Computational Chemistry: Why, What & How?

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Computational Chemistry: Why?

What are the advantages and disadvantages of using computers in molecular sciences?

**Advantages:**
- Cost (1 cpu hour = £0.02).
- Feasibility (synthetic route to compound may be unknown).
- Safety (no COSHH forms!).
- May obtain extra information that you cannot ‘see’ experimentally.

**Disadvantages:**
- Limitations in accuracy.
- Tendency to use computer as a ‘black box’ (i.e. not understand what it is doing).

**Goal of this talk** is to help you to understand a) what sort of problem can be accurately addressed and b) how you can get involved.
Computational Chemistry: What?

**Atomistic modelling** (cf structure-based design)

We know the atomic coordinates, aim to use the laws of physics to calculate molecular properties.

*Lauren Nelson, Chris Ringrose, Ben Cree*

Quantitative structure-activity relationships *(QSAR)*
(cf ligand-based design)

Molecular properties may be too difficult to compute accurately, so we may wish to identify correlations between chemical structures and properties.

*Rachael Pirie, Matthew Roberts*
Structure-Activity Relationships

Quantitative structure-activity relationships (QSAR) (cf ligand-based design)

How to compute chemical/structural similarity is an open research question.

We are looking at techniques from differential geometry to efficiently describe and compare 3D shape.

*Rachael Pirie, Dr Stuart Hall*
Atomistic Modelling

Classical mechanics
fast to run, large system sizes, not very accurate

Quantum mechanics
very accurate, small system sizes, very expensive to run
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Not much available in this space!
Foundations of Quantum Mechanics

In the famous double-slit experiment, electrons (or other sources) are fired through two closely-spaced parallel gaps and detected on a screen.

In the wave picture, the sources interfere (like water waves) and produce a series of light and dark patches. In the classical picture, only two light patches are expected.

http://weelookang.blogspot.co.uk/2011/10/ejs-open-source-double-slit-diffraction.html
 Remarkably, even when the electrons are sent through one at a time, we see a wave-like interference pattern on the screen.

 Profoundly, this tells us that the electrons are not acting like classical particles (e.g. snooker balls). We cannot know exactly where the electron is, only where it is likely to be found (its probability density).

 If we are going to model how electrons behave in molecules (the foundations of all of chemistry), we need a theory that is different to the equations of classical mechanics….quantum mechanics.
Foundations of Quantum Mechanics

The Nobel Prize in Physics 1933 was awarded jointly to Erwin Schrödinger and Paul Dirac “for the discovery of new productive forms of atomic theory”.

Assuming that matter (e.g., electrons) could be regarded as both particles and waves, in 1926 Erwin Schrödinger came up with a wave equation that accurately calculated the energy levels of electrons in atoms.

“The Nobel Prize in Physics 1933”. Nobelprize.org. Nobel Media AB 2014. Web. 25 Jan 2018. <http://www.nobelprize.org/nobel_prizes/physics/laureates/1933/>
Solution of the Schrödinger equation tells us the wave function (where the electrons are) and the total energy of a configuration of atoms.

Goal is to find $E$ and $\psi$, such that action of $\hat{H}$ on $\psi$ returns $E\psi$. 
Quantum Mechanics: Case Study
CDK2 Inhibition

What experimental observation motivated this investigation?

Compelling evidence has recently been found to support a therapeutic role for pharmacological CDK2 inhibition in cancer treatment.

Compound 27 has the potential to form hydrogen bonding interactions with CDK2, as well as $\pi-\pi$ stacking.

However only shows 10% inhibition of CDK2 at 100 $\mu$M concentration.

It was hypothesised that a purine-phenyl twist of 40 degrees would optimise packing.
CDK2 Inhibition

What QM methods are used?

QM dihedral scans with a range of $R_1$, $R_2$ substituents are run.

What useful information does QM provide?

29: $R_1 = \text{Me}$, $R_2 = \text{H}$

Optimal ortho substitution was synthesised and shown to be relatively potent against CDK2 ($IC_{50} = 18 \, \mu\text{M}$).
Biological Applications

Transition state searching in enzymes

Protein-ligand binding in metalloproteins

Optical spectroscopy in a light-harvesting protein

Classical force field parameterisation for drug discovery

“Applications of Large-Scale DFT in Biology”, D. J. Cole & N. D. M. Hine
Journal of Physics: Condensed Matter, 28, 393001 (2016)
Molecular Mechanics

Molecular mechanics (MM) is widely used in computer-aided drug design, protein folding, protonation states, protein-surface interactions, allosteric mechanisms, photochemistry, QM/MM....

