Chemical Structure Standardisation

The ChEMBL database and the standardiser tool

Francis Atkinson
Chemogenomics Group
EMBL-EBI

What is ChEMBL?

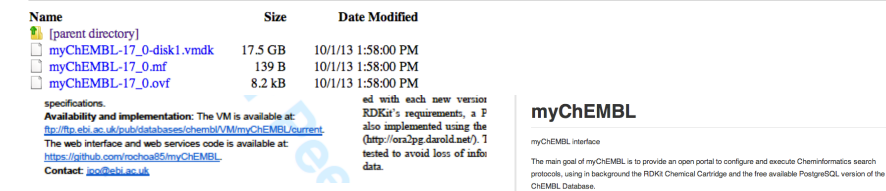
- A freely-available source of bioactivity data
 - Secure on-line access
 - Downloadable in a variety of formats
- Core is data from key MedChem journals
 - J. Med. Chem, Bioorg. Med. Chem. Lett.
 - Manually extracted by CRO
- Supplemented by...
 - Subset of data from PubChem
 - Deposited datasets
 - NTD consortia, GSK kinase set (PKIS) etc.
- Accelrys Direct chemistry cartridge for Oracle

myChEMBL

- Postgres/RDKit version of ChEMBL
- packaged for distribution as VM



Index of /pub/databases/chembl/VM/myChEMBL/releases/myChEMBL-17_0/



Heterogeneous sources of structures

- Chemical Literature
 - Redrawn by CRO from depictions in articles
- PubChem
- Other bioactivity databases
- Data depositions
- Typically received as MDL molfiles

Multiple representations

Hypervalent vs. charge-separated

$$R - N \qquad R - N^{+} \qquad \qquad O^{-}$$

Charged vs. neutral salts

- Chemical cartridge will treat these as different
 - Could cause SSS to fail if alternative depiction used as query
 - Standardization is required

Standardisation

 The FDA have published a compound depiction <u>SOP</u> for their Substance Registration System

This guide is used to standardize the entry of substances into the Food and Drug Administration (FDA)

Substance Registration System (SRS). The primary purpose of this guide is to prevent duplicate entries of a single substance. Conventions for drawing structures and for organizing the characteristics of substances are included. The guide also provides limited aesthetic guidelines for the structures as they are intended to be shared with other databases and may be used in professional publications.

- Used as basis for ChEMBL standardisation rules
 - with some modifications, e.g. ...

8. Salts Formed by the Reaction of Ammonia with Acids

A salt formed by the reaction of ammonia with an acid is represented as the ammonium ion and the conjugate base of the acid.

Example: AMMONIUM LACTATE

ChEMBL would treat these the same as other salts, *i.e.* they would be neutralized

Correct Incorrect

Examples of rules: charges

Consistent rules for drawing salts

- Compounds to be charge-neutral if possible
 - quaternary N an exception if counterion unknown

Example of rules: functional groups

Charge-separated preferred over hypervalent

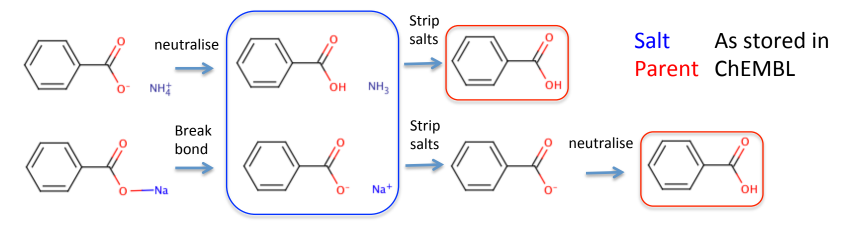
Stylistic rules for sugars, peptides, steroids etc.

Salt stripping

- Salt/solvate components should not affect the biological activity of a compound
 - There can be exceptions
 - e.g. salt form may affect solubility
 - However, any 3D or QSAR modelling should be done using only the bioactive 'parent'
- It is desirable to be able to view compounds with a common parent together
- It can be appropriate to aggregate data for the parent compound

Salt stripping (2)

- Counterions and solvent molecules are removed using a custom 'salt dictionary'
 - Acids with inorganic cations may require a further round of charge-neutralization...



- The parent is registered as a separate structure
 - The parent and original salt are linked via the 'molecule hierarchy' table

Curator intervention

- Structures with permanent charges
 - i.e. counterion not recorded
 - Zero charge used as check on neutralization steps

- Zwitterions
 - Naïve neutralization could introduce errors

- These types automatically routed for inspection
 - Possible manual standardisation

Inorganics & organometallics

- Handled poorly by current chemoinformatics tools...
 - multicenter bonding
 - coordination complexes
 - Non tetrahedral stereochemistry
 - Cannot distinguish cis- and trans-platins



ferrocene?!

