

ACD/I-Lab

Version 8.0 for Microsoft Windows

User's Guide

***Calculating Chemical Properties and
Searching Commercial Databases
On-line via ChemSketch***

Advanced Chemistry Development, Inc.

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Before You Begin

Thank you for purchasing ACD/I-Lab.

About This Guide

This guide provides a comprehensive description of all of the options available in ACD/MS Fragmenter, it is designed for either online use or to be printed and used as a "hard copy" version.

The screen shots shown throughout this guide have been taken with a relatively small window size.

The colors and other properties of the window elements described throughout this guide correspond with the default Windows Display Properties.

This reference guide is provided in electronic form, readable with Adobe Acrobat software. If you cannot locate an index topic you need please do a text string search for the relevant word or phrase, or related words.

Important Note that some of the I-Lab services described in this guide may be missing on your list! For I-Lab: Intranet Edition, the number of available services depends on the ACD/Labs package installed; and for Public I-Lab, on the login type you currently use.

Mouse Conventions

You may perform several actions during your work with this software; the following specific words are used to describe them:

- **Point to** means move the mouse pointer  to an item.
- **Click** means point to an item, and press the left mouse button.
- **Right-click** means point to an item, and press the right mouse button.
- **Double-click** means point to an item, and quickly press the left mouse button twice.
- **Drag** means point to an item, press and hold down the left mouse button while you move the item.
- **Select** means highlight or make an interface element active by either clicking it or dragging over it (other actions are possible if specified in documentation). If used in "select the check box", it means that the check box should be marked with a tick (as opposed to "clear the check box" when the check box should be cleared, without a mark).

For More Information...

To see the latest in ACD/Labs software and services, please visit our Web site at

<http://www.acdlabs.com>

Our Web site is being accessed at the rate of tens of thousands of "hits" per day. There's a reason for this: much is offered through our Web site. As of Summer 2006, we offer free ChemSketch 10.0, a free ISIS 3D Add-in, free ChemDraw extensions, and a free 2-week demo key for "Interactive Laboratory" sessions where you can run test calculations using Java applets without purchasing software. There are TechSmith Camtasia-based movies which show the operation of many of our software packages (especially ChemSketch) available for download.

We are constantly updating the information on our Web site. The Web site will tell you at which scientific conferences you can visit the ACD/Labs booth. You can browse the Frequently Asked Questions page or drop in and "chat" on our newsgroup, which can also be reached via our web page.

If you would like to stay informed of the latest developments in chemical software at ACD/Labs, please be sure to sign up for e-mail broadcasts at our Web site page:

<http://www.acdlabs.com/feedback/mailing.html>

If you would like to participate in the ACD/Labs forums, please access:

<http://forum.acdlabs.com>

How to Contact Us

We are accessible through our Web site, phone, fax, and regular mail, but by far the most popular way to contact us is via electronic mail. Questions on pricing, sales, availability, and general issues should be directed to:

info@acdlabs.com

Technical and scientific support issues should be addressed by visiting:

<http://support.acdlabs.com>

Please tell us the name of the software purchaser; the product name, version number, build number, and license ID of the product you are contacting us about (from the **Help** menu, choose **About** to find this information); as well as a description of the problem you are having. If applicable, please tell us the name of the distributor from whom you purchased the software.

Online Updates

All PC-based ACD/Labs software contains the capability to have software updates delivered online. You will need the registration numbers of the software and an Internet connection from the same computer on which the software is installed. The updates are small fixes, for example, bringing the actual version number of a program from 10.00 to 10.01. For more information, refer to the document located in the ACD/Labs documentation folder, \\DOCS\UP_CLNT.PDF, or contact our technical support department.

1. Introduction

1.1 What is ACD/I-Lab

ACD/I-Lab is the Internet-based service that allows you to get instant access to ACD/Labs chemical databases and property predictions programs via the Internet.

1.1.1 ACD/Public I-Lab

Public access to the latest versions of I-Lab services is available from the ACD/Labs website. Public I-Lab includes both fee-based and free services (the number of available services depends on your login type: authenticated user, guest, or sight license). I-Lab registration and membership are free. An account on the Interactive Lab can be set up from <http://www.acdlabs.com/ilab> or via ChemSketch; a 2-week demo period can be arranged for I-Lab. Users with ChemSketch 6.0 and later can access I-Lab automatically if run on a PC with Internet connection.

Note Some of the I-Lab options are available for ACD/I-Lab: Intranet Edition only. In this guide, these are marked as *“For I-Lab: Intranet Edition only!”*

1.1.2 ACD/I-Lab: Intranet Edition

I-Lab can also be packaged for use in your Corporate Intranet environment. Using I-Lab behind a corporate firewall allows you to use the corporate security infrastructure to keep I-Lab transactions secure. In addition to a standard set of I-Lab services, I-Lab: Intranet Edition allows you to improve prediction results by training predictors with your own experimental data (for more information, refer to Section 3.6.1).

Note Some of the I-Lab options are available for ACD/Public I-Lab only. In this guide, these are marked as *“For Public I-Lab only!”*

1.2 What's New in 8.0?

1.2.1 HNMR DB

The expanded DAT file now contains over 1,384,000 experimental chemical shifts and 449,000 coupling constants for over 165,000 structures.

The internal ACD/Labs database contains very recent entries. Over 3300 entries are from articles published in the year 2001.

1.2.2 CNMR DB

The expanded DAT file now contains about 2,017,000 experimental chemical shifts and 81,000 coupling constants for over 165,000 structures.

The internal ACD/Labs database contains very recent entries. Over 3300 entries are from articles published in the year 2001.

1.2.3 FNMR DB

ACD/FNMR database contains more entries (13,800 records) and represents more classes of structure types than previously. About 50 entries are from references as recent as 2001.

1.2.4 PNMR DB

The internal ACD/PNMR database contains more entries (22,600 records) and represents more classes of structure types than previously. Over 450 entries are from references as recent as 2000, with 22 from 2001.

1.2.5 Connection to ACD/I-Lab: Intranet Edition

Training capability for Aqueous Solubility calculation is improved. ACD/Log*P* Accuracy Extender can be used to improve the accuracy of Log*D*, Aqueous Solubility and Adsorption Coefficient/BCF prediction.

1.2.6 Redesigned Options

The following options have been redesigned in ACD/I-Lab version 8.0:

- The **I-Lab Options** dialog box is redesigned: basic method of Server authentication is supported.
- Improved I-Lab services interface (support for the Stereoconfiguration options for IUPAC Name and Index Name).

2. Quick Start

2.1 Objectives

This chapter will give you the basics of how to work with I-Lab (short for “Interactive Lab”) via the ChemSketch interface. You should have some familiarity with the ChemSketch interface to follow exercises. If you encounter any problems with drawing structures, refer to the *ACD/ChemSketch User's Guide*.

In this chapter you will learn:

- What the prerequisites for accessing I-Lab are;
- What options are necessary for connecting to I-Lab if you are behind a firewall or proxy server;
- How to access I-Lab as a guest;
- How to use I-Lab services with IUPAC Name (Free) as an example;
- What the Task Manager is; and
- How to log out.

2.2 Connecting to I-Lab

2.2.1 Prerequisites

In order to use ACD/I-Lab resources with ChemSketch, you should have:

- A direct Internet connection from the PC where you have ChemSketch installed:
 - *For Public I-Lab:* Access to <http://www.acdlabs.com> and <http://www2.acdlabs.com> addresses, at least.
 - *For I-Lab: Intranet Edition:* Access to the host where the I-Lab: Intranet Edition server is installed.
- ACD/I-Lab Add-on for ChemSketch installed.
- *For Public I-Lab only!* An e-mail address.

If the I-Lab Add-in is already in place, when you start up ChemSketch you will see the standard ChemSketch interface with the additional **I-Lab** menu on the menu bar:



and the additional **I-Lab Login** button on the left part of the status bar:

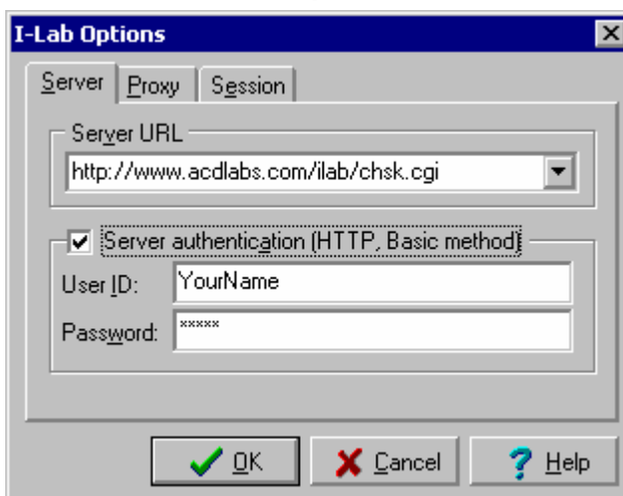


For Public I-Lab: If you do not see the **I-Lab** menu, for example, if you have ChemSketch freeware or have not selected this option during the installation of your purchased copy, you can download the Add-in for free from http://www.acdlabs.com/download/ilab_addon.html. Install the Add-in **in the same directory** where ChemSketch is installed and the next time you start ChemSketch you should see the **I-Lab Login** button and the **I-Lab** menu.

2.2.2 Connection Options

If your computer is on a computer network protected by a *firewall* or *proxy server*, then there are some extra settings required before you can use ChemSketch to access I-Lab.

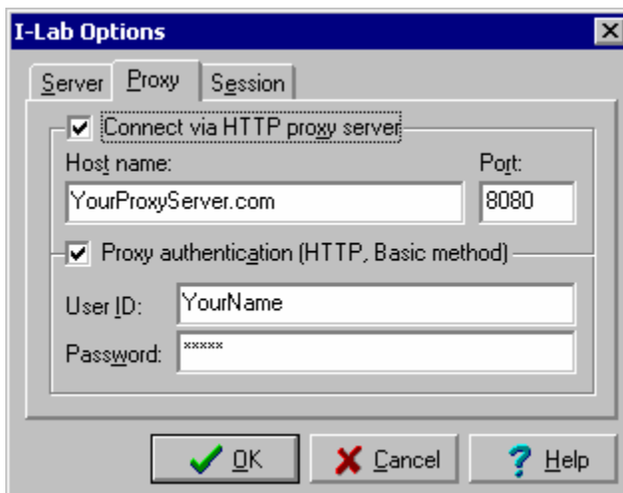
1. From the **I-Lab** menu shown above, choose **Options**:



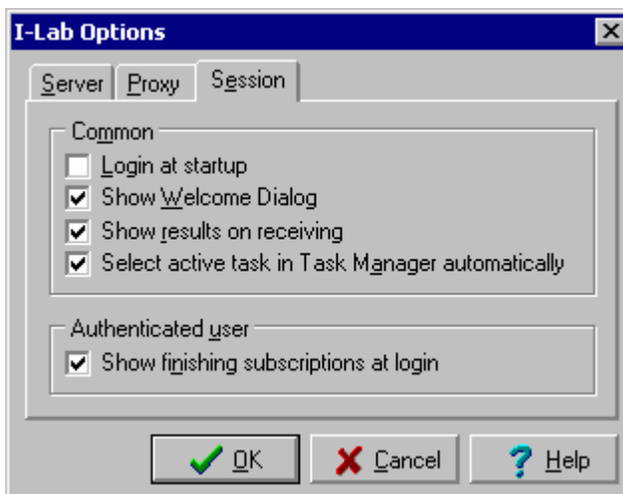
Note In the dialog box image above, the Server URL is set to Public I-Lab URL. If I-Lab: Intranet Edition is installed in your company, ask your I-Lab Administrator for details.

2. In the **Server URL** box, type the I-Lab server URL. Depending on the server currently selected, the **I-Lab** menu will contain a different number of commands (the account-related commands are not displayed if you have selected the I-Lab: Intranet Edition server).
3. To access I-Lab via HTTP server authentication, type your user ID and password in the appropriate boxes. If ACD/I-Lab: Intranet Edition is installed at your company, ask your I-Lab Administrator for the user ID and password.

4. Click the **Proxy** tab and specify your preferences:



5. Fill in the **Host Name of HTTP Proxy:** and **Port:** boxes for your network. This information can be obtained from your network administrator.
6. To access I-Lab via a secured proxy server, type your user ID and password in the appropriate boxes, as they are required for user authentication.
7. Click the **Session** tab, and specify your preferences for I-Lab sessions. Some of the options may be quite useful.
- For example, you can clear the **Show results on receiving** check box if you prefer to perform several calculations without being interrupted and then view the results in the **Task Manager** (from the **I-Lab** menu, choose **Task Manager**).



Note For more information on the **I-Lab Options** dialog box, click **Help** to display the I-Lab Help.

2.2.3 Guest Access

For Public I-Lab only!

The fastest way to have a look at how I-Lab works is to use the Guest Access. In this case, you don't have to register. As a Guest User, you can view the list of available services but can only use the IUPAC Name Free service.

1. On the **I-Lab** menu, click **Login**.
–OR–
Click **I-Lab Login** on the status bar (see above).
2. In the dialog box that appears, click **Guest Login**.
3. Wait for some time (depending on the speed of your Internet connection) until the Welcome message appears.
4. Click **OK** to close the dialog box (or you may click **News** to view the Web page with the latest news from I-Lab).

Now you can view the available services (click **I-Lab** on the status bar or choose the **I-Lab** menu) and use the IUPAC Name Free service.

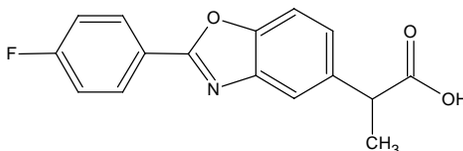
2.3 Generating Name with the IUPAC Name Free Service

Note This chapter provides an overview of the IUPAC Name Free service and is meant for Public I-Lab. However, if you have got ACD/IUPAC Name installed with I-Lab: Intranet Edition, you can follow these instructions to generate an IUPAC name.

As soon as you are logged in, you may start calculations. In this section we are going to describe how to use our free service—IUPAC Name Free.

First, you should draw a structure for which the name should be generated.

