GETTING STARTED

CHEMKIN RELEASE 3.6

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Literature Citation for the CHEMKIN Collection:
The CHEMKIN Collection Release 3.6 should be cited as:

Reaction Design cautions that some of the material in this manual may be out of date. Updates will be available periodically on Reaction Design’s web site. In addition, on-line help is available on the program CD. Sample problem files can also be found on the CD and on our web site at www.ReactionDesign.com.
Overview of Manual

This manual has been designed to get you up and running as quickly as possible with CHEMKIN Release 3.6. The organization of the manual is:

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1. What’s New in CHEMKIN Release 3.6

The CHEMKIN Collection is a powerful system for solving complex chemical modeling problems. CHEMKIN is composed of standalone Applications, as well as subroutine libraries and utilities that can be used to create new problem solving environments. CHEMKIN facilitates the formulation, solution, and interpretation of problems involving gas-phase and heterogeneous (gas-surface) chemical kinetics. A wide range of chemical process modeling problems can be addressed including: combustion, catalysis, corrosion, plasma etching and chemical vapor deposition.

Many new features and enhancements have been incorporated by Reaction Design into Release 3.6, including:

- **Transient stirred-reactor capability.** You can now run AURORA in either transient or steady-state mode. This allows simulation of engine cycles, pulsed plasma, reactor startup, as well as many other applications. The transient capability includes sensitivity analysis for species and temperature, for both gas and surface reactions.
- **Options in AURORA, PLUG, SENKIN, and PREMIX allow you to specify profiles that vary as a function of time or distance** through keyword input. These options include profiles of varying temperature, heat loss, pressure, volume, and power. They eliminate the need for programming and compiling of user subroutines.
- **The CHEMKIN Graphical Post-processor now allows you to import more than one solution set simultaneously, and allows generation of plots from more than one solution set.** This allows plotting of simulation results against imported experimental data, for example, as well as plotting of one simulation result against another when one grid is more refined.
- **OPPDIF now includes sensitivity analysis** for reaction and heat-of-formation sensitivity.
- **All of the Command-Line FORTRAN Post-processors have been re-structured to output text files as comma-delimited columns of data** that may be easily imported into the Graphical Post-processor as well as into Excel or other third-party analysis packages. These FORTRAN post-processor routines read and process the binary solution files, are distributed as source-code, and allow users to tailor post-solution analyses to their own specific needs.

Getting started with CHEMKIN is easy; the installation takes just a few minutes. This manual guides you through the installation process and helps you navigate through the graphical user interfaces. Additional details about the CHEMKIN Applications and Utilities are provided in a complete set of user manuals. The distribution CD for the CHEMKIN Collection contains all of the program documentation, as well as a set of sample problems for each of the Applications.
2. System Requirements

Before you start, take a minute to verify that you have what you need to install and run CHEMKIN:

1. A suitable computer and operating system. CHEMKIN Release 3.6 will run on the following platforms:

<table>
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<th>Operating System</th>
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2. 60 Megabytes of hard disk space.

3. 64 MB of RAM (128 MB recommended). **Note:** You may need more memory if you plan to run very large problems (e.g. problems that involve more than 100 species).

4. A valid license file for the CHEMKIN software

5. Access to the documentation for the applications that you intend to run.

To use CHEMKIN, you should be familiar with the basics of the operating system on your computer. You should know how to use the mouse, how to use menus, manage files and how to enter text. If you are unsure about any of these techniques, we recommend that you consult the documentation that came with your computer.
3. Installation and Quick Start

3.1 Overview of the Installation Process

Installation of the CHEMKIN Collection involves three basic steps:

1. Install the software from the CD-ROM (or web download).
2. Obtain and install a CHEMKIN license file from Reaction Design.
3. Run one or more of the samples problems to make sure that the software is installed correctly.

Each of these steps are described in more detail in the following sections. The instructions are divided according to platform. Section 3.2 describes installation procedures for all Windows-based PCs, while Section 3.3 describes installation procedures for all UNIX platforms.

3.2 PC Installation and Licensing

3.2.1 Running the Setup Program

To install CHEMKIN on your Win32 computer (Windows NT, 95, 98, or 2000), perform the following steps:

1. Insert the CHEMKIN distribution CD-ROM into your CD-ROM drive. After a short delay, the autoplay program starts. Note: If the autoplay program does not start automatically on your computer, double click on your “My Computer” icon and browse to your CD-ROM. Inside the subfolder install, locate the chemkin36_pc_setup.exe file and double-click on this.
2. The program will first prompt you to extract the needed files. Click “Setup” to continue.
3. The Welcome dialog introduces you to the Setup program and informs you of several restrictions and cautions. Click “Next” to continue.
4. The next dialog contains some more information about the Release. Click “Next” to continue with the installation.
5. Choose the Destination Location. This is the location on your hard disk where you would like to install CHEMKIN. The default is “c:\chemkin36”, but you may select any drive or folder name. To choose a different location from the default, click “Browse” and select a location from the Choose Directory dialog. Once you have made your choice (or if you decide to keep the default), click “Next” to continue the installation. Click “Cancel” to leave the installation program and return to Windows.

6. Choose the Setup Type. This allows you to customize what parts of CHEMKIN get installed. We recommend choosing “Typical”. If you choose “Compact,” you will not install most of the online documentation. Once you have made your selection, click “Next” to continue.

7. Choose the short-cut labels for your Start→Programs menu. Modifying the settings here changes the way the short-cuts are organized in your Programs menu. Click “Next” to continue. At this point the Setup program will begin installing files on your computer’s hard disk.

8. After the files have been installed successfully, the setup dialog will ask you to reboot your computer. Choose “Yes” to allow the program to reboot, or “No” to reboot manually later. The reboot step is necessary to register a default license-file location in your system registry.

3.2.2 Modifying your “Path”

We recommend that you modify your “path” variable to include the “c:\chemkin36\bin” directory, particularly if you plan to run from the command line, rather than using the Application User Interface. The following steps will add “c:\chemkin36\bin” to your path. Note that the instructions are different for Windows NT and Windows 2000 than for Windows 95 and Windows 98 operating systems.

**Windows NT or Windows 2000:**

1. Right-click on your computer icon on your desktop and choose “Properties” from the system menu.

2. **NT:** Select the “Environment” tab.
   
   **2000:** Select the “Advanced” tab. Select “Environment Variables”

3. **NT:** Under the “User Variables for <your user name>”, browse for and select the “path”.
   
   **2000:** Under the “User Variables for <your user name>”, browse for and select the “PATH” and choose “Edit”.

4. In the “Value” field, scroll to the end of the text box and add a semicolon “;” followed by the full path to your CHEMKIN “bin” directory (e.g., c:\chemkin36\bin). Click “Set” and then “OK”.


Windows 95 and Windows 98:

1. Open your `autoexec.bat` file, located in the root of your “C” disk, using your favorite text editor (e.g. Notepad).

2. Add the following line near the top of the file (substituting the correct path to your CHEMKIN “bin” directory if you have not installed in the default location):

   ```
   set PATH=%PATH%;c:\chemkin36\bin
   ```

3. Save the file and exit the editor.

4. Restart your computer for the change to take effect.

3.2.3 Uninstalling CHEMKIN

After you have installed CHEMKIN with the above steps, it is possible to remove it from your system if necessary. Open the “Add/Remove Programs” folder in the Control Panel (under “Settings” from the “Start” menu). Select “CHEMKIN 3.6” from the “Install/Uninstall” tab and click “Add/Remove”. Note: If you have added files to or modified files in the CHEMKIN installation, the Uninstall process will not automatically remove these files.

3.2.4 PC Licensing Overview

You must obtain and install a set of license keys in order to run the CHEMKIN Collection software. This section provides instructions on how to obtain and install a Reaction Design license file for the CHEMKIN Collection after you have purchased the software. For purchasing information, contact by email: CHEMKIN@ReactionDesign.com.

Assumptions:

We assume the following, for purposes of providing instructions on obtaining and installing a license on a PC:

1. You have already installed CHEMKIN Release 3.6 on your computer.

2. The default installation directory for your PC is `c:\chemkin36`. If you have installed in a different location, please make the necessary replacements in the instructions that follow.
Required Tasks:

1. Request a license from Reaction Design, providing information about your computer.

2. Install the license received from Reaction Design.

3.2.5 Request a License from Reaction Design

To obtain a license from Reaction Design, we require that you send us certain information about the computer on which you will be running CHEMKIN. Here we provide detailed instructions for obtaining that information. If you have purchased a Network (Simultaneous Active User) License, the following steps should be run on your Network License Server.

For CHEMKIN 3.6, we have added a new license utility for the PC. A short-cut to this utility is installed in your Windows Start → Programs Menu. Follow the steps below to send your computer’s identifying data to Reaction Design:

1. With your mouse, click on:
   
   Start → Programs → CHEMKIN Collection 3.6 → Licensing → Request A New CHEMKIN License
   
   The “RequestLicense” utility should open a dialog box.

2. From the initial dialog box, choose “Get a New License” and click OK

3. In the “Get A License” dialogue box, fill in your Personal Information.

4. Choose one of the following:

   a. **Email Info:** (To email the information gathered to Reaction Design)

   b. **Web Form:** (To post the information gathered via a web browser)

   c. **Save:** (To save the information gathered to a file to be retrieved later). Use this if your computer is not connected to the internet. The information is saved as c:\chemkin36\licenses\request.txt. Email the contents of this file to CHEMKIN@ReactionDesign.com.

Upon receipt of your computer information, Reaction Design will generate a set of license keys, specific to your computer and to the selection of CHEMKIN modules that you have purchased. These license keys will be sent to you by email. The next two sections describe how to install the license keys on your PC. If you have purchased a Node-locked license, follow the instructions in Section 3.2.6; if you have purchased a Network license, proceed to Section 3.2.7.
3.2.6 Installing a Node-locked (or DEMO) License

Once you have received an email with license keys from Reaction Design, you will need to place the license keys in the correct location.

- If the license file arrives as an email attachment, simply save the attachment as a file named “chemkin.lic” in the “licenses” sub-folder of your CHEMKIN installation (e.g. save as “c:\chemkin36\licenses\chemkin.lic”).
- If the license keys are embedded in the email text, simply cut and paste this information into an empty text file, using your favorite editor (e.g. Notepad). Save this file as “c:\chemkin36\licenses\chemkin.lic”.
  Be sure that Notepad does not inadvertently add an extra “.txt” extension onto the end of the file name. To view the file extension, you can open an MS-DOS Prompt, change directories to the “c:\chemkin36\licenses” folder and type “DIR”;
  the full file name of the license file should be listed.

