parKVFinder Guide

Welcome to the **parallel KVFinder** (*parKVFinder*) wiki, this page was built to help you get started with our cavity detection software.

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Please read and cite the original paper paper title> (<doi>).

parKVFinder Installation Guide

Follow these steps to install parKVFinder software on Linux or macOS:

1. Download parKVFinder source code.

Get the latest parKVFinder release from here.

On the terminal, go to the directory containing **parKVFinder.zip** file.

```
$ unzip parKVFinder.zip
$ cd parKVFinder
```

Or, clone the parKVFinder repository into a new directory using git.

```
$ git clone https://github.com/LBC-LNBio/parKVFinder.git
$ cd parKVFinder
```

2. Compile parKVFinder from source.

parKVFinder supports GCC (GNU Compiler Collection) versions later than 6. If necessary, refer to this guide for installing GCC versions later than 6 on Linux and macOS.

\$ make

3. Create a symbolic link to run parKVFinder consistently.

\$ make link

Note: make link create a soft symbolic link for parKVFinder in your /usr/local/bin directory, using the following command sudo ln -s `pwd`/parKVFinder /usr/local/bin/parKVFinder.

4. Export KVFinder_PATH variable and add to ~/.bashrc file or similar configuration file.

```
$ export KVFinder_PATH=`pwd`
$ echo "export KVFinder_PATH=`pwd`" >> ~/.bashrc
$ source ~/.bashrc
```

Note: Depending on the operating system, other configuration files are used instead of \sim /.bashrc. For example, in macOS \sim /.bash profile are used.

GCC Installation Guide

The installation procedures for GCC (GNU Compiler Collection) are different for Linux and macOS. First check the GCC version on your operating system.

```
$ gcc --version
```

parKVFinder supports GCC versions later than 6. Otherwise, you must install a newer GCC version.

Linux

To install the standard available version of GCC on your operating system, type:

```
$ sudo apt install gcc
```

Note: check the GCC version again to make sure you have installed a version later than 6.

If the default version GCC is not later than 6, you can install directly any later version by typing:

```
# GCC version 6
$ sudo apt install gcc-6
# GCC version 7
$ sudo apt install gcc-7
# GCC version 8
$ sudo apt install gcc-8
# GCC version 9
$ sudo apt install gcc-9
```

Then, there are two possible paths to prepare GCC to compile the parKVFinder source code.

• Create a symbolic link to your custom GCC

```
$ sudo ln -s /usr/bin/gcc-X /usr/local/bin/gcc
```

Note: \mathbf{X} is the version of GCC you have installed.

• Replace gcc to gcc-X in Makefile within the parKVFinder directory

macOS

The macOS native C compiler is **Clang LLVM compiler**, which is not supported by parKVFinder.

First, you need to install:

- *Homebrew* package manager;
- Command Line Tools (CLT) for Xcode (xcode-select --install) or Xcode.

To install the standard available version of GCC on your *Homebrew* package manager, type:

```
$ brew install gcc
```

Note: check the GCC version, using brew info gcc, to make sure you have installed a version later than 6. Also, *Homebrew* installation will not override Clang LLVM compiler link.

If the default version GCC is not later than 6, you can install directly any version by typing:

```
# GCC version 6

$ brew install gcc@6

# GCC version 7

$ brew install gcc@7

# GCC version 8

$ brew install gcc@8
```

GCC version 9
\$ brew install gcc@9

Then, there are two possible paths to prepare GCC to compile the parKVFinder source code.

• Create a symbolic link to your custom GCC

\$ sudo ln -s \$(brew --prefix)/bin/gcc-X /usr/local/bin/gcc

 $Note: \mathbf{X}$ is the version of GCC you have installed.

• Replace gcc to gcc-X in Makefile within the parKVFinder directory

\$ sed -i -e 's/gcc/gcc-X/' Makefile

 $Note:\,\mathbf{X}$ is the version of GCC you have installed.