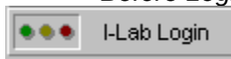
1. Make sure you are in Structure mode and, using the ChemSketch tools, draw the structure of interest. For example,



2. If there are several drawn structures in one page, select the one you want to name.
3. Make sure you are logged in. Check the left part of the status bar. If the Public I-Lab URL is set in the **I-Lab Options** dialog box (see Section 2.2.2 above), it should contain the **I-Lab** button and the “traffic lights” should appear dimmed:

For Public I-Lab:

Before Login:

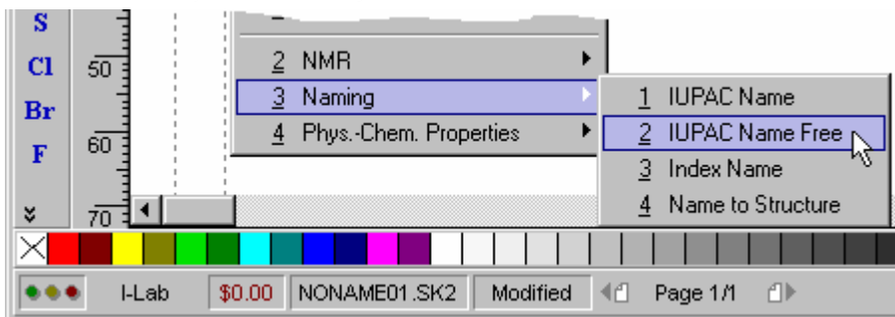


After Login:

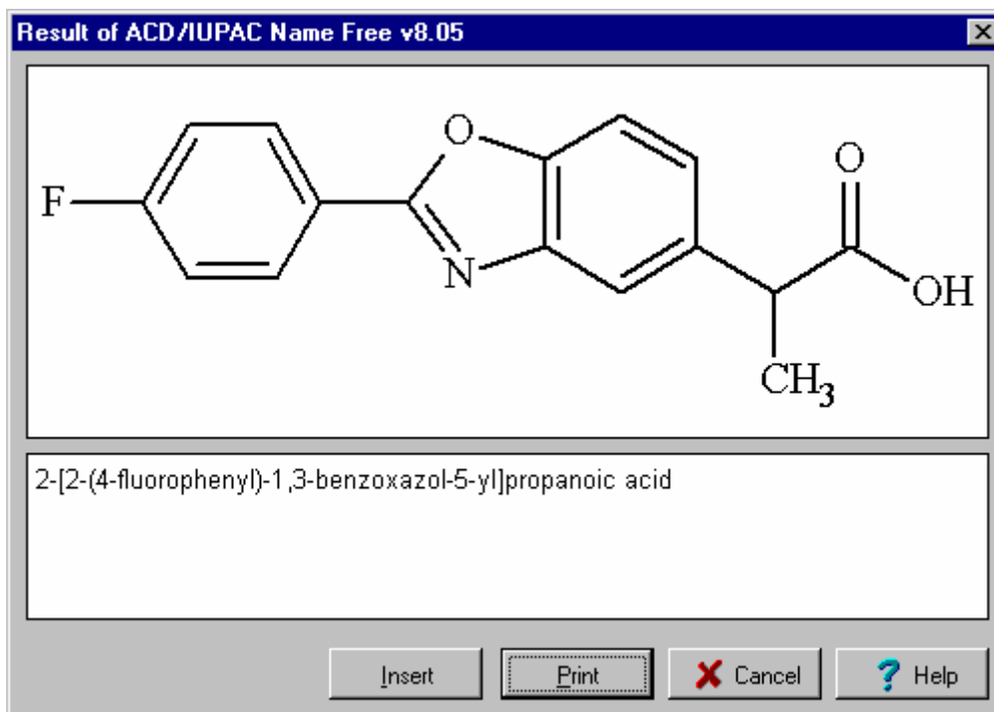


Note The box showing the amount of money you have on your account (if displayed with black) or the amount you owe (with yellow) is not displayed if you are connected by a Site License or as a Guest User or if you are connected to I-Lab: Intranet Edition.

4. Click **I-Lab** on the status bar at the bottom of the ChemSketch window, and on the shortcut menu that appears, point to **Naming**, and then choose **IUPAC Name Free** (or **IUPAC Name** for I-Lab: Intranet Edition). Alternatively, from the **I-Lab** menu, choose **IUPAC Name Free**.



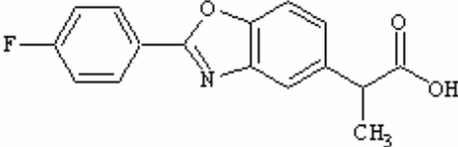
5. A dialog box appears, informing you that no additional parameters are needed. (For other properties, this dialog box may contain other information or options.)
6. Click **Yes**. As soon as your request is processed, the dialog box with the results appears:



Note If the **Show results on receiving** check box is not selected in the **ACD/I-Lab Options** dialog box (from the **I-Lab** menu, choose **Options**), the results dialog box is not displayed, but you can view the results in the **Task Manager (I-Lab** menu).

7. To create a report on the generated name, click **Insert**. The drawn structure, the generated name, and the I-Lab calculation information are laid out on a separate page in the ChemSketch window:

I-Lab Service: **ACD/IUPAC Name Free v8.05**
 Date: **12/Jul/2006** Time: **17:52:52** User:



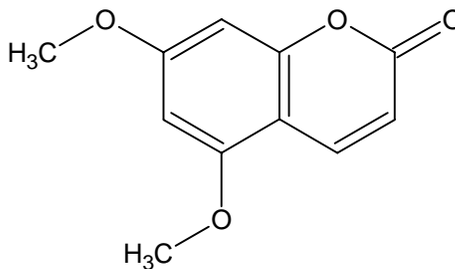
2-[2-(4-fluorophenyl)-1,3-benzoxazol-5-yl]propanoic acid

ACD/IUPAC Name Free v8.05 service is provided by **ACD, Inc.** If you need to quote this information, please include the following: "The result of ACD/IUPAC Name Free v8.05 was obtained using the ACD/I-Lab service".

For more information about **ACD/IUPAC Name Free v8.05** for Windows, please contact ACD (www.acdlabs.com).

Note To refer to an I-Lab calculation in an article, a phrase is suggested in the output page.

8. On the top toolbar, click **New Page**  and draw another structure, for example, this one:



9. Generate its name following steps 4–7.

2.4 Viewing Results (Task Manager)

All of the responses from I-Lab are collected in one place—Task Manager. Using it, you can view the results obtained during the current session.

1. From the **I-Lab** menu, choose **Task Manager**. The dialog box with all of the queries you have made for the current session appears:

ACD/I-Lab Task Manager (Date: 12/Jul/2006; User: Site License User)

Task Name: ACD/IUPAC Name Free
 Charge: free
 Sent Time: 17:51:57
 Received Time: 17:52:06
 Calculation Time: 00:00:09

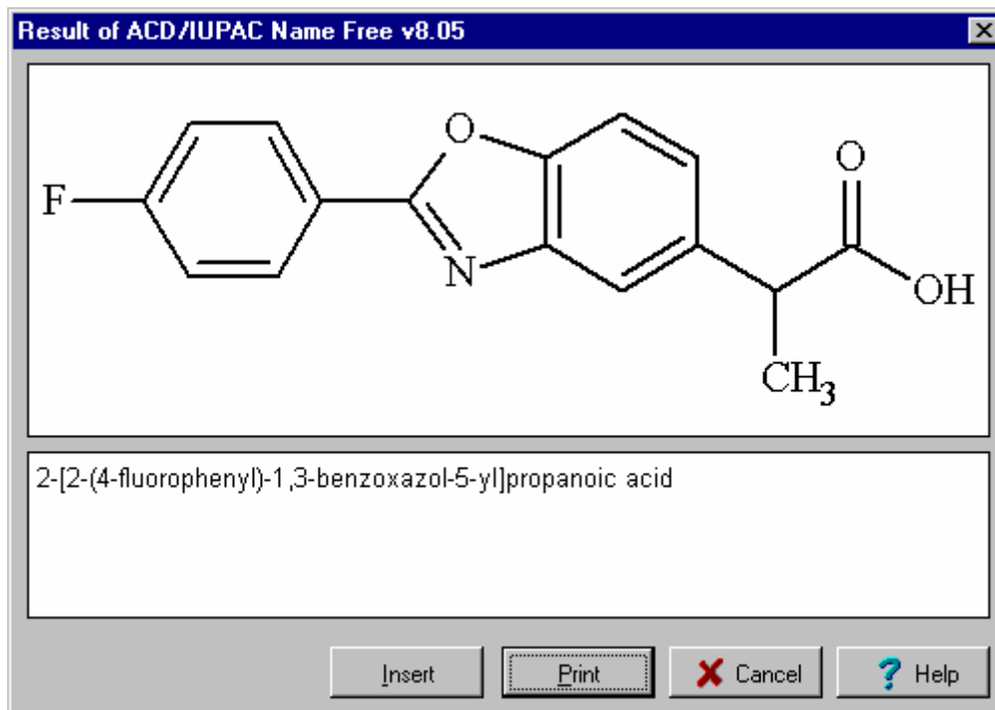
Request Result Delete

Task Name:	Status:	Time:
ACD/IUPAC Name Free v8.05	Results are Received	17:52:06

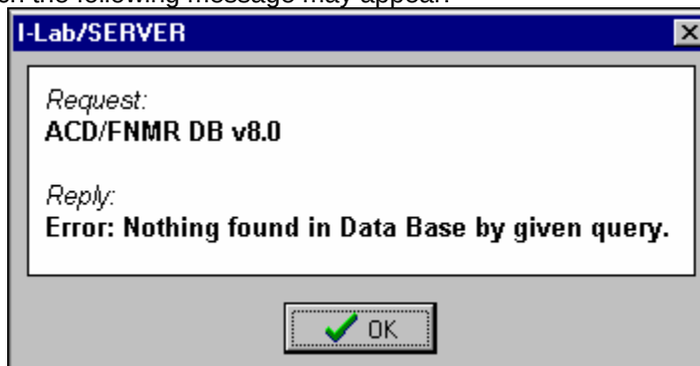
Options Close Help

Note The way the results are received can be controlled by the **Show results on receiving** check box in the **I-Lab Options** dialog box (on the **I-Lab** menu, click **Options**). If you select this check box, the dialog box with the results of the calculation, predictions, or search will appear as they are received. If you keep this check box cleared, the program will just place the received results into the **Task Manager** without informing you about this.

2. To view the results of the query in the **Result of ACD/IUPAC Name Free** dialog box, highlight it in the list, and then click **Result**. For example, for the first IUPAC Name Free query, the following dialog box appears:



Note If the request could not be processed and I-Lab returned an error, it is also reflected in the Task Manager. For such queries, the **Result** button is replaced with the **Error** button. Clicking this button displays the message received from the I-Lab server. For example, if nothing could be found in the CNMR database for your search the following message may appear:



If your request contained any parameters (for example, frequency for HNMR Predictor service), clicking **Request** displays the original request parameters.

3. To close the **Task Manager**, click **Close**.

Important The results are stored during the current session only, so as soon as you log out from I-Lab, the Task Manager contents are cleared.

2.5 Logging Out

To log out, from the **I-Lab** menu, choose **Logout**.

Please note the following:

- When you quit the ChemSketch window, you are logged out automatically. But if ChemSketch has been abnormally terminated (e.g., because of a system crash), your account will be logged off after a certain period of inactivity.
- The list of tasks in the Task Manager is deleted when you log out.

3. Using I-Lab via ChemSketch

3.1 Objectives

In this chapter you will find a description of the I-Lab features available in the ChemSketch interface shown in more detail than in the *Quick Start* chapter. You will learn:

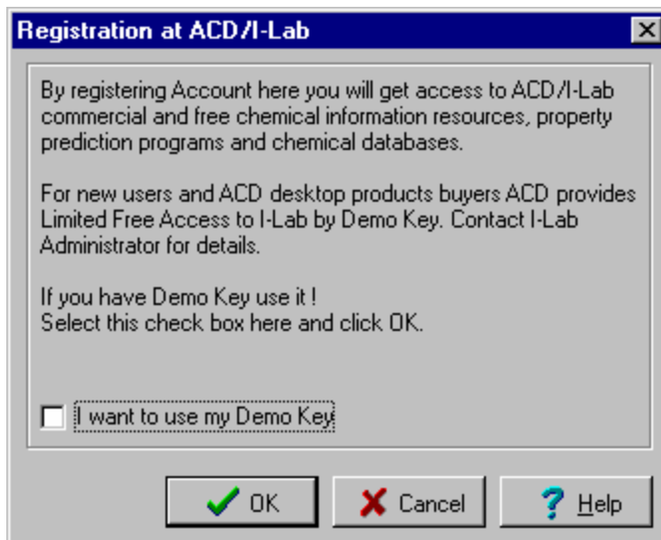
- How to register on I-Lab and activate your I-Lab account;
- How to obtain the demo key;
- How to log in;
- How to calculate various properties;
- How to search databases; and
- How to use subscriptions.

3.2 Registering

For Public I-Lab only!

To be able to work with the commercial services of I-Lab and to be able to use the Demo Key you should first register at I-Lab and then set up your own account.

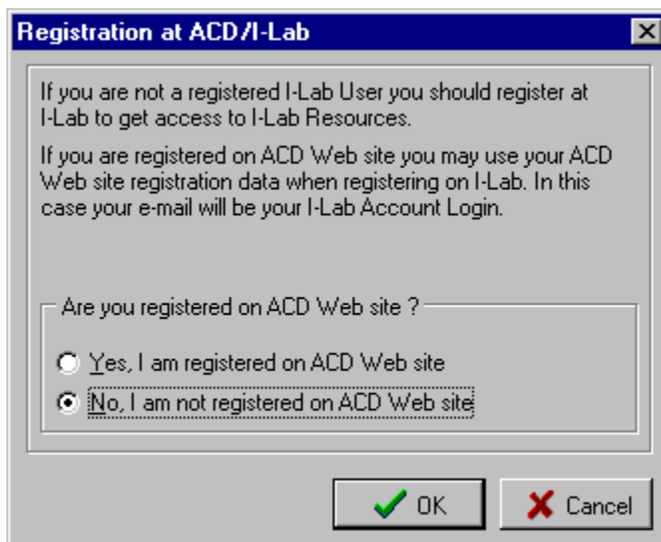
1. If you are logged in to I-Lab as a Guest, from the **I-Lab** menu, choose **Log Out** to log out.
2. From the **I-Lab** menu, choose **Register at I-Lab**. The following dialog box appears:



3. If you do not have the Demo Key available (for more information, refer to Section 3.3), clear the **I want to use my Demo Key** check box and click **OK**.

Note If you want to use your Demo Key, select the corresponding check box. In this case the registration procedure described below will be slightly different: an additional dialog box asking you to provide the Demo Key will appear.

4. Read the license agreement that appears next and, if you agree with its terms, choose **Agree**.
5. The next dialog box will ask you if you are already registered at the ACD/Labs Web site. Choose **No**.



6. Click **OK**. The **ACD/I-Lab Account Registration Form** appears where you should specify your details. Items marked with a red dot are required fields:

The screenshot shows the 'ACD/I-Lab Account Registration Form' dialog box. At the top, it says 'To finish/update registration fill the form and press OK'. The form has the following fields:

- Title: Dr. (dropdown)
- First Name: Mary
- Last Name: Brown (marked with a red dot)
- Login: brown (marked with a red dot)
- Password: [masked with asterisks] (marked with a red dot)
- E-mail address: maru.brown@mail.com (marked with a red dot)
- Position: [empty]
- Phone: 123-45-67-89
- Company: [empty]
- Fax: [empty]
- Country: Australia (dropdown)
- Province: [empty]
- City: Melbourne (marked with a red dot)
- Address: Queen St, 8 (marked with a red dot)
- Postal Code: [empty]
- Referrer: E-mail Newsletter (dropdown)

At the bottom, there is a message: 'Fields marked by [red dot] are required !'. Below this are three buttons: OK (with a green checkmark), Cancel (with a red X), and Help (with a blue question mark).

7. When you click **OK**, you are asked to re-enter your password and are shown the information you have just entered. Check the details and if everything is correct, click **OK**.
8. An e-mail message will be automatically sent to the e-mail address you provided. The message will have the subject "Your ACD/I-Lab activation key" and will contain the activation key, as well as remind you of your login and password.