MM can complement experiment in many ways. At the simplest level, we can use molecular dynamics to ‘animate’ the system.

Simulations tend to be limited only by finite sampling and accuracy of the force field.
Force Field

\[ E_{Total} = \sum_{\text{Bonds}} K_r (r - r_0)^2 + \sum_{\text{Angles}} K_\theta (\theta - \theta_0)^2 + \sum_{\text{Torsions}} \frac{V_n}{2} \left[ 1 + \cos(n\Phi - \gamma) \right] \]

Bonded (Intramolecular) Parameters

\[ + \sum_{\text{Non-Bonded}} 4\varepsilon_{ij} \left\{ \left( \frac{\sigma_{ij}}{r_{ij}} \right)^{12} - \left( \frac{\sigma_{ij}}{r_{ij}} \right)^6 \right\} + \frac{q_i q_j}{r_{ij}} \]

Non-Bonded (Intermolecular) Parameters

An open and collaborative approach to better force fields

https://github.com/qubekit/QUBEKit
In lead optimisation studies, we are typically interested in optimising the target-ligand binding affinity.

In other words, we need to find the free energy difference between a small molecule (L1) in solution and bound to the protein (R).

Free energy perturbation (FEP) theory provides a rigorous means to compute the binding free energy.
If we have two similar molecules, then often we only need to compute the relative binding free energy $\Delta \Delta G$.

$$\Delta \Delta G = \Delta G_2 - \Delta G_1$$
The total free energy change around a closed loop is zero:

\[ \Delta \Delta G = \Delta G_2 - \Delta G_1 = \Delta G_B - \Delta G_A \]

Free energy changes computed using Zwanzig equation:

\[ \Delta G_A = -kT \ln \left< \exp \left[ -\frac{(U_{L2} - U_{L1})}{kT} \right] \right>_L \]

We can use FEP to transform molecule L1 into molecule L2 in the protein and in water. Conformational sampling performed using force field.
Aurora A kinase plays a central role in cell division. It is oncogenic and over-expressed in various tumour types.

Inhibition of Aurora A leads to cell death in dividing cells and it is a potential drug target in cancer.

It has recently been shown that an allosteric surface site can be targeted. Interruption of the protein-protein interaction with TPX2 reduces kinase activity.
Used FEP to investigate small substitutions around the phenyl ring. Replica exchange with solute tempering (REST) enhanced sampling of ligand and L178.

Cole et al., Chem. Commun. 2017, 53, 9372
Cole, Tirado-Rives & Jorgensen, JCTC 2014, 10, 565
Computer-Aided Drug Design

\[
\begin{array}{cccccc}
X & Y & Z & \Delta \Delta G^a & IC_{50}^b & K_i^b \\
1 & H & H & 1.05 & 289 & 62.5 \\
2 & F & H & 0.00 & 75.9 & 16.5 \\
3 & F & H & -0.94 & 36.0 & 7.8 \\
4 & Cl & H & -0.73 & ND & ND \\
5 & Cl & H & -0.89 & 20.5 & 4.4 \\
6 & Br & H & -0.49 & 25.6 & 5.5 \\
7 & CF_3 & H & 0.11 & 26.5 & 5.7 \\
8 & CH_3 & H & 1.12 & ND & ND \\
9 & F & CH_3 & -0.49 & 42^c & 8.7^c \\
10 & Br & CH_3 & -0.90 & 11.1^c & 2.3^c \\
\end{array}
\]

\(^a\) kcal mol\(^{-1}\). \(^b\) \(\mu\)M. \(^c\) This work (see the ESI).

RMSE = 0.32 kcal/mol

D. J. Cole et al., *Chem. Commun.* 2017, 53, 9372
M. Janecek et al., *Scientific Reports* 2016, 6, 28528
Atomistic Modelling

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Not much available in this space!
Gaussian Approximation Potential

Gaussian approximation potential (GAP) writes the potential energy as a generic function of atomic coordinates. Trained using QM energies and forces.

\[ E(\mathcal{A}) = \sum_{\mathcal{B} \in M} x_{\mathcal{B}} K(\mathcal{A}, \mathcal{B}) \]

- Target energy of conformation A (represented by vector of interatomic distances)
- Squared exponential kernel (similarity function between A and B)
- Unknown coefficients (regularised least squares regression)

Bartók, Payne, Kondor, Csányi, Phys. Rev. Lett. 2010, 104, 136403
Bartók, Csányi, Int. J. Quantum Chem. 2015, 115, 1051-1057
Gaussian Approximation Potential

Small molecule GAP trained using QM energies/forces from ~3000 configurations.