You are now ready to run CHEMKIN; Skip the following sections and go directly to Section 3.4

3.2.7 Installing a Network (Simultaneous Active User) License

In network licensing, more than one user may be running CHEMKIN Applications by accessing a centrally located license file that resides on a Network License Server. The license allows one or more simultaneous active users to run CHEMKIN Applications. Clients of the Network License Server should have their own local installation of CHEMKIN, but obtain a license from the Server while the CHEMKIN Application is running. The instructions here describe what needs to be done on both the Network License Server and on the Client machines.

On the Network License Server:

1. Install the license as described in Section 3.2.6 (assuming that the Server is a PC).

2. Edit the license file using your text editor of choice:
   a. Change the text “servername” in the line:
      
      SERVER servername

      to the name (e.g. “MYCOMPUTER”) or IP Address (e.g. “192.168.0.1”) of the Network License Server.

   b. On the DAEMON line, verify that the PATH to reaction.exe is correct; the default location is: c:\chemkin36\licenses\reaction.exe
3. **Launch LMTOOLS** by selecting with your mouse from the **Start → Programs** menu. Click on **Start → Programs → CHEMKIN Collection 3.6 → Licensing → Manage CHEMKIN Licenses**. This should start the LMTOOLS utility window.
   
   a. Click on the **"Configuration using Services"** option (radio button).
   
   b. Click on **"Configure Services"** tab.
   
   c. Browse for the location of lmgrd.exe. With default installation, this will be: 
      
      c:\chemkin36\licenses\lmgrd.exe
   
   d. Browse for the license file location. This will be the location used in Step 1 above.
   
   e. It is recommended that you keep a log file. In the log path, type in: 
      
      c:\chemkin36\licenses\debug.log
   
   f. Check the boxes **"Use Services"** (NT) and **"Start server at Power Up"**.
   
   g. Press the **"Save Service"** button and click **"Yes"** to save the settings.
   
   h. Select the **"Start/Stop/Reread"** tab.
   
   i. Click **"Start"** to start the daemon service on the Server.
   
   j. Choose **"Exit"** from the **"File"** pull-down menu.

**On each Network Client:**

1. Place a copy of the modified Network license file in the licenses sub-directory of the client CHEMKIN 3.6 installation; by default this will be:
   
   **Windows/PC Client:**
   
   c:\chemkin36\licenses\chemkin.lic
   
   **UNIX Client:**
   
   $HOME/chemkin36/licenses/chemkin.lic
   
2. Now the client will be able to check out a license through the Server daemon.

You are now ready to run CHEMKIN; Skip the following sections and go directly to Section 3.4
3.3 UNIX INSTALLATION AND LICENSING

This section provides detailed instructions for installing CHEMKIN on any one of our supported UNIX platforms.

3.3.1 Mounting the CHEMKIN Distribution CD-ROM

Insert the CHEMKIN distribution CD-ROM into a mounted CD-ROM drive. You must then mount the disk as a file system on your computer. **Note:** on most systems you must be root to mount a CD-ROM. Specific instructions for mounting the CD on supported platforms are as follows:

**Compaq Tru64 UNIX (Digital Alpha):**

Mount the CD using Rock Ridge format by entering:

```
/usr/sbin/mount -t cdfs -r -o rrip <CD-ROM-device> /cdrom
```

replacing `<CD-ROM-device>` with the actual name of the CD-ROM drive on your system. A typical name is `/dev/rz4c`

**Hewlett-Packard HP-UX:**

Mount the CD by entering:

```
mount -r <CD-ROM-device> /cdrom
```

replacing `<CD-ROM-device>` with the actual name of the CD-ROM drive on your system. A typical name is `/dev/dsk/c1t2d0`

**SGI IRIX:**

IRIX automatically mounts the CD when you insert it into the CD-ROM drive. It is mounted as:

```
/CDROM
```

**Notes:** the installation instructions below assume that the CD-ROM is mounted at `/cdrom`. You will therefore need to replace this with the actual path above (all upper case) as you follow the instructions below.

**SUN Solaris 2.6:**

Solaris includes a “Volume Manager” that automatically mounts the CD when you insert it into the CD-ROM drive. The CD will be mounted as:

```
/cdrom/Chemkin36
```

**Notes:** the installation instructions below assume that the CD-ROM is mounted at `/cdrom`. You will therefore need to replace this with the actual path above as you follow the instructions below.
3.3.2 Executing the Installation Script

If you are installing CHEMKIN into a public directory, make sure that you have root or similar privileges before running the installation script. Once you have the correct permissions, open a UNIX shell and execute the installation script with the following commands (Note: for SGI and SUN users, you must substitute the appropriate mount point for /cdrom below, as described in Section 3.3.1):

```
cd /cdrom/install
/bin/sh /cdrom/install/chemkin36_unix_setup.sh
```

The installation script will echo information and prompt you for input as it proceeds through the following steps:

1. **Verify the current directory and the username.** Check that the directory printed is the install directory at the CD-ROM mount point. Check that the user login has the correct privileges for installing to your desired destination. If this information is correct, type “y” and <RETURN>.

2. **Choose the Destination Location.** This is the location on your hard-disk where you would like to install CHEMKIN and is the location where the install script will create a “chemkin” subdirectory. The default is the directory identified by the environment variable, $HOME, or, if $HOME is not set on your computer, the default is $HOME. The prompt asks if you want to use the default. To accept the default, type “y” and <RETURN>. To change the Destination Location, type “n” and <RETURN>, then type in the full path of your preferred destination and hit <RETURN> again.

3. **Create any needed directories.** If the directory you selected for the Destination Location does not already exist, you will be prompted to allow the install script to create the new directory. Type “y” and <RETURN> to continue with the install.

4. **Verify the platform for the installation.** The install script will determine which platform you are currently running on and will ask you whether this is the correct platform information. If the information is correct, type “y” and <RETURN> to continue. If you want to install onto a different platform than the one specified, type “n” and <RETURN>. You will then be asked to choose from the list of available platforms and enter a corresponding number (e.g. “1” for Compaq (Alpha)). Type the number and <RETURN>.

5. **Verify all the information entered** thus far before installing the CHEMKIN software. If the information is correct, type “y” and <RETURN>.
6. **Install the files.** The script will copy compressed tar files from the CD-ROM to your hard disk, 
uncompress and untar them into the Destination Location, and then delete the tar files. When the 
installation is complete, information on setting environment variables needed to run CHEMKIN 
are displayed on your screen, using the UNIX `more` utility. These instructions are also given 
below and in Section 3.3.4.

### 3.3.3 Setting the CHEMKIN_DIR Environment Variable

The UNIX version of the CHEMKIN software requires you to set an environment variable that points to the 
top-level directory where CHEMKIN was installed. Although you could set the environment variables 
locally in a UNIX shell every time you run CHEMKIN, we recommend that you instead include the 
environment settings in your `.cshrc` (for C-shell users) or your `.profile` (for K-shell users) file, which 
will result in the variables being set automatically every time you open a new shell. This section gives 
you detailed instructions for modifying these start-up files.

The following instructions assume that you have installed CHEMKIN in the directory `$HOME`, such that the 
top-level CHEMKIN directory is `$HOME/chemkin36`. If you have installed into a different location, or if 
the `$HOME` environment variable is not set for your user account, you will need to replace “$HOME” in the 
following instructions with the actual (absolute) path.

1. Modify your `.cshrc` file (for C-shell users) or your `.profile` file (for K-shell users) to include 
the environment variable `CHEMKIN_DIR`. The command(s) that need to be added are shown 
below for the two shell environments:

   **C-shell:**
   ```
   setenv CHEMKIN_DIR $HOME/chemkin36
   ```

   **K-shell:**
   ```
   CHEMKIN_DIR=$HOME/chemkin36
   ```
   ```
   export CHEMKIN_DIR
   ```

2. To verify that the `CHEMKIN_DIR` is set correctly, open a new shell and type:

   ```
   echo $CHEMKIN_DIR
   ```

   This should result in an echo of the directory path to the top-level `chemkin36` directory. If not, 
verify that you have performed step 1 correctly; you may need to log-out and log back in to your 
computer for the changes to take effect.
3.3.4 Modifying your “Path”

We recommend modifying your “path” variable to include the $HOME/chemkin36/bin directory (or the equivalent, if you installed elsewhere from $HOME). This will simplify the commands that you need to type to start the CHEMKIN Application User Interface, the CHEMKIN Graphical Post-processor, or to run programs from the command line. The path variable can be modified in your .login or .cshrc files for C-shell users, or in your .profile for Bourne or Korn shell users with the following added line(s):

**C-shell:**
```
set path=( $path $HOME/chemkin36/bin )
```

**K-shell:**
```
PATH=$PATH:$HOME/chemkin36/bin:
```

```
export PATH
```

**Note:** if you installed CHEMKIN in a different directory than $HOME, you will need to replace the actual (full) path in the above commands.

3.3.5 UNIX Licensing Overview

You must obtain and install a set of license keys in order to run the CHEMKIN Collection software. This section provides instructions on how to obtain and install a Reaction Design license file for the CHEMKIN Collection, once you have purchased the software. For purchasing information, contact by email: CHEMKIN@ReactionDesign.com.

**Assumptions:**

We assume the following for purposes of providing instructions on obtaining and installing a license on a UNIX workstation:

1. You have already installed CHEMKIN Release 3.6 on your computer.

2. The default installation directory for your UNIX workstation is $HOME/chemkin36. If you have installed in a different location, please make the necessary replacements in the instructions that follow.

**Required Tasks:**

1. Request a license from Reaction Design, providing information about your computer.

2. Install the license received from Reaction Design.
3.3.6 Request a License from Reaction Design

Once you have purchased a license, you will be asked to provide the “HOSTID” for the machine on which CHEMKIN will be installed, so that we can generate the needed license file. You can get this information by changing directories to the licenses subdirectory of the CHEMKIN installation and running the command (in a UNIX shell):

```
lmutil lmhostid
```

This returns the statement “The Flexlm hostid of this machine is “xxxx”, where “xxxx” will be a unique identifier or HOSTID for your machine. If the result is “ffffffff”, it probably means you don’t have a network card installed on your machine. In this which case, issue the command

```
lmutil lmhostid -vsn
```

This returns the statement ‘The Flexlm host ID of this machine is “DISK_SERIAL_NUM =101717e5”.’