3.3 Demo Keys

For Public I-Lab only!

If you like what you've seen during Guest access and IUPAC Name Free calculation, consider moving on to the next step: a demo key.

Demo keys allow you to try out the I-Lab services for a limited period of time. During this demo period, you will have access to all of the I-Lab services free of charge. After the demo period expires, your I-Lab account will be charged on a per transaction basis.

Demo keys are for a two-week (14 days) period. A distributor or sales representative may request a demo key on behalf of a client for longer periods of time.

Demo keys must be generated by the I-Lab Administrator. When a demo key request comes in, it requires human intervention. It is not an automated process such as the generation of the account activation key.

Only one demo key is issued per account per year. Typically the demo keys are given to newly opened accounts. The demo keys must be requested. They are not applied automatically.

Demo keys only have to be applied once! They do not have to be entered every time to gain demo access. Once it has been successfully applied to the account, the user just has to login normally.

If there are problems applying a demo key to an I-Lab account, the I-Lab Administrator can manually set up the demo period for the account. To contact the I-Lab Administrator, send an e-mail to ilab@acdlabs.com.

3.3.1 To Obtain the Demo Key

1. Make sure you are not logged in to I-Lab. If you are, from the **I-Lab** menu, choose **Log Out**.
2. From the **I-Lab** menu, choose **Request Demo Key**. Before requesting the Demo Key, you should register at I-Lab. As you click the **Request Demo Key** command, the ACD/Labs Request Demo Key browser is activated where you can type your Login and Password in the corresponding boxes. Click **Send Request** to apply and **Reset the Form** to change the settings.
3. In a few seconds, you will receive a Demo Key.

3.4 Activating Your I-Lab Account

For Public I-Lab only!

Once you have registered and received the activation key you must activate your account.

1. From the **I-Lab** menu, choose **Activate Account** to display the following dialog box:



2. Type the login and activation key you received in the e-mail message, in the **ACD/I-Lab Activation Account** dialog box. Click **OK** to start activation.

Tip Use CTRL+C and CTRL+V to copy and paste the activation key from your e-mail message directly into this dialog.

3. You should get a confirmation that the activation key was applied successfully. Your I-Lab account is now ready to be used. *This is the only time you need to use the activation key.*

3.5 Logging In

As soon as you register and activate your account, you can log in.

1. From the **I-Lab** menu, choose **Login** or click **I-Lab Login** on the status bar.
2. *For Public I-Lab:* In the dialog box that appears, type your login name and password.
For I-Lab: Intranet Edition: You are instantaneously logged in to I-Lab (skip the note and step 3 below).

Note In case you want to log in as a guest, simply click the corresponding button without providing any login or password.

3. Click **OK**. The confirmation dialog box will appear when you are successfully logged in to I-Lab.

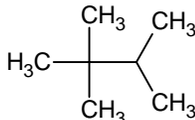
Note If connection is not possible, an error message will appear in the dialog box. You may have to try again later.

4. You are ready to select any one of the available calculations. Choose the service you would like from the **I-Lab** menu. Keep in mind that a **Confirm your request** dialog box will appear (if the **Ask confirmation for calculation request** check box is selected in the **I-Lab Options** dialog box—see Section 2.2.2), in which you can select **No**, if you decide not to proceed.

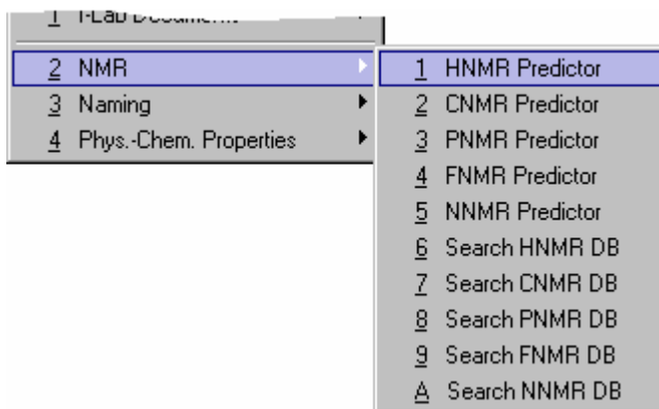
3.6 Calculating Properties

3.6.1 Predicting a Spectrum

1. Draw a structure for which you want to predict a spectrum, for example, the following one:

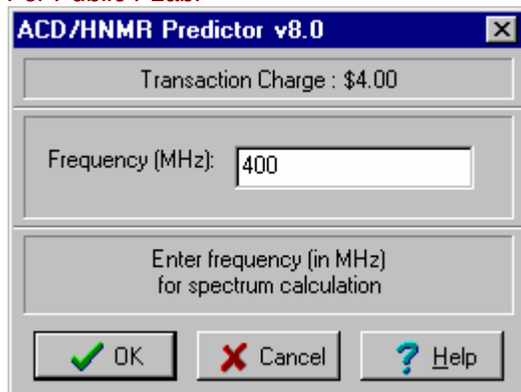


2. If there is more than one structure drawn in the workspace, select the required one.
3. On the **I-Lab** menu, point to **NMR**, and then choose the type of spectrum you want to calculate: HNMR, CNMR, PNMR, or FNMR. For example, if you want to calculate the HNMR spectrum, choose **HNMR Predictor**:

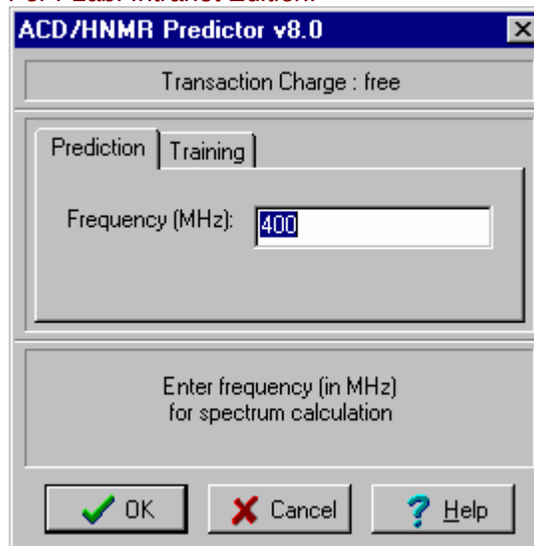


4. The dialog box containing the price for the transaction and parameters that can be specified appears. Note that some predictions do not require any additional parameters. In case of the HNMR spectrum calculation, the following dialog box appears:

For Public I-Lab:

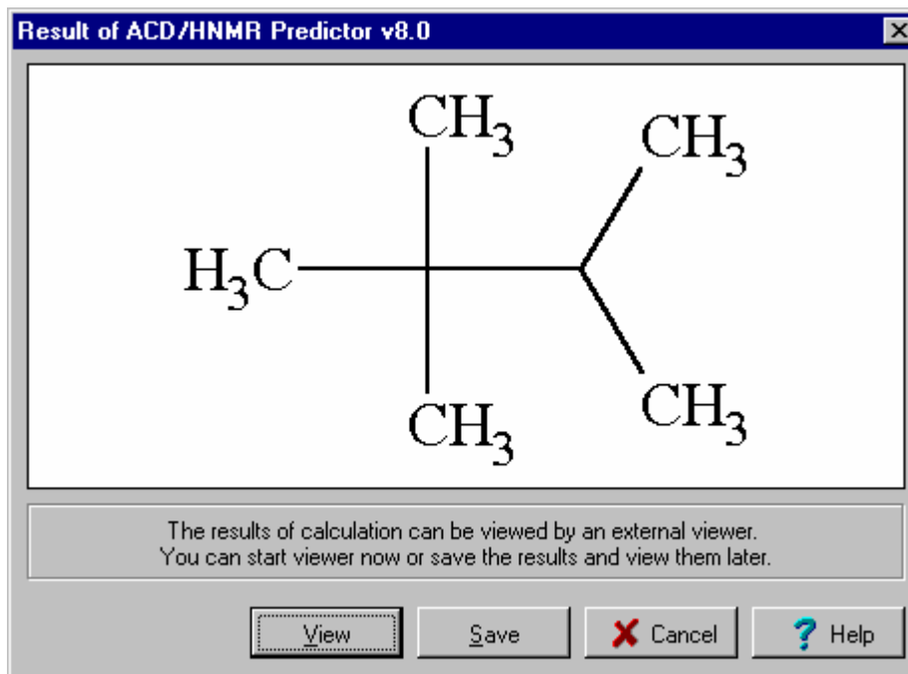


For I-Lab: Intranet Edition:



As you see, the dialog box that appears if you are running ACD/I-Lab: Intranet Edition contains two tabs: the **Prediction** tab where you can specify the appropriate frequency, and the **Training** tab that will be described in Section 3.6.1.1 below.

5. Click **Yes** to complete the request. As soon as the results arrive, they are displayed in the following dialog box:



Note If the **Show results on receiving** check box is not selected in the **ACD/I-Lab Options** dialog box (from the **I-Lab** menu, choose **Options**), this dialog box is not displayed, but you can find the results in the **Task Manager (I-Lab** menu).

6. To view the resultant spectrum, click **View**. To save it as a file in .HSP format, click **Save**.

Important To be able to view the spectrum you should have CNMR and/or HNMR Viewer installed. These programs can be downloaded for free from our Web site. More information about this software can be found at: http://www.acdlabs.com/download/c_h_view.html.

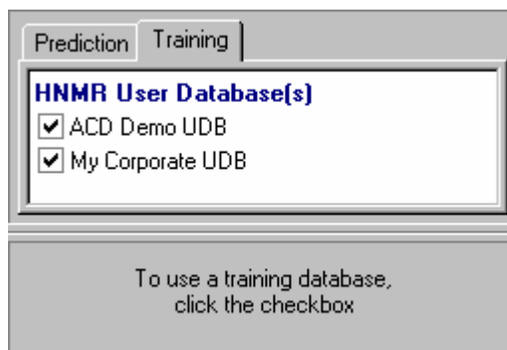
In the case of PNMR and FNMR spectra, the results are represented in the form of the table and can be printed out or placed into the ChemSketch pad as a report.

More details on predictors can be found in Chapter 4, *ACD/I-Lab Services*.

3.6.1.1 System Training

For I-Lab: Intranet Edition only!

ACD/I-Lab: Intranet Edition allows you to select (an) additional user database(s) that will be used along with the internal database for predictions:



ACD/I-Lab: Intranet Edition contains a "self-training" system, which increases the accuracy of prediction, the more you add data to your own user's database. If you are not satisfied with the accuracy of spectra calculated for a new class of compounds, you can create and/or update with the help of your system administrator your own database with the experimental or literature chemical shifts. The next time you use this database and make calculations for any new compound, the program automatically uses the information from both this database, as well as system databases, so that the accuracy of calculations is increased.

I-Lab Intra can improve the accuracy of your predictions by training the algorithm with your own data. The following services support the training feature: CNMR, HNMR, LogP, pK_a, LogD, Adsorption Coefficient, and Bioconcentration Factor.

The training feature is available after I-Lab is installed. It contains sample databases. I-Lab Intra includes a simple administrative interface allowing you to add custom training databases. Adding as well as enabling/disabling custom databases is the administrator's privilege.

3.6.2 Calculating PhysChem Properties

1. Draw or import the structure for which you want to calculate a property. If there is more than one structure in the drawing area, select the required one by clicking near it with the **Select** tool active.

Note For the Naming service (IUPAC Name and Index Name), you can select several structures to be named.

2. From the **I-Lab** menu, select the required property (e.g., Boiling Point). The dialog box appears. Depending on the property selected it may contain parameters that are necessary for the property to be calculated (e.g., Pressure, Pressure and Temperature Units for Boiling Point). You can also find information on the price of the calculation displayed at the top of this dialog box.
3. As soon as the parameters are specified (if any), click **OK**.
4. When the properties are calculated and the results are received, you can go to the **Task Manager** dialog box (from the **I-Lab** menu, choose **Requests Manager**) to view the results.
Note If the **Show results on receiving** check box in the **ACD/I-Lab Options** dialog box is selected, the **Result** dialog box will be displayed as soon as the results are received.
5. In the Task Manager, if you have several tasks in the list, click the required one. Then click **Result** to open the dialog box containing the query structure and the calculations. (If the property could not be calculated for any reason, the **Result** button is replaced with the **Error** button by clicking on which you can view the reason of error).
6. To insert the results into the ChemSketch window, click **Insert**.

3.7 Searching a Database

3.7.1 Similar Structure Search

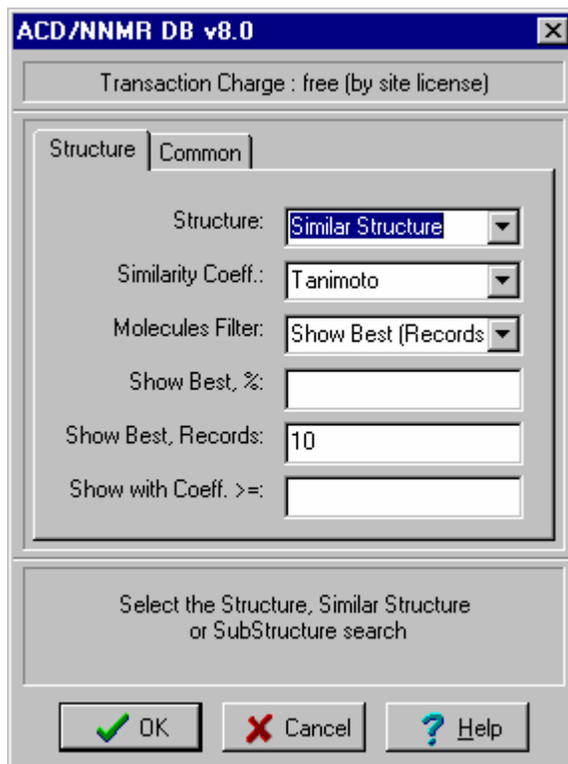
One of the services provided by ACD/I-Lab is the capability to search chemical databases for various structures and parameters. Version 7.0 and later allows you to perform a so-called similar structure search. The output from the search is a ranked list in which the structures that are assumed to be the most similar to the target structure are located at the top of the list. These most similar structures will be those that have the greatest probability of being of interest to the user, given an appropriate measure of intermolecular structural similarity. The question is how to assess the degree of similarity—for that purpose, so called similarity coefficients are used. For more information on the similarity coefficients, see Section 3.7.1.1 below.

1. In the ChemSketch window, click **Table of Radicals** on the rightmost toolbar, and then select benzene.
2. Click in the workspace to draw the corresponding structure:



3. If there is more than one structure on the current page, select the required one using the **Select/Move** tool  on the Structure toolbar.
4. On the **I-Lab** menu, point to **NMR**, and then choose **Search NNMR DB**. The **ACD/NNMR DB** dialog box appears.

5. In the **Structure** box, select **Similar Structure**, and in the **Similarity Coeff.** box, select **Tanimoto**. For more detail on the similarity coefficients, see Section 3.7.1.1 below.



Note All other boxes are optional. It is very convenient to restrict your search conditions by setting a molecular filter. You can specify a percentage or a number of records best matching your query in the **Show Best (%)** and **Show Best (Records)** boxes. Alternatively, you can set the threshold for the selected similarity coefficient in the **Show with Coeff.** box, and the records with a similarity coefficient exceeding the set value will appear in the search results list.

