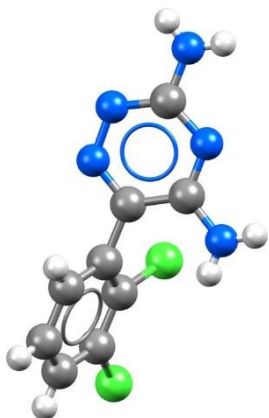


Introduction to the Conformer Generator in Mercury (CONF-001)

2024.1 CSD Release



CCDC
advancing structural science

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Introduction

The CSD Conformer Generator provides the ability to both minimise molecular conformations and generate diverse conformer subsets based on CSD data. The methodology starts from an input 3D molecular structure with all hydrogen atoms present, which is optionally minimised in the first step. Subsequently, conformations are sampled based on CSD-derived rotamer distributions and ring templates. A final diverse set of conformers, clustered according to conformer similarity, is returned. Each conformer is locally optimised in torsion space. There are three methods for generating conformers with the CSD portfolio; via the Mercury interface, the command line utility and the CSD Python API. This workshop focuses on generating conformers through the Mercury interface.

Before beginning this workshop, ensure that you have a registered copy of CSD-Materials, CSD-Discovery or CSD-Enterprise installed on your computer.

Learning Outcomes

This tutorial will guide you through the process of generating conformers in Mercury. At the end of this workshop, you will be able to:

- Sketch a structure in Mercury.
- Generate a set of conformers.
- Visualize multiple structures at once.

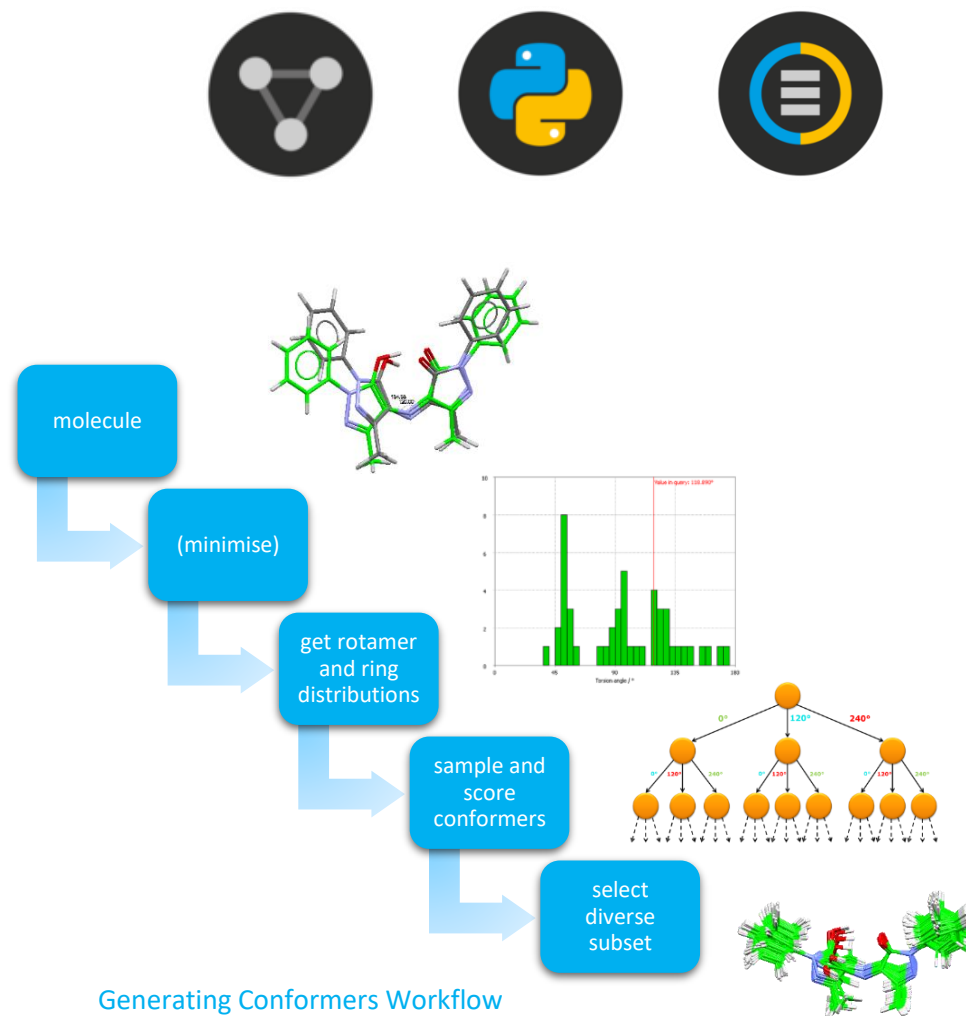
Note: This handout should take approximately **25 minutes** to be completed.

Pre-required Skills

Familiarity with the Mercury interface is important; you can access the Visualization in Mercury self-guided workshop [here](#) or review the basics of visualization in Mercury [here](#). Knowledge of sketching a structure in ConQuest is expected; a [review of sketching in ConQuest](#) is at the end of this document.

Materials

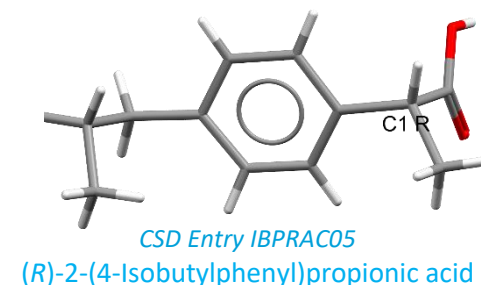
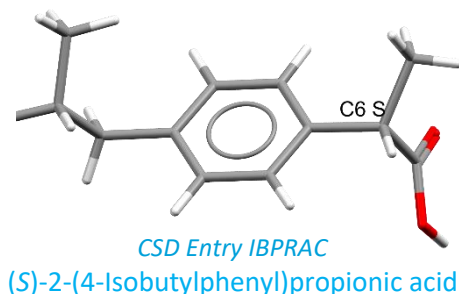
There are no additional materials for this workshop.




