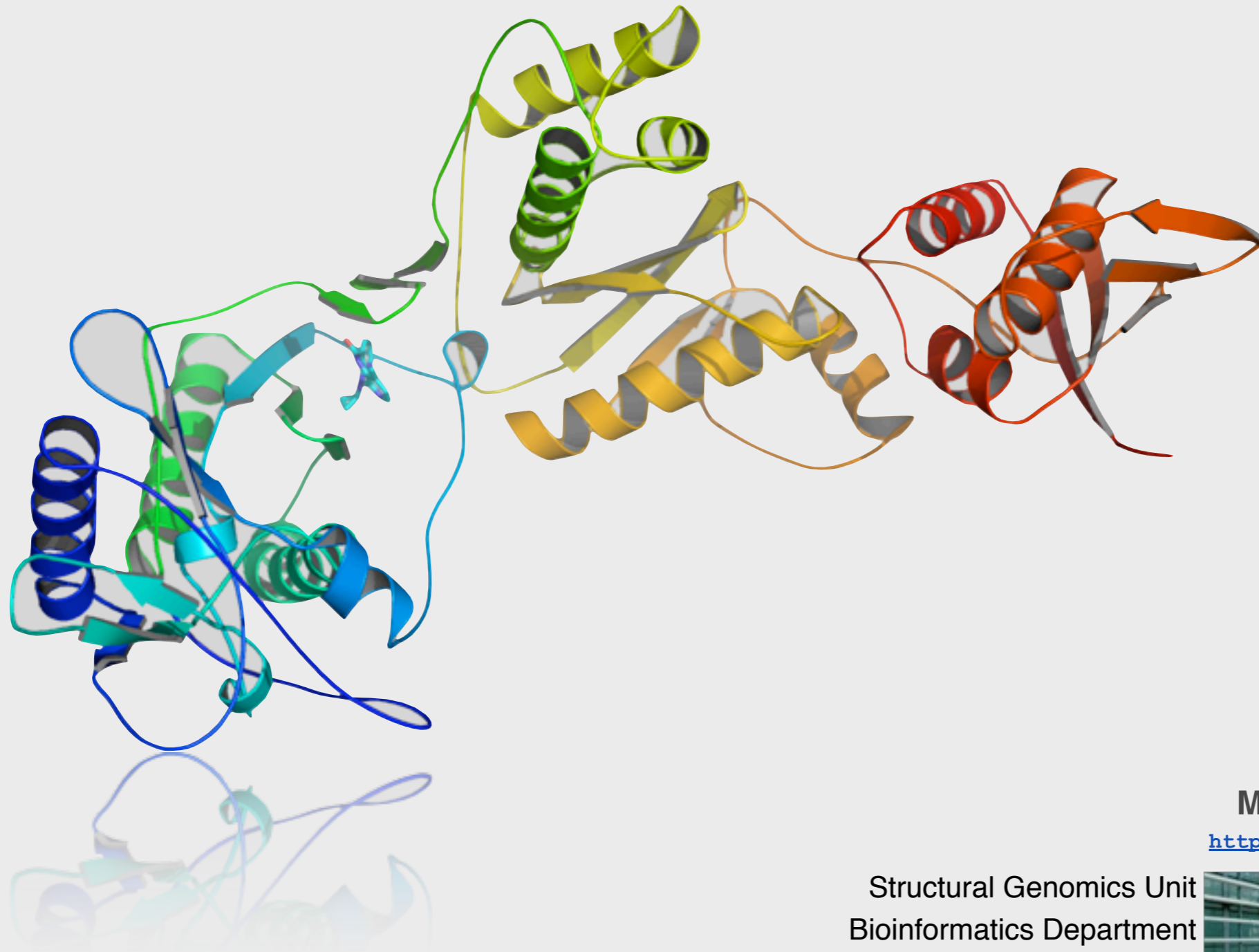


Exercise. SNP-based drug resistance to Nevirapine drug against the HIV reverse transcriptase