Please check if the filter selected in the **Molecules Filter** list is correct.

6. In the **Molecules Filter** list, select **Show Best (Records)**, and then type "10" in the **Show Best (Records)** box.

Note You can set additional restrictions on the **Common** tab.

7. Click **OK**.

8. As soon as the results are received, the **Results of ACD/NNMR DB v8.0** dialog box appears containing the 10 records best matching the query:

Result of ACD/NNMR DB v8.0

Formula: C₇H₉N

Molecule Name: 1-phenylmethanamine

Molecule Weight: 107.15

Nominal Mass: 107

Average Mass: 107.1558

Exact Mass: 107.0735

Chemical Shifts:

Nucl.	Atom No.	Chem. Shift (ppm)	R
15N	8	-357.30	
15N	8	-356.00	

Structures: 10

1. C7H9N
2. C7H5N
3. C7H6N
4. C7H10N
5. C13H12N

Buttons: Previous, Next Search, Insert, Print, Close

Note If the **Show results on receiving** check box in the **ACD/I-Lab Options** dialog box is not selected, the results are not displayed, but can be found in the **Task Manager** dialog box (**I-Lab** menu).

9. Browse through the entries by clicking the lines in the **Structures** list.
10. If you want to insert any entry into the ChemSketch window, highlight it in the list and click **Insert**.

3.7.1.1 Similarity Coefficients

The **similarity coefficient** should be used to provide a quantitative measure of structural relatedness between a pair of structural representations. The similarity coefficient determines a numerical measure of similarity (or conversely, the distance) between two objects, each characterized by a common set of attributes. Now we are going to review the coefficients that have found widespread use in chemical information systems.

Each structure is represented as a binary vector containing n attributes.

Let S_{AB} and D_{AB} be similarity and distance between the structures A and B respectively.

Let $X_A = \{x_{1A}, x_{2A}, \dots, x_{jA}, \dots, x_{nA}\}$ and $X_B = \{x_{1B}, x_{2B}, \dots, x_{jB}, \dots, x_{nB}\}$ be binary vectors describing the structures A and B respectively.

$X_{iA} = 1$ ($X_{iA} = 0$) if the structure object number i is present (absent) at the structure A . Structure objects are generated automatically depending on DB structures type.

Let S_{AB} and D_{AB} be similarity and distance between the structures A and B respectively.

Let:

$$a = \sum_{j=1}^n x_{jA} \quad b = \sum_{j=1}^n x_{jB} \quad c = \sum_{j=1}^n x_{jA}x_{jB}$$

Let m be the value of the **Minimum Coefficient** list. The following table describes how Web DB searches records by similarity for each item of the **Similarity Coefficient** list:

Coefficient	Parameter	Condition
Tanimoto	$S_{AB} = \frac{c}{a+b-c}$	$S_{AB} \geq m$
Dice	$S_{AB} = \frac{2c}{a+b}$	$S_{AB} \geq m$
Cosine	$S_{AB} = \frac{c}{\sqrt{ab}}$	$S_{AB} \geq m$
Based on Hamming Distance	$D_{AB} = 1 - \frac{a+b-2c}{n}$	$D_{AB} \geq m$
Based on Euclidian Distance	$D_{AB} = 1 - \frac{\sqrt{a+b-2c}}{n}$	$D_{AB} \geq m$

If the condition is fulfilled, the record containing structure B will be shown as a result of the search.

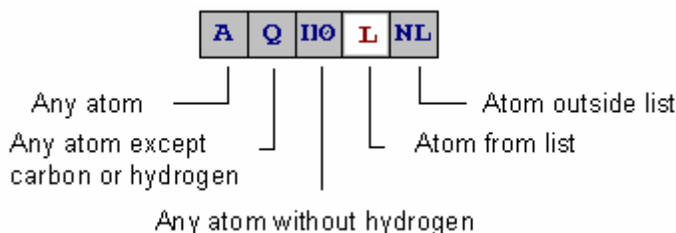
The program shows records in descending order by the coefficient.

Reference: Willet, Peter, John M. Barnard, and Geoffrey M. Downs. "Chemical Similarity Searching." J. Chem. Inf. Sci. 1998, 38, 983–996.

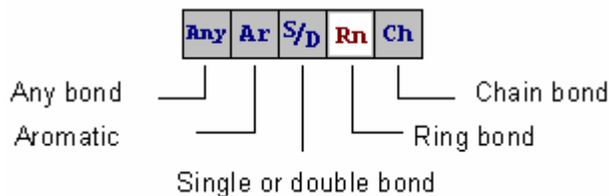
3.7.2 Substructure Search

If you want to search for a "general" structure, use the Query buttons to modify the drawn structure according to search criteria. These buttons are located at the top of the left vertical toolbar in the ChemSketch window as **A** and **Any**. Click the triangle on the lower right corner of the **A** and **Any** buttons to get all the query atoms and query bonds. They allow you to do the following:

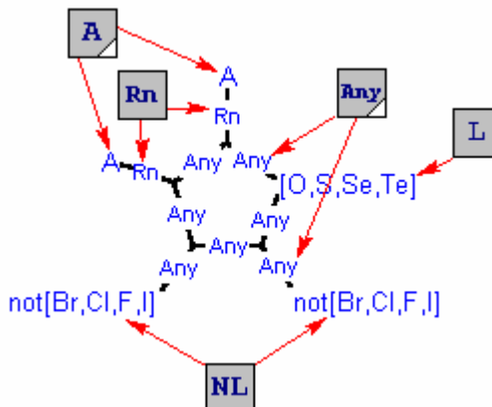
Query Atoms



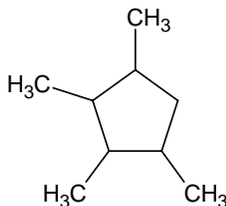
Query Bonds



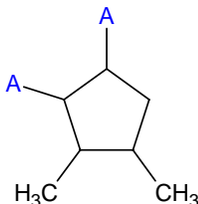
For example, a search according to the following query structure will result in all condensed furans, thiophenes, and their derivatives with no halogen substituents:



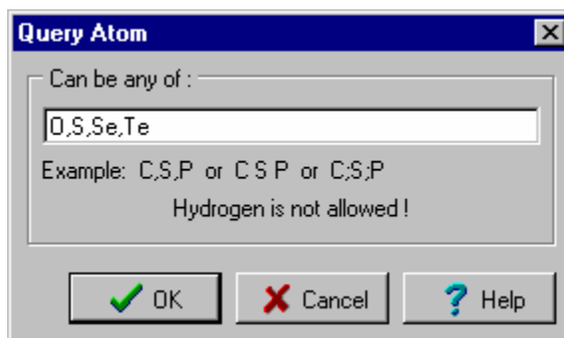
1. First draw the following structure:



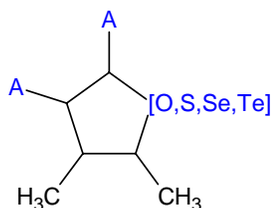
2. Click **Any Atom**  and then on the two atom groups as shown:



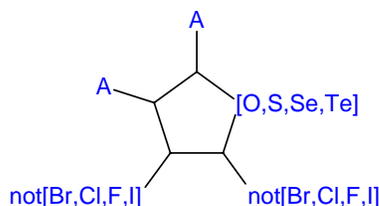
3. Now click the right bottom triangle of **Any Atom** and, from the set of buttons that appears, click **Atom from List** , then click on the right most group of atoms. In the **Query Atom** dialog box, type **O,S,Se,Te**:



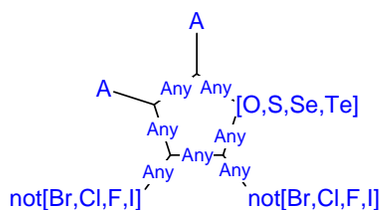
4. Click **OK**. You should obtain the following:



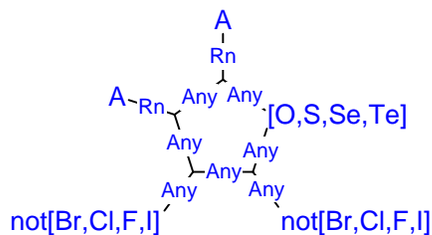
5. Click **Atom Not in List**  and specify the list of elements that should not be present for other groups of atoms:



6. Now we are going to define bonds. Click **Any Bond**  and then on the bonds of the structure to get the following:

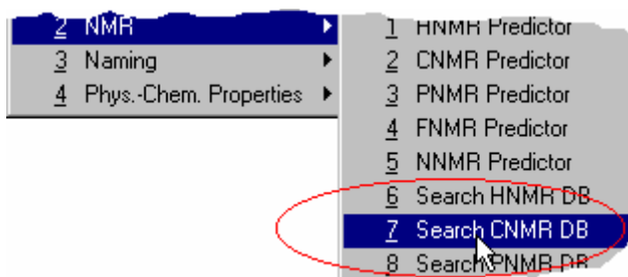


7. Choose **Ring Bond**  and click the bonds to define them:

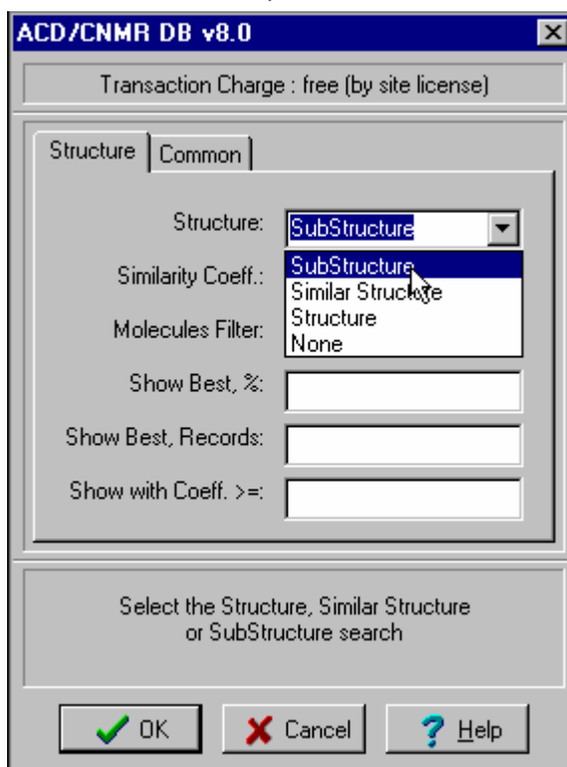


8. If there is more than one structure in the drawing area, select the required one by clicking near it with the **Select/Move** tool  active.

9. On the **I-Lab** menu, point to **NMR**, and then select the database you want to search through:



10. The dialog box appears where you can specify your query. If there is a structure drawn in the workspace, the dialog box contains the **Structure** option that allows you to specify whether you want to search by the whole structure or substructure. If you have used the query buttons, you must select the **SubStructure** option, otherwise an error will be returned:

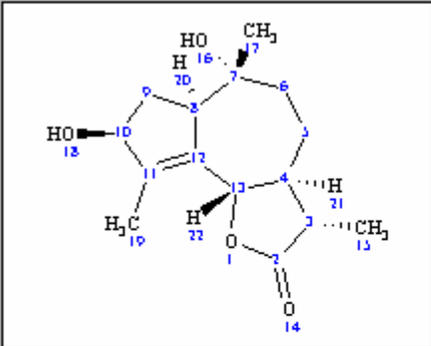


Note You can restrict your search by specifying molecular formulae, molecular weight, chemical shifts, name substring, references, or all of them.

If you don't want to use the structure as a criterion for the search but only want to use other properties, select the **None** option in the **Structure** box.

11. After you specify all of the parameters, click **OK**. As soon as the results are received, the dialog box containing the first 15 found entries is displayed:

Result of ACD/CNMR DB v8.0



Formula: C₁₅H₂₂O₄

Molecule Name:
6,8-dihydroxy-3,6,9-trimethyl-3a,4,5,6,6a,7,8,9b-octahydro[5-b]furan-2(3H)-one

Molecule Weight: 266.33

Nominal Mass: 266

Average Mass: 266.3382

Exact Mass: 266.1518

Chemical Shifts:

Atom No.	Chem. Shift (ppm)	Refs.
2	177.80	1

Structures: 15

1. C15 H22 O4
2. C15 H24 O4
3. C21 H38 O4 Si2
4. C21 H38 O5 Si2
5. C21 H38 O5 Si2

Previous Next Search Insert Print Close

Note If the **Show results on receiving** check box in the **ACD/I-Lab Options** dialog box is not selected, the results are not displayed, but can be found in the **Task Manager** dialog box (**I-Lab** menu).

12. Browse through the entries by clicking on the lines in the **Structures** list.
13. If the desired entry is not among the first 15 found ones, click **Next Search**.

Important Obtaining the next set of entries is considered as a new transaction and is charged as well.

14. If you want to insert any entry into the ChemSketch window, highlight it in the list and click **Insert**.

3.8 Subscribing for Services

For Public I-Lab only!

If you plan to do a series of predictions or searches, ACD/I-Lab contains a great money-saving service—subscriptions. If you subscribe to any of the services (search, prediction, or calculation), you can use the corresponding modules as many times as you wish during the specified period for a flat fee. To subscribe to any of the available services:

1. On the **I-Lab** menu, point to **Account Management**, and then choose **Subscription Wizard**. The first step—**How to subscribe**—gives short instructions on how to use the wizard. Read it carefully and click **Next**.

2. The second step—**Service selection**—lists all of the services available via I-Lab. Choose the one you require and click **Next**.
3. The third step—**Subscription selection**—shows the subscriptions that contain the selected service. The upper part of the dialog box lists the subscriptions and the lower part displays the list of services available in the selected subscription. Choose the subscription in the upper part and click **Next**.
4. The fourth step—**Duration selection**—allows you to choose among the types of the selected subscription that differ by their duration. As usual, there are daily and monthly subscriptions. Click the required subscription in the list. In the upper part you can see the price. Click **Next**.
5. At the last step—**Confirmation**—the summary of the selected subscription is displayed and you should confirm your choice by clicking **Finish**.
6. Wait until the **Active subscriptions** dialog box appears informing you of the services you are subscribed for. If you wish to subscribe for more services, click **Next**. If you don't, click **Cancel**.

Note If the **Show finishing subscriptions at login** check box is selected in the **ACD/I-Lab Options** dialog box (**I-Lab** menu), you will be informed of the current subscriptions upon each login.

4. I-Lab Services

4.1 Objectives

In this chapter you will find a list of services available through I-Lab with the corresponding description and statistical information.

4.2 HNMR Predictor

ACD/HNMR Predictor uses parameters gleaned from its internal database with over 1,384,000 experimental chemical shifts and 449,000 experimental coupling constants for 165,000 structures. To quantitate intramolecular interactions in new organic structures and to predict their chemical shifts, HNMR Predictor module uses a unique algorithm based on intramolecular interaction parameters for over 3,000 structural fragments, which have been carefully derived from experimental data using ACD/Labs' specially developed HNMR data processing approaches.