Once you have obtained a unique identifier, please send the following information to Reaction Design (email to CHEMKIN@ReactionDesign.com) to receive your license file:

1. The complete result of the “lmutil lmhostid” command.
2. Your CHEMKIN License Number (located on your “License Specifications Form”).
3. The Licensee Contact for your organization (located on your “License Specifications Form”).

Upon receipt of your computer information, Reaction Design will generate a set of license keys, specific to your computer and to the selection of CHEMKIN packages that you have purchased. These license keys will be sent to you by email. The next sections describe how to install the license keys on your UNIX workstation.

3.3.7 Installing a Node-locked (or DEMO) License

After requesting a license, should receive by email a specific license to run CHEMKIN on your system. The file may have either UNIX or PC line-endings, depending on where and how you receive your email; if you need to transfer the license file from one machine to another, be sure to use ftp in ascii mode to guarantee the line endings get converted correctly. Once you have received an email with license keys from Reaction Design, you will need to place the license keys in the correct location and set an environment variable to point to the license path, as described below:
Installation and Quick Start

1. Save the received license keys into a text file named “chemkin.lic” and copy the file into the directory $HOME/chemkin36/licenses.

2. Set the environment variable called REACTION_LICENSE_FILE to point to the absolute path of the installed license file. The specific instructions depend on the type of UNIX shell that you use:

   **C-shell:**
   
a. Add the following line to your .cshrc file:
   ```
   setenv REACTION_LICENSE_FILE $HOME/chemkin36/licenses/chemkin.lic
   ```
b. Run the command:
   ```
   source .cshrc
   ```

   **K-shell or Bourne Shell:**
   
a. Add the following lines to your .profile file:
   ```
   REACTION_LICENSE_FILE=$HOME/chemkin36/licenses/chemkin.lic
   export REACTION_LICENSE_FILE
   ```
b. Run the command:
   ```
   . .profile
   ```

   **Note:** if the environment variable REACTION_LICENSE_FILE has already been set for another product (i.e., another product from Reaction Design), simply append the license path to the existing one. For example, if the existing definition is:
   ```
   REACTION_LICENSE_FILE=$HOME/simulator/flight.lic,
   ```
   then modify it to be:
   ```
   REACTION_LICENSE_FILE= \n
   $HOME/chemkin36/licenses/chemkin.lic:$HOME/simulator/flight.lic
   ```

You are now ready to run CHEMKIN; Skip the following sections and go directly to Section 3.4.

### 3.3.8 Installing a Network (Simultaneous Active User) License

In network licensing, more than one user may be running CHEMKIN Applications by accessing a centrally located license file that resides on a Network License Server. The license allows one or more simultaneous active users to run CHEMKIN Applications. Clients of the Network License Server should have their own local installation of CHEMKIN, but obtain a license from the Server while the CHEMKIN Application is running. The instructions here describe what needs to be done on both the Network License Server and on the Client machines.

**On the Network License Server:**

1. Install the license as described in Section 3.3.7 (assuming that the Server is UNIX machine).
2. Edit the license file using a text editor of your choice:
   a. Change the text “servername” in the line:
      
      SERVER servername ... 
      
      to the name (e.g. “MYCOMPUTER”) or IP Address (e.g. “192.168.0.1”) of the Network License Server.
   b. On the DAEMON line, verify that the PATH to reaction is correct; the default location is: $HOME/chemkin36/licenses/reaction

3. Start the Flexlm License Manager, lmgrd:

   (WARNING: DO NOT RUN AS ROOT)
   a. cd $HOME/chemkin36/licenses.
   b. Type (using full path names to files):
      
      lmgrd -c $HOME/chemkin36/licenses/chemkin.lic \\
      -l $HOME/chemkin36/licenses/chemkin.log
   c. Confirm that the daemon is actually running, using the following command. The command should show a “lmgrd” process in the returned list:
      
      ps -ef | grep lmgrd
      
      If the daemon lmgrd is not listed, check your log file for errors (e.g. “chemkin.log”). If you need further assistance, send email to support@ReactionDesign.com.

On each Network Client:

1. Place a copy of the modified Network license file in the “licenses” sub-directory of the client CHEMKIN 3.6 installation; by default this will be:
   Windows/PC Client:
   
   c:\chemkin36\licenses\chemkin.lic
   
   UNIX Client:
   
   $HOME/chemkin36/licenses/chemkin.lic

2. Now the client will be able to check out a license through the Server daemon.

   You are now ready to run CHEMKIN; Skip the following sections and go directly to Section 3.4
3.4 RUNNING THE SAMPLE PROBLEMS

Once CHEMKIN 3.6 and an appropriate license have been installed, you can launch the CHEMKIN Application User Interface and run any of the sample problems with just a few mouse clicks.

3.4.1 Launching the CHEMKIN Application User Interface

The way to launch the CHEMKIN Application User Interface depends on your platform:

**Windows/PC:**

In this case, the install program will have placed an icon labeled “CHEMKIN Collection 3.6” on your desktop. Just double-click the “CHEMKIN Collection 3.6” icon to launch the Application User Interface. You can also access either the CHEMKIN Application User Interface or the CHEMKIN Graphical Post-processor from the Windows Start → Programs → CHEMKIN Collection 3.6 menu.

**UNIX:**

If you set your “path” variable as instructed in Section 3.3.4 above, then you can open a UNIX shell and simply type:

```bash
chemkin <RETURN>
```

to launch the Application User Interface. If you did not set the path, you will have to include the full path to the CHEMKIN “bin” directory in front of the above command.

An interface as shown in Figure 1 should appear on your screen. **Note:** this manual includes illustrations from the PC version only; the interface will look slightly different on other platforms. However, the functionality is the same and the instructions provided here are the same for all computer systems.
Figure 1. Startup window for the CHEMKIN Application User Interface.

From here you can choose an Application, as shown in Figure 2.

Use the pull down menu or click on the Application button to start.

Figure 2. Selecting an Application to run in the Application User Interface.
3.4.2 Running a Sample Problem

For demonstration purposes, select the SPIN Application. The SPIN Application Interface, shown in Figure 3, should appear on your screen.

When CHEMKIN is first installed, the software is pre-set to start the user in a directory containing one of the sample problems for the chosen Application. For SPIN, the “Input” and “Output” directory text boxes should be set to the location of the CHEMKIN sample called “rotating_disk_cvd”. The names of the input and output files are also pre-set to the correct files in this sample directory. You can “Edit” any of the input files to see their contents. The pre-processors can be run separately by clicking on the individual “Chem”, “Surf” or “Tran” buttons, or you can run all of the needed preprocessors and the Application sequentially by clicking on the run “All” button.

![Chemkin Collection, Release 3.6](image)

**Figure 3. The SPIN Application User Interface**
Click the run “All” button now. Messages that describe the progress of the executing programs should appear in the Message Box (See No. 6 in Figure 4). If the CHEMKIN installation was successful and the License File has been installed correctly, the messages should report successful completion for the three pre-processors and the SPIN Application.

**Note for Windows:** On PCs, CHEMKIN 3.6 first creates MS-DOS prompt windows to run each of the CHEMKIN programs. You will see this sequence of windows flash on the screen as programs execute. These events are normal.
To view the results of the SPIN run, you may look at the output files using the “View” buttons (See No. 5 in Figure 4). You may also wish to view the results graphically. The next section describes the use of the CHEMKIN Graphical Post-Processor.

3.4.3 Launching the CHEMKIN Graphical Post-Processor

From the Application User Interface, you can start the CHEMKIN Graphical Post-Processor by simply clicking on the “Post-Process” button (See No. 7 in Figure 4). You can also launch the Graphical Post-Processor independently of the Application User Interface, as follows:

**Windows/PC:**

From the Windows desktop, go to: **Start → Programs → CHEMKIN Collection 3.6** and select “CHEMKIN Post-processor” to launch the Graphical Post-Processor.

**UNIX:**

If you set your “path” variable as instructed in Section 3.3.4, then you can open a UNIX shell and simply type:

```
chemkin_post <RETURN>
```

to launch the Graphical Post-Processor. If you did not add the CHEMKIN “bin” directory to your path, you will have to include the full path to that directory before “chemkin_post” in the above command.

From the Application User Interface, click “Post-Process” now. As the post-processor starts up, it will look for a solution file (save.bin) in the Output Directory indicated in the Application User Interface or in the user Preferences (see Section 6.4.1 for more information on setting Preferences). **(Note:** If a valid save.bin file is not found, a sample plot (sine/ cosine) will be displayed.) For the SPIN sample problem, the solution file includes both the solution data itself as well as sensitivity data for the gas-phase reactions, the surface reactions, and the heat-of-formation of species. The Data Selection window, shown in Figure 5, allows you to select which solution components will be imported for plotting purposes.
1. **Solution sets available for plotting**

2. **Species fraction or sensitivity filter options**

3. **Selection of plot variables by value or by species name (✓)**

4. **Plot results**

**Figure 5.** Data Selection window for importing data from the SPIN sample into the Graphical Post-Processor.

**General Note:** Any time there is more than one solution “subset”, such as sensitivity data or multiple solutions, or when the number of variables in the solution exceeds 20, the program first provides the user with an opportunity to down-select the number of arrays that will be imported into the Graphical Post Processor, with the Data Selection window. For smaller solution files, the program will go directly to the default plot.

By default, all of the species data for the SPIN solution will be read into the Graphical Post-Processor. **Click “OK”** to read the current selections into the post-processor. **Figure 6** shows the default plot that you should now see on your screen.
Figure 6. Default plot generated by the Graphical Post-Processor for the SPIN sample.

By default, the Graphical Post-Processor chooses the first two arrays to plot as “X” and “Y” in the initial plot. The dependent variable (in this case “Distance”) is typically the first array. To plot a different set of variables, use the “Plot” pull-down menu and choose “XY Plot”. This will give you a menu of the available arrays, as shown in Figure 7. To generate a new plot, leave the “X” variable the same, but select the other two velocity components from the list of available arrays, as shown in Figure 7.
Use the pull down menu to select X and Y variables for plotting

You can now change the look of the plot by double-clicking on the X-axis, the Y-axis, the Plot Title, or the Legend, and adjusting the settings in the properties panels. You can also drag-and-drop the Plot Title or the Legend for more optimal placement. Figure 8 shows the results of changing some of the settings for the new velocity plot.