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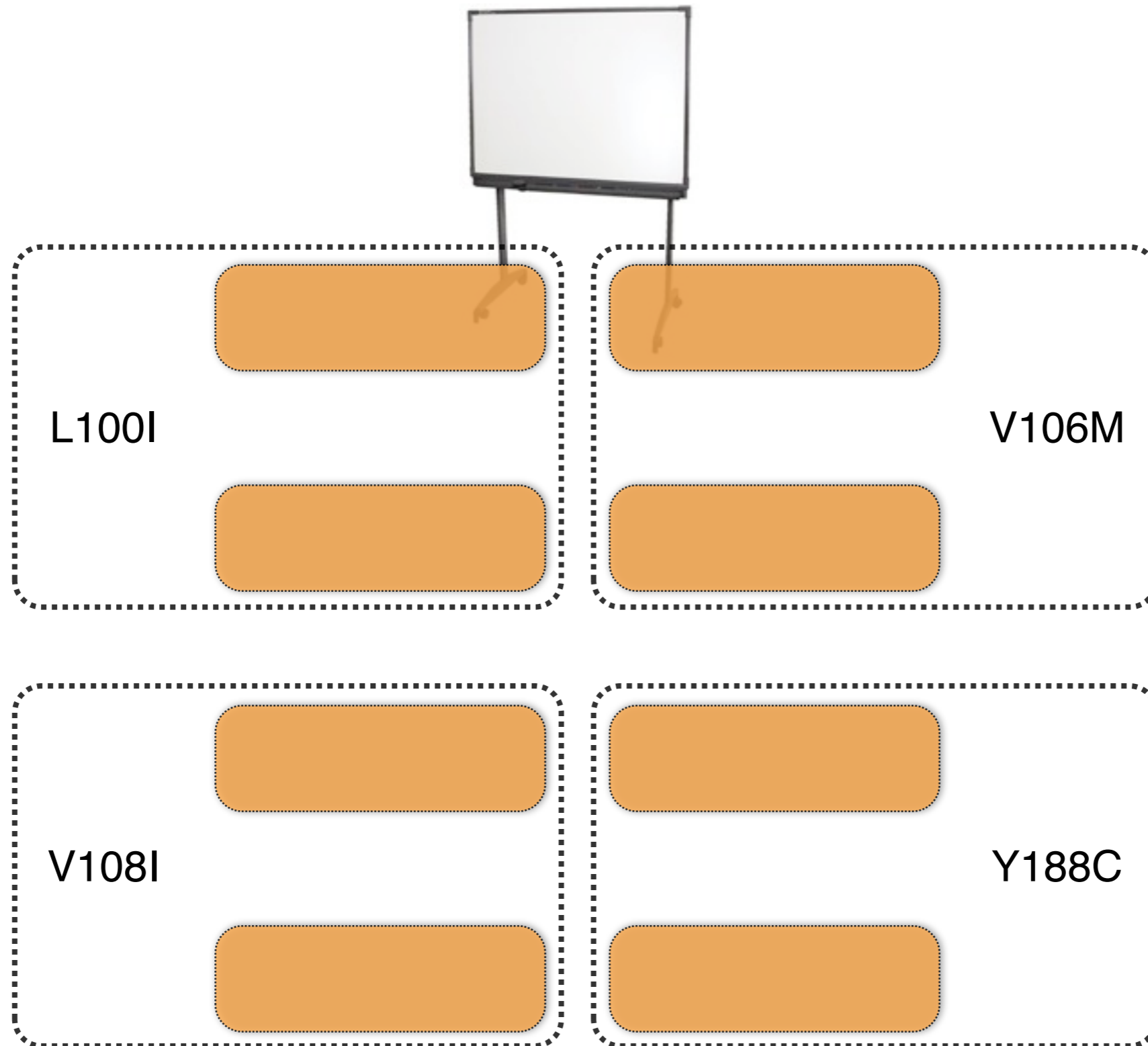
Problem