ACD/HNMR:

- Calculates accurate HNMR spectra under any basic frequency. The most accurate spectra are obtained for neat liquid samples or solutes in non-polar non-aromatic solvents. Calculated chemical shifts and coupling constants are provided with 95% confidence intervals so that you know the reliability of the calculated values.
- Takes into account second order interactions (strong coupling effects) and long-range coupling constants.
- Simulates double resonance and plots integral curves.
- Allows you to calculate exact spectra for any strongly coupled system with up to 8 magnetically non-equivalent nuclei (spin 1/2) or more if some nuclei are magnetically equivalent.

You can specify the frequency for the spectrum to be calculated.

If you are using I-Lab via ChemSketch, the results of HNMR spectrum prediction can be viewed from a special dialog box that appears automatically as soon as the results are received from the server (if the corresponding check box is selected in the **ACD/I-Lab Options** dialog box) or that can be opened from the **Task Manager** by clicking **Result**. The results are represented in the form of a spectrum that can be viewed with the external viewer (HNMR Viewer) that is included into the ChemSketch installation package by default.

4.3 CNMR Predictor

CNMR Predictor quickly and precisely calculates ^{13}C NMR spectra for almost any drawn organic structure. The calculated chemical shifts are always provided with the confidence limits, so that you always know how far the calculated values can be trusted.

In most cases, the CNMR Predictor module calculates chemical shifts with an accuracy of 3 ppm or better. For this purpose, the module uses a large internal data base with over 2,017,000 experimental chemical shifts and 81,400 coupling constants for 165,000 structures. To quantitate intramolecular interactions in new organic structures and to predict their chemical shifts, CNMR Predictor module uses a unique algorithm based on the intramolecular interaction parameters for over 3,000 structural fragments, which have been carefully derived from the experimental data using the specially developed CNMR data processing approaches. It calculates the spin-spin interaction of carbon nuclei with the magnetic nuclei of other elements, proceeding from the natural ratio of magnetic isotopes. Calculated chemical shifts and coupling constants are provided with 95% confidence intervals so that you know the reliability of the calculated values.

ACD/CNMR recognizes spectral differences between the following types of isomeric structures:

- Cis-trans isomers of alkenes;
- Cis-trans and endo-exo isomers of cyclic structures; and
- Syn-anti isomers of amides, oximes, hydrazones, and nitrosoamines.

The results of CNMR spectrum prediction can be viewed from a special dialog box that appears automatically as soon as the results are received from the server (if the corresponding check box is selected in the **ACD/I-Lab Options** dialog box) or that can be opened from the **Task Manager** by clicking **Result**.

The results are represented in the form of a spectrum that can be viewed with the external viewer (CNMR Viewer) that is included into the ChemSketch installation package by default.

4.4 PNMR Predictor

ACD/PNMR Predictor accurately calculates ^{31}P NMR chemical shifts and coupling constants for many typical chemical structures containing P. It uses parameters gleaned from its internal ^{31}P database, which contains over 28,200 chemical shifts and 19,400 coupling constants for over 22,600 structures.

The resultant table of chemical shifts includes the following information:

- The type of nucleus giving rise to the nuclear magnetic resonance (here, it is ^{31}P);
- The number of the atom in the structure which gives rise to the predicted shift;
- The value of the predicted shift; and
- The uncertainty of the predicted shift, based on 95% confidence limits for the structure fragment.

The table of coupling constants displays:

- The type of nucleus giving rise to the nuclear magnetic resonance coupling (here, it is ^{31}P);
- The numbers of the two atoms in the structure which give rise to the coupling;
- The value of the predicted coupling; and
- The uncertainty of the predicted coupling, based on 95% confidence limits for the structure fragment. Note that this is usually larger than the uncertainty of the predicted chemical shift.

4.5 FNMR Predictor

ACD/FNMR Predictor accurately calculates ^{19}F NMR chemical shifts and coupling constants for many typical chemical structures containing F. It uses parameters gleaned from its internal ^{19}F database, which contains over 29,300 chemical shifts and 20,600 coupling constants for over 13,800 structures.

The resultant table of chemical shifts includes the following information:

- The type of nucleus giving rise to the nuclear magnetic resonance (here, it is ^{19}F);
- The number of the atom in the structure which gives rise to the predicted shift;
- The value of the predicted shift; and
- The uncertainty of the predicted shift, based on 95% confidence limits for the structure fragment.

The table of coupling constants displays:

- The type of nucleus giving rise to the nuclear magnetic resonance coupling (here, it is ^{19}F);
- The numbers of the two atoms in the structure which give rise to the coupling;
- The value of the predicted coupling; and
- The uncertainty of the predicted coupling, based on 95% confidence limits for the structure fragment. Note that this is usually larger than the uncertainty of the predicted chemical shift.

4.6 NNMR Predictor

ACD/NNMR Predictor accurately calculates ^{15}N NMR chemical shifts and coupling constants for many typical chemical structures containing N. It uses parameters gleaned from its internal ^{15}N database, which contains over 18,500 chemical shifts and 4400 coupling constants for over 8,000 structures.

The resultant dialog box displays the query structure and the table of calculated chemical shifts and coupling constants.

The table of chemical shifts includes the following information:

- The type of nucleus giving rise to the nuclear magnetic resonance;
- The number of the atom in the structure which gives rise to the predicted shift;
- The value of the predicted shift; and
- The uncertainty of the predicted shift, based on 95% confidence limits for the structure fragment.

The table of coupling constants display:

- The type of nucleus giving rise to the nuclear magnetic resonance coupling;
- The numbers of the two atoms in the structure which give rise to the coupling;
- The value of the predicted coupling.

The uncertainty of the predicted coupling, based on 95% confidence limits for the structure fragment. Note that this is usually larger than the uncertainty of the predicted chemical shift.

4.7 Search CNMR DB

This tool allows you to search the ^{13}C NMR database of over 2,017,000 observed chemical shifts for 165,000 structures.

Each database entry contains:

- Chemical structure
- Formula
- Name
- Molecular weight
- Chemical shifts
- References

You can perform searches according to the following parameters:

- Chemical substructure
- Structure
- Similar structure
- Formula
- Molecular weight
- Chemical shifts
- Name substring
- References

4.8 Search HNMR DB

This tool allows you to search the ^1H NMR database of over 1,384,000 experimental chemical shifts and 449,000 experimental coupling constants for over 165,000 structures.

Each database entry contains:

- Chemical structure
- Formula
- Name
- Molecular weight
- Chemical shifts
- Coupling constants
- References

You can perform searches according to the following parameters:

- Chemical substructure
- Structure
- Similar structure
- Formula
- Molecular weight
- Chemical shifts
- Coupling constants
- Name substring
- References

4.9 Search FNMR DB

This option allows you to search ACD/FNMR database that includes more than 13,800 fluorine-containing compounds with experimental shifts and coupling constants.

Each database entry contains:

- Chemical structure
- Formula
- Name
- Molecular weight
- Chemical shifts
- Coupling constants
- References

You can perform searches according to the following parameters:

- Chemical substructure
- Structure
- Similar structure
- Formula
- Molecular weight
- Chemical shifts
- Coupling constants
- Name substring
- References

4.10 Search PNMR DB

This option allows you to search the ACD/PNMR database that includes more than 22,600 phosphorus-containing compounds with experimental shifts and coupling constants.

Each database entry contains:

- Chemical structure
- Formula
- Name
- Molecular weight
- Chemical shifts
- Coupling constants
- References

You can perform searches according to the following parameters:

- Chemical substructure
- Structure
- Similar structure
- Formula
- Molecular weight
- Chemical shifts
- Coupling constants
- Name substring
- References

4.11 Search NNMR DB

This option allows you to search the ACD/NNMR database that includes more than 8,200 nitrogen-containing compounds with experimental chemical shifts and coupling constants.

Each database entry contains:

- Chemical structure
- Formula
- Name
- Molecular weight
- Chemical shifts
- Coupling constants
- References

You can perform searches according to the following parameters:

- Chemical substructure
- Structure
- Similar structure
- Formula
- Molecular weight
- Chemical shifts
- Coupling constants
- Name substring
- References

4.12 IUPAC Name

This module generates the systematic name for general organic compounds, according to guidelines specified by the International Union of Pure and Applied Chemistry (IUPAC).

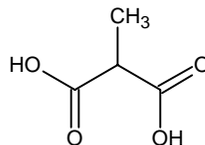
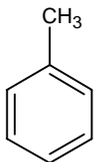
Note that the IUPAC Name Free service (with a set of restrictions on naming) is also available for Public I-Lab.

When you choose this property from the menu, the dialog box appears where you can specify the options described in the following sections.

4.12.1 Common Tab

4.12.1.1 Use Retained Replacements

This allows you to specify whether or not to replace pure systematic parts of the name by its commonly agreed upon name. For example, for the two structures shown, highly different names can be generated:

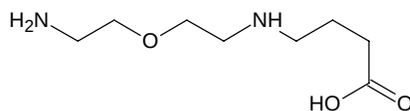


Yes	toluene	methylmalonic acid
No	methylbenzene	methylpropanedioic acid

4.12.1.2 Advanced Enclosing Marks

This allows you to specify which enclosing marks should be used: just parentheses or parentheses, brackets, braces, etc. in nesting order. The latter is preferred for clarity, but some text editors cannot handle these.

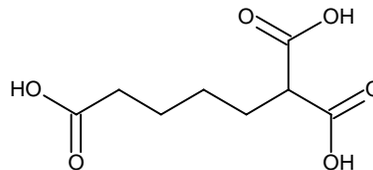
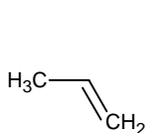
For example:



Yes	4-[[2-(2-aminoethoxy)ethyl]amino]butanoic acid
No	4-((2-(2-aminoethoxy)ethyl)amino)butanoic acid

4.12.1.3 Forward Locants Position

This allows you to specify the position of locants: either before the whole name or immediately before the corresponding part of the name. For example:

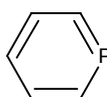


Yes	1-propene	1,1,5-pentanetricarboxylic acid
No	prop-1-ene	pentane-1,1,5-tricarboxylic acid

4.12.2 Old/New Tab

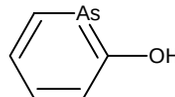
4.12.2.1 Hantzsch-Widman New Stems

This permits you to specify the usage of the extended Hantzsch-Widman system stems based on either *Recommendation 1982 (Pure Appl. Chem. 1983, 55, 409-416)* or *Nomenclature of Organic Chemistry, 1979 ed. (Blue book)*. For example:



Yes
No

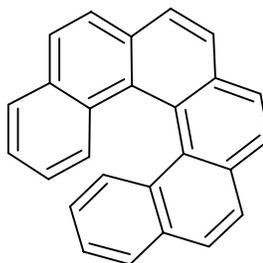
phosphinine
phosphorin



arsinin-2-ol
arsenin-2-ol

4.12.2.2 Extended Fused List

Specifies whether to use the additional trivial or semi-systematic retained names according to *A Guide to Nomenclature of Organic Compounds (Recommendations 1993)* or to use the more limited list based on *Nomenclature of Organic Chemistry, 1979 ed. (Blue book)*. For example:

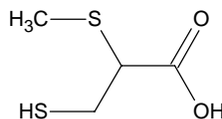


Yes
No

hexahelicene
benzo[c]naphtho[1,2-g]phenanthrene

4.12.2.3 New Functional Groups Names

This allows the user to specify the usage of suffixes and prefixes based on either *A Guide to Nomenclature of Organic Compounds (Recommendations 1993)* or *Nomenclature of Organic Chemistry, 1979 ed. (Blue book)*. For example:



Yes
No

2-(methylsulfanyl)-3-sulfanylpropanoic acid
3-mercapto-2-(methylthio)propanoic acid

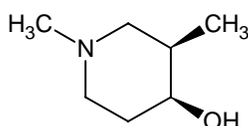
4.12.3 Structure Tab

The third set of preferences to be decided upon is the **Structure**. It specifies the preferred stereo conventions which will be used in name generation.

Note In order for the stereochemistry of a chiral center of a sketched-in molecule to be decided, the active centers must be designated with stereo-bonds.

4.12.3.1 Stereoconfiguration

Commonly, only relative configurations of chiral centers are known or a substance is a racemic mixture of stereoisomers. The options of this section allow you to choose whether to specify these situations in generated names or to ignore the configuration of chiral centers and double bonds.



Ignore	1,3-dimethyl-4-piperidinol
Absolute	(3 <i>R</i> ,4 <i>S</i>)-1,3-dimethyl-4-piperidinol
Racemic	<i>rac</i> -(3 <i>R</i> ,4 <i>S</i>)-1,3-dimethyl-4-piperidinol
Relative	<i>rel</i> -(3 <i>R</i> ,4 <i>S</i>)-1,3-dimethyl-4-piperidinol

For naming MDL SDfiles and molfiles or ISIS .DB files, if you select the **Take into account MDL chiral flag** option, all of the structures in those files that have such a flag will be named as having an absolute configuration and those that do not as having a relative one.

4.12.3.2 Stereo Wedge Direction

This allows you to choose between two possible procedures of stereo bond analysis.

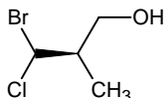
If **yes** is chosen, the “up” and “down” wedges which specify stereo centers will be analyzed with regard to their directions (depending on which atom a narrow end is attached).

If no is chosen, the sign (“up” or “down”) of the stereo wedge doesn’t depend on the direction.

For example:

Yes	(2<i>R</i>)-2-chlorobutane	(2<i>S</i>)-2-chlorobutane
No	(2<i>R</i>)-2-chlorobutane	(2<i>R</i>)-2-chlorobutane

If "no" is chosen, stereo bonds are not allowed between two chiral centers. For example:



Yes (2R,3R)-3-bromo-3-chloro-2-methylpropan-1-ol

No Refused to name

Note Avoid drawing stereo bonds represented as $\text{H}_3\text{C} \blacksquare \text{CH}_3$ or $\text{H}_3\text{C} \text{|||||} \text{CH}_3$, where the direction of the stereo bond is not explicitly shown. Such bonds will be replaced by a usual stereo bond, e.g., $\text{H}_3\text{C} \blacktriangleleft \text{CH}_3$ or $\text{H}_3\text{C} \blacktriangleright \text{CH}_3$, $\text{H}_3\text{C} \text{||||} \text{CH}_3$ or $\text{H}_3\text{C} \text{|||||} \text{CH}_3$, and named correspondingly.

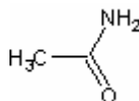
4.12.3.3 Name Preferred Tautomeric Form

This option allows you to choose between two possible procedures for tautomeric analysis.

If **Yes** is selected, the name for preferred tautomeric form of the drawn structure will be generated.

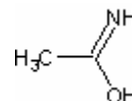
If **No** is selected, the name for the drawn tautomeric form of the structure will be generated.

For example:



Yes

acetamide



No

acetamide

ethanimidic acid

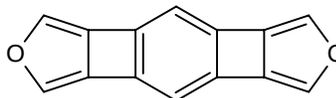
4.12.4 Custom Tab

The fourth set of preferences deals with refusals and utilization of natural product names.

4.12.4.1 Fused Multiparent Systems

Although ACD/Name can handle a broad variety of compounds, improvements to the algorithm are ongoing. For some compounds, the current name is not the most preferable IUPAC name. The **Refuse to Name** preference group allows one to specify what naming should be skipped.