Generating Conformers Workflow

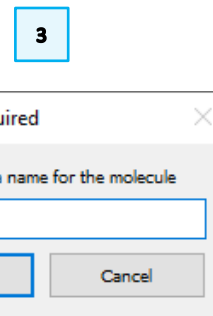
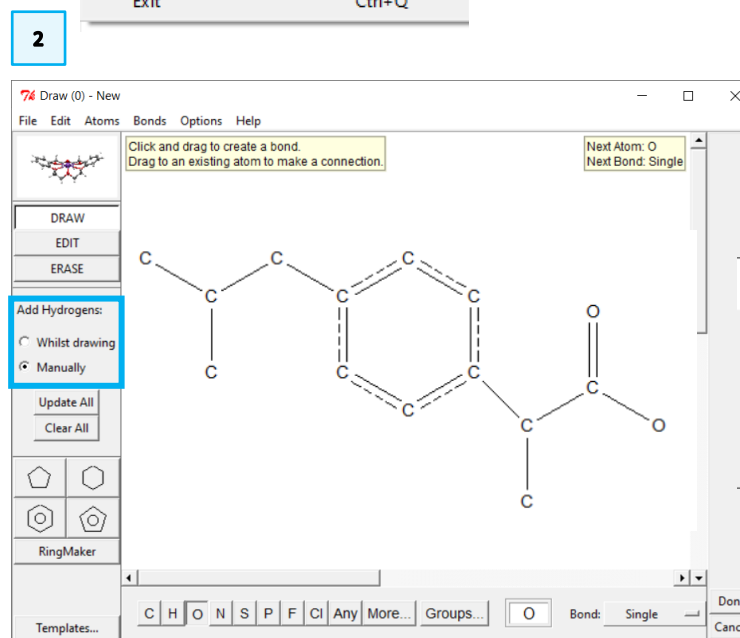
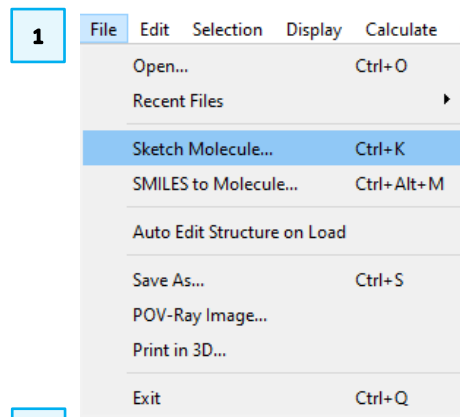
Example 1. Generating Conformers in Mercury.

In this example, we will go through the steps of setting up a conformer generation run in Mercury. The CSD Conformer Generator requires as input 3D structures with all hydrogen atoms present. We will be generating conformers for both stereoisomers of 2-(4-isobutylphenyl)propionic acid (ibuprofen).



You can either sketch your structure in Mercury, import a .cif file or load one of the structures from the CSD by typing the refcode in the Structure Navigator. We will first sketch the structure of ibuprofen, but you can use one of the ibuprofen refcodes: IBPRAC or IBPRAC05.

1. Start Mercury by double-clicking on the icon . Click on *File > Sketch Molecule* to bring up the sketcher.
2. The sketcher interface is the same used in ConQuest. If you need a refresher on how to sketch structures in ConQuest, check out the [summary at the end of this handout](#). Sketch the structure of ibuprofen shown to the right.
3. Once you are done sketching the molecule, click **Done**. Your settings might add H atoms by default, making your sketched molecule look a little different. We will edit the structure later in Mercury. A pop-up window will appear asking for a name for the structure. Provide a name for the structure and click **OK**.
4. You will now return to the main Mercury window with your sketched structure displayed.