Errors on 900 configurations from Monte Carlo simulations:

GAP RMS error = 1 kcal/mol

OPLS RMS error = 4 kcal/mol

DJ Cole, L Mones, G Csányi, Faraday Discuss., 2020, in press
Implemented in MCPRO molecular modelling software.

Intermolecular interactions computed using OPLS force field (similar to QM/MM).

Simulated in 3 environments, bound to two proteins and in water.

Good agreement with crystal structure for kinase, but propose alternative structure for hydrolase that is consistent with x-ray data.

DJ Cole, L Mones, G Csányi, *Faraday Discuss.*, 2020, in press
Free energy calculations

It would be preferable to compute the free energy of binding using the GAP, but this is still too expensive at the moment.

Instead we can use FEP to compute the GAP correction to MM free energy of binding ($\Delta\Delta G$):

$$\Delta\Delta G / \text{kcal/mol}$$

| Kinase  | 1.0 |
|--------|-----|
| Hydrolase | 2.0 |

$$E_L = (1 - \lambda)E_{GAP} + \lambda E_{MM}$$

DJ Cole, L Mones, G Csányi, *Faraday Discuss.*, **2020**, in press
Computational Chemistry: How?

Start by learning how to code!

A programming language is an interface between humans and computers.

The **python** programming language is very popular at the moment as it has a very intuitive syntax:

```python
In [46]: raining = True
day = "Saturday"
temperature = 22

if day == "Saturday" and temperature > 15 and not raining:
    print("Go for a walk")
else:
    print("Stay indoors")

Stay indoors
```

Lots of online courses (e.g. udemy, codecademy), welcome to ask me or Rachael for CHY1610 material.
Computational Chemistry: How?

Quantitative structure-activity relationships (QSAR) (cf ligand-based design)

TeachOpenCADD

A teaching platform for computer-aided drug design (CADD) using open source packages and data

⚠️ We are introducing major changes to the TeachOpenCADD repository very soon. Before making them available, we are cutting a 1.3.0 version with the current changes on master. This will be last release in the v1.x series. Subscribe to the repository releases (top-right menu: Watch > Releases) to stay tuned!

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volkamerlab.org

TeachOpenCADD Jupyter Notebooks

DOI 10.1186/s13321-019-0351-x
DOI 10.5281/zenodo.2600902
.launch binder

TeachOpenCADD KNIME Workflows

DOI 10.1021/acs.jcheminf.9b00662
DOI 10.5281/zenodo.3626897
KNIME Hub TeachOpenCADD-KNIME

Rachael Pirie, Matthew Roberts

https://github.com/volkamerlab/teachopencadd

“launch binder” run tutorials on the cloud from anywhere
Computational Chemistry: How?

Atomistic modelling (cf structure-based design)

Bit harder to get into without training, though part of Ben’s project is to build & deploy tutorials/workflows like the TeachOpenCADD one.

Otherwise come and talk to us!

Lauren Nelson, Chris Ringrose, Ben Cree
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Mike Payne (Cambridge)
Julien Michel (Edinburgh)
Open Force Field Initiative
\[
\hat{H}\psi = E\psi
\]

\[
E_U[\hat{n}^{I\sigma}] = \sum_{I\sigma} U^{I} \frac{1}{2} \text{Tr}[\hat{n}^{I\sigma}(1 - \hat{n}^{I\sigma})]
\]

\[
n_a(r) = \frac{w_a(r)}{\sum_k w_k(r)} n(r)
\]

\[
[k_{AB}] = -\begin{bmatrix}
\frac{\partial^2 E}{\partial x_A \partial x_B} & \frac{\partial^2 E}{\partial x_A \partial y_B} & \frac{\partial^2 E}{\partial x_A \partial z_B} \\
\frac{\partial^2 E}{\partial y_A \partial x_B} & \frac{\partial^2 E}{\partial y_A \partial y_B} & \frac{\partial^2 E}{\partial y_A \partial z_B} \\
\frac{\partial^2 E}{\partial z_A \partial x_B} & \frac{\partial^2 E}{\partial z_A \partial y_B} & \frac{\partial^2 E}{\partial z_A \partial z_B}
\end{bmatrix}
\]

https://blogs.ncl.ac.uk/danielcole/
https://github.com/cole-group/

@ColeGroupNCL

Thank you for your attention