Session 4: Introduction to in silico docking

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This document should also be available online at http://sbcb.bioch.ox.ac.uk/users/greg/teaching/docking-2012.html.

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- 1. Introduction

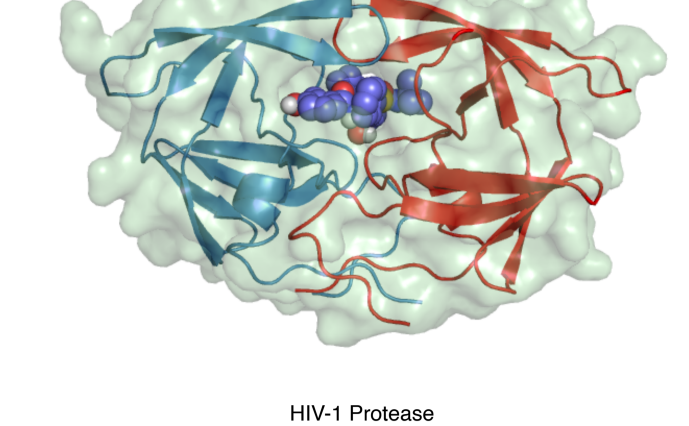
Why is docking important?

The purpose of this practical is to give a flavour of how one could dock small flexible molecules to a protein structure. Such in silico methods are extremely useful for both finding potential binding sites and also to discover

A case study: HIV-1 protease

and/or engineer new molecules that could bind to a known site. This is a multi-billion dollar industry. Virtual screening and blind docking are often employed in an attempt to discover new medicines.

The HIV-1 protease [1] is an enzyme that is vital for the replication of HIV. It cleaves newly formed polypeptide chains at appropriate locations so that they form functional proteins. Hence, drugs that target this protein could be vital for suppressing viral replication. A handful of drugs - called HIV-1 protease inhibitors (saguinavir, ritonavir, nelfinavir, etc.) [2] - are currently commercially available that inhibit the function of this protein, by binding in catalytic site that binds the polypeptide.



We will first look at the protein structure (PDB ID: 1HSG) [3] used in the molecular dynamics practical in a bit more detail. We will then try to dock couple of drug molecules into the binding site to see how well docking can

What does this practical cover?

reproduce the binding pose. If time permits, we will generate the electrostatic surface of the protein to study how the drug interacts with the protein. Make sure the tools work

Anything that starts with the symbol % should be run in a Linux terminal/command prompt and PyMOL> should be run in a PyMOL command line.

i. All the files you need for this practical can be found at http://sbcb.bioch.ox.ac.uk/users/greg/teaching/docking-2012.tar.gz. Download and save it to your Desktop. (Note: To do this, you might have to right click the link, select "Save Link As..." and make sure the folder in

"Save in folder" is Desktop. Click Save) ii. On your Desktop, right click the downloaded file docking-2012.tar.gz and select Extract here. This should create a directory called dock-prac on your Desktop. iii. Open a Terminal and make sure you have access to the following commands. Some of these may open up a graphical user interface (GUI). Close it and proceed to the next step. If you get an error or "command not

found" inform a demonstrator. % cd ~/Desktop % vina --version % apbs --version % pdb2pqr.py --help