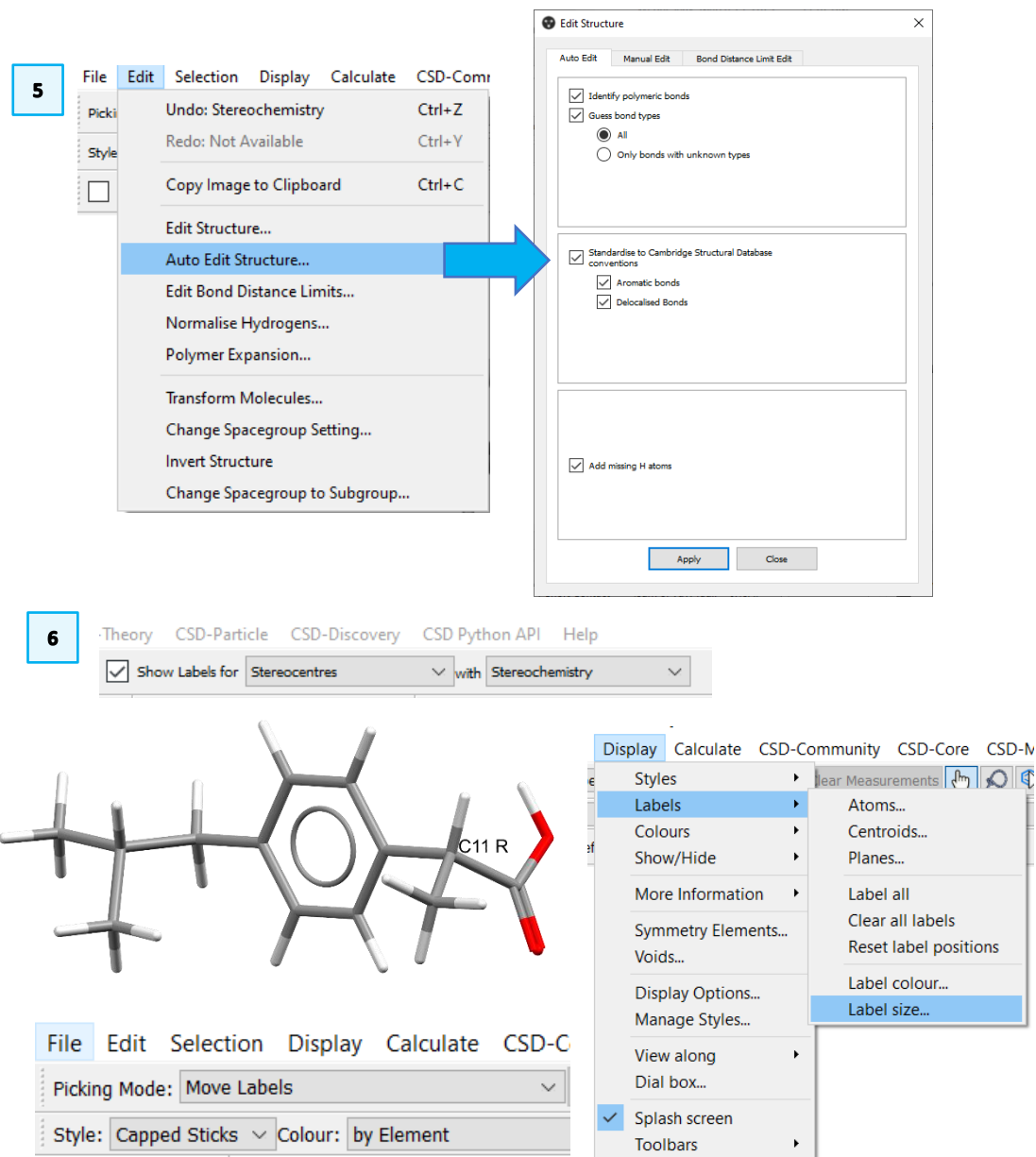
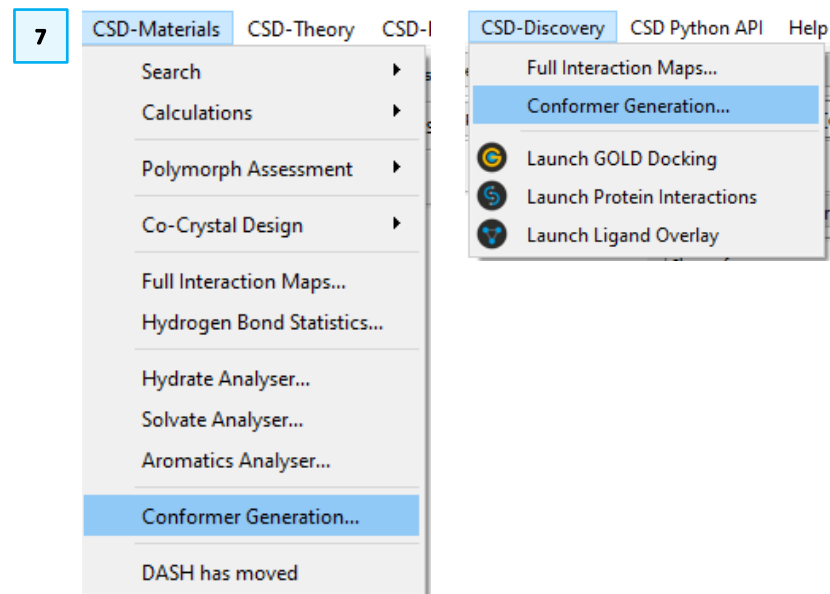


5. We first need to make sure that all bond types in our structure are correct. We can do so by using the Edit Structure option. Click *Edit > Auto Edit Structure*. Leave the default options ticked and click **Apply**. If there are any changes made, it will be shown in the pop-up.
6. 2-(4-Isobutylphenyl)propionic acid has a chiral centre, however we were unable to specify chirality when we sketched the structure. We can view the configuration that our structure has. Tick the **Show Labels for** box, select *Stereocentres* with *Stereochemistry*. The structure shown here is the *R* enantiomer but yours could be different.

Tip: You can change the label size from the Display menu; *Display > Labels > Label size*. Change the *Picking Mode* to *Move Labels*.