PyMOL Plug-in Installation Guide

PyMOL is required if you wish to use parKVFinder with a graphical user interface (parKVFinder PyMOL Tools). If necessary, refer to this guide for installing PyMOL on Linux and macOS.

parKVFinder PyMOL Tools is available to use parKVFinder with PyMOL v1.8 and v2.

However, plug-in requires the installation of toml and future modules and native python do not have them installed. So you need to install them:

```
# PyMOL v1.8 (python 2)
$ pip install toml future
# PyMOL v2 (python 3)
$ pip3 install toml future
# In an conda environment via Anaconda Cloud
$ conda install -c conda-forge toml future
```

Note: pip or Anaconda package management system installation is required.

Then, to install the parKVFinder PyMOL Tools on PyMOL, follow these steps:

- 1. Open PyMOL.
- 2. Go to Plugin menu \rightarrow Plugin Manager.
- 3. The Plugin Manager window will open, go to the Install New Plugin tab.
- 4. Under Install from local file Group, click Choose file...
- 5. The **Install Plugin** window will open, go to the **tools** directory at **parKVFinder** directory and select **parKVFinder_PyMOL_tools.py**.
- 6. The Select plugin directory window will open, select /home/<user>/.pymol/startup and click OK.
- 7. The **Confirm** window will open, click **OK**.
- 8. The Sucess window will appear, confirming that the plug-in has been installed.
- 9. Restart PyMOL.
- 10. parKVFinder PyMOL Tools is ready to use.

PyMOL Installation

The installation procedures for PyMOL are different for Linux and macOS.

Linux

The easiest way is to use your distribution package manager.

```
$ sudo apt install pymol
```

Note: PyMOL version varies according to your distribution version.

Or, you can install PyMOL through Anaconda package management system.

PyMOL 1.8

An open source version of PyMOL 1.8 is available from Anaconda. However, a conda environment must have python 2 installed.

Follow these steps:

1. Create a new environment and activate it:

```
$ conda create --name pymol1.8 python=2.7
$ conda activate pymol1.8
```

2. Install PyMOL 1.8 via Anaconda Cloud:

```
$ conda install -c mw pymol
```

3. Try PyMOL 1.8:

\$ pymol

PyMOL 2

A version of PyMOL 2 is available from Anaconda. However, a conda environment must have python 3 installed.

Follow these steps:

1. Create a new environment and activate it:

```
$ conda create --name pymol2 python=3.7
$ conda activate pymol2
```

2. Install PyMOL 2 via Anaconda Cloud:

```
$ conda install -c schrodinger pymol
```

3. Try PyMOL 2:

\$ pymol

macOS

In macOS, PyMOL requires XQuartz installation.

Install Xquartz via *Homebrew*.

\$ brew cask install xquartz

Or.

An XQuartz installer is provided here.

PyMOL 1.8

An open source version of PyMOL 1.8 is available from Anaconda. However, a conda environment must have python 2 installed.

Follow these steps:

1. Create a new environment and activate it:

```
$ conda create --name pymol1.8 python=2.7
$ conda activate pymol1.8
```

2. Install PyMOL 1.8 via Anaconda Cloud:

```
$ conda install -c mw pymol
```

3. Try PyMOL 1.8:

\$ pymol

PyMOL 2

A version of PyMOL 2 is available from Anaconda. However, a conda environment must have python 3 installed.

Follow these steps:

1. Create a new environment and activate it:

```
$ conda create --name pymol2 python=3.7
$ conda activate pymol2
```

2. Install PyMOL 2 via Anaconda Cloud:

```
$ conda install -c schrodinger pymol
```

3. Try PyMOL 2:

\$ pymol

If the options above does not work, you can try an installation via *Homebrew* package manager. A brief description of how to install open source PyMOL for macOS is provided here.

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).



Whole protein detection

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

On PyMOL, open **parKVFinder PyMOL Tools** under **Plugin** tab. The objects on the scene will be listed on the **Input PDB** list box, 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 list box.



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.



