# **CryptoSite Documentation**

Release master

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CryptoSite is a computational tool for predicting the location of cryptic binding sites in proteins and protein complexes. Contents:

# CHAPTER 1

### Installation

#### 1.1 In the Sali Lab

If you are working in the Sali lab, you don't need to build and install CryptoSite - it is already set up for you as a module. Just run module load cryptosite to load it.

### **1.2 Dependencies**

All dependencies listed below are expected to be found in standard system paths. This may require setting PYTHONPATH, PATH and/or LD\_LIBRARY\_PATH environment variables. Note that Linux is the only platform for which all these dependencies are available, and so is the only platform on which CryptoSite currently functions.

- Python 2.6 or later (Python 3 should also be OK).
- MODELLER plus the SOAP-Protein library. The soap\_protein\_od.hdf5 file needs to be placed into MODELLER's modlib directory.
- MUSCLE.
- DSSP. It is expected that the mkdssp binary is in the PATH.
- fpocket (version 2).
- PatchDock.
- IMP.
- NCBI BLAST+ plus a local copy of the UniProt database for it to search against. To make this local database, gunzip the uniprot\_trembl.fasta.gz and uniprot\_sprot.fasta.gz files available from the UniProt website and combine them into a single uniprot text file.
- USEARCH version 8.1 or later.
- Biopython.
- NumPy and SciPy.

- scikit-learn. Note that precisely version 0.12 is needed other versions won't work.
- AllosMod is needed to run part of the protocol.
- nose is also needed to run the test suite (recommended but not essential).

In the Sali lab, running module load modeller muscle dssp fpocket patch\_dock imp blast+ usearch will get all of these dependencies.

## 1.3 Building

Use make test to test the library, and make install to install it. In most cases you will need to tell make where to install (if running on a Linux cluster, CryptoSite will need to be installed on a network-accessible filesystem) and where your local copy of UniProt is, with something like make PREFIX=/shared/cryptosite UNIPROT=/ database/uniprot install. See Makefile.include for all make variables that can be configured.

# CHAPTER 2

### Basic usage

Once installed, the CryptoSite protocol can be run by means of a command line tool (cryptosite). Each component of the protocol is also a Python package, which can be called directly from other Python software (via import cryptosite).

#### 2.1 Overview

Running the basic protocol consists of these steps:

- 1. Create a set of input files specifying the structure to probe, and basic CryptoSite parameters.
- 2. Run cryptosite setup to calculate structural features and prepare input files for AllosMod.
- 3. Run AllosMod given the set of input files.
- 4. Calculate additional features using the AllosMod output (cryptosite soap, cryptosite pockets, cryptosite am\_bmi, cryptosite analysis).
- 5. Gather together all features into a single file (cryptosite gather).
- 6. Predict cryptic binding sites with an SVM using the complete set of features as input (cryptosite predict).
- 7. Optionally visualize the results in Chimera (cryptosite chimera).

# chapter $\mathbf{3}$

Indices and tables

- genindex
- modindex
- search