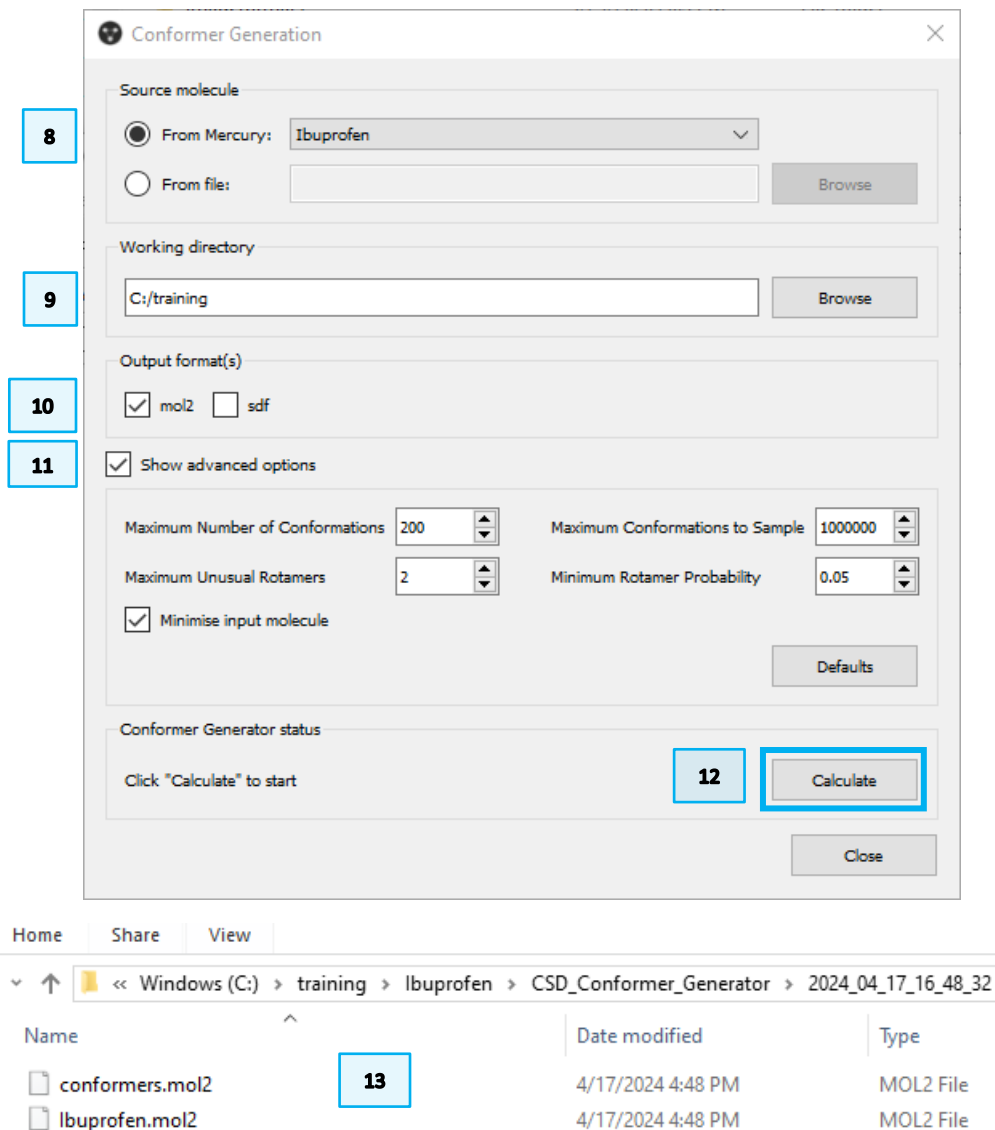
We are now ready to set up the Conformer Generator. The CSD Conformer Generator can be accessed through the **CSD-Materials** and **CSD-Discovery** menus within Mercury.

7. From the CSD-Materials or CSD-Discovery menu, click on *Conformer Generation...* to open the Conformer Generation window.



8. In the **Conformer Generation** window in the *Source molecule* section, you can specify a CSD entry, an in-house database entry, or another structure as the input molecule for conformer generation. The default selected molecule is the present molecule in the main display window. Alternatively, you can select a file on the disk as the input by selecting *From file:* option and clicking on the **Browse** button.
9. The working directory can be selected under *Working Directory* section of the **Conformer Generation** window. You can stick with the default here or Click the **Browse** button and navigate to the folder you want the results saved to. A folder will be created in this directory named after the identifier of the input molecule used. Both a copy of the input molecule and the generated conformers will be written to the output folder.
10. Conformers may be generated in *.mol2* or *.sdf* format by choosing the relevant checkboxes under *Output format(s)*. The default *.mol2* format will be used in this tutorial.
11. Tick the **Show advanced options** to view the defaults. You can change the values here if necessary but we will stick with the defaults.
12. Click the **Calculate** button to generate conformers.
13. The calculation should take less than a minute. You can navigate to the folder you selected in step 9, the conformers you have generated are placed in `\path\to\structure\CSD_Conformer_Generator\YYYY_MM_DD_hh_mm_ss\conformers.mol2`.

Replace `\path\to\` with the actual path to the folder selected in step 9. Please note that `YYYY_MM_DD_hh_mm_ss` is the date and time when the calculation was run. You will have a different folder name corresponding to your own calculation date and time.



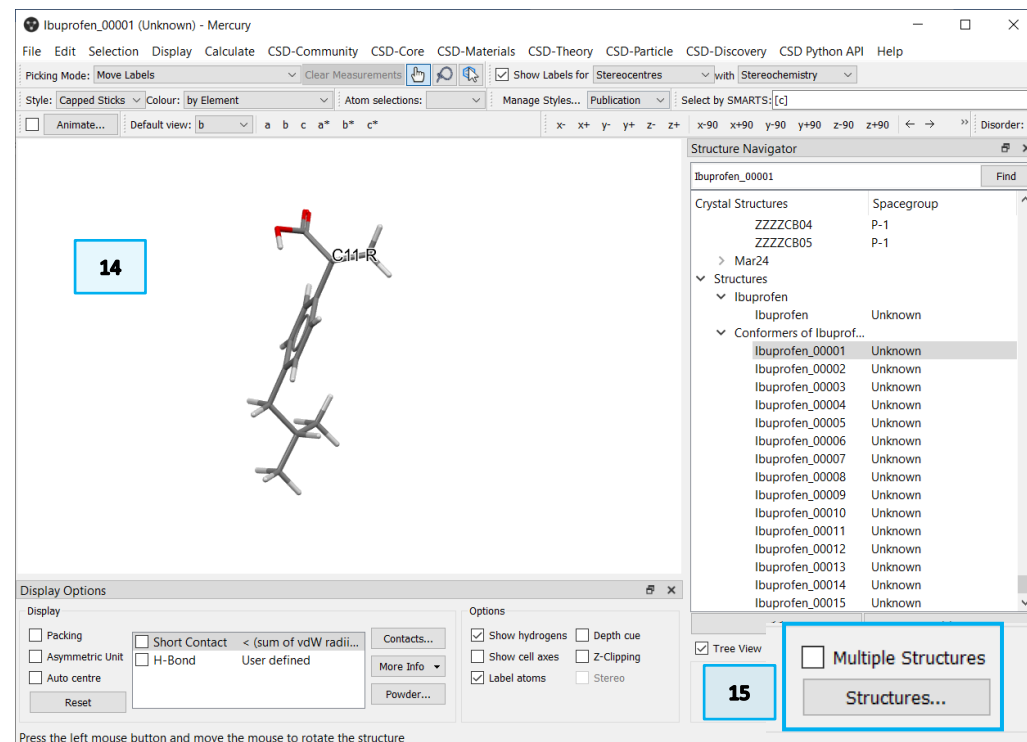
14. The conformers are also populated in the **Structure Navigator** as new entries.