We 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, we can select cavities in the **Interface Residues** list





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:

```
# Copy adenosine
select resn ADN
create adenosine, sele
delete sele
```

Also, copy the adenosine cavity (KAF) to a new object (adnsite) to compare the cavity ceiling from the previous execution.

```
# Copy adenosine cavity
select resn KAF and output.KVFinder.output
create adnsite, sele
delete sele
```

Finally, prepare the new PyMOL scene.

```
# Prepare PyMOL scene
color magenta, adnsite
disable
enable (adnsite, adenosine)
```



Adjusting Probe Out

As mentioned above, adjusting the Probe Out size changes the level of the cavity ceiling. So let's go back on the **Main** tab and change the **Probe Out** size to 8.0 Å. Run parKVFinder again.



Again, copy the adenosine cavity (KAF) to a new object (adnsitePO).

Copy new adenosine cavity
select resn KAF and output.KVFinder.output
create adnsitePO, sele
delete sele

Finally, prepare the PyMOL scene.

Prepare PyMOL scene
disable
enable (adenosine, adnsite, adnsitePO)



Note that the adenosine cavity detected with the 4 Å Probe Out (magenta) has a lower ceiling than that detected with the 8 Å probe (white). Therefore, by increasing the size of the Probe Out, the cavity ceiling is also raised.

Adjusting Removal Distance

Besides adjusting the Probe Out size, we can also adjust the Removal Distance to change the cavity ceiling. So let's go back to the **Main** tab and change the **Removal Distance** to 1.2 Å and the size of **Probe Out** back to 4.0 Å. Run parKVFinder again.



Again, copy the adenosine cavity (KAH) to a new object (adnsiteRD).

Copy new adenosine cavity
select resn KAH and output.KVFinder.output
create adnsiteRD, sele
delete sele

Finally, prepare the PyMOL scene.

Prepare PyMOL scene
disable
enable (adenosine, adnsite, adnsiteRD)



Note that the adenosine cavity detected with the 2.4 Å Removal Distance (magenta) has a lower ceiling than that detected with the 1.2 Å (white). Therefore, by decreasing the Removal Distance, the cavity ceiling is also raised.

Furthermore, changing the cavity ceiling by varying Probe Out and Removal Distance also affects cavity segregation.

Note: Usually the Removal Distance adjustment is less time consuming than the Probe Out adjustment for similar effects.

Steered detection

An important feature of parKVFinder is the steered detection of cavities. We continue our tutorial illustrating two distinct methods of cavity segmentation.

Box adjustment mode

Box adjustment mode explores closed regions with a custom box, which can be drawn via the GUI.

On the **Search Space** tab, select **Box Adjustment** option under **Search Procedure** group. This will enable a **Box Adjustment** frame, which handles the custom box in PyMOL viewer.



The custom box is drawn based on the (sele) object in the PyMOL viewer.

Then, select the adenosine ligand. This can be made on the PyMOL viewer by clicking on the ligand structure or using select resn ADN PyMOL command.

Click on **Draw Box** Button. This will create a custom box that limits the search space. It is fully customizable, but we will not change it for now.



On the Main tab, change Removal Distance back to 2.4~Å and rerun parKVFinder.



Now, let's customize the box parameters to segment the binding site of our target protein.

Each axis is associated with one color (red with X, green with Y and blue with Z). The adjustment is made by the arrows or directly setting the value in the entry on the **Search Space** tab in the **Box Adjustment** group. We can also adjust the box angles by the same procedure. After altering the values, just click on **Redraw Box** button to redraw the box object using the new values.

Then, on the **Search Space** tab, reduce **Maximum X** to 1.0 Å and click **Redraw Box**. Rerun parKVFinder.



Lastly, click on **Delete Box** button to delete the custom box.

Ligand adjustment mode

A last feature is to limit the search around a structure. In this last example, let's do a whole protein prospection again, but limiting the search space around ligands.

First, on the **Search Space** tab, select **Whole Protein** option under **Search Procedure** group. This will disable the previous enabled **Box Adjustment** frame.