TO **STUDY** THE EFFECT IN BINDING
OF KNOWN **SNPs** OF HIV REVERSE
TRANSCRIPTASE

TOOLS

- ◆ AnnoLyze (DBAli)
- ◆ PubChem and DrugBank
- ◆ MODELLER
- ◆ Vina and AutoDockTools
- ◆ PyMol

Organization

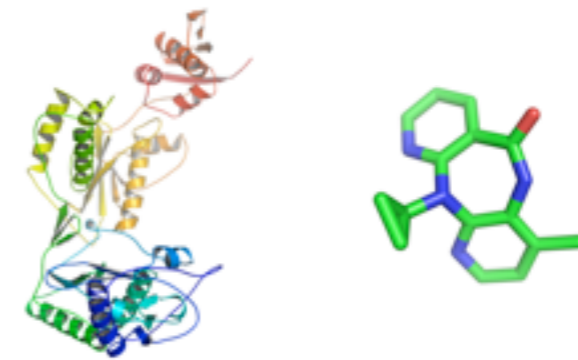


input data and files

Mutation paper



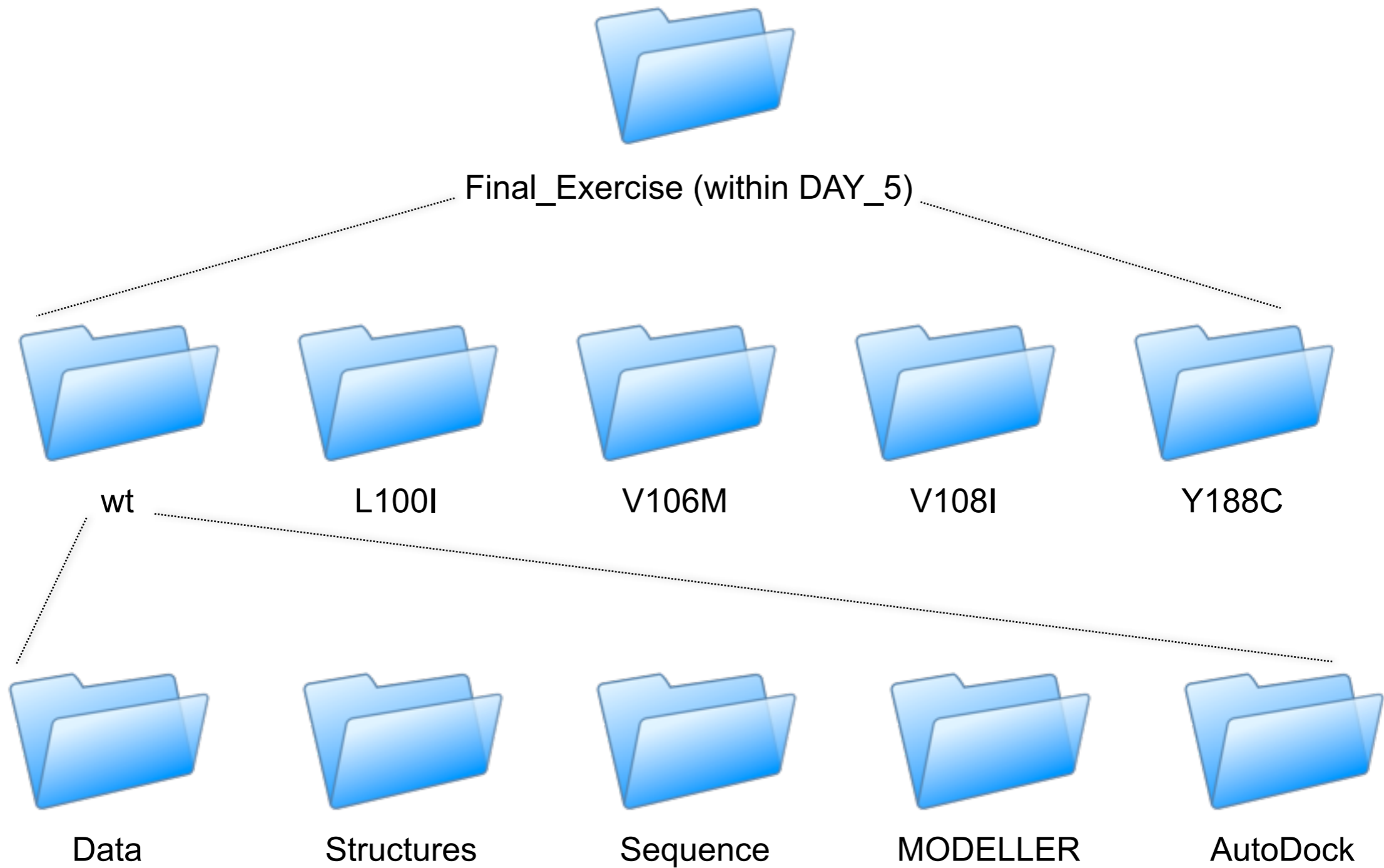
Structure files



Sequence and alignment files

```
>1vruA
PISPIETVPVKLPGMDGPKVKQWPLTEEKIKALVEICTEMEKEGKISKIGPENPNTPVFAIKKKDSTKWRKLVDFREL
NKRTQDFWEVQLGIPHPAGLKKKKSVTVLVDGDYFVPLDEDFRKYTFTIPSIINNETPGIRYQYVNLPGWKGSPAIF
QSSMTKILEPFRKQNPDIIVYQYMDLLVYVSDLEIGQHRTKIEELRQLLRWGLTTPDKKHQKPEPFLWNGYELHPDKWT
VQPVLPEKDSWTVNDIQKLVGKLNWASQIYPGKIVRQLCKLLRGTALTEVIPLTEAELELAENREILKEPVHGVYYD
PSKDLIAEQKQGGQWYIYQEPFNKLTGKYARMRGAHTNDVKQLTEAVQKITTESIVIWGKTPKFKLPIQKETWET
WWTEYWQA TWIPEWEFVNTPPVLKLVQYLEKEPIVGAETFYVDGAANRETKL GKAGYVYVNRGRQKVVTLDTTNQKTELQ
AIYLALQDSGL EVNIVTDSQYALGIIQAQPPDQSESELVNQIEQLIKKEKVVYLAWVPAHKGIGGNEQVDKLVSAGIRKVL
```

Folder organization



Recipe

LIGAND

1. Go to PubChem and look at Nevirapine (NPV). Smile it!. (<http://pubchem.ncbi.nlm.nih.gov>)
2. Divided by groups:
 - a) Get similar compounds with a Tanimoto score larger than 95%. Download the SDF files.
 - b) Do a sub-structure search based on the SMILES. Download the SDF files.
 - c) Do a sub-structure search + filter by molecular weight (200-600Da). Download the SDF files.
 - d) Do a super-structure search + filter by molecular weight (200-400Da). Download the SDF files.

BINDING SITE

1. Run AnnoLyze for the chain 1vruA. (<http://www.dbali.org>)
2. Get predicted binding site to Nevirapine (NVP ligand).
3. Calculate a central point to the ligand using PyMol (see *conf.txt* file under data folder).

COMPARATIVE PROTEIN STRUCTURE PREDICTION

1. Model the 3D structure of the wild-type using its own structure.
2. Model the point mutation for your group.

DOCKING OF SMALL MOLECULES

1. Dock the NVP ligand to the wild-type structure.
2. Dock the NVP ligand to the wild-type model.
2. Dock the NVP ligand to the mutant.

PRESENTATION

1. How would you explain the differences between the wild-type and the point mutant?

and...

A price for the one that can design a point mutation that “stabilizes” the ligand-protein interaction. That is that can find a mutation that gives lower energy scores.