% pymol -c % adt -h 2. Exercise 1: Visualising the protein

% cd ~/Desktop/dock-prac

PyMOL> select indinavir, resn MK1

PyMOL> set sphere scale, 0.4

PyMOL> set depth cue, 0

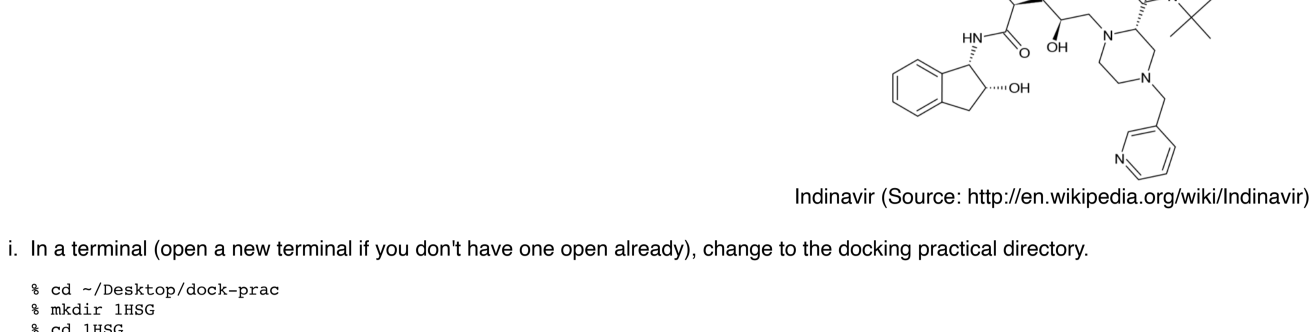
PyMOL> png protein.png

PyMOL> ray

% mkdir 1HSG % cd 1HSG

be clear now.

In this practical, we will use the structure of the HIV-1 protease used in the molecular dynamics practical - PDB ID: 1HSG. This is a 2Å resolution X-ray crystal structure of HIV-1 protease with a bound drug molecule indinavir. We will use pymol to view the protein, the binding site and the drug molecule. pymol options/controls that you learned in the homology modelling practical will be handy here. An introduction to PyMOL can be found here.



ii. Download the protease structure in PDB format (PDB ID: 1HSG) from the Protein Data Bank (http://www.pdb.org). Search using the PDB ID 1HSG. On the results page, click Download Files and select PDB File (Text).

iv. Lets start by hiding everything. Click the **H** (for hide) next to **all** in PyMOL's **Object Control Panel** (the panel on the right with buttons **A**ctions, **H**ide, **S**how, **L**abel and **C**olour) and select **everything**. The screen should

Save it in ~/Desktop/dock-prac/1HSG. If the internet is down you can get a copy from ~/Desktop/dock-prac/backup/data/1HSG.pdb. (Note: Once again you might have to right click, select Save Link As... and browse to

/home/biocomp/Desktop/dock-prac/1HSG. Make sure you set the filename Name to 1HSG.pdb. Click Save) iii. Load the structure into pymol. You should see the protein structure displayed as lines and water molecules as little red crosses. % pymol 1HSG.pdb &

resn MK1 selects the residue MK1, which is indinavir. You should now have an object in the object control panel called (indinavir). Display it as sticks. Click anywhere in the display screen to unselect the selected atoms. vii. Now, rotate (left mouse button), zoom (right mouse button) and move (middle mouse button) the molecule to get an idea where the binding site is. You will need to know where it is for the next exercise. viii. Water molecules have the residue name HOH. Select and display all water molecules as red spheres. If you think the spheres are too big, type

vi. Let's select the ligand indinavir and show it as sticks. In the PyMOL command line (with a "PyMOL > ") type

v. Now show (S) the protein (1HSG) using the cartoon representation and colour (C) by chain to show that it is a homodimer.

xi. Q: Can you think of a way in which the ligand could enter the binding site? xii. If you would like to save a nice image of the protein, ray trace it and save it as an image file. PyMOL> bg color white

ix. Q: Water molecules normally have 3 atoms. Why do we see just one atom per water molecule?

x. Q: There is a conserved water molecule in the binding site. Can you identify this water molecule?

xiii. Close PyMOL by typing guit in PyMOL command window or clicking the x in the PyMOL Tcl/Tk GUI window. xiv. Using eog, you can visualise the image you just generated. % eog protein.png

Docking algorithms require each atom to have a charge and an atom type that describes its properties. However, the PDB structure lacks these. So, we have to prep the protein and ligand files to include these values along with the atomic coordinates. Furthermore, for flexible ligand docking, we should also define ligand bonds that are rotatable. All this will be done in a tool called AutoDock Tools (adt).

Prepare the protein

3. Exercise 2: Preparing the protein and ligand for docking

i. The PDB file (1HSG.pdb) contains protein, ligand and water oxygen atoms. First we have to extract just the protein atoms, which are lines that start with the keyword ATOM. Each protein chain is terminated with a line that starts with **TER**. (If you would like to confirm this, open 1HSG.pdb in a text editor and scroll through the text.)

% adt & iii. Load the protein using File > Read Molecule. Select 1HSG protein.pdb. Click Open.

v. Q: Can you locate the binding site visually?