Note that the first conformer generated is now shown in the Mercury window. You can scroll through the conformers by clicking on each entry or the down arrow on your keyboard. A couple of things to note here are (a.) the stereochemistry is conserved in all conformers and (b.) it is not easy to see the differences between the conformers if viewing one by one. We will load multiple conformers at once and superimpose them.

15. Tick the **Multiple Structures** box in the lower right corner of the Mercury window. Click on however many conformers you want and the input molecule to view them at once. For some users, the structures might be superimposed but if they are not, you can use the view options to change that.

16. From the *Default view:* menu, select *Custom*. You may need to untick and tick again the Multiple Structures box. Now when you click on multiple structures, they should be superimposed along the same axis.

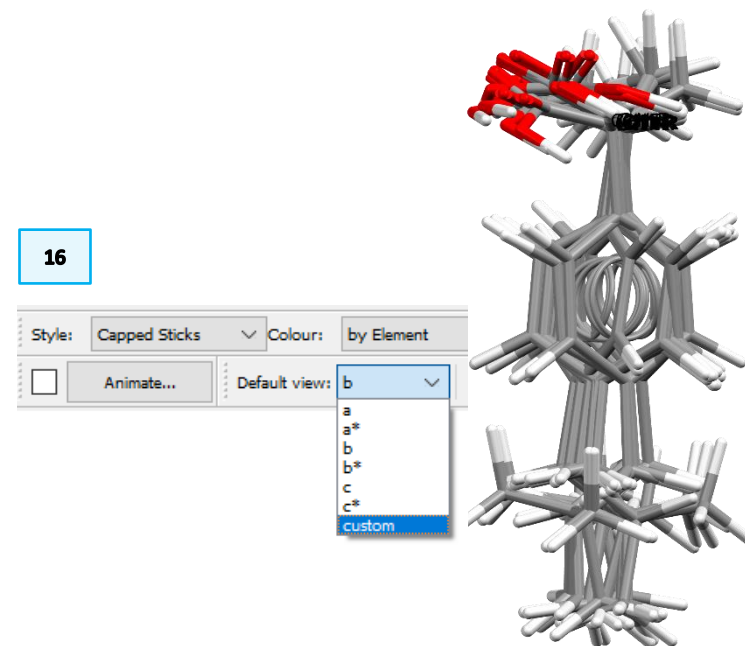
17. You can change the colour of each conformer displayed to make the differences more visible. Click on **Structures...** in the lower right corner to bring up the *Multiple Structures* menu. Change the colour by clicking on **by Element** box and choose a colour. You can also hide some of the entries from the display.



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Multiple Structures					
Actions, e.g. packing, will be applied to Active structure(s) only					
Rotation is around					
<input checked="" type="radio"/> Global rotation centre <input type="radio"/> Local rotation centres					
Delete All		Visible	Active	Movable	Colour
1 Delete	Ibuprofen	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	Yellow
2 Delete	Ibuprofen_00001	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	Red
3 Delete	Ibuprofen_00002	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	by Element
4 Delete	Ibuprofen_00003	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	by Element
5 Delete	Ibuprofen_00004	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	by Element
6 Delete	Ibuprofen_00005	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	by Element

☐ Move the structure that is nearest the mouse cursor

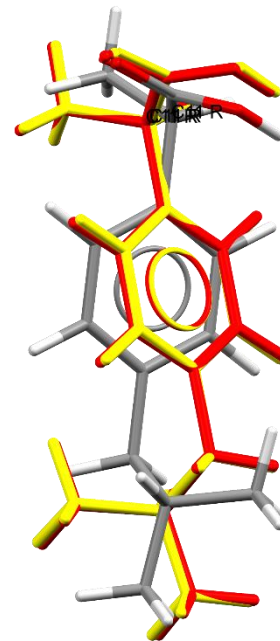


18. The colour of the entries should change in the display window.

Conclusion

In this example, we sketched the structure of ibuprofen and generated a set of conformers using the Conformer Generator module in Mercury. As mentioned, it is possible to generate conformers from structures already in the CSD. You can generate conformers for the other enantiomer using either refcode IBPRAC for an *S* enantiomer or IBPRAC05 for an *R* enantiomer. You can also try changing some of the default settings to see if this changes your results.

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Example 2. Generating Conformers via the CSD Python API Script in Mercury.