- Charge-balancing, salt-stripping etc. are difficult
- Cannot be properly searched for by cartridge
- Not put through main standardisation process
 - May be redrawn for clarity in some cases
 - e.g. approved drugs such as the platins
- Structures now excluded (post-CHEMBL17)
 - ~3200 structures affected
 - Bioactivities are retained

Implementation

- Rules implemented using Pipeline Pilot
 - Run by ChEMBL's chemical curator
 - Some manual intervention in difficult cases
- Protocols not currently publically available
 - Tightly coupled with data-loading pipeline
 - Subject to change as new issues identified
- Pipeline Pilot is commercial software
- We have a interest in an FOSS equivalent...
 - e.g. for loading proprietary data into myChEMBL

standardiser

- Tool to pre-process structures for modelling
 - A component of the eTOX predictive modelling system
- Molecular representation can affect results...
 - Descriptors, forcefields, QM
- Need to standardise representation
 - Common to both training and application
- 'Inspired by' ChEMBL curation & InChI preprocessing
 - Only interested in parent (bioactive) component
 - Standardise tautomers
 - *i.e.* hydroxy-pyridine -> pyridone
 - Not tautomer canonicalization
 - No manual intervention
- Implemented in Python & RDKit
 - i.e. all components are open-source
 - Runs under Python2.6+ (2.6 is a requirement for eTOX project)

Procedure

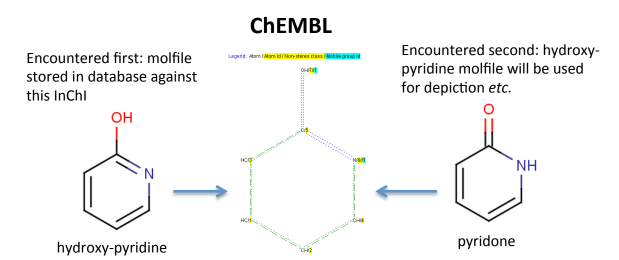
- Break bonds to Group I or II metals
- Neutralize charges by adding/removing protons
- Apply standardization rules
- Neutralize any charges exposed by rules
- Discard any salt/solvate components
- Return standardized parent

Limitations

- Rule set could be expanded
 - e.g. will review RCS / CVSP rules
- No attempt to handle inorganics
 - Impractical with current tools
- No attempt to produce 'canonical' tautomer
 - Could flag tautomeric molecules for inspection?
- re-charging not handled

Key differences to ChEMBL

- ChEMBL makes no attempt to standardise tautomers
- InChI codes are used for registration (i.e. assignment of database identifiers/keys)
 - i.e. two molecules are the same if they have the same InChI
- For any molecule, the first-encountered tautomer will always be used
 - to generate images, for searching and in downloadable SDF files etc.
- Thus, for example: if the first time a molecule encountered it is shown as the hydroxypyridine, this tautomer will be stored (as a molfile) and will be used for the depiction of this molecule even where an alternative tautomer is encountered in a later document
- By contrast, standardiser *does* attempt to standardise tautomers
 - e.g. the pyridone is preferred, as it is likely to be the lower-energy tautomer



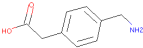
standardiser

InChI=1S/C5H5NO/c7-5-3-1-2-4-6-5/h1-4H,(H,6,7)

Example

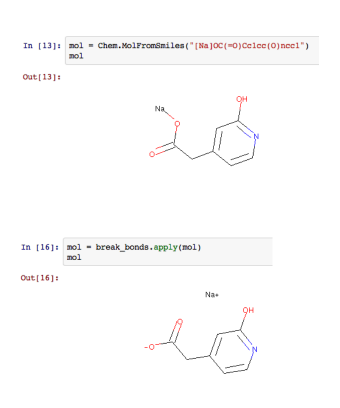
- Provided as a Python package
 - Compatible with python2.6 (for etoxws VM)
- A simple driver program for batch processing is included
 - Accepts SDF or SMILES input

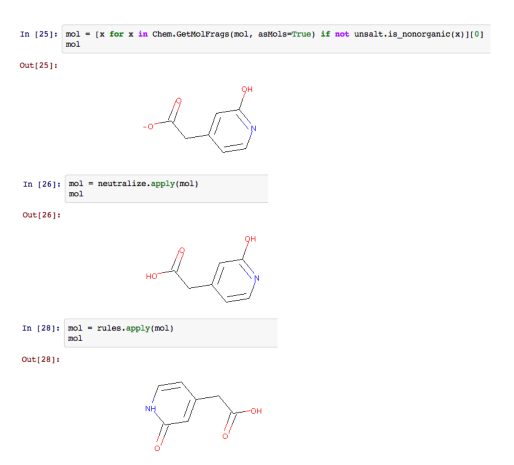
```
In [2]: from standardise import standardise
Examples
 In [3]: mol = Chem.MolFromSmiles("[Na]OC(=0)Cclccc(C[NH3+])ccl.clnnn[n-]1.0")
 Out[3]:
 In [4]: parent = None
          try:
              parent = standardise.apply(mol)
          except standardise.StandardiseException as e:
             logging.warn(e.message)
 Out[4]:
```