% egrep "^(ATOM|TER)" 1HSG.pdb > 1HSG protein.pdb

ii. Lauch AutoDock Tools (ADT) using the command adt

Note: In ADT, you can translate the molecule by clicking and holding down the right mouse button while moving the mouse, rotate by clicking and holding down the middle button and zoom in/out by using the scroll wheel of the mouse. iv. Bonds and atoms are shown in white. For better visualisation, colour the structure by atom type - Color > By Atom Type. Click All Geometries and then OK.

vi. Crystal structures normally lack hydrogen atoms. Q: Why? However, hydrogen atoms, or more specifically polar hydrogen atoms are required for appropriate treatment of electrostatics during docking. Add hydrogen

vii. Now we need to get ADT to assign charges and atom type to each atom in the protein. We do this with Grid > Macromolecule > Choose.... Choose 1HSG_protein in the popup window and click Select Molecule. ADT will merge non-polar hydrogens, assign charges and prompt you to save the macromolecule. Click Save. This will create a file called 1HSG_protein.pdbqt in the current folder. Open this in a text editor and look at the last two columns - these should be the charge and atom type for each atom. Q: Look at the charges. Does it make sense? viii. Now, we need to define the 3D search space were ligand docking will be attempted. Remember the binding site that you observed in one of the earlier steps. Ideally, if we do not know the binding site, we will either define a box that encloses the whole protein or perhaps a specific region of the protein. In this case, to speed up the docking process, we will define a search space that encloses the known binding site.

Spacing (angstrom) of 1Å (this is essentially a scaling factor). So set this dial to 1.000. So that we all get consistent results, let us set the (x, y, z) center as (16, 25, 4) and the number of points in (x, y, z)-dimension as (30, 30, 30). Make a note of these values. We will need it later. Close the Grid Options dialog by clicking File > Close w/out saving. x. That is all we need to do with the protein file. Delete it from the display using Edit > Delete > Delete Molecule and select 1HSG protein. Click Delete Molecule and CONTINUE. Prepare the ligand

ix. To define the box, use Grid > Grid Box.... This will draw a box with opposite faces coloured in red, green and blue. Fiddle with the dials and see how you can enclose regions of the protein. In this instance we will use a

i. Like the protein, the ligand lacks hydrogen atoms. We need to add hydrogen atoms and also define rotatable bonds that will be used for flexible docking. ii. First, extract the ligand atoms from the PDB. As mentioned in Exercise 1, the ligand residue name for indinavir in the PDB file is MK1 and the lines start with the keyword HETATM for heteroatoms. In a terminal type % grep "^HETATM.*MK1" 1HSG.pdb > indinavir.pdb

iii. Load the ligand structure into ADT using File > Read Molecule and select indinavir.pdb

iv. Again, colour by atom type. Color > By Atom Type. Click All Geometries and then OK. v. Now we have to add polar hydrogen atoms. Add all hydrogen atoms initially. Non polar hydrogens will be merged in the next step. Edit > Hydrogens > Add. Select All Hydrogens and click OK. vi. Define this as the ligand in ADT so that ADT assigns partial charges and sets rotatable ligand bonds using Ligand > Input > Choose.... Select indinavir and click Select Molecule for AutoDock4. You should see a message that confirms that non-polar hydrogens have been merged, charges added and rotatable bond detected. Click OK. The ligand will now have only polar hydrogens.

atom to the structure using - Edit > Hydrogen > Add. Click OK. You should see a lot of white dashes where the hydrogens were added.

vii. To check the rotatable bonds detected by ADT, go to Ligand > Torsion Tree > Choose Torsions.... You should see 14 rotatable bonds. viii. Q: What does the three colours - green, magenta and red - mean? Does it make sense? ix. Click **Done** when you are done. x. Save the ligand file in PDBQT format. Do this using **Ligand > Output > Save as PDBQT...**. Click **Save** to save the file as indinavir.pdbqt.

xi. Quit ADT using **File > Exit** and then **OK**. Prepare a docking configuration file

i. Before we can perform the actual docking, we need to create an input file that defines the protein, ligand and the search parameters. We will create the input file in a text editor.