This concludes the “Quick Start” instructions. You can exit the program by choosing “Quit” from the “File” pull-down menu or by simply closing the window(s). For more information on the CHEMKIN Collection or about Reaction Design, please see Chapters 4 and 5. For more information on using the
CHEMKIN Application User Interface or the CHEMKIN Graphical Post-Processor, see Chapters 6 and 7 respectively. If you would like to run CHEMKIN from the command line, rather than through the Application User Interface, you should refer to Chapter 8. Chapter 9 provides information on available documentation, as well as on how to get help if anything has gone wrong in your installation, or if you are having difficulty running any of the programs.
Figure 8. Example plot generated in the Graphical Post-Processor, where the plot attributes have been modified from the default settings by double-clicking on the plot features (e.g. axis, legend, and title).
4. Introduction to the CHEMKIN Collection

4.1 About Reaction Design

CHEMKIN Release 3.6 is part of Reaction Design's continuing commitment to expanding the capabilities of the original system developed at Sandia National Laboratories. In 1997, Sandia National Laboratories selected Reaction Design as the exclusive worldwide licensee for its CHEMKIN Collection. Today under this agreement, Reaction Design provides support and new developments for hundreds of industrial and academic users worldwide.

Reaction Design was founded in 1995 to provide software simulation and modeling tools to help process engineers create more efficient and environmentally friendly manufacturing processes. The company is committed to the ongoing development of a comprehensive and easy-to-use set of software simulation tools that incorporate both chemical reaction and transport models. Reaction Design engineers can also provide consulting services and software-based solutions to the customer's specific chemical process problems. These software products and consulting services help industrial customers develop better products at a lower cost and with a smaller impact on the environment. Reaction Design focuses its development, consulting and marketing efforts on four chemistry-intensive areas:

**Combustion:** Process heat and power generation, incinerators, jet and automotive engines, gas turbines and fuel design.

**Chemical Processing:** Chemical and pharmaceutical, oil refining, paper, high performance materials, corrosion and lubrication, food flavors and fragrances.

**Microelectronics:** Integrated circuit manufacturing, semiconductor processing equipment and reactive precursors, protective and optical coatings.

**Environment Protection:** Pollutant treatment and atmospheric chemistry.

More information about Reaction Design products and services is available through our web site at the address: www.ReactionDesign.com.
4.2 OVERVIEW OF CHEMKIN

The CHEMKIN software architecture represents a modular approach that separates problem-specific information from problem-independent software. For a reacting flow simulation, Figure 9 illustrates this approach. The Application represents the general model description, independent of what chemical species are included in a specific problem. The general model description typically includes conservation equations for mass, momentum, energy, and species. The upper ovals in Figure 9 contain the input data that make the problem specific, i.e. species identity, species properties, reaction paths, and reaction rates. The CHEMKIN software utilities provide the interface between this problem-specific information and the problem-independent Application. The Application may itself be broken down into modules that perform different functions, such as residual evaluation, matrix manipulation, solution printing, or post-processing. Additional problem-specific information that is non-chemical may also be input directly by the user, such as reactor pressure or inlet flow rate.

Figure 9. Illustration of the CHEMKIN modular approach to problem solving.
Included in the CHEMKIN distribution are sample problems for each Application, as well as detailed
documentation describing each model. In addition to the CHEMKIN Applications, described in Section 4.4,
the CHEMKIN Collection includes Utilities that facilitate user development of modular application
programs. The Utilities are described in Section 4.3 below.

4.3 THE CORE UTILITIES OF THE CHEMKIN COLLECTION

The following are brief descriptions of the components of the Core Utilities included in the CHEMKIN
Collection:

**CHEMKIN**
Pre-Processor (Interpreter) and Subroutine Library for the analysis of
gas-phase chemical and plasma kinetics.

**SURFACE CHEMKIN**
Pre-Processor (Interpreter) and Subroutine Library for the analysis of
heterogeneous chemical kinetics at gas-solid interfaces.

**Thermodynamic Database**
Compilation of polynomial fits to temperature for species enthalpy,
specific heat, and entropy.

**TRANSPORT**
Pre-processor and Subroutine Library for the evaluation of gas-phase,
multi-component transport properties including diffusion coefficients,
viscosities, and thermal conductivities of species and their mixtures.

**TRANSPORT Database**
Compilation of molecular parameters used in the calculation of transport
properties.

**TWOPNT**
A module for solving two-point boundary-value problems using a
modified Newton iteration method.

**EQUIL**
A program for predicting the equilibrium state of systems containing
ideal gas mixtures or ideal solutions of gases and liquids.
### 4.4 CHEMKIN APPLICATIONS

The following are brief descriptions of the Applications available in the CHEMKIN Collection:

<table>
<thead>
<tr>
<th>Application</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>AURORA</strong></td>
<td>A program for predicting the time-dependent or steady-state properties of a well mixed or perfectly stirred reactor for thermal or plasma chemical systems. AURORA allows inclusion of surface chemistry on multiple materials in the reactor and optionally performs time-dependent sensitivity analysis of solution results.</td>
</tr>
<tr>
<td><strong>CRESLAF</strong></td>
<td>A program for modeling laminar, chemically reacting, boundary-layer flow in cylindrical or planar channels. Surface chemistry on the inner walls of the channel may be included.</td>
</tr>
<tr>
<td><strong>OPPDIF</strong></td>
<td>A program for modeling opposed-flow diffusion flames. OPPDIF includes homogenous, gas-phase kinetics with sensitivity analysis.</td>
</tr>
<tr>
<td><strong>PLUG</strong></td>
<td>A program for the analysis of plug-flow reactors with gas-phase and surface chemistry.</td>
</tr>
<tr>
<td><strong>PREMIX</strong></td>
<td>A program for modeling steady, laminar, one-dimensional, pre-mixed flames. PREMIX includes homogenous, gas-phase kinetics with sensitivity analysis.</td>
</tr>
<tr>
<td><strong>SENKIN</strong></td>
<td>A program for predicting the time-evolution of homogenous, gas-phase kinetics with sensitivity analysis.</td>
</tr>
<tr>
<td><strong>SHOCK</strong></td>
<td>A program for predicting chemical behavior behind incident and reflected shock waves.</td>
</tr>
<tr>
<td><strong>SPIN</strong></td>
<td>A program for modeling one-dimensional, rotating-disk or stagnation-flow chemical-vapor-deposition reactors. SPIN includes homogenous gas-phase and heterogeneous gas-surface kinetics with sensitivity analysis.</td>
</tr>
<tr>
<td><strong>SURFTHERM</strong></td>
<td>A program for analyzing thermochemical and kinetic data in gas-surface chemical reaction mechanisms.</td>
</tr>
</tbody>
</table>

There are several advantages of the modular approach to chemically reacting flow simulations. One is that a user may “mix and match” Applications and chemistry descriptions. Users who are interested in the determination of dominant reaction mechanisms, for example, are free from having to repeatedly develop computational tools with large systems of chemical reactions. The CHEMKIN input-file standard also facilitates communication and sharing of reaction sets between researchers. An analyst who must
perform calculations for several different chemical mixtures is free from concern over software programming. This also leads to more robust software that is applied to a wide variety of problems.

4.5 CONTENTS OF THE CHEMKIN COLLECTION 3.6 PACKAGES

CHEMKIN can be purchased in several different “packages” tailored to the needs of individual customers. The following is a brief description of the packages that are currently available:

Full Package

Utility Set
CHEMKIN Interpreter and Gas Phase Subroutine Library, SURFACE CHEMKIN Interpreter and Subroutine Library, Thermodynamic Database, TRANSPORT Preprocessor and Database, TWOPNT, and EQUIL

Applications
AURORA, CRESLAF, OPPDIF, PLUG, PREMIX, SENKIN, SHOCK, SPIN, and SURFTHERM

Chemistry Package

Utility Set (See above)

Applications
AURORA, CRESLAF, PLUG, SENKIN, and SURFTHERM

Combustion Package

Utility Set (See above)

Applications
AURORA, OPPDIF, PLUG, PREMIX, and SENKIN

Microelectronics Package

Utility Set (See above)

Applications
AURORA, CRESLAF, SPIN, SENKIN, and SURFTHERM

Customer Designed Packages

Utility Set (See above)

Applications
Customer choice
4.6 HOW DOES CHEMKIN WORK?

The CHEMKIN modular approach described in Section 4.2 provides a broad overview of the CHEMKIN philosophy. Here we describe more specifically how the CHEMKIN Collection provides the necessary interface between general Applications and chemistry-specific information. The flow chart in Figure 10 illustrates components of the CHEMKIN software for a typical application. The major software components can be categorized as preprocessors, subroutine libraries, application models, and databases. The flow chart illustrates the communication between these components for a typical application model.

Figure 10. Schematic showing the relationship between the CHEMKIN Utilities and a CHEMKIN Application, including input and output files.
Pre-processors include the CHEMKIN Interpreter, the SURFACE CHEMKIN Interpreter, and the TRANSPORT Fitting Program. The pre-processors read information that is either supplied directly by the user, or is obtained from one of the CHEMKIN databases, or both. The user-input data tells the pre-processors what species to consider and what reactions can occur between these species. The pre-processors parse and interpret the user information, compile species property information from the databases, and calculate any additional chemistry-specific information that may be needed by the Application and that are appropriate to determine a priori. This information is then stored in “Linking Files” that are later accessed by the CHEMKIN Subroutine Libraries when called by the Application. The CHEMKIN Interpreter must always precede the other pre-processors as it provides information that is used by the others through its Linking File.