For example:



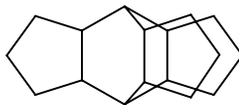
IUPAC name: benzo[1'',2'':3,4;4'',5'':3',4']dicyclobuta[1,2-c:1',2'-c']difuran

ACD name: furo[3''',4''':3'',4'']cyclobuta[1'',2'':4',5']benzo[1',2':3,4]cyclobuta[1,2-c]furan

4.12.4.2 Complex Bridged Fused Systems

If this check box is selected, bridged fused systems containing more than one trivalent or two bivalent acyclic bridges will be refused. If this check box is cleared, all bridged fused systems will be named as Von Baeyer polycycles.

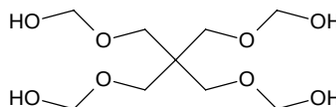
For example:



pentacyclo[5.5.5.0^{2,6}.0^{8,12}.0^{13,17}]heptadecane

4.12.4.3 Multiplicative Nomenclature

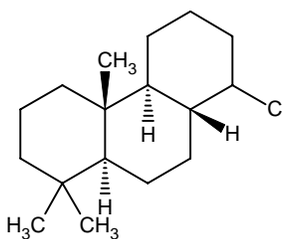
For example:



IUPAC name: [[2,2-bis[(hydroxymethoxy)methyl]propane-1,3-diyloxy]bis(oxy)]dimethanol
ACD name: {3-(hydroxymethoxy)-2,2-bis[(hydroxymethoxy)methyl]propoxy}methanol
 (*non-preferred name*)

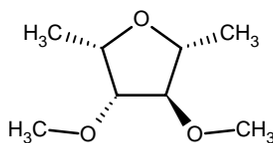
The rest of the preference allows you to specify whether to use retained natural product and biochemical names for several classes of compounds. This version allows you to use or avoid retained names for several dozens of basic natural product parent structures (steroids, alkaloids, terpenes, and some others), carbohydrates, common aminoacids, and peptides.

4.12.4.4 Steroids, Alkaloids, and Terpenes



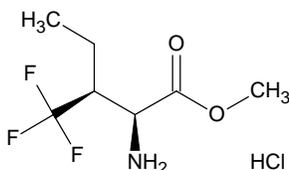
Yes 14-chloropodocarpene
No (4a*R*,4b*S*,8a*R*,10a*S*)-8-chloro-1,1,4a-trimethyltetradecahydrophenanthrene

4.12.4.5 Carbohydrates



Yes 2,5-anhydro-1,6-dideoxy-3,4-di-O-methyl-D-glucitol
No (2*R*,3*R*,4*R*,5*S*)-3,4-dimethoxy-2,5-dimethyltetrahydrofuran

4.12.4.6 Aminoacids and Peptides



Yes methyl 3',3',3'-trifluoro-L-isoleucinate hydrochloride
No methyl (2*S*,3*S*)-2-amino-3-(trifluoromethyl)pentanoate

4.12.4.7 Ignore Selected Settings if Cannot Name Otherwise

If this option is selected, all chosen settings in this tab will be ignored if the program cannot name the structure according to selected preferences. In this case "non-preferred name" notice will be added in the parentheses at the end of the name

4.13 IUPAC Name Free

This is a restricted version of the commercial ACD/IUPAC Name service.

Restrictions:

- This free version can treat structures with no more than 50 atoms, including hydrogen.
- Only the following atoms are allowed: C, H, N, P, O, S, F, Cl, Br, I, Li, Na, and K.
- No preferences can be specified for name generation.

4.14 Index Name

This module generates the systematic name for general organic compounds, according to guidelines specified by the Chemical Abstracts Service (CAS).

When you choose this property from the menu, the dialog box appears where you can specify the options described in the following subsections.

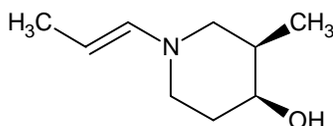
4.14.1 Structure Tab

The first set of preferences to be decided upon is the **Structure**. It specifies the preferred stereo conventions which will be used in name generation.

Note In order for the stereochemistry of a chiral center of a sketched-in molecule to be taken into account, the active centers must be designated with stereo bonds.

4.14.1.1 Stereoconfiguration

Commonly, only relative configurations of chiral centers are known or a substance is a racemic mixture of stereoisomers. The options of this section allow you to choose whether to specify these situations in generated names or to ignore the configuration of chiral centers and double bonds.



Ignore	4-piperidinol, 3-methyl-1-(1-propenyl)-
Absolute	4-piperidinol, 3-methyl-1-[(1E)-1-propenyl]-, (3R,4S)-
Racemic	4-piperidinol, 3-methyl-1-[(1E)-1-propenyl]-
Relative	4-piperidinol, 3-methyl-1-[(1E)-1-propenyl]-, (3R,4S)- <i>rel</i> -

For naming MDL SDfiles and molfiles or ISIS .db files, if you select the **Take into account MDL chiral flag** option, all of the structures in those files that have such a flag will be named as having an absolute configuration and those that do not as having a relative one.

4.14.1.2 Stereo Wedge Direction

This allows the user to choose between two possible procedures of stereo bond analysis.

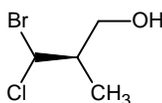
If **Yes** is selected, the “up” and “down” wedges which specify stereo centers will be analyzed with regard to their directions (depending on which atom a narrow end is attached).

If **No** is selected, the sign (“up” or “down”) of the stereo wedge doesn’t depend on the direction.

For example:

Yes	butane, 2-chloro-, (2R)	butane, 2-chloro-, (2S)	butane, 2-chloro-, (2R)
No	butane, 2-chloro-, (2R)	butane, 2-chloro-, (2R)	butane, 2-chloro-, (2R)

Note If "ignore" is selected, stereo bonds are not allowed between two chiral centers.
For example:



Take into account 1-propanol, 3-bromo-3-chloro-, (2R,3R)-
Ignore Refused to name

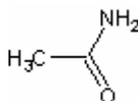
4.14.1.3 Name Preferred Tautomeric Form

This option allows you to choose between two possible procedures for tautomeric analysis.

If **Yes** is selected, the name for the preferred tautomeric form of the drawn structure will be generated.

If **No** is selected, the name for the drawn tautomeric form of the structure will be generated.

For example:

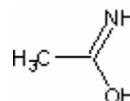


Yes

acetamide

No

acetamide



acetamide

ethanimidic acid

4.15 Name to Structure

ACD/Name to Structure generates chemical structures from the names of general organic and biochemical compounds, within the scope of the possibilities and restrictions described below. It is an independent program unit.

ACD/Name to Structure enables you:

- To generate chemical structures for the following name types:
 - Systematic names of most classes of general organic compounds,
 - Many derivatives of more than 150 basic natural product parent structures,
 - Semisystematic and trivial names of common organic compounds, and
 - Incorrect names that do not strictly follow Organic Nomenclature rules;
- To treat chemical names both in their uninverted and inverted form (CAS Index names);
- To report warnings about the ambiguity of a chemical name;
- To designate the stereo configuration of chiral centers and double bonds;
- To assigns numbers to atoms in the generated structure;
- To search the trivial or semitrivial names in ACD/Dictionary and generates the structure of some derivatives of such structures; and
- To treat the names in formatted and unformatted text.

When you choose this property, from the **I-Lab** menu, choose **Naming**, the dialog box appears where you should type the name from which you want to generate a structure and specify whether the structure should be searched for in ACD/Dictionary or not.

ACD/Dictionary is an additional module that contains a database of more than 85,340 systematic and non-systematic names of the most frequently used chemicals and biologicals. The entries cover more than 200 therapeutic areas.

The resultant dialog box displays the query name and the structure generated or found in the Dictionary in accordance with the specified settings.

4.16 Boiling Point

This module calculates the boiling point values for compounds, from 0.001 torr to 7,600 torr. The predictions are normally accurate to within $\pm 5^{\circ}\text{C}$ for structures with fewer than two polar groups, and to within $\pm 10^{\circ}\text{C}$ for structures with more polar groups.

To determine constants and relationships, the algorithm uses an internal database which contains boiling points for 10,000 compounds—6,000 of which have a boiling point at 760 mmHg. Additive increments of atoms and groups, force increments of interactions between groups and atoms are calculated from this database. These database values are also used to evaluate and fine-tune the specialized algorithms for ACD/Boiling Point. *Your compound does not have to be in the database in order to have its boiling point calculated!*

Prior to calculation, you should select units for pressure and temperature, and specify pressure at which the boiling point value is to be calculated.

The resultant dialog box contains structure, boiling point value, and pressure at which it was calculated. All of this data can be printed at once by clicking **Print** or you may wish to edit the default report page in ACD/ChemSketch and then print it (click **Insert** to start editing).

4.16.1 Limitations

Restrictions are:

- A maximum of 255 atoms excluding hydrogen can be calculated; and
- Only C, H, O, S, P, N, F, Cl, Br, I, Se, Si, Ge, Pb, Sn, As, and B are permitted elements.

4.16.2 What is Boiling Point?

The boiling point of a pure substance is in principle a non-additive property. It has been observed experimentally that in homologous sets, the dependence of boiling point on the number of CH_2 - groups obeys, approximately, the following non-linear function:

$$n_c = a_0 + \frac{a_1}{bp - a_2} \quad (1)$$

where n_c is the number of CH_2 - groups in the structure, bp is the observed boiling point, and a_i are empirically-determined constants. Note, however, that the additive algorithm cannot be applied to prediction of the boiling point.

We have made a detailed comparison of the behavior of different macroscopic properties, such as the index of refraction (nd_{20}), density (d_{20}), and surface tension (γ), for different homologous sets, with respect to boiling point. We observed that all of these properties can be approximately described by Equation (1). These three properties are further related by different relationships between two out of four macroscopic properties (molar volume, molecular weight, molar refractivity, and the parachor):

$$nd_{20} = f(MR, MV)$$

$$d_{20} = f(MW, MV)$$

$$\gamma = f(\text{parachor}, MV).$$

via the following well-known functions:

$$f(d) = \frac{1}{d} = \frac{MV}{MW} \quad (2)$$

$$f(n) = \frac{n^2 + 2}{n^2 - 1} = \frac{MV}{MR_r} \quad (3)$$

$$f(\gamma) = \frac{1}{\gamma^{1/4}} = \frac{MV}{P_r} \quad (4)$$

The noteworthy discovery, made by senior scientists at ACD/Labs, is that there is a function of boiling point with respect to other additive molecular properties, which is a linear (additive) function too. (Moreover, such an approach can also be used for prediction of the dielectric constant for organic compounds—an added feature of ChemSketch).

We express this function as:

$$K = f(MV, BP) \quad (5)$$

For different homologous sets, to a good approximation, the linear relationship is obeyed:

$$K = f(MV, BP) = c_0 + c_1 \cdot n_c \quad (6)$$

where the c_i are empirically determined.

From this, we can obtain two linearly predicted properties (using additive algorithms): our function K , and the molar volume (MV), which are similar in construction to our algorithm for prediction of $\log P$. Of course there are some differences: for example, the boiling point algorithm is different and more specialized for homologous sets and for other classes of compounds. Once K and MV are obtained, the boiling point is easily calculated.

4.17 LogP

ACD/LogP DB calculates the partition coefficient, $\log P$, for almost any drawn organic structure, quickly and accurately.

ACD/LogP DB contains a searchable database containing over 18,412 chemical structures with references to experimental $\log P$ values.

In most cases, ACD/LogP DB calculates $\log P$ values with an accuracy ± 0.3 or better. Such an accuracy has been demonstrated for many different classes of compounds (including acridines, amino acids and peptides, amphenicoles, barbituric acids, benzimidazoles, benzotriazoles,

carbamates, furans, furazans, furoxans, imidazoles, indoles, β -lactams, nucleosides and nucleotides, phenothiazines, phosphoroorganics, polyhalogenated aromatic and aliphatic compounds, purines, pyrazines, pyrazoles, pyridazines, pyridines, pyrimidines, quinolines, quinoxalines, saccharides, steroids, thiophenes, and many others).

The resultant dialog box displays the following data:

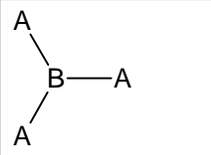
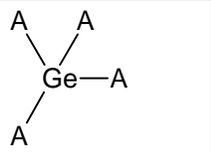
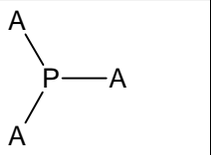
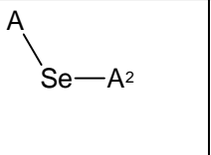
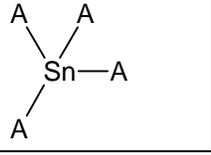
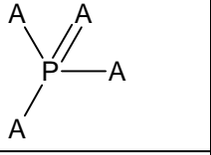
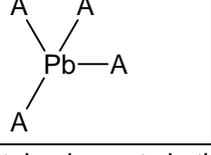
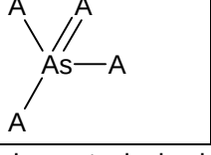
- If there are any literature references containing information on the obtained value, the references are placed at the top of the list.
- Next is the calculated $\log P$ value.

All of this data can be printed at once by clicking **Print** or you may wish to edit the default report page in ChemSketch and then print it (click **Insert** to start editing).

4.17.1 Limitations

The general ACD/Log P algorithm does not calculate the $\log P$ values for the following chemical structures:

- Charged structures other than the zwitterionic alpha-amino acids and their peptide derivatives and other than the non-ionic derivatives of IV-valent Nitrogen (+) bonded to Oxygen (-) or bonded to Nitrogen (-) with double bond (note that such compounds can be calculated, if their $\log P$ values are included in the user database that is used for system training);
- Structures containing atoms other than C, H, O, S, N, and F in possible chemical surroundings or structures containing atoms P, Cl, Br, I, Se, Si, Ge, Pb, Sn, As, and B that are not within the chemical surroundings shown below (note that A denotes any atom out of C, O, S, N, F, or any group listed below¹):

				Cl—A
			Se=A	Br—A
				I—A

- Structures that contain elements in their non-typical valence;
- Structures with coordinating bonds; and
- Structures containing more than 255 atoms excluding hydrogen.

¹ This group cannot be a part of a cycle unless otherwise stated.

² This fragment can be a part of an aromatic cycle.

ACD/Log P does not take into account the specific features of different geometric isomers, stereoisomers, conformers, isotopes, and structures with non-covalent bonds.

It predicts log P values so that in most cases the reliable experimental measurements lie within the calculated \pm log P interval. However, it is still possible that some new chemical structures might possess new specific structural features, such as far-range non-covalent bonding, intra-molecular shielding, or inter-molecular association. In such cases the discrepancy between a newly measured experimental value and the calculated \pm log P interval might occur.

For calculation of the partition coefficients of a partially dissociated mixture, we recommend to use ACD/Log D . This combines knowledge of the extent of acid-base ionization with partition coefficient calculations.

4.17.2 What is Log P ?

The hydrophobicity (also known as the lipophilicity) is a very important physicochemical property used in different areas of chemistry, medicine, and pharmacology. It is a very important indicator of transport and permeation through membranes, interaction with biological receptors and enzymes, toxicity, and biological potency.

An octanol-water partition coefficient, log P , is a measure of compound hydrophobicity; once you have obtained reliable log P values for a series of compounds, you are able to estimate many of their properties as listed above, provided these properties correlate with log P .