% gedit & iii. The input file should look something like:

ii. If you have used Unix/Linux before, open your favourite text editor - vi, emacs, nano - or use a GUI based editor called gedit.

receptor = 1HSG protein.pdbqt ligand = indinavir.pdbqt $num\ modes = 50$

out = all.pdbqt

center x = XXcenter y = XXcenter z = XXsize x = XXsize y = XXsize z = XXseed = 2009This defines your protein (receptor), ligand, number of docking modes to generate (num_modes). All the docked modes will be collated in a file defined by out (all.pdbqt). You should replace the XX with the center of your 3D search space ($center_x/y/z$) and the size of the box ($size_x/y/z$) that you defined in the "Prepare the protein" section. iv. Save the file as config.txt in the folder containing the protein and ligand .pdbqt files.

i. For this practical, we will use a program called **Autodock Vina** [4] for docking. Autodock Vina is a fast docking alogorithm that requires minimal user intervention. We will run it from a terminal. ii. Make sure you are in the folder containing 1HSG protein.pdbgt, indinavir.pdbgt and config.txt iii. Run vina to perform the docking. We will keep a log of all the program output in a file log.txt % vina --config config.txt --log log.txt

This will take a few minutes depending on how fast your computer is. While you wait, if you are interested, read [1] for a review on HIV-1 protease structure, function and drug discovery. iv. Once the run is complete, you should have a two files allipdbgt, which contains all the docked modes, and logitxt, which contains a table of calculated affinities based on AutoDock Vina's scoring function [4]. The best docked mode, according to AutoDock Vina, is the first entry in all.pdbgt. v. Lets visualise the docks and compare it to the crystal conformation of the ligand. Load the protein, the extracted ligand and the docks into pymo1.

5. Exercise 4: Docking nelfinavir into HIV-1 protease

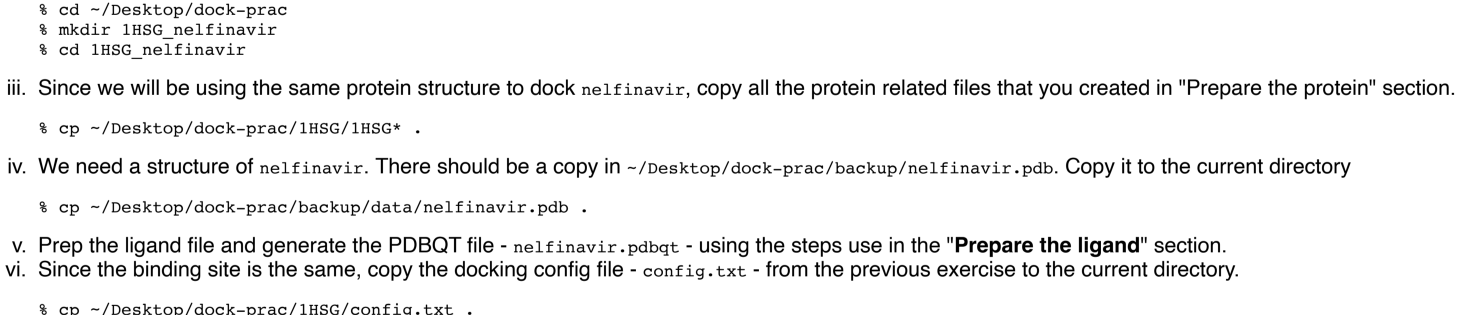
4. Exercise 3: Docking indinavir into HIV-1 protease

% pymol 1HSG protein.pdb indinavir.pdb all.pdbgt & vi. Begin by hiding everything. Now, display the protein in cartoon representation, indinavir in stick representation and the docked conformations as sticks as well. vii. Cycle through the docks by clicking the playback control buttons on the lower right hand corner of the window.

viii. Q: Qualitatively, how good are the docks? ix. Q: Is the crystal binding mode reproduced? If not or if there is a difference, what could be the reason? Is it the best conformation according to AutoDock Vina? x. Close PyMOL.

i. In the previous exercise, AutoDock Vina should have found a near native docked conformation for the drug indinavir. In this exercise, we will try to dock another drug nelfinavir into the protein structure 1HSG broadly along the lines of how one would dock a small molecule into a target structure. This approach could be refined and extended to the "virtual screening" approach.