1. Load the ibuprofen structure in Mercury or select the entry from the **Structure Navigator**.
2. From the CSD Python API menu select *Analysis > generate_conformers.py*
3. There will be a window showing the progress, the script should take less than 30 seconds to be completed. The output folder where the files are written is displayed in this window.
4. Once the script is done running, a **Data Analysis** window will open with the conformers generated and torsion angles in each molecule. Clicking on each entry will place the conformer in the Mercury display window. You can choose to plot some of the angles or export the parameters.
5. To export the entries into a .csv, click on *File > Export* in the **Data Analysis** window. This will open the **Export Spreadsheet** window where you can select items you want saved to the spreadsheet.
6. If you have experience editing Python scripts and want to view this script, it is in path\to\CCDC\ccdc-software\mercury\scripts\Analysis replacing path\to\ with your actual install location.

Data Analysis
File Options

ibuprofen_generate_conformers Spreadsheet 1

File Tools Descriptors Display Selection Plots Statistics

Find identifier Find next

Identifier	NAME	Molecule	...rsion C10-C8-C7-C2	...rsion C9-C8-C7-C2	...rsion C3-C2-C7-C8	...rsion C1-C2-C7-C8	...rsion O15-C12-C11-C14	...rsion O13-C12-C11-C14	...rsion O15-C12-C11-C5	...rsion O1
ibuprofen_generate_conformers ibuprofen_0001 0	ibuprofen_0001	ibuprofen_0001	173.0000	-65.2721	90.2759	-90.0000	19.0000	-160.8796	-103.6171	
ibuprofen_generate_conformers ibuprofen_0002 1	ibuprofen_0002	ibuprofen_0002	173.0000	-65.2721	-89.7241	90.0000	19.0000	-160.8796	-103.6171	
ibuprofen_generate_conformers ibuprofen_0003 2	ibuprofen_0003	ibuprofen_0003	62.0000	-176.2721	90.2759	-90.0000	19.0000	-160.8796	-103.6171	
ibuprofen_generate_conformers ibuprofen_0004 3	ibuprofen_0004	ibuprofen_0004	62.0000	-176.2721	-89.7241	90.0000	19.0000	-160.8796	-103.6171	
ibuprofen_generate_conformers ibuprofen_0005 4	ibuprofen_0005	ibuprofen_0005	173.0000	-65.2721	90.2759	-90.0000	-134.9140	45.2063	102.4689	
ibuprofen_generate_conformers ibuprofen_0006 5	ibuprofen_0006	ibuprofen_0006	173.0000	-65.2721	-89.7241	90.0000	-134.9140	45.2063	102.4689	
ibuprofen_generate_conformers ibuprofen_0007 6	ibuprofen_0007	ibuprofen_0007	62.0000	-176.2721	90.2759	-90.0000	-134.9140	45.2063	102.4689	
ibuprofen_generate_conformers ibuprofen_0008 7	ibuprofen_0008	ibuprofen_0008	62.0000	-176.2721	-89.7241	90.0000	-134.9140	45.2063	102.4689	
ibuprofen_generate_conformers ibuprofen_0009 8	ibuprofen_0009	ibuprofen_0009	173.0000	-65.2721	90.2759	-90.0000	20.8311	-159.0485	-101.7859	
ibuprofen_generate_conformers ibuprofen_0010 9	ibuprofen_0010	ibuprofen_0010	173.0000	-65.2721	-89.7241	90.0000	20.8327	-159.0470	-101.7844	
ibuprofen_generate_conformers ibuprofen_0011 10	ibuprofen_0011	ibuprofen_0011	62.0000	-176.2721	90.2759	-90.0000	20.8311	-159.0485	-101.7859	
ibuprofen_generate_conformers ibuprofen_0012 11	ibuprofen_0012	ibuprofen_0012	62.0000	-176.2721	-89.7241	90.0000	20.8311	-159.0485	-101.7859	
ibuprofen_generate_conformers ibuprofen_0013 12	ibuprofen_0013	ibuprofen_0013	173.0000	-65.2721	90.2759	-90.0000	152.0016	-27.8781	29.3845	
ibuprofen_generate_conformers ibuprofen_0014 13	ibuprofen_0014	ibuprofen_0014	173.0000	-65.2721	-89.7241	90.0000	152.0016	-27.8781	29.3845	
ibuprofen_generate_conformers ibuprofen_0015 14	ibuprofen_0015	ibuprofen_0015	62.0000	-176.2721	90.2759	-90.0000	152.0107	-27.8689	29.3937	
ibuprofen_generate_conformers ibuprofen_0016 15	ibuprofen_0016	ibuprofen_0016	62.0000	-176.2721	-89.7241	90.0000	152.0107	-27.8689	29.3937	
ibuprofen_generate_conformers ibuprofen_0017 16	ibuprofen_0017	ibuprofen_0017	169.4741	-68.7979	64.7680	-115.5079	19.0000	-160.8796	-103.6171	
ibuprofen_generate_conformers ibuprofen_0018 17	ibuprofen_0018	ibuprofen_0018	169.4240	-68.8480	-115.2000	64.5241	19.0000	-160.8796	-103.6171	
ibuprofen_generate_conformers ibuprofen_0019 18	ibuprofen_0019	ibuprofen_0019	62.0000	-176.2721	60.2759	-120.0000	19.0000	-160.8796	-103.6171	

2

CSD Python API Help

- Analysis
 - conformers_similarity.py
 - generate_conformers.py
- Reports
- Searches
- user_support.py
- welcome.py
- aromatic_ring_model
- scripts
- scatter_info.py
- Options...
- CSD Python API Documentation
- CSD Python API GitHub Script Repository

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ibuprofen

The python interpreter is: C:/Users/.../CCDC/ccdc-software/csd-python-api/miniconda/python.exe

The working directory is: C:/training/Ibuprofen/generate_conformers/2024_04_18_16_27_03

Output files will be written in: C:/training/Ibuprofen/generate_conformers/2024_04_18_16_27_03

Script completed...
12 seconds

5

Data Analysis
File Options

ibuprofen_00001_generate_conformers Spreadsheet 1

File Tools Descriptors Display Selection Plots Statistics

Export...