4.17.3 Description of Algorithm

The ACD/Labs algorithm performs the following procedures in the course of calculating log P :

1. Splits the structure into fragments.
2. Searches for identical fragments in the **internal databases**:
 - The database of **Fragmental Increments** contains well-characterized increments for over 500 different functional groups. These differ from each other by their chemical structure (e.g., amide, carboxy, ester, *etc.*), attachment to the hydrocarbon skeleton (aliphatic, vinylic, or aromatic), cyclization (cyclic or non-cyclic), and aromaticity (non-aromatic, aromatic, or fused aromatic).
 - The database of Carbon Atom Increments contains well-characterized increments for different types of carbons that are not involved in any functional group. They differ from each other by their state of hybridization (sp, sp², or sp³), number of attached hydrogens, branching (primary, secondary, tertiary, or quaternary), cyclization (cyclic or non-cyclic), and aromaticity (non-aromatic, aromatic, or fused aromatic).
 - The database of the Intramolecular Interaction Increments contains well-characterized increments for over 2,000 different types of pair-wise group interactions. They differ from each other by the type of the interacting terminal groups (see the differences among functional groups above), and the length and type of the fragmental system in-between the interacting groups (aliphatic, aromatic, and vinylic).
 - Searches for identical fragments in the data sources specified for system training. You can regulate the usage of each data field by its status: **Included/Excluded** in training, and statistical significance as **High/Low**.
3. If some fragments are not found in neither of the above mentioned databases, the program estimates their increments (as well as increments of inter-fragmental interactions) using Secondary Algorithms.

Lastly, the algorithm estimates the probability of tautomeric and ionic equilibria, calculation error and displays the results.

4.17.3.1 For I-Lab: Intranet Edition ACD/LogP Accuracy Extender

The training capability for log P /log D calculation is improved in version 7.0. As well, the LOGPUSER.CLC file can be created with ACD/Log P Accuracy Extender. You can further use this file to improve the accuracy of log P /log D prediction. (For more information, refer to Section 3.6.1.1.)

4.18 Log D

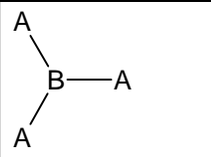
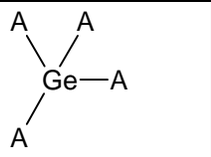
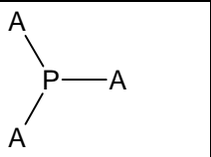
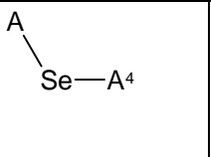
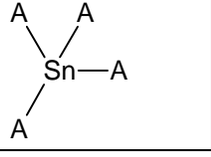
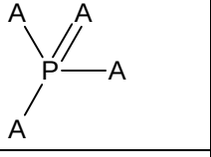
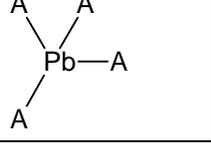
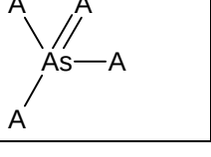
This module calculates the apparent partition coefficient D for dissociative systems—log D values at various pH (from 0 to 14).

The resultant dialog box displays a list of log D values at pH from 0.0 to 14.0 in a single step. All of this data can be printed at once by clicking **Print**, or you may wish to edit the default report page in the ChemSketch and then print it (click **Insert** to start editing).

4.18.1 Limitations

The ACD/Log D algorithm does not calculate the log D values for the following chemical structures:

- Structures containing atoms other than C, H, O, S, N, and F in possible chemical surroundings or structures containing atoms P, Cl, Br, I, Se, Si, Ge, Pb, Sn, As, and B that are not within the chemical surroundings shown below (note that A denotes any atom out of C, O, S, N, F, or any group listed below³):

				Cl—A
			Se=A	Br—A
				I—A

- Structures containing atoms in non-typical valence states (see the Elements Table in the ACD/ChemSketch for information on individual atomic valences);
- Structures containing non-covalent (co-ordination) bonds;
- Structures containing more than 255 heavy atoms or 10 ionization centers; and
- Structures containing more than 10 exchangeable protons.

³ This group cannot be a part of a cycle unless otherwise stated.

⁴ This fragment can be a part of an aromatic cycle.

Additionally, the ACD/LogD algorithm does not take into account the specific features of different geometric isomers, stereoisomers, conformers, and isotopes. The algorithm does not correct its prediction for long-range non-covalent bonding, intra-molecular shielding, or inter-molecular association.

Due to the ability to “train” the log*P* portion of the algorithm, however, certain effects which cannot be accounted for explicitly, can be adjusted for with a user database that contains other compounds of a related nature experiencing similar effects.

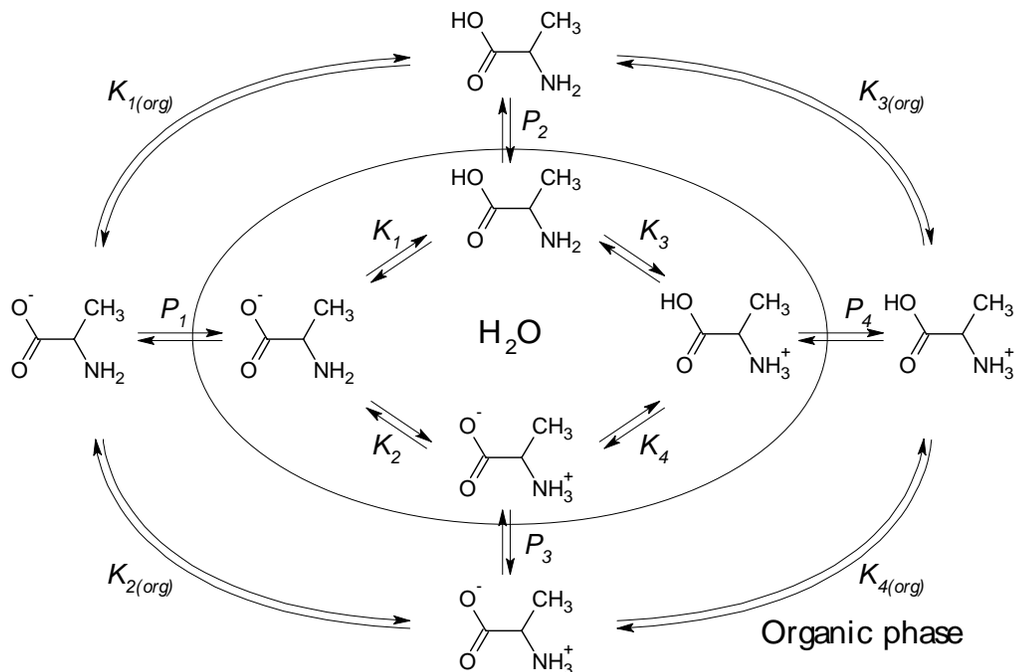
4.18.2 What is LogD?

The hydrophobicity (also known as the lipophilicity) is a very important physicochemical property used in different areas of chemistry, medicine, and pharmacology. It is a very important indicator of transport and permeation through membranes, interaction with biological receptors and enzymes, toxicity, and biological potency. In environmental sciences the hydrophobicity is often used to predict solubility (S), the bioconcentration factor (BCF), and the organic carbon adsorption coefficient (K_{oc}).

The octanol-water partition coefficient, log*P*, is a measure of compound hydrophobicity, which in many cases correlates well with the above chemical and biological properties. However, the log*P* can be accurately calculated only for uncharged substances. If the chemical compound contains one or more ionogenic groups (i.e., functional groups which can easily form ions), it may exist as a mixture of different ionic forms. The composition of this mixture depends strongly on pH.

In such cases the apparent partition coefficient *D* for dissociative systems (or log*D*) gives a more appropriate description than log*P* of the complex partitioning equilibria.

The following scheme presents an example of the complex equilibrium which exists for a solution of alanine in a water-octanol mixture:



K_i —the ionization constant of the *i*-th microspecies

$K_{i(org)}$ —the ionization constant of the *i*-th microspecies in octanol

P_i —the partition coefficient of the *i*-th microspecies

So, if such complex equilibria exist, then the partitioning of chemicals between water and organic phase must be a function of:

- (i) The extent of ionization, and
- (ii) The partition constants.

of the numerous different microspecies shown in the diagram above.

These problems have already been tackled, and to a good degree of success, by two ACD/Labs algorithms:

- (i) The acid-base ionization coefficient, pK_a , can be predicted normally to within ± 0.2 pK_a units for most functional groups by ACD/ pK_a .
- (ii) The octanol-water partition coefficients can be predicted normally to within ± 0.3 $\log P$ units by ACD/ $\log P$ DB.

Thus, it only makes sense to combine the capabilities of both "prediction engines" in one simple-to-use prediction of $\log D$: ACD/ $\log D$ Suite.

4.18.3 Definition of $\log D$

The partition coefficient for dissociative mixtures, $\log D$, is defined as follows:

$$\log D = \log \left(\frac{\sum a_i^{org}}{\sum a_i^{H_2O}} \right) \quad (I)$$

where $a_i^{H_2O}$ —the concentration of the i -th microspecies in water

a_i^{org} —the concentration of the i -th microspecies in organic phase

For correct solution of a proposed scheme of equilibrium we use the master equation approach. We construct a system of equations where the concentrations of all microspecies (both in organic and aqueous phase) are represented by a set of independent linear equations.

To calculate the concentrations all microspecies at any pH we need to solve the following system of equations:

$$a_i^{H_2O} = \frac{a_j^{H_2O} \times [H^+]}{K_i} \quad (II)$$

$$a_i = a_i^{H_2O} + a_i^{org}$$

$$\sum a_i = 1$$

$$a_i^{org} = a_i^{H_2O} \cdot P_i$$

where K_i —the ionization constant of the i -th microspecies $K_i = 10^{-pK_a}$

P_i —the partition coefficient of the i -th microspecies

a_j —the total concentration of the j -th microspecies having one H^+ less than the i -th

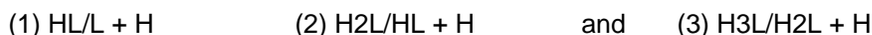
a_i — the total concentration of the i -th microspecies in both H_2O and octanol

Once the concentrations are known, the ratio in equation (I) and thus the $\log D$ value can easily be determined.

4.19 pK_a

This module calculates acid-base ionization constants (pK_a values) for organic and some inorganic chemical compounds in the system at the equilibrium state under standard conditions (25°C and zero ionic strength in aqueous solutions), the algorithm of this calculation mimics the experimental order of protonation of the drawn molecule and determines the pK_a values which can be experimentally measured in aqueous solution. Each calculation is provided with its ±95% confidence limits and, when available, literature references. The accuracy of calculations for simple structures is usually better than ±0.2 pK_a units (for complex structures it is better than ±0.5 pK_a units).

The resultant dialog box displays the query structure and the calculated pK_a values. The calculated pK_a values of a stepwise equilibrium are reported. The following scheme gives an estimate of the meaning of each of the calculated constants for the reactions:



More details can be found in the sections that follow.

4.19.1 Limitations

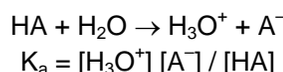
The ACD/pK_a algorithm will refuse to predict pK_a values for structures which:

- Contain more than 255 atoms (note that the program refuses to predict pK_a for some cyclic compounds having less than 255 atoms due to the fact that the program uses a cycle-breaking algorithm that increases the number of atoms);
- Do not contain an ionization center;
- Contain atoms of non-typical valence;
- Contain atoms other than C, H, O, S, P, N, F, Cl, Br, I, Se, Si, Ge, Pb, Sn, As, and B;
- Contain two or more fragments;
- Contain more than 20 ionization centers;
- Contain d-block or f-block metal atoms; or
- Contain textual abbreviations which cannot be transformed to structural fragments.

4.19.2 What is ACD/pK_a?

The pK_a is a measure of the tendency of a molecule or ion to keep a proton, H⁺, at its ionization center(s). It is related to ionization capabilities of chemical species. The more likely ionization occurs, the more likely a species will be taken up into aqueous solution, because water is a very polar solvent (dielectric constant, ε²⁰ = 80). If a molecule does not readily ionize, then it will tend to stay in a non-polar solvent such as cyclohexane (ε²⁰ = 2) or octanol (ε²⁰ = 10). In biological terms, pK_a is thus an important concept in determining whether a molecule will be taken up by aqueous tissue components or the lipid membranes. It is also closely related to the concepts of pH (the acidity of solution) and log*P* (the partition coefficient between immiscible liquids).

The equilibrium acid ionization constant, K_a, expresses the ratio of concentrations for the reaction:



where, by convention, it is assumed that the concentration of water is constant, and it is absorbed into the K_a definition.

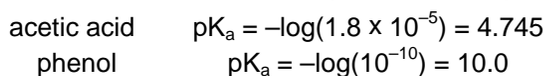
The acid ionization constant varies by orders of magnitude. For example at 25°C:



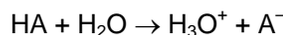
It is easier to refer to such extreme numbers on a logarithmic scale and, again by convention, “p” is used to denote the negative logarithm (base 10):

$$pK_a = -\log(K_a)$$

The K_a values of the compounds above are then easily converted to pK_a values:

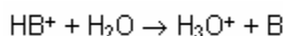


There is an essential difference between interpreting the pK_a values for molecules vs. ions. A molecule which loses a proton ionizes:



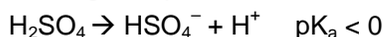
and so a low pK_a value denotes good aqueous solubility.

An ion which loses a proton, however, de-ionizes:



and so a high pK_a value denotes good aqueous solubility.

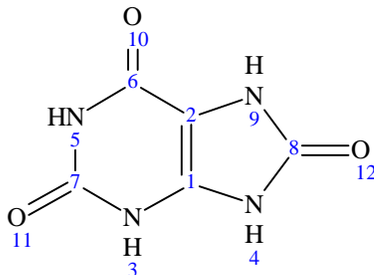
Note that there is no intrinsic reason to rule out pK_a values less than 0 or greater than 14. For example, sulfuric acid, H_2SO_4 , has a negative pK_a for the loss of its first proton:



although normally, experiment can only measure pK_a between 1 and 13.

4.19.3 Understanding pK_a Calculation Results

For example, if you calculate pK_a for uric acid:



the obtained results can be interpreted in the following way:

Ionic form: H4L

It means that the ionic form of the specified structure has 4 protons that can dissociate in the molecule according to the ACD/Labs algorithm.

$$pKa_1(HL/L+H; 4) = 17.05 \pm 0.20$$

It means that if the molecule contains 3 protons less than the query molecule (compare Ionic Form: H4L and the left part of the equation in the brackets), for the current dissociation step, the pK_a value is 17.05. The number of the atom dissociating is defined in the brackets after the semicolon (4 in this case) in accordance with the numbering in the corresponding MOL file. The subscript after "pKa" simply denotes the running number of the pK_a value.

$pKa2(H2L/HL+H; 3) = 14.19 \pm 0.20$

...

$pKa3(H3L/H2L+H; 5) = 9.51 \pm 0.20$

...