Nelfinavir (Source: http://en.wikipedia.org/wiki/Nelfinavir)



xx. You may have observed that the two HIV-1 protease structures are in different orientations in space. To make sense of the docking results, you have to align the two structures. You can use the PyMOL align command

viii. Now you should have everything required by AutoDock Vina to generate a series of docking modes. % vina --config config.txt --log log.txt This will take a few minutes to run... ix. Load the protein and the docked structures and show the protein as cartoon and the docks as sticks?

ii. Start by creating a new directory for this docking run. Your current directory should now be: ~/Desktop/dock-prac/1HSG.

xiii. Q: Based on the literature, what is common among this class of HIV-1 protease inhibitors? xiv. Q: How would you check whether the conformation makes sense? xv. Q: How would you look at it from a "virtual screening" perspective? xvi. Fortunately, in this instance, there are crystal structures of nelfinavir bound to HIV-1 protease inhibitor that you can use to validate your docking results.

to do this. In the PyMOL command line, type:

Colour it by element and pick any colour combination.

% mkdir ~/Desktop/dock-prac/1HSG APBS % cd ~/Desktop/dock-prac/1HSG APBS

7. Concluding remarks

% cp ~/Desktop/dock-prac/1HSG/1HSG_protein.pdb . % cp ~/Desktop/dock-prac/1HSG/indinavir.pdbqt .

v. Launch the PyMOL APBS plugin using Plugin > APBS Tools.

PyMOL> align 10HR, 1HSG_protein

% pymol 1HSG protein.pdb all.pdbqt &

xi. Q: Is it docked in the binding site?

x. Q: Qualitatively, how good are the docks?

% cd ~/Desktop/dock-prac % mkdir 1HSG nelfinavir % cd 1HSG_nelfinavir

% cp ~/Desktop/dock-prac/1HSG/1HSG* .

% cp ~/Desktop/dock-prac/1HSG/config.txt .

% cp ~/Desktop/dock-prac/backup/data/nelfinavir.pdb .

vii. You have to change one entry in the configuration file. Make the change.

xii. Q: Is the best docked conformation of nelfinavir similar to indinavir?

xvii. Use the Protein Data Bank and download the PDB file for 10HR. Save it in the current directory - ~/Desktop/dock-prac/1HSG_nelfinavir. (If the internet is down, there should be a copy in ~/Desktop/dockprac/backup/data/10HR.pdb). xviii. Load this structure into PyMOL using **File > Open...**. Select **10HR.pdb** and click **OK**. xix. Hide all representations of 10HR using H and then everything. Now, display 10HR as cartoon and colour by chain.

10HR should now be overlaid on **1HSG_protein** Note: A copy of the aligned structure can be found in ~/Desktop/dock-prac/backup/data/10HR_aligned.pdb if the align command does not work. xxi. Q: What is the RMSD of the alignment? Is it qualitatively and quantitatively good? xxii. You may have observed that moving the structure around the window is a bit difficult since the origin of the view has been altered when you loaded 10HR.pdb. To reset it, try:

PyMOL> reset xxiii. **nelfinavir** has the residue name **1UN** in **10HR**. Select and display it as sticks. PyMOL> select nelfinavir, 10HR and resn 1UN

xxiv. Q: Is the "best mode" generated by AutoDock Vina similar to the crystal conformation of nelfinavir? If not, why not?

xxv. Q: How about the "second best mode"? Why do you think this is the case? (Hint: Look at the affinities reported in the log file)

xxvi. Q: In this instance we assumed that chain A of 1HSG corresponds to chain A of 1OHR. Is this true always? xxvii. Let us align chain A of 10HR to chain B of 1HSG. PyMOL> align 10HR and chain A, 1HSG protein and chain B

xxix. Close PyMOL. **Exercise 5: Generating the electrostatic surface of the protein**

xxviii. Q: Compare the first two AutoDock Vina docks with the crystal conformation. What does this tell you? What could be a reason for this observation?