Subroutine libraries include the CHEMKIN Gas-phase Library, the SURFACE CHEMKIN Library, and the TRANSPORT Library. Each of these libraries includes an initialization routine that reads the information in the Linking File, stores the information in arrays and returns these storage arrays to the Application for use in subsequent calls to other library routines. The CHEMKIN Gas-phase Library contains routines that provide information about the size of the chemistry set; information about the elements, species, and reactions; values of physical constants; equation-of-state calculations and units conversions; thermodynamic properties; calculations of chemical production rates; and reaction equilibrium constants. The SURFACE CHEMKIN Library contains routines that perform similar functions for the surface chemistry set. The TRANSPORT Library contains routines that evaluate gas-phase species and gas-mixture transport properties including ordinary diffusion coefficients, thermal diffusion coefficients, thermal conductivities, and viscosities. The transport properties can be evaluated using either mixture-averaged or multicomponent formulations.

Currently, the CHEMKIN Collection includes databases of fundamental species data for calculation of transport and thermodynamics properties. Details on the species data available can be found in the Thermodynamic Data and TRANSPORT manuals. Users may also augment or replace database entries by providing their own sets of data.

We have compiled a list of journal articles and other publications that have referenced CHEMKIN Collection software since 1985. These citations may be useful in discovering how other people are using the CHEMKIN Collection in research and industrial applications. The citation bibliography is included on our website (www.ReactionDesign.com) and is updated periodically.
5. Problem Solving with the CHEMKIN Collection

5.1 How to Use CHEMKIN 3.6 to Solve Chemical Modeling Problems

The CHEMKIN software can be used in several different ways, depending on your own areas of expertise and your modeling needs. The most straightforward way to use the CHEMKIN software is by direct application of the models included in the CHEMKIN Collection. A brief summary of the Applications is shown in Table 1.

<table>
<thead>
<tr>
<th>Application</th>
<th>Brief Description of the CHEMKIN Application</th>
</tr>
</thead>
<tbody>
<tr>
<td>AURORA</td>
<td>Well stirred thermal and plasma reactors with gas and surface reactions.</td>
</tr>
<tr>
<td>CRESLAF</td>
<td>Laminar, boundary layer flows in cylindrical or planar channels</td>
</tr>
<tr>
<td>EQUIL</td>
<td>Chemical equilibrium of an ideal gas or solution mixture</td>
</tr>
<tr>
<td>OPPDIF</td>
<td>Diffusion flame between two opposing nozzles</td>
</tr>
<tr>
<td>PLUG</td>
<td>Plug flow reactor with gas and surface chemistry</td>
</tr>
<tr>
<td>PREMIX</td>
<td>Steady, laminar one-dimensional pre-mixed flames</td>
</tr>
<tr>
<td>SENKIN</td>
<td>Sensitivity analysis of homogeneous gas-phase kinetics</td>
</tr>
<tr>
<td>SHOCK</td>
<td>Chemical dynamics behind incident and reflected shock waves</td>
</tr>
<tr>
<td>SPIN</td>
<td>Rotating-disk/ stagnation flow chemical vapor deposition reactors</td>
</tr>
<tr>
<td>SURFHERM</td>
<td>Analysis of thermochemical and kinetic data in gas-surface reaction mechanisms</td>
</tr>
</tbody>
</table>

Additional applications can be built using the sub-components or utilities of the CHEMKIN Collection. This mode requires a much higher level of modeling expertise, but offers more flexibility in describing a specific system of interest to you. Table 2 presents a brief description of the utilities.
### Table 2  CHEMKIN Utilities and Supporting Databases

<table>
<thead>
<tr>
<th>CHEMKIN Utility/Database</th>
<th>Brief Description of the CHEMKIN Utility/Database</th>
</tr>
</thead>
<tbody>
<tr>
<td>CHEMKIN</td>
<td>Library of analysis tools for gas-phase chemical and plasma kinetics</td>
</tr>
<tr>
<td>SURFACE CHEMKIN</td>
<td>Library of analysis tools for chemical kinetics at gas-solid interfaces</td>
</tr>
<tr>
<td>Thermodynamic Property Database</td>
<td>Temperature variation of species enthalpy, specific heat, and entropy</td>
</tr>
<tr>
<td>TRANSPORT</td>
<td>Library of tools for evaluation of multi-component transport properties</td>
</tr>
<tr>
<td>TRANSPORT Property Database</td>
<td>Molecular properties used in calculating transport properties</td>
</tr>
<tr>
<td>TWOPNT</td>
<td>Algorithms for solution of two-point boundary value problems</td>
</tr>
</tbody>
</table>

Each of the Application and Utility manuals describe in much more details how to build or extend the capabilities of the existing programs.

### 5.2 Steps in Using CHEMKIN 3.6

There are six basic steps involved in using CHEMKIN to solve a typical chemical process modeling problem. Depending on the particular Application, some of the steps are not required; these are indicated by ( ) around the step.

1. **Prepare the chemistry input file(s)**
   - Gas-phase chemistry
     - (Surface chemistry)

2. **Assemble the thermodynamic and transport databases**
   - Gas-phase thermodynamics
     - (Surface thermodynamics)

   - (Gas-phase species transport data)
3. **Prepare the Application-specific input data**

   (Geometry)

   (Process conditions)

   (Solution method options)

4. **Run the Pre-Processing programs**

   Chem  -  The CHEMKIN gas-phase reaction mechanism interpreter

   (Tran)  -  The TRANSPORT property fitting program

   (Surf)  -  The SURFACE CHEMKIN reaction mechanism interpreter

5. **Run the Application**

6. **Post-process the results**

Table 3 summarizes the input data requirements for each of the CHEMKIN Applications. The specific details of the data formats, and information needs are described in the user manuals for the Applications and the CHEMKIN Utilities.

<table>
<thead>
<tr>
<th>Application</th>
<th>Gas-phase Chemistry</th>
<th>Surface Chemistry</th>
<th>Application-specific Input</th>
<th>Thermodynamic Database</th>
<th>Transport Properties Database</th>
</tr>
</thead>
<tbody>
<tr>
<td>AURORA</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✔</td>
</tr>
<tr>
<td>CRESLAF</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>EQUIL</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>OPPDIF</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>PLUG</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>PREMIX</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>SHOCK</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>SENKIN</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>SPIN</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>SURFTHERM</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
</tbody>
</table>

An illustration of some of the steps is presented in Figure 11.
Problem Solving with the CHEMKIN Collection

1. CHEMKIN 3.6 
   Startup Panel

2. Application Panel

3. Chemistry Input Data

4. Application-specific Input Parameters

5. CHEMKIN Graphical Post-processor

Figure 11. A schematic representation of the information flow in running a typical CHEMKIN Application.
5.3 WHERE TO GET THE NEEDED INPUT DATA?

A typical question is, “I can't find data for certain species in the Thermodynamic or Transport database. What should I do?” First, check the complete species list in the Thermodynamic Database and Transport manuals. These data were originally supplied to Reaction Design by Sandia. While the original data has not been officially updated, Reaction Design continues to assemble information for additional species that will be added to future releases. If you are unable to locate certain species in the Thermodynamic or Transport databases there are several other sources for developing the needed information including:

1. Original Sources: These are organizations that generate or critically evaluate data. One of the most important is the National Institute of Standards and Technology (NIST).

2. Secondary and Tertiary Sources: These are compilations of data from original sources and are often published in handbooks and journals. For chemical kinetics a good starting point is the J. Physical and Chemical Reference Data.


4. Web Sites: The Reaction Design web site contains links to many different sources of thermochemical and transport property information.

Many other sources of data are described in Maizell (1998), How to Find Chemical Information: A guide for Practicing Chemists, Educators and Students, John Wiley & Sons, Inc., New York.
6. The CHEMKIN Application User Interface

This Section provides a more complete description of the elements of the CHEMKIN Application User Interface. The information here includes descriptions of menus and buttons as well as quick reference “Hints” on program features. We recommend that you first read through Section 3.4 so that you are familiar with starting the Application User Interface and you are sure that your software is installed correctly.

6.1 Selecting Input and Output Files

The Application User Interface is designed to help you organize your input and output files, and to remind you of the steps and input data required to run each Application. For an illustration of the Application User Interface features, see Figure 4 in Chapter 3. These features are described in more detail here.

Directories

You should first set the working directories where you will store the input and output files. By default, the program assumes that the output directory is the same as the Input Directory.

Hints:

• The directory must already exist.
• You can make the Output Directory independent of the Input Directory by modifying the user Preferences (Choose “Preferences” from the “File” pull-down menu).
• You can change the default directory that the program assumes at startup in the user Preferences (Choose “Preferences” from the “File” pull-down menu).

Input File Names

The input files that are used by the pre-processors and the Application must reside in the Input Directory. By default, the program will suggest filenames in the text boxes. To select a different filename, you can either type the new name in the text box or click “Browse” to select a new file. You can also click on “Edit” to modify the selected file or to create a new file.
Introduction to the CHEMKIN Application User Interface

Hints:

• Since the text box is small on some platforms, using the "Browse" button may be more reliable than typing to ensure that you have the correct input file selected.

• You may change the default editor invoked by the “Edit” buttons by modifying the user Preferences (Choose “Preferences” from the “File” pull-down menu).

• Remember: your input files must exist in the specified Input Directory.

Output File Names

The Pre-processors and the Application program will create output files in the specified Output Directory. By default, this is the same directory as the Input Directory. Also by default, the Application User Interface will suggest filenames in the text boxes. To select a different filename, you can either type the new name in the text box or click “Browse” to select a new file. Once you have clicked on “Browse” you may still need to type in the filename, since this file typically does not yet exist. **If you choose an existing file, the file will be overwritten when the program is run.** After the programs have been run, you can view the output files by clicking on their “View” button.

Hints:

• Since the text box is small on some platforms, using the “Browse” button may be more reliable than typing to ensure that you have the correct output filename entered.

• You may change the default editor invoked by the “View” buttons by modifying the user Preferences (Choose “Preferences” from the “File” pull-down menu).

• Remember: the output files will always be created in the specified Output Directory.

Database Files

Unlike the Input and Output Files, the Database Files are specified by providing the full directory path to the required files. These files typically will not reside in the Input Directory or the Output Directory, although it is OK if the directory paths happen to be the same. By default, the database files selected are the standard Thermodynamic Database file and the Transport Database file distributed with CHEMKIN. To change to a different file, click “Browse” and navigate to the desired file. You can also click on “View” to open the selected file in an editor.

Hints:

If the Transport Database is not required for a specific Application, then the selection features will become inactive.
6.2 Running the Programs

As described in Chapter 5, solving a problem using CHEMKIN requires running one or more pre-processors as well as running the Application program itself. The right-hand column of the Application User Interface allows you to run these programs one at a time or, alternatively, to run all needed pre-processors and the Application by clicking the run “All” button. In general, the order in which the pre-processors is run is important: all required pre-processors must be run before the Application, and the “Chem” pre-processor should always be run first. The two options are described in more detail below.

Run “Chem”

The “Chem” button runs the CHEMKIN Interpreter using the specified gas-phase chemistry input file, as well as the Thermodynamic Database File. The CHEMKIN Interpreter creates a Linking File that is required by the other pre-processors and by the Application. For more information about the CHEMKIN Interpreter program and input, refer to the CHEMKIN user manual.

Run “Surf”

The “Surf” button runs the SURFACE CHEMKIN Interpreter using the specified surface chemistry input file, as well as the Thermodynamic Database File. The SURFACE CHEMKIN Interpreter creates a Linking File that is required by the Application. For more information about the SURFACE CHEMKIN Interpreter program and input, refer to the SURFACE CHEMKIN user manual.

Run “Tran”

The “Tran” button runs the TRANSPORT Fitting Routine using the specified Transport Database File. The TRANSPORT program creates a Linking File that is required by the Application. For more information about the TRANSPORT program and input, refer to the TRANSPORT user manual.

Run the Application

Each Application User Interface panel has a button labeled with the Application name and aligned with the Application Input and Output File selection features. This Application button runs the Application program using the specified Input File. The Application will create (or overwrite) the Output File. In addition to the text Output File, all of the Applications (except for SURFThERM) generate a binary solution file named save.bin. The save.bin file is read by the Graphical Post-Processor to plot the solution results, as described briefly in Chapter 7 and in more detail in Chapter 7.

Running “All”

The “All” button automatically runs the required sequence of pre-processors and then runs the Application, as described by the individual buttons above.
Hints:

When running Chem and Surf, the program will create a local copy of the Thermodynamic Database File in the Output Directory, and rename the file “therm.dat”. If there is already a therm.dat file in that directory, you will be prompted whether or not to overwrite the existing file. Similarly, running Tran will try to create a local tran.dat file, and will prompt you for a response if an existing file is found. You can allow the program to always overwrite the therm.dat and tran.dat files found in the Output Directory, by modifying your user Preferences. (Choose “Preferences” from the “File” pull-down menu).

6.3 INFORMATIONAL MESSAGES AND ERROR REPORTS

The Message Box

The Message Box, located at the bottom of the Application User Interface panel, helps you to follow the progress of your work. The messages tell you what command is being executed, when it has finished executing, and whether or not the process has completed successfully.

Error, Warning and Information Dialogs

In addition to the advisory messages in the Message Box, you may also see pop-up Information, Warning, or Error Dialog boxes. These Dialogs inform you that some operation did not complete properly, that you need to make a choice between one or more options, or that an operation about to occur may result in some unwanted effects (such as overwriting files). The message may also provide some suggested action that you can undertake to correct or avoid the problem. In order to continue working, you must click on a button in the box (“OK”, “Yes”, “No”, etc.).

6.4 THE “FILE” MENU

The “File” pull-down menu is located at the left-hand end of the menu bar on the Application User Interface. There are four options available under this menu in Release 3.6: “Save Session”, “Recover Session”, “Preferences” and “Quit”. The “Quit” option simply exits the CHEMKIN Application User Interface. The other three options are discussed further below. Figure 12 demonstrates the selection of “Preferences” from the “File” menu.
Select User Preferences from File menu

Figure 12. Selection of Preferences from the File pull-down menu.
6.4.1 Preferences

The Preferences option provides you with some control over the settings for the CHEMKIN Application User Interface, as well as the settings for the CHEMKIN Graphical Post-Processor. The Preferences selection panel is shown in Figure 13.

**Note:** For a few of the preference options, you will need to restart the CHEMKIN Application User Interface or the CHEMKIN Graphical Post-Processor before the changes take effect.

**Hints:**

If the path to the directory containing your Editor program is contained in your environment “path”, then only the edit command must be entered in the Preferences Editor box (for example, “wordpad” on Windows). If not, then you must include the full directory path to the editor program for the editor command to work properly (for example, “c:\Program Files\Accessories\wordpad”).

6.4.2 Saving and Restoring Sessions

Once you have set up a problem in the Applications User Interface panel, you can save your directory, filename, and database settings using the “Save Session” option in the “File” menu. A dialog will prompt you for a filename and a directory location for the Session File.

A previous session’s settings can be recovered by selecting the “Recover Session” option in the “File” menu. A file browser allows you to select the previously created Session File.

By default, you will be prompted to save a session file every time you quit the Application User Interface, or when you change from one Application to another. This prompting can be disabled in the Preferences menu.

**Hint:**

When you recover a Session File, you should be in the same Application User Interface panel (e.g. SPIN), as you were when you created the Session File.
Introduction to the CHEMKIN Application User Interface

1. Default working directories for input/output and data files

2. Options for running CHEMKIN Applications

3. Text editor for viewing or editing files

4. Limit arrays to improve plotting speed

5. Units for species fractions in plotting

Figure 13. The Preferences selection panel for the Application User Interface and for the Graphical Post-Processor
7. The CHEMKIN Graphical Post-Processor

This Section provides a more complete description of the elements of the CHEMKIN Graphical Post-Processor. The information here includes descriptions of menus and buttons as well as quick reference “Hints” on program features. We recommend that you first read through Section 3.4 so that you are familiar with starting the Graphical Post-Processor and you are sure that your software is installed correctly.

7.1 The “File” Pull-down Menu

The “File” pull-down menu is located at the left-hand end of the menu bar that runs across the top of the Graphical Post-Processor window. There are six operations available in CHEMKIN Release 3.6: “Open”, “Print”, “Import”, “Export”, “Preferences” and “Quit”. The “Quit” option simply exits the Graphical Post-Processor. The other five options are discussed below.

7.1.1 Open

The “Open” option allows you to open a new CHEMKIN binary solution file, which the program expects to have a “.bin” suffix. Selecting the “Open” option puts you in a file browser on your computer, where you can choose an existing binary solution file. Once the file is selected, the program will attempt to read the data on the file. If there are any difficulties reading the file, a message will appear in the Message Box at the bottom of the Graphical Post-Processor window.

As described in Section 3.4 when you open a new binary solution file, the program will either open a Data Selection window, or will directly create a default plot from the data that was read. The Data Selection window is opened whenever there are more than 20 variables in the solution or when there is more than one solution sub-set.

**Hints:**

When reading in a new save.bin file, you will be prompted to keep or clear existing data sets in the Graphical Post-Processor. Keeping more than one solution data set open in the Post-processor allows you to plot one solution against another.
7.1.2 Import

You can import x-y data from plain text files into the Graphical Post-Processor using the “Import” option in the “File” menu. The import utility expects to find columns of data, delimited by tabs, commas, or spaces. Selecting the “Import” option opens a dialog box where you can choose the delimiter for the columns in the file, the number of header lines to skip before reading the data, and whether or not to read the first line (after the skipped lines) as character-string labels for the columns. If your selections do not appear to agree with the format of the first couple of lines of data in the file, you will receive an warning message. If you click “OK” to read the data in and the import utility encounters errors reading the specified format, you will receive a message in the Message Box, and the data will not be imported. If the format is read correctly, the first two columns found in the file will be used to create a default plot.

7.1.3 Export

To give you more flexibility in analyzing and post-processing your data, you can export all of the data arrays currently stored in the Graphical Post-Processor into a file of text columns. The columns in the exported text files may be delimited by tabs, commas, or spaces. Such formats can be easily imported into spreadsheet programs, as well as other visualization and analysis software. When you select the “Export” option from the “File” menu, you will be asked to choose the delimiter for the exported data columns, and then the name and location of the data file that will be written. If there are any errors writing the file, a message will appear in the Message Box at the bottom of the Graphical Post-Processor window.

7.1.4 Print

Any plot generated on your screen can be printed to a color or black-and-white printer using the “Print” utility. This requires that you already have a print driver installed and a default printer set on your computer. You can also print to a Postscript file, by selecting the file option on the printer dialog.

7.1.5 Preferences

As described in Chapter 6, Section 6.4.1, the “Preferences” option provides you with some control over the settings for the CHEMKIN Application User Interface, as well as the settings for the CHEMKIN Graphical Post-Processor. The main setting for the CHEMKIN Graphical Post-Processor is a choice of whether the species composition data are imported from CHEMKIN solution files as mole fractions or mass fractions. The default is to convert the data to mole fractions. Note: if you change the settings in the Preferences dialog, you will need to restart the CHEMKIN Graphical Post-Processor before the changes take effect.
7.2 The “View” Pull-Down Menu

Two options are available from the “View” pull-down menu, “Show/Hide Legend” and “Show/Hide Title”. These options allow you show the Plot Title or Legend, if you have previously selected to “Hide” them in the Plot Title or Legend Attributes Dialog. Alternatively, if you have previously selected to “Show” the Plot Title or Legend (the default), you can use the “View” menu to “Hide” these items.

7.3 The “Plot” Pull-Down Menu

Although we plan to have more plot options in future CHEMKIN releases, the current release only allows creation of X-Y plots. Choosing the “X-Y Plot” option in the “Plot” menu opens a window from which you can select the data arrays to plot as “X” (horizontal axis) or “Y” (vertical axis) variables in the Graphical Post-Processor.

7.4 Changing Plot Attributes

By using your mouse to double-click or drag-and-drag items in the plot window, you can change the appearance of the plot as displayed in the Graphical Post-Processor. Table 4 summarizes what items can be accessed by the mouse and what attributes can be changed.
### Table 4 Plot Attributes That Can Be Changed

<table>
<thead>
<tr>
<th>Item</th>
<th>Mouse Action</th>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>X-Axis / Y-Axis</td>
<td>double-click</td>
<td>Axis Label Text</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Axis Label Font and Font Size</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Linear/Log Axis Scaling</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Axis Minimum/Maximum values</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Tick Direction, Number, and Size</td>
</tr>
<tr>
<td>Plot Title</td>
<td>double-click</td>
<td>Title Text</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Font and Font Size</td>
</tr>
<tr>
<td>Plot Title</td>
<td>click-and-drag</td>
<td>Title Position on Plot</td>
</tr>
<tr>
<td>Legend</td>
<td>double-click</td>
<td>Hide/Show Legend</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Hide/Show Outline</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Hide/Show Legend Title</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Legend Title Text</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Legend Title Font and Font Size</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Legend Item Labels</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Legend Item Font and Font Size</td>
</tr>
<tr>
<td>Legend</td>
<td>click-and-drag</td>
<td>Legend Position on Plot</td>
</tr>
</tbody>
</table>

### 7.5 Plotting Solutions Against External Data or Other Solutions

A new feature of CHEMKIN Release 3.6 is the ability to import more than one solution set at one time into the CHEMKIN Graphical Post-processor. To demonstrate this feature, let’s assume that we have some experimental temperature data that we want to compare to the temperature profile calculated in the “freely propagating flame” sample problem for PREMIX. The procedure below describes how to import data and compare to a Chemkin Application solution. The same procedure can be used to plot one solution against another, except in that case we would use “Open” from the file menu instead of “Import” to obtain the second data set.

**Temperature Data for Comparison:**

First, you will need to save the “experimental” data in a text file format that can be imported into the CHEMKIN Graphical Post-processor. Example contents of a (fictitious) data file are shown below:
To follow along with this example, the above data needs to be saved in a file named “mydata.csv,” in the PREMIX “freely propagating flame” sample directory.

Run the PREMIX Sample Problem

Next, run the “freely_propagating_flame” sample problem, which is located in a subfolder of the samples “premix” directory. Follow the same approach as described in Section 3.4.2 for the SPIN sample. When the PREMIX Application has run successfully, launch the Graphical Post-Processor. Select “SOLUTION_3_VS_DISTANCE” in the Data Select panel, to select the last, most-refined solution. The default plot will be Temperature vs. Distance.

Plot Imported Data Against a Solution in the Graphical Post-Processor

With the temperature plot displaying in the Graphical Post-Processor, select “Import” from the File menu. You will first see the “Keep or Clear” dialog window shown in Figure 14. Click on the “Premix Solution Data” to indicate that you want to keep this set (or click on the “KEEP ALL” button) and “OK”. Then, browse to the “mydata.csv” text file. In the Import dialog box, select “Comma” as the delimiter and check the “Read column titles” box, and click “OK”. The default plot that appears will be a plot from the imported data only. Now, go to the Plot⇒XY-Plot menu. You will next see a window that shows both the PREMIX Solution and the Imported Data Sets. For each, click on the “Select XY” button and make select Temperature for “Y” and Distance for “X”, as shown in Figure 15. The result is shown in Figure 16 after modifying the plot attributes.
1. Choose Import from File menu
2. Opt to Keep the existing solution
3. Open file with external data
4. Import comma-delimited values

Figure 14. Importing external data to plot against CHEMKIN solution results. Note: the File browser will look different on different platforms.
Figure 15. Selection of plot variables for multiple data sets displayed in same plot.
Figure 16. Comparison Plot of PREMIX Solution and Imported Data.

7.6 Units of Solution Variables

The binary solution files (save.bin) produced by the CHEMKIN Applications include values for solution variables, (e.g. velocities, temperatures, pressure, and species fractions), which have physical units (e.g. [cm/s], [K], [dyne/cm²], and mole fractions). In some cases, the variable names assigned to the arrays indicate the units (e.g. a variable labeled “Pressure_In_Torr” would indicate that the date has the units of Torr instead of the default). In all other cases, you can assume that the units are the standard units for the “cgs” system (centimeters, grams, seconds). Table 5 below indicates the cgs units for some of the common solution variables:
### Table 5 CGS Units for Common Solution Variables

<table>
<thead>
<tr>
<th>Variable</th>
<th>CGS Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time</td>
<td>s</td>
</tr>
<tr>
<td>Distance</td>
<td>cm</td>
</tr>
<tr>
<td>Velocity</td>
<td>cm/s</td>
</tr>
<tr>
<td>Area</td>
<td>cm²</td>
</tr>
<tr>
<td>Pressure</td>
<td>dyne/cm²</td>
</tr>
<tr>
<td>Density</td>
<td>g/cm³</td>
</tr>
<tr>
<td>Temperature</td>
<td>K</td>
</tr>
<tr>
<td>Mass flow rate</td>
<td>g/s</td>
</tr>
<tr>
<td>Volume flow rate</td>
<td>cm³/s</td>
</tr>
<tr>
<td>Deposition/Etch Rate</td>
<td>mole/cm²-s</td>
</tr>
<tr>
<td>Molecular weight</td>
<td>g/mole</td>
</tr>
<tr>
<td>Species Fraction</td>
<td>Determined by user Preferences (see Section 7.1.5)</td>
</tr>
</tbody>
</table>

For variables not listed in Table 5, see the specific Application user manual for information about units of variables.
8. Using CHEMKIN from the Command Line

We recognize that there will always be some users who prefer to run CHEMKIN by typing commands from a command line (in a MS-DOS prompt or a UNIX shell, for example), rather than using the CHEMKIN Application User Interface. Also, if you are writing your own applications or your own post-processing routines, you will need to be familiar with running CHEMKIN from a command line. This section describes the steps for running a sample problem, the makefiles and build scripts available in the CHEMKIN distribution, as well as the FORTRAN post-processors that are available as templates for writing your own post-processing routines. For most uses of command-line programming, you will need to have a FORTRAN compiler installed on your platform. For Windows/PC users, the required compiler is Digital Visual Fortran version 5.0D or later.

Note for Windows: To open an MS-DOS prompt in Windows, go to your Start menu, select Programs, and then select MS-DOS Prompt (Windows) or Command Prompt (Windows NT).

8.1 Running the CHEMKIN Sample Problems

The CHEMKIN distribution includes one or more sample problems for each Application. Generally, each sample consists of a set of input files, a “make” file that builds and runs the sample, and an HTML file that describes what type of problem the sample demonstrates. For the purposes of describing the commands in this section, we will refer to the root of the CHEMKIN installation as c:\chemkin36 for PCs and ~/chemkin36 for UNIX. If you changed the default location during installation, you will need to substitute the actual path where CHEMKIN is installed in place of these references.

8.1.1 Build All Samples

To build and run all the samples in the release, change directories to the samples subdirectory of the CHEMKIN installation (c:\chemkin36\samples for PCs or ~/chemkin36/samples for UNIX) and type:

```bash
buildsamples
```
**Note for UNIX:** On UNIX, you may not have the current directory (.) in your “path”. If typing buildsamples does not work in your UNIX shell, try prefixing the command with your current directory, like this:

```
./buildsamples
```

This command will launch a batch script that will build and run all of the samples by changing into each sample directory in turn and running the make file found there. Under the samples directory, there are subdirectories named for each Application. Within these subdirectories, there are different sample problems, such as “gas_psr” or “rotating_disk_cvd”, depending on the Application. Within these sample-problem directories, you can look at the sample input and output files with a standard text editor.

### 8.1.2 Clean All Samples

You can undo the buildsamples command by running cleansamples in the same manner as described in Section 8.1.1. This command removes all of the output and intermediate files from the sample sub-directories.

**General Note:** It is often desirable to run cleansamples to reduce the use of disk space.

### 8.1.3 Using the Make files

As mentioned above, each of the sample sub-directories contains a “make” file that will compile and link programs if necessary, run pre-processors, run the Application, and run a command-line post-processor if required. The make files obtain information about the build dependencies and machine-specific commands from files located in the “include” directory in the root of the CHEMKIN installation (c:\chemkin36\include for PCs or ~/chemkin36/include for UNIX). The include file is named chemkin_make_pc.inc on the PC and chemkin_make_unix.inc on UNIX.

**General Note:** The make files are designed to be used in the directory level where they are located, since they determine the location of the Include File relative to this directory. If you want to copy a makefile to another directory for your own use, you may need to modify the makefile before it will work properly in its new location.

As an example, we will demonstrate here how to run the AURORA sample problem called “gas_psr”. First, change directories to the AURORA gas_psr sample directory and then run the “make” utility as follows:
Using CHEMKIN from the Command Line

Windows/PC:

```
cd c:\chemkin36\samples\aurora\gas_psr
nmake -f gas_psr_pc.mak > gas_psr_pc.log
```

UNIX:

```
cd ~/chemkin36/samples/aurora/gas_psr
make -f gas_psr_unix.mak > gas_psr_unix.log
```

Note that the results of the make utility are redirected to a log file so that you can review any warnings or error messages that occur during the build process.

**Notes for Windows:**

- Notice that the “make” utility on the PC is called `nmake` instead of `make`
- You must have Digital Visual Fortran installed on your PC for the nmake command to be recognized.

8.1.4 Running Step by Step

In case you prefer not to use the makefiles provided or you want to build your own scripts, this section describes how to run each command directly on the command line. Here we will repeat the above example, but show the commands that you will need to type to the terminal for each step.

**General Note:** For all of the CHEMKIN pre-processors and Applications, all input and output files can now be specified using command-line arguments (e.g. “-i ” precedes an input file name). A complete list of the command-line argument options is given in Section 8.1.5.

1. **Change directories** to the AURORA gas_psr sample directory:

   Windows/PC:
   
   ```
cd c:\chemkin36\samples\aurora\gas_psr
   ```

   UNIX:
   
   ```
cd ~/chemkin36/samples/aurora/gas_psr
   ```
2. **Run the CHEMKIN Interpreter:**

   **Windows/PC:**
   ```
   ..\..\..\bin\chem -i chem.inp -o chem.out -d c:\chemkin36\data\therm.dat
   ```

   **UNIX:**
   ```
   ../../../bin/chem -i chem.inp -o chem.out -d ~/chemkin36/data/therm.dat
   ```

   Here, the command-line argument “-i” precedes the name of the gas-phase chemistry input file, while the command-line argument “-o” precedes the diagnostic output file name. The location of the thermodynamics database file is given by “-d”. In addition to the `chem.out` file, the CHEMKIN Interpreter creates an output file named `chem.asc`. In this example, the Linking File will be created by default in the local directory. The Linking File is required by other CHEMKIN pre-processors and by the Application. You should always check for errors by reviewing the contents of `chem.out`, before proceeding.

   **General Note:** For Applications that require TRANSPORT data (not required by AURORA) or SURFACE CHEMKIN (optional input for AURORA), you will also need to run the TRANSPORT and SURFACE CHEMKIN pre-processors. These additional pre-processors must be run after the CHEMKIN Interpreter and before the Application.

3. **Run the Application**, in this case the AURORA program:

   **Windows/PC:**
   ```
   ..\..\..\bin\aurora -i aurora.inp -o aurora.out -b save.bin
   ```

   **UNIX:**
   ```
   ../../../bin/aurora -i aurora.inp -o aurora.out -b save.bin
   ```

   In addition to the `aurora.out` file, the AURORA program creates a binary solution file called `save.bin`. This file is required by the CHEMKIN Graphical Post-Processor as well as the FORTRAN post-processor described below. You should always check for errors by reviewing the contents of `aurora.out`, before proceeding.
4. **Run the FORTRAN Post-Processor** for AURORA, called **AURORA_POST**:

Windows/PC:

```
..\..\..\bin\ aurora_post < aurora_post.inp > aurora_post.out
```

UNIX:

```
../../../bin/ aurora_post < aurora_post.inp > aurora_post.out
```

In the above commands, the "<" and ">" are used to redirect standard input and output, respectively, from and to the specified files. The **AURORA_POST** program reads the binary solution file, `save.bin`. In addition to the `aurora_post.out` output file, which contains diagnostics information, the post-processor creates one or more files named `*.csv`, where the "*" is a prefix determined by the input in the `aurora_post.inp` file. These files contain rows of comma-separated values representing the **AURORA** solution results (such as species mass fractions). In this particular sample, the number of rows is determined by the number of continuations (keyword **CNTN**) included in the **AURORA** input file. For transient problems, the number of rows will be the number of time steps.

**General Note**: All of the Applications in the **CHEMKIN** Release 3.6 have sample FORTRAN post-processors. The release includes full source code for these routines. The source code is located in the "post_processors" directory at the root of the **CHEMKIN** installation. These post-processors provide examples of how to read the binary solution files for exporting **CHEMKIN** solution data into custom or third-party applications.

### 8.1.5 Command-Line Argument Options

Table 6 provides a list of the available command-line argument flags for each of the **CHEMKIN** Pre-processor and Application programs. Table 6 also indicates whether each of the input or output files is always needed (A) or whether it is optional (O). For all required files, default file names are provided if files are not specified on the command line. These default file names are listed in the last column in Table 6. The command-line arguments allow specification of full pathnames to input and output files, in addition to the file names themselves. By default, any un-specified path is assumed to be the local working directory. Examples of using the command-line arguments are provided in Section 8.1.4.
Table 6 Command-Line Argument Options for CHEMKIN Pre-processors and Applications

<table>
<thead>
<tr>
<th>Flag</th>
<th>File Description</th>
<th>chem</th>
<th>surf</th>
<th>tran</th>
<th>aurora</th>
<th>creslaf</th>
<th>equil</th>
<th>oppdif</th>
<th>plug</th>
<th>premix</th>
<th>sentin</th>
<th>shock</th>
<th>spin</th>
<th>surftherm</th>
<th>Default File Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>-i</td>
<td>Input</td>
<td>A</td>
<td>A</td>
<td>O</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td></td>
<td></td>
<td>xxx.inp</td>
</tr>
<tr>
<td>-o</td>
<td>Output</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td></td>
<td></td>
<td>xxx.out</td>
</tr>
<tr>
<td>-d</td>
<td>Data</td>
<td>O</td>
<td>O</td>
<td>A</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td></td>
<td></td>
<td>yyy.dat</td>
</tr>
<tr>
<td>-r</td>
<td>Restart</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>O</td>
<td>O</td>
<td>--</td>
<td>O</td>
<td>--</td>
<td>O</td>
<td>--</td>
<td>--</td>
<td></td>
<td></td>
<td>rest.bin</td>
</tr>
<tr>
<td>-b</td>
<td>Binary Solution</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td></td>
<td></td>
<td>save.bin</td>
</tr>
<tr>
<td>-c</td>
<td>CHEMKIN Gas-phase Linking</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>A</td>
<td>chem.asc</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-s</td>
<td>Surface Linking</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>A</td>
<td>A</td>
<td>O</td>
<td>--</td>
<td>A</td>
<td>--</td>
<td>--</td>
<td>A</td>
<td>A</td>
<td></td>
<td>surf.asc</td>
</tr>
<tr>
<td>-t</td>
<td>Transport Linking</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>A</td>
<td>--</td>
<td>A</td>
<td>--</td>
<td>A</td>
<td>--</td>
<td>--</td>
<td>A</td>
<td>O</td>
<td></td>
<td>tran.asc</td>
</tr>
<tr>
<td>-v</td>
<td>Recover</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>A</td>
<td>--</td>
<td>--</td>
<td>A</td>
<td>--</td>
<td>--</td>
<td>A</td>
<td>--</td>
<td>--</td>
<td></td>
<td>recov.bin</td>
</tr>
<tr>
<td>-p</td>
<td>Initial Profile</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>O</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td></td>
<td>cres.pro</td>
</tr>
<tr>
<td>-l</td>
<td>Extra Output #1</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>O</td>
<td>A</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>--</td>
<td>A</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

8.2 SETTING UP YOUR OWN PROBLEM

To begin applying CHEMKIN Applications to your own problems, we recommend that you set up a working directory where you will store your input and output files for the application you wish to run. We also recommend that you add the CHEMKIN “bin” directory (e.g. c:\chemkin36\bin on a PC or ~/chemkin36/bin for UNIX) to your local “path” definition, so that you don't have to specify the full path of the executables every time you run them. The procedure for modifying path variables is described in Chapter 3, Sections 3.2.2 (for PCs) and Section 3.3.4 (for UNIX).

1 In the following list “xxx” is used in place of the name of the program executable (e.g. “chem” would have an input file named “chem.inp” and an output file named “chem.out”). The text “yyy” is either “tran” or “therm” for the Transport and Thermodynamics database files, respectively.
2 The Extra Output #1, 2, etc., allow growth for additional output files. At present, the –1 covers: “last.s” for AURORA, “ldas.bin” for CRESLAF, and “flux.out” for SPIN.
You can then follow a similar procedure as when running the sample problems in the step-by-step mode, but this time you will have to create your own input files. For this discussion we will use SENKIN as the target Application. The following are the steps for running SENKIN for your own problem.

1. **Create the input files** that you want to use to describe your reactor and process conditions. Use the text editor you are most comfortable with and save the files to your working directory. (See the Application and Utility user manuals for more guidance in creating the input data.)

2. **Open a MS-DOS Prompt or UNIX shell** and change directories to this working directory.

3. **Run the necessary pre-processors.** For the SENKIN Application, you will only need to run the CHEMKIN Gas-phase Interpreter. In the example below, we show how to run with your own thermodynamic database file located in your local directory. You can then run the CHEMKIN Interpreter by typing the following, assuming that you set your path as suggested above and have defined the chemistry in a file named `mychem.inp`:

   ```
   chem -i mychem.inp -o mychem.out -d mytherm.dat
   ```

   Make sure check the `mychem.out` file before proceeding. For other applications, you may also need to run the TRANSPORT pre-processor (which requires the `tran.dat` file to be copied or linked to the local directory) and/or the SURFACE CHEMKIN Pre-processor. To determine which pre-processors and input files are needed, see Table 3 in Chapter 5.

4. **Run the Application** with your input files. For example, to run SENKIN, you would type:

   ```
   senkin -i mysenkin.inp -o mysenkin.out
   ```

   Again, you would want to **check the mysenkin.out file to see if the program ran successfully**, before attempting to post-process the results.

5. **Run the SENKIN post-processor:**

   ```
   senkin_post < mysenkin_post.inp > mysenkin_post.out
   ```

The SENKIN post-processor (like all of the FORTRAN post-processors) produces a number of text files that contain comma-separated values according to the options requested in the `mysenkin_post.inp` file. These text files can easily be imported into the CHEMKIN Graphical Post-Processor, or into other analysis tools, such as spread-sheet programs.
9. Getting Help and Support

9.1 Online Documentation

CHEMKIN 3.6 provides a complete set of documentation in the form of HTML pages and PDF documents focused precisely on the details of building and running the CHEMKIN software. Included in the CHEMKIN installation are PDF documents for each of the user manuals for the CHEMKIN Utilities and Applications. You can view the PDF documents using Adobe Acrobat Reader. The HTML pages installed with CHEMKIN describe the installation directory structure and the sample problems available for each Application. You can view the HTML files using any internet browser (such as Internet Explorer or Netscape Navigator). The best method for getting started with the online documentation depends on your platform:

Windows/PC:

In this case, the install program will have placed short-cuts to the HTML pages and to the PDF documents in your desktop Start menu. Selecting a PDF document from the Start menu is illustrated in Figure 17 below.

Figure 17. Locating the CHEMKIN user manual PDFs and the HTML Documentation from the Windows Desktop Start menu on a PC.

UNIX:

All of the PDF documents are located in: $HOME/chemkin36/docs. The top-level of the HTML-based online documentation is: $HOME/chemkin36/index.html. Assuming that you have the executable for Adobe Acrobat Reader in your path, you may type:

    acroread getting_started.pdf <RETURN>
to open a PDF of this document in the Acrobat Reader. Alternatively, most web browsers include a plug-in for Acrobat Reader, so that you can open a PDF directly from the web browser. For example, if you have Netscape installed on your computer and in your path, then you could type:

```
netscape getting_started.pdf <RETURN>
```

to open the PDF from within Netscape. Similarly you can use the web browser from the command line to open the top-level "index.html" for browsing the HTML documentation.

9.2 FREQUENTLY ASKED QUESTIONS

Reaction Design provides a set of answers to Frequently Asked Questions (FAQs) that we have compiled through our interaction with customers. The up-to-date list of FAQs is maintained in the Customer Support section of our website (www.ReactionDesign.com).

9.3 CONTACTING TECHNICAL SUPPORT

Reaction Design is committed to the highest level of technical support. If the answer to your question is not found in the Frequently Asked Questions (FAQ) section, please email your question to Support@ReactionDesign.com. You may also contact Reaction Design's Technical Support, by Fax: (858) 550-1925, or by Phone: (858) 550-1920.

If you call or send email, you should include the following information in your support request:

1. Your CHEMKIN License Number.
2. The name and email address of the Licensee, if different from yourself.
3. Any input or output files for the problem you are running.
4. Any pertinent error or informational messages you have received from the CHEMKIN software.
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