Export selected identifiers...

Close

Export Spreadsheet

127 entries from spreadsheet

Available Items (Right-click for options)

- NAME
- Molecule
- Torsion C10-C8-C7-C2
- Torsion C9-C8-C7-C2
- Torsion C3-C2-C7-C8
- Torsion C1-C2-C7-C8
- Torsion O15-C12-C11-C14
- Torsion O13-C12-C11-C14
- Torsion O15-C12-C11-C5
- Torsion O13-C12-C11-C5

Selected Items (Right-click for options)

Up Down

CSV (*.csv) Browse

OK Cancel

Conclusion

In this example, we have generated conformers for a single molecule in Mercury through a script in the CSD Python API menu. The Data Analysis window includes the torsion angles in the molecule for each conformer. You can generate plots of the torsion angles to view structures that may be outliers.

Summary

We have used the conformer generator feature in Mercury to generate conformers of ibuprofen. After this workshop you should now be able to:

- Sketch a structure in Mercury.
- Generate conformers for a given structure.
- Use different display options for viewing multiple structures.
- Analyse results using the Data Analysis window.

Next Steps

The CSD Conformer Generator User Guide can be found [here](#) or from the Documentation and Resources section of our website. There are two other methods for generating conformers: via the CSD Python API and the command line. If you are interested in exploring those, instructions can be found in the *How to Generate Ensembles of Conformers using the CSD Conformer Generator* (CONF-002) on this [page](#).

<https://www.ccdc.cam.ac.uk/community/training-and-learning/workshop-materials/csd-discovery-workshops/>

Feedback

We hope this workshop improved your understanding of the *CSD Conformer Generator* and you found it useful for your work. As we aim to continuously improve our training materials, we would love to hear your feedback. Click on [this link](#) to a survey (link also available from workshops webpage), it will take less than 5 minutes to complete. The feedback is anonymous. You will be asked to insert the workshop code, which for this self-guided workshop is CONF-001. Thank you!

Data Analysis

File Options

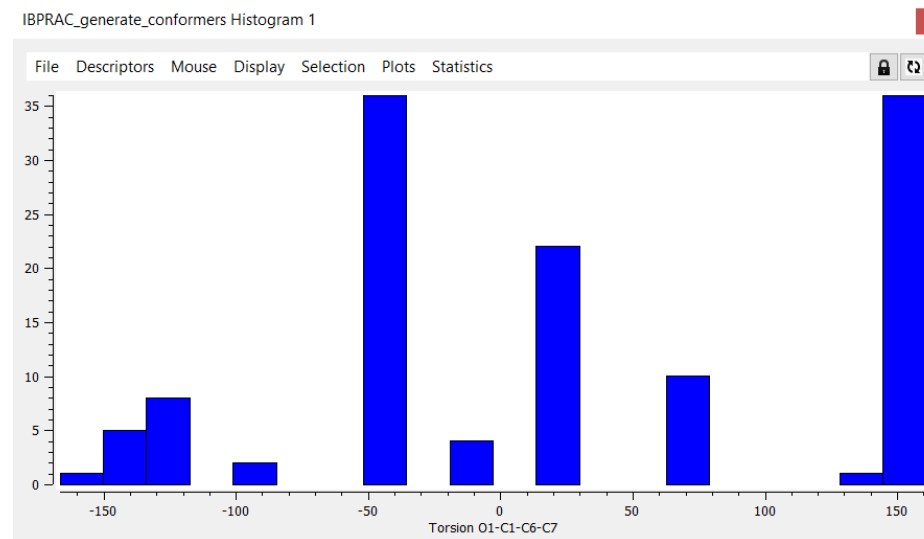
IBPRAC_generate_conformers Spreadsheet 1

File Tools Descriptors Display Selection Plots Statistics

Find identifier

Find next

Identifier	File	Torsion 01-C1-C6-C7
IBPRAC_generate_conformers IBPRAC_0001 0	001	161.0071
IBPRAC_generate_conformers IBPRAC_0002 1	002	161.0071
IBPRAC_generate_conformers IBPRAC_0003 2	003	161.0071
IBPRAC_generate_conformers IBPRAC_0004 3	IBPRAC_0004 IBPRAC_0004	161.0071
IBPRAC_generate_conformers IBPRAC_0005 4	IBPRAC_0005 IBPRAC_0005	-42.9929
IBPRAC_generate_conformers IBPRAC_0006 5	IBPRAC_0006 IBPRAC_0006	-42.9929



Basics of Mercury Visualization

Mercury is the CCDC's visualization software to view 3D structures of small molecules, generate images, and animations of molecules.

In the following we will see some of the basics of navigation and visualization in Mercury that you will find helpful to support your analysis.