By now you might have realised that electrostatics play a very important role in docking molecules. The purpose of this exercise is to generate and view the surface of the HIV-1 protease structure and to see how a ligand interacts with this. i. As we found out earlier, PDB structures do not contain partial charges for the atoms in the structure. Partial charges are crucial for electrostatic calculations. Hence, we need to add that to a PDB file. ii. Create a new directory for this exercise and copy the protein structure - 1HSG protein.pdb into this folder.

iii. PyMOL has a nice plugin for computing and viewing electrostatic surfaces generated using a program called Adaptive Poisson-Boltzmann Solver (APBS) [5]. iv. Open 1HSG protein.pdb in PyMOL. % pymol 1HSG protein.pdb &

vi. On the tab titled Main make sure Use PyMOL generated PQR and PyMOL generated Hydrogens and termini is selected. This adds hydrogens and assigns partial charges and atomic radii to each atom in the PDB file. vii. Switch to the Configuration tab and click Set grid. This defines a grid that encloses the protein (Note: Grid is not displayed on the screen). Set the System Temperature to 300 and set the ion concentration (+1) and (-1) to 0.15, which is equivalent of 0.15M cations and cations. viii. Click Run APBS. This should take a while. During this period, the APBS plugin window will probably be unresponsive and the Run APBS button will appear depressed.

PyMOL> load pymol-generated-PE0.dx x. We now want to display the electrostatic surface of the protein. Start with the default positive and negative threshold. Click Show in Molecular Surface. Make the positive threshold higher and negative threshold lower, click **Update** and observe what happens to the protein surface. xi. Load the ligand indinavir into PyMOL and display it in stick representation.

PyMOL> load indinavir.pdbqt xii. Q: Show/Hide the electrostatic surface and compare with the exposed protein atoms. Does it match up? xiii. Q: Does anything on the protein surface (in the binding site) stand out? xiv. Q: Why is the base of the binding site largely negative? How is it related to catalysis in this case? xv. Q: Does the protein surface charge (or lack thereof) correspond to complementary regions in the ligand?

ix. Once the calculations are done, switch to the Visualization tab and click Update. If this does not load and you get an Unable to open file error, type the following in the PyMOL shell.

In this practical, we have looked at how small molecules could be docked into a protein. This approach is widely used to detect binding sites and also to screen a library of small molecules to find potential drugs that could bind to a known binding site. Scoring functions play an important role in identifying a "good dock" and as such remains an area of active research. Several other considerations are worth noting including water molecules and ions in the binding site, flexibility of binding site residues, etc. Some docking programs (including AutoDock Vina) allow you to define a subset of flexible sidechains. Whilst this permits binding site rearrangement to accommodate distinct ligands, the computational search space increases many fold. Defining water molecules that are important in the binding site also remains an area of active research. In this practical, we have considered only protein-ligand docking. Protein-protein docking is also widely used, but not considered here due to the significantly higher search space that has to be considered. As the number of published and

1. Brik A, Wong CH. "HIV-1 protease: mechanism and drug discovery". Org. Biomol. Chem. (2003) 1:5–14.

4. Trott O, Olson AJ. "AutoDock Vina: improving the speed and accuracy of docking with a new scoring function, efficient optimization and multithreading", J. Comput. Chem. (2009) Online in advance of print. 5. Baker NA, Sept D, Joseph S, Holst MJ, McCammon JA. "Electrostatics of nanosystems: application to microtubules and the ribosome." Proc. Natl. Acad. Sci. (2001) 98:10037-10041.

http://autodock.scripps.edu **AutoDock Tools** http://vina.scripps.edu AutoDock Vina http://www.poissonboltzmann.org/apbs **APBS**

Appendix II: Basic Linux commands provides a "long" list of all files in the current directory in reverse order of time. ls -lrt change directory to the directory 'dir' cd dir

delete (remove) 'file' rm file rename file to newfile mv file newfile cat file

print the current working directory on the screen pwd print the contents of file to the screen

9. Appendix I: Useful links/resources PyMOL http://www.pymolwiki.org/index.php/Main Page

print the contents of file to the screen but with more navigation possible.

unpublished protein structures continue to grow, in silico docking will continue to play an important role in the drug discovery process.

inhibitor of the HIV proteases." J. Biol. Chem. (1994) 269:26344-26348.

more file

8. References 2. Wensing AM, van Maarseveen NM, Nijhuis M. "Fifteen years of HIV Protease Inhibitors: raising the barrier to resistance." Antiviral Res. (2009) Online in advance of print. 3. Chen Z, Li Y, Chen E, Hall DL, Darke PL, Culberson C, Shafer JA, Kuo LC. "Crystal structure at 1.9-A resolution of human immunodeficiency virus (HIV) II protease complexed with L-735,524, an orally bioavailable