# **Tutorial**

On this tutorial, we are going to demonstrate some features of the parKVFinder, including the graphical user interface (parKVFinder PyMOL Tools) and the command-line interface.

All files used on this tutorial can be found under input directory, on the parKVFinder directory.

# parKVFinder PyMOL Tools

First, load **input/1FMO.pse** into PyMOL viewer, which loads two objects in your scene. The **1FMO** is a subunit of a protein kinase A and the **ligs\_1FMO** is an adenosine (ADN) and a peptide kinase inhibitor (PKI).



### Default parameters

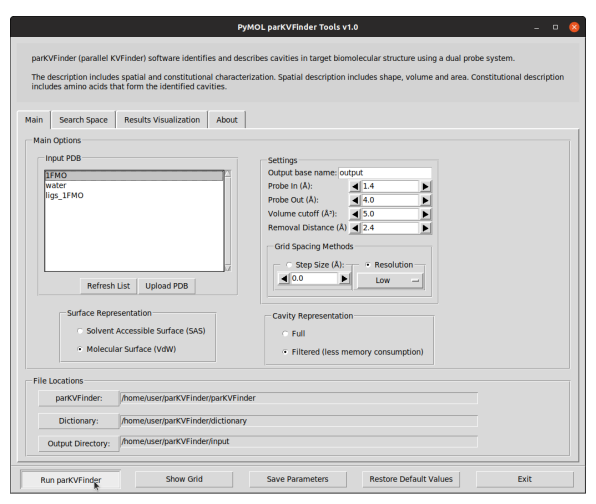
The default parameters are designed to make a simple and fast whole protein prospection.

On PyMOL, open **parKVFinder PyMOL Tools** under **Plugin** tab. The objects on the scene will be listed on the **Input PDB** listbox, on the **Main** tab. If not, press the **Refresh List** 

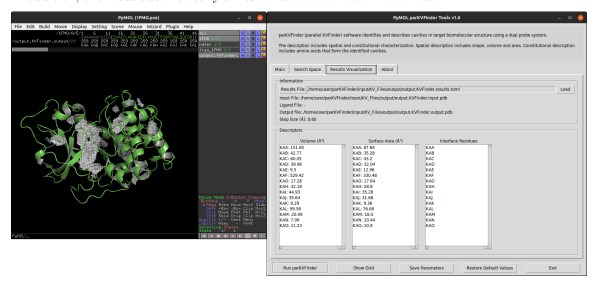
The **Input PDB** selection sets which object will be analyzed by parKVFinder. Select **1FMO** on the listbox.



To run parKVFinder with the default parameters, just click **Run parKVFinder** button or press **Enter**.

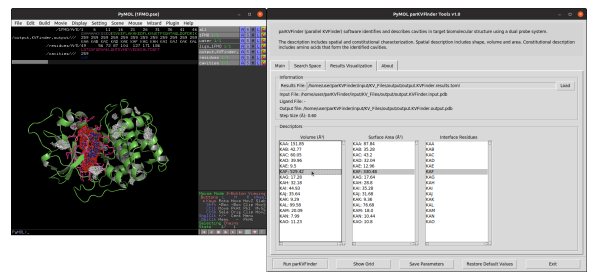


After execution is complete, cavities PDB is loaded into PyMOL viewer as <Output Base Name>.KVFinder.output object and the results file is loaded on the **Results Visualization** tab. In addition, the focus automatically shifts to **Results Visualization** tab.



You can select cavities in the **Volume** or **Surface Area** lists to highlight them on a new object called **cavities**, helping to identify each cavity. Also, you can select cavities in the **Interface Residues** 

list to highlight the residues around the cavities on a new object named residues.



## Changing cavity ceiling

parKVFinder is all about parameter customization. One of parKVFinder's most powerful assets is the ability to manually set the cavity ceiling. parKVFinder works with a double probe system. A smaller probe, called Probe In, and a bigger one, called Probe Out, that defines two molecular surfaces with different molecular accessibility. The space left between these surfaces is considered cavities.

Let's show the effect of varying Probe Out and Removal Distance on the cavity ceiling.

First, we should copy the adenosine to a new object using the following PyMOL commands:

```
select resn ADN create adenosine, sele delete sele
Also,
```

#### Exploring enclosed regions with box adjustment mode

### Ligand adjustment mode

## Command line interface

parKVF inder has a command-line interface, which can be useful for molecular dynamics and high-throughput analysis. It also handles the same parameters available in parKVF inder PyMOL Tools, except for box rotations in box adjustment mode.