Tutorials of the calculations of Synthetic Accessibility, Lipinski's rule of five and PAINS by MolAICal

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1. Introduction

In this tutorial, the calculations of synthetic accessibility, Lipinski's rule of five and PAINS are introduced. The synthetic accessibility (SA) of compounds can be used to evaluate the novel growth ligands generated by *de novo* method. The Lipinski's rule of five also known as the rule of five (RO5) is a rule to estimate drug-like or determine if a chemical compound has pharmacological or biological activity that would be likely orally active drug¹. Pan-assay interference compounds (PAINS)² are the compounds which often show the false positive results in the biological assay.

2. Materials

2.1. Software requirement

1) MolAICal: https://molaical.github.io

2.2. Example files

1) All the necessary tutorial files are downloaded from:

https://github.com/MolAICal/tutorials/tree/master/009-SA Ro5 Pains

3. Procedure

3.1. SA calculation

You can calculate the SA by SMILES sequence directly.

#> molaical.exe -tool sa -i "FC(F)(F)c1cc(ccc1)N5CCN(CCc2nnc3[C@H]4CCC[C@H]4Cn23)CC5"

Or you can calculate SA of ligands in one file that contains many SMILES sequences.

```
#> cd "009-SA Ro5 Pains/SA"
```

#> molaical.exe molaical.exe -tool sa -i SmilTest.smi

3.2. RO5 calculation

The RO5 of single ligand can be calculated as below:

```
#> cd "009-SA Ro5 Pains/ro5"
```

#> molaical.exe -tool ruleoffive -f mol2 -n zinc 1879871.mol2

If you want to calculate Lipinski's rule of five for many ligands in bulk, you can do it below steps:

1. Using command "ls > mol2List.dat" in Linux console, or "dir /b > mol2List.dat" in DOS console of Window. Open generated file "mol2List.dat" and delete no useful characters. Make sure the file "mol2List.dat" only contains the ligand names.

2. Run command as below:

#> molaical.exe -tool ruleoffive -f mol2list -i mol2List.dat -o result.dat

It will generate the file named "result.dat" which contains RO5 values in bulk. More detail about RO5, please see the manual of MolAICal.

3.3. PAINS calculation

The single ligand with SMILES or mol2 format can be calculated as below:

```
#> cd "009-SA_Ro5_Pains/pains"
#> molaical.exe -tool pains -f smi -n "c1ccccc1N=Nc1ccccc1"
#> molaical.exe -tool pains -f mol2 -n ZINC00154323.mol2
```

If you want to calculate PAINS for many ligands in bulk, you can do it below steps:

For SMILES format:

#> molaical.exe -tool pains -f smilist -i painsTest.txt -o smiResult.txt

For Mol2 format:

```
#> cd "009-SA Ro5 Pains/pains/mol"
```

- 1. Using the command "ls > mol2List.dat" in Linux console, or "dir /b > mol2List.dat" in DOS console of Window. Open generated file "mol2List.dat" and delete no useful characters. Make sure the file "mol2List.dat" only contains the ligand names.
- 2. run command as below:

```
#> molaical.exe -tool pains -f mol2list -i mol2List.dat -o mol2Result.txt
```

It will generate the file named "mol2Result.txt" which contains PAINS information in bulk.

Note: recommend the SMILES format for PAINS calculation.

References

- Lipinski, C. A., Lombardo, F., Dominy, B. W. & Feeney, P. J. Experimental and computational approaches to estimate solubility and permeability in drug discovery and development settings. *Adv Drug Deliv Rev* **46**, 3-26 (2001).
- Dahlin, J. L. *et al.* PAINS in the assay: chemical mechanisms of assay interference and promiscuous enzymatic inhibition observed during a sulfhydryl-scavenging HTS. *J Med Chem* **58**, 2091-2113 (2015).