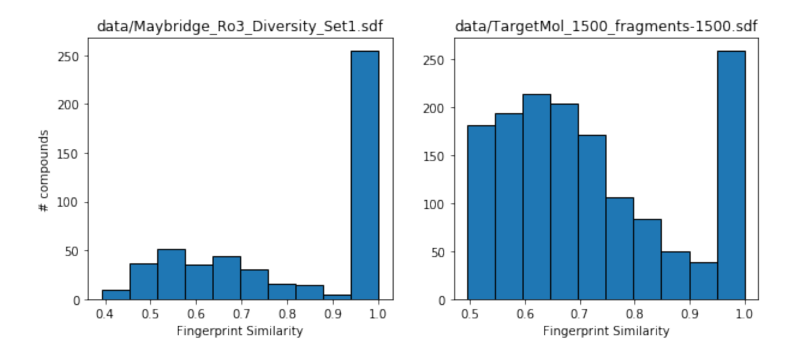
5.2 Comparing nearest-neighbor similarity between two libraries

- Example use case: would combining two different compound libraries lead to substantial redundancy?
- In the example shown below, each compound fingerprint in one library is compared to each compound fingerprint in another and the highest pair-wise similarity score is recorded for each molecule

- Here, scoring ranges from 0.0 (no similarity) to 1.0 (structurally identical)
- E.g. if a compound is associated with a score of 1.0, this means the same compound exists in both libraries

```
In [13]: def calc_nearest_neighbors(fpsA, fpsB):
             Calculate lists of nearest-neighbor similarities between to compound collections
             molsA = load_library(libraryA)
             molsB = load_library(libraryB)
             fpsA = []
            for x in range(len(molsA)):
                 try:
                     fpsA.append(FingerprintMols.FingerprintMol(molsA[x]))
                 except:
                     pass
             fpsB = []
            for x in range(len(molsB)):
                 try:
                     fpsB.append(FingerprintMols.FingerprintMol(molsB[x]))
                 except:
                     pass
            nnA = np.zeros(len(fpsA))
            for x, tq in zip(range(len(fpsA)), tqdm(range(len(fpsA)), desc='calculating A...')):
                 similarity = 0.
                for y in range(len(fpsB)):
                     tmp = DataStructs.FingerprintSimilarity(fpsA[x], fpsB[y])
                     if tmp > similarity:
                         similarity = tmp
                     nnA[x] = similarity
            nnB = np.zeros(len(fpsB))
             for x, tq in zip(range(len(fpsB)), tqdm(range(len(fpsB)), desc='calculating B...')):
                 similarity = 0.
                 for y in range(len(fpsA)):
                     tmp = DataStructs.FingerprintSimilarity(fpsB[x], fpsA[y])
```

```
if tmp > similarity:
                         similarity = tmp
                     nnB[x] = similarity
             return nnA, nnB
         def plot_nearest_neighbors(libraryA, libraryB):
             plot nearest-neighbor similarities between two compound collections
             nnA, nnB = calc_nearest_neighbors(join('data', libraryA), join('data', libraryB))
             fig, ax = plt.subplots(1, 2, figsize=(10, 4))
             ax[0].hist(nnA, edgecolor='k')
             ax[0].set_title(libraryA)
             ax[0].set_ylabel('# compounds')
             ax[0].set_xlabel('Fingerprint Similarity')
             ax[1].hist(nnB, edgecolor='k')
             ax[1].set_title(libraryB)
             ax[0].set_xlabel('Fingerprint Similarity')
             ax[1].set_xlabel('Fingerprint Similarity')
             plt.show()
             plt.close()
In [14]: libraryA = 'data/Maybridge_Ro3_Diversity_Set1.sdf' # Note: can change this to any other structure file
         libraryB = 'data/TargetMol_1500_fragments-1500.sdf' # Note: can change this to any other structure file
         plot_nearest_neighbors(libraryA, libraryB)
```



6 2D depictions of molecular fingerprints:

- The dimensionality of fingerprint bitmaps (here we used 2048-bit Morgan fingerprints) can be reduced using statistical techniques like Principle component analysis (PCA) to allow for visualization and comparison of library diversity.
- Here we fit a pca model on a set of ~348 million fragments (MW \leq 350 g/mol) downloaded from the ZINC database
- The model (loaded here as 'ipca.pkl') is then used to project 2048 bit Morgan fingerprints calculated from compound libraries onto two dimensions for visual comparison of diversity.

```
In [15]: ipca = pickle.load(open('ipca.pkl', 'rb'))
```

```
def decompose_mols(library):
    calculate Morgan fingerprints for each molecule in a collection and
   reduce 2048 bit fingerprints to 20 principle components using a pre-fit incremental PCA model
   mols = load_library(library)
   bits = []
   for x in mols:
        try:
            bits.append(AllChem.GetMorganFingerprintAsBitVect(x, 2))
   X = np.zeros((len(bits), len(bits[0])))
   for i, b in enumerate(bits):
       X[i] = [int(x) for x in b]
   x = ipca.transform(X)
   return x.T
def plot_pcs(libraryA, libraryB):
   plot 2D molecular fingerprint depictions to compare two compound collections
   XA, XB = decompose_mols(libraryA), decompose_mols(libraryB)
   xmin, xmax = np.min([np.min(XA[0]), np.min(XB[0])]), np.max([np.max(XA[0]), np.max(XB[0])])
   ymin, ymax = np.min([np.min(XA[1]), np.min(XB[1])]), np.max([np.max(XA[1]), np.max(XB[1])])
   fig, ax = plt.subplots(1, 2, figsize=(10, 4))
   ax[0].scatter(XA[0], XA[1], c=gaussianKDE(XA[0], XA[1]), cmap='jet', s=5, alpha=0.8)
   ax[0].set_ylabel('PC2')
   ax[0].set_xlabel('PC1')
   ax[0].set_xlim([xmin, xmax])
   ax[0].set_ylim([ymin, ymax])
   ax[0].set_title(libraryA)
```

```
ax[1].scatter(XB[0], XB[1], c=gaussianKDE(XB[0], XB[1]), cmap='jet', s=5, alpha=0.8)
ax[1].set_xlabel('PC1')
ax[1].set_xlim([xmin, xmax])
ax[1].set_ylim([ymin, ymax])
ax[1].set_title(libraryB)
plt.tight_layout()
plt.show()
plt.close()
```