$pKa4(H4L/H3L+H; 9) = 5.60 \pm 0.30$

This line means that if the molecule contains as many protons as the specified molecule, for the current dissociation step the pK_a value is equal to 5.6. The dissociating atom stands under number 9.

$pKa5(H5L/H4L+H; 4) = -1.82 \pm 0.20$

If the molecule contains 1 proton more than the drawn structure, the pK_a value for this step is equal to -1.82 . Atom 4 is dissociating, but this time it turns from the positively charged into a neutral one. As compared to the first pK_a value ($pK_a = 17.05$), this atom became negatively charged.

4.20 $\Delta H(vap)$

This module calculates enthalpy of vaporization values.

Prior to calculation, you should select the units in which the enthalpy of vaporization is to be calculated.

The resultant dialog box contains structure, enthalpy of vaporization value, and pressure at which it was calculated. All of this data can be printed at once by clicking **Print** or you may wish to edit the default report page in ChemSketch and then print it (click **Insert** to start editing).

4.20.1 Limitations

Restrictions are:

- A maximum of 255 atoms excluding hydrogen can be calculated; and
- Only C, H, O, S, P, N, F, Cl, Br, I, Se, Si, Ge, Pb, Sn, As, and B are permitted elements.

4.20.2 What is Enthalpy of Vaporization?

The following equations are used to calculate the enthalpy of vaporization at different temperatures:

$$\phi = 4.19 + (n - 1) \cdot 0.2343 + 0.0021576 \cdot (BP_{760} [in^\circ C] + 60) \quad (1)$$

where

n = pre-assigned number related to the class of compound, which can be estimated according to the dependence of functional groups present in the compound. For example, for ethanol and amyl alcohol (non aromatic OH-containing) $n=7$, for acetic acid $n=4$, for different amines $n=3$, etc.

and

BP_{760} —the boiling point at 760 mm Hg (normal boiling point)

$$\Delta H_{vap} = 2.303 \cdot R \cdot BP \cdot \phi \quad (2)$$

where

R = molar gas constant

BP = boiling point

ϕ = result of equation (1)

4.21 Vapor Pressure

This module calculates vapor pressure values.

Prior to calculation you should select units for pressure and temperature, and specify temperature at which the vapor pressure value is to be calculated.

The resultant dialog box contains structure, vapor pressure value, and temperature at which it was calculated. All of this data can be printed at once by clicking **Print** or you may wish to edit the default report page in ChemSketch and then print it (click **Insert** to start editing).

4.21.1 Limitations

Restrictions are:

- A maximum of 255 atoms excluding hydrogen can be calculated; and
- Only C, H, O, S, P, N, F, Cl, Br, I, Se, Si, Ge, Pb, Sn, As, and B are permitted elements.

4.21.2 How is it Calculated?

The following equations are used to calculate vapor pressure at different temperatures:

$$\log(VP) = 2.808 - \frac{\phi \cdot (BP_{760} - t)}{273.1 + t - 0.15 \cdot (BP_{760} - t)} \quad (3)$$

where t = temperature (in °C); BP_{760} = boiling point at 760 mm Hg (normal boiling point); ϕ = result of equation (1)

$$VP = 10^{\log(VP)} \quad (4)$$

4.22 Aqueous Solubility

This module calculates accurate aqueous solubility values at any pH under the standard conditions (and zero ionic strength). The accuracy of calculations for simple structures is usually better than 0.2–0.5 logarithmic units (for complex structures it is better than 0.5–1.0 logarithmic units).

The aqueous solubility of a chemical may be defined as the maximum amount of the chemical that will dissolve in pure water at a specified temperature. Above this concentration, two phases will exist if the organic chemical is a solid or a liquid at the system temperature: a saturated aqueous solution and a solid or liquid organic phase.

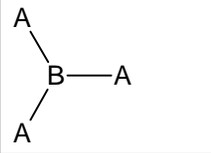
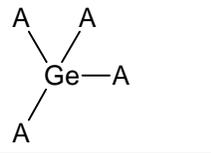
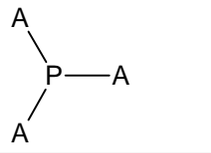
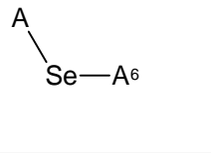
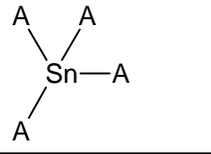
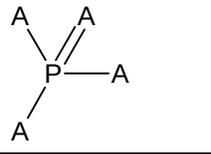
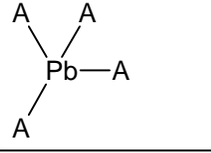
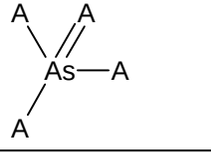
ACD/Solubility prediction has the following features:

- Compounds subdivided into classes with separate prediction equations for each, including new parameters for internal hydrogen bonds.
- The following parameters are used in correlation equations: boiling point, $\log P/\log D$, MW, MolVol, HydBonding (number of H-acceptors and H-donors), and refractive index.
- Prediction is not sensitive to the melting point presence for many classes of compounds. Even information such as whether the compound is a solid or liquid at room temperature is useful in the prediction.

4.22.1 Limitations

The general ACD/Solubility algorithm does not calculate the solubility values for the following chemical structures:

- Charged structures containing IV-valent Nitrogen (+);
- Structures containing atoms other than C, H, O, S, N, and F in possible chemical surroundings or structures containing atoms P, Cl, Br, I, Se, Si, Ge, Pb, Sn, As, and B that are not within the chemical surroundings shown below (note that A denotes any atom out of C, O, S, N, F, or any group listed below⁵):

				Cl—A
			Se=A	Br—A
				I—A

- Structures that contain elements in their non-typical valence;
- Structures with coordinating bonds; and
- Structures containing more than 255 atoms excluding hydrogen (note that the program refuses to predict solubility for some cyclic compounds having less than 255 atoms due to the fact that the program uses cycle-breaking algorithm that increases the number of atoms).

The program does not take into account the specific features of different geometric isomers, stereoisomers, conformers, isotopes and structures with non-covalent bonds.

⁵ This group cannot be a part of a cycle unless otherwise stated.

⁶ This fragment can be a part of an aromatic cycle.

It predicts solubility values so that in most cases the reliable experimental measurements lie within the following solubility interval: 0.2–0.5 logarithmic units for simple compounds and 0.5–1.0 logarithmic units for complex compounds. However, it is still possible that some new chemical structures might possess new specific structural features, such as far-range non-covalent bonding, intra-molecular shielding, or inter-molecular association. In such cases the discrepancy between a newly measured experimental value and the calculated solubility interval might occur.

Note There certainly exist some structures that formally meet the aforementioned limitations, but cannot be calculated by the current algorithm.

4.23 Search pK_a DB

Using this module you can search through the ACD/ pK_a database containing almost 16,000 structures with thousands of experimental pK_a values under different temperatures and ionic strengths in purely aqueous solutions.

You can perform searches according to various parameters:

- Chemical substructure
- Structure
- Similar structure
- Formula
- Molecular weight
- pK_a values
- Reference
- Temperature
- Ionic strength

Each database entry contains the following data:

- Name
- Chemical structure
- Formula
- Molecular weight
- Ionic form
- pK_a values
- Corresponding references

4.24 Search LogP DB

Using this module you can search through the ACD/LogP database containing more than 18,400 chemical structures with references to experimental logP values.

You can perform searches according to various parameters:

- Chemical substructure
- Structure
- Similar structure
- Formula
- Molecular weight
- LogP values
- Reference
- Name

Each database entry contains the following data:

- Chemical structure
- Formula
- Name
- Molecular weight
- LogP values
- Corresponding references

4.25 BCF and Adsorption Coefficient

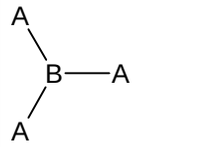
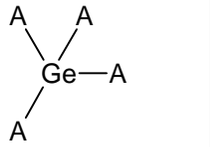
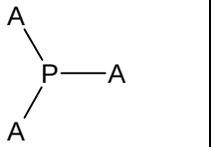
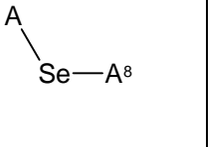
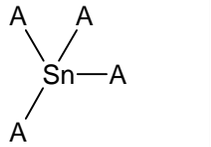
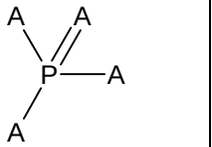
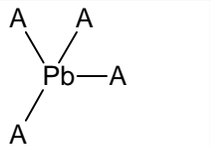
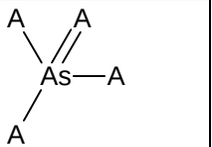
This module calculates BCF (Bioconcentration Factor) at various pH (from 0 to 14) and K_{oc} (Organic Carbon Adsorption Coefficient) at various pH (from 0 to 14).

The resultant dialog box displays a list of BCFs at pH from 0.0 to 14.0 with a single step and a list of K_{oc} values at pH from 0.0 to 14.0 with a single step. All of this data can be printed at once by clicking **Print**, or you may wish to edit the default report page in the ChemSketch and then print it (click **Insert** to start editing).

4.25.1 Limitations for BCF

The algorithm does not calculate the BCF for the following chemical structures:

- Structures containing atoms other than C, H, O, S, N, and F in possible chemical surroundings or structures containing atoms P, Cl, Br, I, Se, Si, Ge, Pb, Sn, As, and B that are not within the chemical surroundings shown below (note that A denotes any atom out of C, O, S, N, F, or any group listed below⁷):

				Cl—A
			Se=A	Br—A
				I—A

- Structures containing atoms in non-typical valence states (see the Elements Table in the ACD/ChemSketch for information on individual atomic valences);
- Structures containing non-covalent (co-ordination) bonds; and
- Structures containing more than 255 heavy atoms or 10 ionization centers.

Additionally, the algorithm does not take into account the specific features of different geometric isomers, stereoisomers, conformers, and isotopes. The algorithm does not correct its prediction for long-range non-covalent bonding, intra-molecular shielding, or inter-molecular association.

4.25.2 What is BCF?

BIOCONCENTRATION—the increase in concentration of a chemical in an organism resulting from tissue absorption levels exceeding the rate of metabolism and excretion.

BIOCONCENTRATION FACTOR (BCF)—used to describe the accumulation of chemicals in organisms, primarily aquatic, that live in contaminated environments.

According to EPA guidelines, "the BCF is defined as the ratio of chemical concentration in the organism to that in surrounding water. Bioconcentration occurs through uptake and retention of a substance from water only, through gill membranes or other external body surfaces. In the context of setting exposure criteria it is generally understood that the terms "BCF" and "steady-state BCF" are synonymous. A steady-state condition occurs when the organism is exposed for a sufficient length of time that the ratio does not change substantially."

⁷ This group cannot be a part of a cycle unless otherwise stated.

⁸ This fragment can be a part of an aromatic cycle.

Bioconcentration factors (BCFs) are used to relate pollutant residues in aquatic organisms to the pollutant concentration in ambient waters. The bioconcentration factor (BCF) is related to biomagnification effects. Many chemical compounds, especially those with a hydrophobic component, partition easily into the lipids and lipid membranes of organisms and bioaccumulate. If the compounds are not metabolized as fast as they are consumed, there can be significant magnification of potential toxicological effects up the food chain. The concern about bioaccumulation and biomagnification comes mainly from experience with chlorinated compounds, especially pesticides and PCBs, and their deleterious effects on vulnerable species, especially birds, frogs, and fish. Only minimal experimental and monitoring information has been gathered on the bioaccumulation properties of many other currently used chemical compounds. In fact, the biomagnification of many widely available chemicals has not been observed or predicted in aquatic systems.

BCF or BAF values are based on U.S. Environmental Protection Agency publications pursuant to Section 304(a) of the Federal Water Pollution Control Act as amended, literature values, or site-specific bioconcentration data. Current EPA guidelines for the derivation of human health water quality criteria use BCFs as well.

BCFs are quantified by various procedures depending on the lipid solubility of the pollutant. For lipid soluble pollutants, the average BCF is calculated from the weighted average percent lipids in the edible portions of fish and shellfish, which is about 3%; or it is calculated from theoretical considerations using the octanol/water partition coefficient. The *BCF* is calculated using the following equation:

$$\log BCF = 0.79 \times \log P - 0.40$$

where $\log P$ is the octanol/water partition coefficient. The "rule of thumb" is that a BCF value greater than 1000 poses a concern for bioaccumulation, particularly if the chemical is considered persistent in aquatic environments. This BCF value corresponds to a $\log P$ of 4.3 or greater.

For non-lipid soluble compounds, the BCF is determined empirically. The assumed water consumption is taken from the National Academy of Sciences publication *Drinking Water and Health* (1977). (Referenced in the Human Health Guidelines.) This value, of 2.0 liters/day, is appropriate as it includes a margin of safety so that the general population is protected. The 6.5 grams per day of contaminated fish and shellfish consumption value is the average per-capita consumption rate of all (contaminated and non-contaminated) freshwater fish and shellfish for the U.S. population.

Although BCF assessments began as aquatic measurements, the exposure of plants and cattle to certain chemicals is also rated in terms of the bioconcentration factors.

4.25.3 Limitations for Adsorption Coefficient

The algorithm does not calculate the K_{oc} for the following chemical structures:

- Structures containing atoms other than C, H, O, S, N, and F in possible chemical surroundings or structures containing atoms P, Cl, Br, I, Se, Si, Ge, Pb, Sn, As, and B that are not within the chemical surroundings shown below (note that A denotes any atom out of C, O, S, N, F, or any group listed below⁹):

⁹ This group cannot be a part of a cycle unless otherwise stated.

				Cl—A
			Se=A	Br—A
				I—A

- Structures containing atoms in non-typical valence states (see the Elements Table in the ACD/ChemSketch for information on individual atomic valences);
- Structures containing non-covalent (co-ordination) bonds; and
- Structures containing more than 255 heavy atoms or 10 ionization centers.

Additionally, the algorithm does not take into account the specific features of different geometric isomers, stereoisomers, conformers, and isotopes. The algorithm does not correct its prediction for long-range non-covalent bonding, intra-molecular shielding, or inter-molecular association.

4.25.4 What is Adsorption Coefficient?

The organic carbon adsorption coefficient, K_{oc} , is crucial for estimating a chemical compound's mobility in soil, and the prevalence of leaching from soil. The adsorption of a compound increases with an increase in organic content, clay content, and surface area of soil. The presence of a chemical compound can also be detected in groundwater, and inference can be made about its residence time in the soil and the degradation period before reaching the water table. The presence of continuous pores or channels in soil will increase the mobility of a chemical compound in the soil.

The Adsorption Coefficient (K_{oc}) may be defined as the ratio of the amount of chemical adsorbed per unit weight of organic carbon (oc) in the soil or sediment to the concentration of the chemical in solution at equilibrium:

$$K_{oc} = \frac{\mu\text{g} \cdot \text{adsorbed} / \text{g} \cdot \text{organic} \cdot \text{carbon}}{\mu\text{g} / \text{mL} \cdot \text{solution}}$$

K_{oc} is calculated using the following equation [1,4]:

$$\log K_{oc} = 0.544 \cdot \log P + 1.377$$

¹⁰ This fragment can be a part of an aromatic cycle.