Modules within package

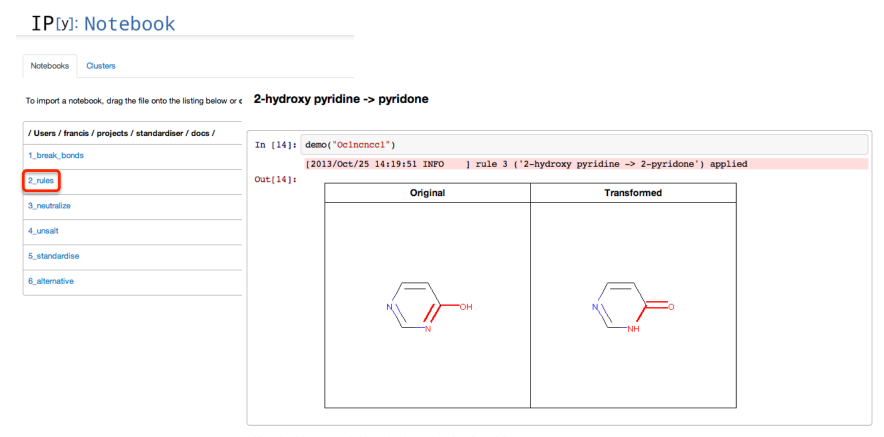
- Modules implementing different steps may be called independently
 - could be incorporated into different workflows





Documentation

- Provided as IPython Notebooks
 - Also available as static web <u>pages</u>

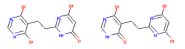


Note that it is the 2-pyrimidone that is produced preferentially.

Futher Documentation

- Also included are some pages describing issues encountered
 - Intended to stimulate debate on future directions of tool

Issue 01 The molecule below is a simplified version of real examples. It illustrates a problem with the original (very simple) version of the algorithm. While the code has been fixed to cope with cases like this, it should be noted that it is not really clear how often problems of this nature would occur in practice, and thus how much the extra complexity needed to deal with them is warranted. It would also be possible to deal with this issue by rewriting the SMARTS transforms to be more specific (i.e. by excluding cases such as the present one). However, tt was felt that this would likely end up being more compolicated than the chosen method, as mutiple complicated SMARTS could become necessary to represent fairly simple transforms. In [2]: mol = Chem.MolFromSmiles("Oclnc(CCc2c(0)ncnc20)nc(0)c1") Out[2]: In [109]: rxn = AllChem.ReactionFromSmarts("[OX2H1:1]-[c:2]:[nX2:3]>>[OH0:1]=[c:2]:[nH1:3]") # 2-hydroxy pyridine -> 2-pyridone In [111]: # Run transform... # NB As the transorm is a one-component reaction, only the first component from each product tuple will exist products = [x[0] for x in rxn.RunReactants((mol,))] # Sanitize product mols... for x in products: Chem.SanitizeMol(x) # Keep unique products (i.e. tautomers) only products = {Chem.MolToSmiles(x): x for x in products}.values() # Depict... Draw.MolsToGridImage(products)



Out[111]:

Futher Documentation

Out[28]:

- The tool will be run on diverse test sets and the results posted
 - Again, designed to detect flaws and promote discussion
 - One example up so far...

Check the bahaviour of the standardiser by running on ChEMBL parent structures

Note that as these are ChEMBL parent structures, they will already have been through ChEMBL's normalization pipeline. The interest here is thus in what changes when the standardiser is run. Recall that ChEMBL uses InChIs to register structures, so tautomer standardisation is unnecessary. By contrast, the goal here is to product structures appropriate for e.g. modelling, so it is desirable to 'fix' certain tautomeric forms. This difference in goals accounts for the bulk of the difference observed.

```
In [2]: %run setup.py
In [3]: chembl parents = pd.read table(open('chembl parents.smi', 'r'), header=None, names=['smiles',
         chembl id'])
         chembl parents.set index('chembl id', inplace=True)
In [4]: standardised = pd.read_table(open('standardised.smi', 'r'), header=None, names=['smiles', 'chem
          standardised.set index('chembl id', inplace=True)
In [25]: len(standardised) # Out of 10000
Out[25]: 9994
In [5]: # Merge standardised with originals...
         merged = chembl parents.join(standardised, how='inner', lsuffix=' old', rsuffix=' new')
In [12]: # Keep only those that changed...
         merged["changed"] = merged.loc[:, "smiles old"] != merged.loc[:, "smiles new"]
         changed = merged[merged.changed == True]
         del changed["changed"]
```

In [28]: not_hydroxy_pyridine mol old mol new chembl id CHEMBL570855 CHEMBL532261 This isn't good. CHEMBL1186335

Future developments

- We would like to open-source this code
 - Tool would improve as more bugs/issues are found
 - e.g. establish wiki for community annotation of rules
 - Need to tidy up code and documentation
- More testing on diverse compound sets
- Incorporate into registration tool for myChEMBL?
 - Longer term project!
 - recall differences between standardisation for modelling and for database registration...