In the **Mercury interface** we find:

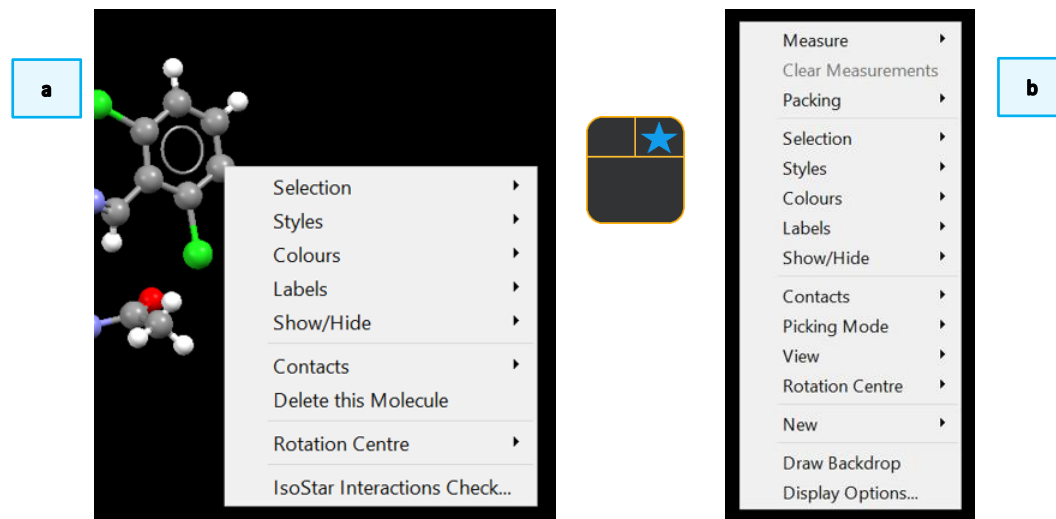
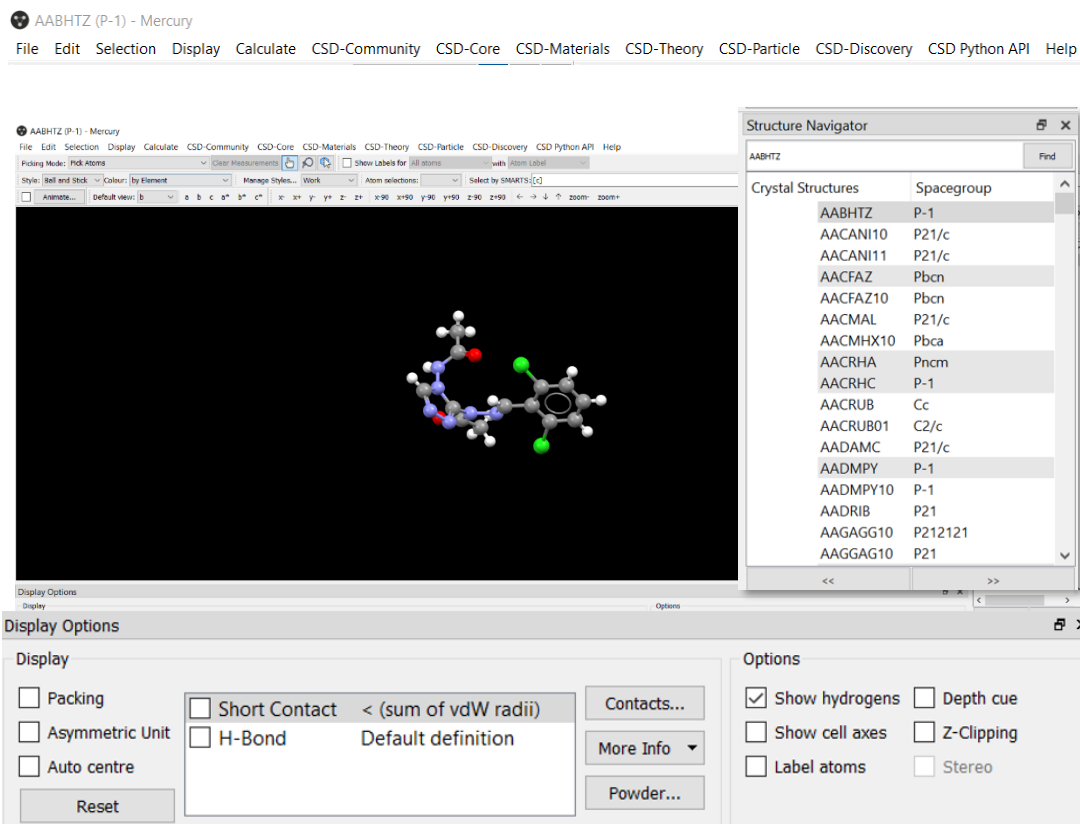
- **At the top:** list of menus from which we can access visualization and analysis options, and other CSD components such as CSD-Materials.
- **On the right-hand side:** the **Structure Navigator**, with the database loaded (depending on your licence). The Structure Navigator allows you to select a refcode to visualize in the main Mercury window.
- **Beneath the main display window:** **Display options toolbar**. You can quickly view a packing diagram, display Hydrogen bonding and detailed information about the molecule using the More Info option.

Using the **mouse to enhance visualization**:

- Left mouse button and move – rotate molecules.
- Middle Mouse wheel – move molecules up and down.
- Right mouse button and move up and down – zoom in and out of molecules.
- Shift + Left mouse button and move - rotate in the plane molecules.
- Ctrl + Left mouse button and move - translate molecules.

Right click:

a) Near a molecule and b) Away from a molecule



Review. Draw Window

All drawing takes place in the central white area of the *Draw* window. In addition to creating 2D chemical structure sketches, the *Draw* window allows for the inclusion of 3D parameters for searching or for filtering.

ConQuest sketching conventions

- Left click in the sketcher to insert the selected atom type
- Left click and drag to sketch two bonded atoms
- Use the **Edit** button to modify properties of or delete atoms, bonds or entire substructures
- Right-click on atoms or bonds to modify their properties
- Use the **Templates...** button to pick from a list of CSD editor devised and drawn substructures
- Use the **More...** button to find less frequently used element types, or generic atom type groups (e.g. halogens), or define custom element combinations (e.g. C or N or O).