Still on the **Search Space** tab, click on the check button **Ligand Adjustment**. This will enable the buttons **Refresh List** and **Upload Ligand**.

Click the **Refresh List** button to display all objects in the scene in the **Ligand PDB** list box. Select the adenosine on the list box and reduce **Ligand Cutoff** to 3.0 Å. Run parKVFinder again.



Now, let's shift focus to the two ligands (adenosine and PKI) in the ligs_1FMO object.

On the **Search Space** tab, select the ligs_1FMO on the **Ligand PDB** list box and increase **Ligand Cutoff** back to 5.0 Å. Back on the **Main** tab, increase **Probe Out** to 10.0 Å and reduce **Removal Distance** to 0.0 Å. Run parKVFinder again.



Command line interface

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

```
/home/user/parKVFinder$ parKVFinder
parKVFinder (parallel KVFinder) software identifies and describes cavities in
target biomolecular structure using a dual probe system.

The description includes spatial and constitutional characterization. Spatial
description includes shape, volume and area. Constitutional description includes
amino acids that form the identified cavities.

Usage: parKVFinder PDB [options],
   where PDB is a path to a target PDB file.

Options:
   -h, --help
        Display this help message.
   -v, --version
```

```
Display parKVFinder version.
--verbose
Print extra information to stdout.
```

Whole Protein detection

Standard execution of the command line interface only requires the PDB file path of the target protein. So let's repeat the execution of 1FMO protein with default parameters.

```
/home/user/parKVFinder$ parKVFinder$ parKVFinder input/1FMO.pdb
[PID 8504] Running parKVFinder for: /home/user/parKVFinder/input/1FMO.pdb
done!
Elapsed time: 0.78 seconds
```

To view the cavities, the cavities PDB file must be loaded into PyMOL.



Custom execution

There a set of options for customizing parKVFinder's command line execution. These options are displayed in a help menu with their default values when applicable.

```
/home/user/parkVFinder$ parkVFinder -h
parKVFinder (parallel KVFinder) software identifies and describes cavities in
target biomolecular structure using a dual probe system.
The description includes spatial and constitutional characterization. Spatial
description includes shape, volume and area. Constitutional description includes
amino acids that form the identified cavities.
Usage: parKVFinder PDB [options],
   where PDB is a path to a target PDB file.
Options:
  -h, --help
     Display this help message.
  -v, --version
     Display parKVFinder version.
     Print extra information to stdout.
General options:
  -p, --parameters [<.toml>]
     Define path to parameters file.
  -d, --dictionary [<dictionary>]
     Define path to a custom dictionary file.
  -r, --resolution <enum>
                           (Low)
     Define resolution mode. Options include: Off, Low, Medium and High.
  -s, --step
                  <real> (0.0)
     Define step size (grid spacing).
  -i, --probe_in <real>
                            (1.4)
     Define probe in size.
  -o, --probe_out <real>
                            (4.0)
     Define probe out size.
  --volume_cutoff <real>
                            (5.0)
     Define cavities volume filter.
  --removal_distance <real> (2.4)
     Define removal distance when comparing probes surfaces.
                        (paramters.toml)
  -t, --template
     Create a parameter file template with defined parameters in current
     working directory.
Box adjustment options:
  -B, --box
     Define a search box mode where parKVFinder will detect cavities.
  --custom box [<file>]
     Define a custom search box based on a file containing the minimum and
     maximum cartesian values of each axis in angstrom.
  --residues box [<file>]
     Automatically set a search box based a file containing a tab-separated
     list of residues.
  --padding
               <real>
                          (3.5)
     Define residues box padding. Adds a length in each box direction.
```

```
Surface options:

-S, --surface <enum> (VdW)

Define a surface representation. Options include: SAS and VdW. SAS specifies solvent accessible surface. VdW specifies van der Waals molecular surface.

Ligand options:

-L, --ligand [<.pdb>]

Define path to ligand PDB file.

--ligand_cutoff <real> (5.0)

Define ligand radius distance cutoff.
```

So let's repeat the previous analysis, but changing Probe Out and Volume Cutoff parameters.

