CalculatePartialChargesWithPsi4

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0.1 Calculate (RESP) partial charges with Psi4

import the following modules

```
[29]: import os
import psi4
import resp
import openbabel as ob
from rdkit import Chem
from rdkit.Chem import AllChem
```

0.1.1 some helper functions

```
[30]: def neutralize_atoms(mol):
          pattern = Chem.MolFromSmarts("[+1!h0!$([*]~[-1,-2,-3,-4]),-1!
       \Rightarrow$([*]~[+1,+2,+3,+4])]")
          at_matches = mol.GetSubstructMatches(pattern)
          at_matches_list = [y[0] for y in at_matches]
          if len(at_matches_list) > 0:
              for at_idx in at_matches_list:
                  atom = mol.GetAtomWithIdx(at idx)
                  chg = atom.GetFormalCharge()
                  hcount = atom.GetTotalNumHs()
                  atom.SetFormalCharge(0)
                  atom.SetNumExplicitHs(hcount - chg)
                  atom.UpdatePropertyCache()
          return mol
      def cleanUp(psi4out_xyz):
          deleteTheseFiles = ['1_default_grid.dat','1_default_grid_esp.dat','grid.

dat','timer.dat']

          deleteTheseFiles.append(psi4out_xyz)
          for fileName in deleteTheseFiles:
              if os.path.exists(fileName):
                  os.remove(fileName)
      def get_xyz_coords(mol):
          if not mol is None:
```

```
num_atoms = mol.GetNumAtoms()
        xyz_string=""
        for counter in range(num_atoms):
            pos=mol.GetConformer().GetAtomPosition(counter)
            xyz_string = xyz_string + ("%s %12.6f %12.6f %12.6f\n" % (mol.
 →GetAtomWithIdx(counter).GetSymbol(), pos.x, pos.y, pos.z) )
    return xyz_string
def calcRESPCharges(mol, basisSet, method, gridPsi4 = 1):
    options = {'BASIS_ESP': basisSet,
               'METHOD_ESP': method,
               'RESP_A': 0.0005,
               'RESP_B': 0.1,
               'VDW_SCALE_FACTORS':[1.4, 1.6, 1.8, 2.0],
               'VDW_POINT_DENSITY':int(gridPsi4)
    }
    resp_charges = resp.resp([mol], [options])[0][1]
    return resp_charges
```

0.1.2 Set some variables and stuff

```
[32]: method = 'b3lyp'
basisSet = '3-21g' #'6-31G**'
neutralize = True
psi4.set_memory('10 GB')
obConversion = ob.OBConversion()
obConversion.SetInAndOutFormats("xyz", "mol2")
singlePoint = True
path = "./data"
```

0.1.3 Read sdf file (3D) into a list

```
[24]: inputFile = "./data/twoCpds.sdf"
molList = Chem.SDMolSupplier(inputFile, removeHs=False)
```

0.1.4 ...or read a SMILES files into a list

```
[38]: SMILESasInput = False

if SMILESasInput:
    molList = []
    inputFile = "./data/twoCpds.smi"
    suppl = Chem.SmilesMolSupplier(inputFile, titleLine = False)

for mol in suppl:
```

```
mol = Chem.AddHs(mol)
AllChem.EmbedMolecule(mol)
try:
     AllChem.MMFFOptimizeMolecule(mol)
except:
     AllChem.UFFOptimizeMolecule(mol)
molList.append(mol)
```

0.1.5 Loop over compounds in list and calculate partial charges

```
[39]: for mol in molList:
          if not mol is None:
              molId = mol.GetProp(" Name")
              print('Trying:', molId)
              if neutralize:
                  mol = neutralize_atoms(mol)
                  mol = Chem.AddHs(mol)
              xyz_string = get_xyz_coords(mol)
              psi_mol = psi4.geometry(xyz_string)
              ### single point calculation
              outfile_mol2 = inputFile[:-4]+".mol2"
              if singlePoint:
                  print('Running singlepoint...')
                  resp_charges = calcRESPCharges(psi_mol, basisSet, method, gridPsi4_
       \rightarrow= 1)
              else:
                  print('Running geometry optimization...')
                  methodNbasisSet = method+"/"+basisSet
                  psi4.optimize(methodNbasisSet, molecule=psi_mol)
                  resp_charges = calcRESPCharges(psi_mol, basisSet, method, gridPsi4⊔
       \rightarrow= 1)
              ### save coords to xyz file
              psi4out_xyz = molId + '.xyz'
              psi_mol.save_xyz_file(psi4out_xyz,1)
              ### read xyz file and write as mol2
              ob_mol = ob.OBMol()
              obConversion.ReadFile(ob_mol, psi4out_xyz)
```

```
### set new partial charges
        count = 0
        for atom in ob.OBMolAtomIter(ob_mol):
             newChg = resp_charges[count]
             atom.SetPartialCharge(newChg)
             count += 1
         ### write as mol2
        outfile_mol2 = path+"/"+molId+"_partialChgs.mol2"
        print("Finished. Saved compound with partial charges as mol2 file: s''
 →% outfile_mol2)
        obConversion.WriteFile(ob_mol, outfile_mol2)
         ### clean up
        cleanUp(psi4out_xyz)
Trying: TXA
Running singlepoint...
Finished. Saved compound with partial charges as mol2 file:
./data/TXA_partialChgs.mol2
Trying: 4piol
Running singlepoint...
Finished. Saved compound with partial charges as mol2 file:
./data/4piol_partialChgs.mol2
```

[]: