# Phase 4.2 Command Line Tutorial: Creating and Searching 3D Databases

This tutorial demonstrates how 3D databases may be created, modified and searched by running command line tools that accompany Phase version 4.2, which is part of the Schrödinger Suite 2015 release.

Phase databases created in this tutorial are stored in the same format as those created using the Maestro graphical interface. Users may therefore employ the graphical interface to modify and search a Phase database created using the tools described in this tutorial.

#### 1.1 Conventions

Shell commands that the user should type will be preceded by the "shell>" prompt and they will appear in fixed font, e.g.,

```
Shell> cd db tutorial
```

Parameters for which the user must supply a value will appear in italics and in brackets, e.g.,

```
-i <inFile>
```

Important information about changes from previous versions of the software will be indented and italicized:

Users who are familiar with the Phase 3.8 command line database tools may recognize that a new option...

## 1.2 Getting Started

In order to run the programs described in this tutorial, Phase 4.2 (or later) must be installed on the machine you are currently using, as well as on any remote hosts on which jobs will be run. You must also define the SCHRODINGER environmental variable so that it points to the directory in which the Schrödinger software is installed.

A Phase 3D database must reside in a filesystem that is uniformly accessible to all machines on the network that will need to access the database, whether for purposes of creation, modification, or searching. Thus if a job is launched from *host1* and run on 10 processors that reside on *host2*, all 10 CPUs on *host2* must have access to the database via the same absolute path. Note that

<sup>&</sup>lt;sup>1</sup> Schrödinger software is frequently installed on a single filesystem that is accessible to multiple hosts, so strictly speaking, not every host must have its own separate installation.

*host1* is not required to have access to the database, although the present tutorial assumes that the database filesystem is mounted locally.

To create a database, you must have a collection of molecules stored in one or more Maestro or SD files, which may be compressed via gzip. Valid file extensions are .mae, .mae.gz, .mae.gz, .sdf, .sd, .sdf.gz, and .sd.gz. Each molecule may be represented by a single low-energy 3D conformer or by multiple 3D conformers. In the former case, you may import the single-conformer molecules into the database and create multi-conformer models later. If you are starting with 2D structures or SMILES strings, you must create low-energy 3D models first, preferably using LigPrep.

Each Maestro or SD file may contain multiple molecules, but if multiple conformers are present, they must be grouped in consecutive CT blocks. In the case of Maestro files, a multi-conformer molecule may be stored as a full CT block ( $f_m_ct$ ) followed by partial CT blocks ( $p_m_ct$ ), or it may be stored as a set of consecutive full CT blocks.

## 1.3 Database Programs

The following *top-level* programs should be present in the directory \$SCHRODINGER:

```
    phase_database
    Performs all tasks related to Phase database creation and management.
    Searches a Phase database, one or more structure files, or a Phase command line project for matches to a pharmacophore hypothesis.
```

These two programs comprise the entire command line interface for Phase database creation and searching.

# 1.4 Creating a New 3D Database

Place a copy of the file db\_tutorial.tar.gz in a directory located in a uniformly accessible filesystem (as discussed in Section 1.2), and extract the file as follows:

```
Shell> gunzip -c db tutorial.tar.gz | tar xf -
```

Move to the db tutorial/directory, which should be created automatically by the extraction:

```
Shell> cd db tutorial
```

For future convenience, we shall define an environment variable TPATH to hold the absolute path to the tutorial directory:

```
csh/tcsh:
Shell> setenv TPATH `pwd`
```

#### sh/bash/ksh:

```
Shell> export TPATH=`pwd`
```

Note that single back-quotes are used to surround the pwd command. Verify that TPATH has been assigned correctly:

```
Shell> echo $TPATH
```

This should print the absolute path to the current directory. Note that if you require more than one terminal session to complete the tutorial, or if you switch shells or windows, you will need to redefine TPATH.

Before running any command line program, it's a good idea to simply type the name of that program to see the usage message. You will be creating the database using phase\_database, so type the following command:

```
Shell> $SCHRODINGER/phase database
```

Observe that a <task> argument must be supplied to indicate what sort of operation is to be performed. To see a shortened help message with options for only a particular task, invoke phase\_database with the flag -help\_<task> (e.g., -help\_import, -help\_revise, etc.). To see a series of examples showing how to perform various tasks, use -help\_examples. Please examine the help messages in detail so that you are familiar with the wide variety of tasks and options supported by phase\_database.

Now create a new database named stdDB.phdb in the directory \$TPATH, and populate it with 100 single-conformer molecules contained in \$TPATH/userFiles/singleConfs.mae.gz:

By default, the assigned job name is database\_<task>, so it will be database\_import when an import is done. This convention can be overridden by supplying the optional third argument <jobName>.

The above command will run the job on the machine you are currently logged into, but you could direct it to be run on another machine by using the option <code>-HOST</code> <code><host></code>, where <code><host></code> is an entry name in the file <code>\$SCHRODINGER/schrodinger.hosts</code>. As discussed in Section 1.2, the database directory must be accessible to <code><host></code> via the absolute path that's specified in the <code>phase\_database</code> command. The database need not be accessible to the machine you are logged into, so if you specify <code>-HOST</code> <code><host></code>, you could launch the job from any machine on your network that can reach <code><host></code> through Schrödinger job control. You may alternatively run <code>phase\_database</code> locally as a foreground process, rather than as a Schrödinger job, by using the flag <code>-NOJOBID</code>.

Note that phase\_database supports only single-CPU import. If you are importing a large number of single-conformer molecules, you may wish to utilize multiple processors by running \$SCHRODINGER/phase multi database.<sup>2</sup>

The import will require only about one second to finish, after which you should see the following job files in the current directory:

File Name	Description
database_import.log	Log file with details of the import process.
database_import.okay	Created only if the job completes successfully.
database_import_errors.out	Error messages, if any errors occur. Empty otherwise.
database_import_new_phase.inp	Phase subset file with newly imported record numbers.

The tail end of the file database import.log should look like the following:

The reported CPU time of course depends on the hardware you are using.

Now verify that the appropriate files have been created in the database directory:

```
Shell> ls -F $TPATH/stdDB.phdb
```

You should see the following files:

<sup>&</sup>lt;sup>2</sup> phase\_multi\_database is for parallel import only and therefore does not support all the tasks/options in phase\_database. If the input files have to be split up by phase\_multi\_database, a non-trivial amount of upfront time is required to count and divide the structures over the appropriate number of files, and the overall speedup will be no more than a factor of approximately 10. Significantly greater speedup can be achieved if a set of pre-split input files are provided.

File Name	Description
database selite	SQLite database with tables that hold global information
database.sqlite	about the Phase database. See below for details.
	Version file. Holds the Phase version number, the method of
database_dbversion	creation (always "CL"), and the storage format ("SQLite"). Do
	not modify.
	Default pharmacophore feature definitions. You may replace
	this file with customized definitions before creating
database_feature.ini	pharmacophore sites (or you could have used the -fd
	<fdfile> option when phase_database was used to create</fdfile>
	the database).
database_history.log	Terse descriptions of all changes to the database.
database_info.log	Verbose descriptions of all changes to the database.
d-+-1 12d-/	Subdirectory under which all database structure files are
database_ligands/	stored.
database_master_phase.inp	List of all database records (do not modify).
database_summary.csv	Human-readable copy of the data in summary table (see
	below).

#### The file database.sqlite contains the following tables:

Table Name	Description
dprop_name_table	Holds the name of the property used to detect/eliminate duplicate
	structures. This is currently hard-coded to s_phase_Unique_SMILES.
dprop_values_table	Holds mol_id and s_phase_Unique_SMILES for each database
	record, where mol_id is the primary key.
	This table will exist only if an "extract" job has been run. If so, it will
nrong tablo	hold mol_id, title, num_confs, has_sites, plus all imported and
props_table	computed properties for each record that was in the database when the
	extract job was run.
	Mappings of the column names in props_table to the CT-level
nrong man tahla	property names in the block database files stored under
props_map_table	database_ligands/. This table always exists, but it will be empty if
	properties have not been extracted.
props_stale_table	Contains a 1/0 value indicating whether or not the database contains
	records that are not in props_table. A value of 1 indicates that an
	"extract" job needs to be run in order to bring props_table up-to-
	date.
summary_table	Holds mol_id, title, num_confs, and has_sites for each database
	record. has_sites will be 1 or 0, depending on whether or not
	pharmacophore sites are stored for a given record.

At this point, we have extracted 100 single-conformer molecules from the file singleConfs.mae.gz, and we have stored them in a single block SQLite file:

```
$TPATH/stdDB.phdb/database ligands/block 1/block struct 1.sqlite
```

Up to 5,000 molecule records will be stored in this file, with block\_2/block\_struct\_2.sqlite being created when molecule 5,001 is added. You can reduce the number of molecules per block during an import by using the option -blimit <maxRec>. This is most relevant when you create a small database and you wish to split conformer and/or site generation over multiple processors. Because it is not safe to allow multiple processors to write to the same block SQLite file, each processor must operate on one or more separate blocks. By reducing the block size, a larger number of blocks will be created, and more processors may be utilized for conformation/site generation.

We could now search this database if we were willing to generate conformers and pharmacophore sites in memory without storing them in the database, a process we refer to as *flexible* searching. In the present case, however, we will be storing conformers and sites in the database. Flexible databases are covered in Sections 1.11 and 1.12.

# 1.5 Adding Molecules to an Existing Database

In this section, a set of multi-conformer molecules is added to the database created in the previous section. If you are resuming the tutorial in a terminal session or window that is different from that used in Section 1.4, you will need to redefine the TPATH environment variable.