/home/user/parKVFinder\$ parKVFinder\$ parKVFinder input/1FMO.pdb --probe_out 8.0 --volume_cutoff 100. [PID 12089] Running parKVFinder for: /home/user/parKVFinder/input/1FMO.pdb done!



If we set the verbose flag, extra information will be printed in stdout.

```
/home/user/parKVFinder$ parKVFinder$ parKVFinder input/1FMO.pdb --probe_out 8.0 --volume_cutoff 100.
[PID 12198] Running parKVFinder for: /home/user/parKVFinder/input/1FMO.pdb
> Setting 'dictionary_name' to default file: /home/user/parKVFinder/dictionary
> Setting 'probe_in' to default value: 1.40
> Setting 'removal_distance' to default value: 2.40
> Setting 'step' (grid spacing) to 0.00.
> Setting 'resolution' to flag: Low
> Setting 'ligand_cutoff' to default value: 5.00
> Chosen 'surface' representation: VdW
> Running parKVFinder for whole biomolecular structure
> Loading atomic dictionary file
> Calculating grid dimensions
> Reading PDB coordinates
> Creating grid
> Filling grid with probe in surface
> Filling grid with probe out surface
> Defining biomolecular cavities
> Calculating volume
> Calculating surface points
> Calculating area
> Retrieving residues surrounding cavities
> Writing cavities PDB file
> Writing results file
done!
Elapsed time: 1.99 seconds
```

Box Adjustment modes

The command line interface takes a different approach to box adjustment mode compared to PyMOL parKVFinder Tools. As the GUI is not available to drawn the custom box, the command line interface set the custom box based on a list of residues (residues_box) or a set of cartesian coordinates (custom_box).

So let's execute parKVFinder using both box approaches.

Residues box

First, we need to define a list of residues in a tab-separated file, in which we specify the residues as the residues number and chain identifier separated by an underscore character (__).

Considering the example file called 1FMO.residues.KVFinder.in, located inside input directory, we select the residues 51E, 71E, 99E and 134E (highlighted in red) to draw the custom box.



/home/user/parKVFinder\$ parKVFinder\$ cat input/1FMO.residues.KVFinder.in 51_E 71_E 99_E 134_E

So let's run parKVFinder for the 1FMO protein using a custom box based on this set of residues. A custom box will be drawn around these residues with a default padding, which can be changed using the --padding <real> option.

/home/user/parKVFinder\$ parKVFinder\$ parKVFinder input/1FMO.pdb -B --residues_box input/1FMO.residues [PID 15230] Running parKVFinder for: /home/user/parKVFinder/input/1FMO.pdb done!
Elapsed time: 0.27 seconds



Note: This method uses the same procedure as PyMOL parKVFinder Tools to draw the custom box in Box Adjustment mode.

Cartesian box

First, we need to define a tab-separated file containing the minimum and maximum cartesian values of each axis in the following order: Xmin, Xmax, Ymin, Ymax, Zmin and Zmax.

Considering the example file called 1FMO.box.KVFinder.in, located inside input directory, to draw the custom box.

```
/home/user/parKVFinder$ parKVFinder$ cat input/1FMO.box.KVFinder.in -20.0 20.0 -20.0 20.0 20.0
```

So let's run par KVFinder for the 1FMO protein using a custom box based on this set of coordinates.

```
/home/user/parKVFinder$ parKVFinder$ parKVFinder input/1FMO.pdb -B --custom_box input/1FMO.box.KVFin [PID 16070] Running parKVFinder for: /home/user/parKVFinder/input/1FMO.pdb done!
Elapsed time: 0.42 seconds
```



Manual

About

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parKVFinder and PyMOL parKVFinder Tools was developed by:

- <author 1> (<e-mail 1>)
- <author 2> (<e-mail 2>)
- ..
- <author N> (<e-mail N>)

<Institution> (<website>).

If you have any further questions, inquires or if you wish to contribute to parKVFinder project, please contact us at <e-mail>.