The file \$TPATH/userFiles/multiConfs.mae.gz contains a set of 36 endothelin ligands with pre-generated conformational models. In order to add these structures to the existing database using phase\_database, we omit the -new flag, and we must supply the flag -multi to indicate that the file being imported contains multiple conformers per molecule. We will also supply a unique <jobName> argument so that the job files from the previous import are not overwritten. Accordingly, add molecules to the database as follows:

```
Shell> $SCHRODINGER/phase_database $TPATH/stdDB.phdb import add_multi \
-i $TPATH/userFiles/multiConfs.mae.gz -multi -verbose
```

The job should finish in a just a few seconds. Once the job has completed, the tail end of the file add multi.log should contain output similar to the following:

```
.
LIGAND_NAME = block_1/mol_134  # Title = "endo-34"  Confs = 8
LIGAND_NAME = block_1/mol_135  # Title = "endo-35"  Confs = 3
LIGAND_NAME = block_1/mol_136  # Title = "endo-36"  Confs = 3
Committing 36 records
Updating top-level database files . . .

A total of 36 records added
Total number of records stored = 136
```

```
Removing restart directory

CPU time = 1.35 sec

phase_database successfully completed

Driver script for phase_database job add_multi finished Current time: Mon Dec 22 16:22:11 2014

Elapsed time = 00:00:02

phase database results for add multi are complete
```

You may also examine the file \$TPATH/stdDB.phdb/database\_info.log to see the full history of changes made to the database.

A new -annotations flag is now supported to speed up detection of stereochemistry during multi-conformer import from Maestro files. If this flag is used in conjunction with "-multi-stereo", stereochemistry will be gotten from mmstereo annotations in the file rather than by detecting from 3D structure. There is also a new -dropped flag, which may be used in conjunction with -unique to get a subset file with record numbers of database compounds that weren't found in the file(s) being imported. Use of "-unique -dropped" allows import of only the new compounds from, e.g., a vendor collection, while detecting compounds that have been dropped or discontinued.

# 1.6 Creating Conformers and Pharmacophore Sites

So far, we have created a database of 136 molecules, storing a single conformer for each of the first 100, and multiple conformers for the last 36. We now demonstrate how to create conformers for the first 100 molecules only, and pharmacophore sites for all 136. Conformers are generated using a rapid torsional sampling technique, followed by application of a soft non-bonded potential to eliminate high-energy structures.

We will be running a phase\_database "revise" job, so type the following command to view the relevant help message:

```
Shell> $SCHRODINGER/phase database -help revise
```

The options <code>-confs</code> <code>all|auto</code>, <code>-sites</code>, and <code>-isub</code> <code>/subset/</code> provide a high level of control and flexibility when creating conformers and pharmacophore sites. You may use <code>-confs</code> <code>all</code> to force conformer and site creation for all molecules, or, if you wish to generate conformers only for molecules that are represented by a single conformer, use <code>-confs</code> <code>auto</code>. Note that sites are automatically generated whenever <code>-confs</code> is supplied. If <code>-confs</code> is omitted, conformer generation will be skipped altogether, and <code>-sites</code> may be used to generate pharmacophore sites for the existing conformers. The <code>-sub</code> option allows you to change the domain of operation from the entire database to any subset of molecules in the database (see Section 1.17 for more on using subsets).

Ensure that the TPATH environment variable is defined as described in Section 1.4, and then launch the phase database "revise" job as follows:

Observe that we are using default settings for conformer generation, so a maximum of 100 conformers will be created and stored for each molecule which isn't already represented by multiple conformers. When the job has successfully finished, the file auto\_confs.okay will appear. The tail end of auto confs.log should look like the following:

```
. . . . All subjobs successfully completed. Updating top-level database files . . . Revision of 136 out of 136 records completed

Average number of conformers per revised record = 45.70

Total number of records stored = 136

Removing restart directory

CPU time = 0.10 sec

phase_database successfully completed

Driver script for phase_database job auto_confs finished

Current time: Mon Dec 22 16:39:15 2014

Elapsed time = 00:01:32

phase database results for auto confs are complete
```

One subtle point worth mentioning is that if you generate conformers more than once for the same database records, you will generally not get the same sets of conformers created each time. This is because you are not starting with the same seed structure. The first time around, the seed structure for a given molecule is whatever structure you imported. That structure is discarded by default when the new conformers are stored in the database, so if you generate conformers for the same molecule again, the seed structure will be the first conformer stored. While the conformers generated with different seeds are usually similar, they are not the same.

## 1.7 Deleting Molecules from a Database

There may come a time when a user needs to delete molecules from a database, and we demonstrate how to do this by running a phase\_database "delete" job and specifying a set of database records for deletion. To see the relevant help message, type the following:

```
Shell> phase database -help delete
```

Observe that in addition to deleting a specific set of records, phase\_database can identify records that contain duplicate structures and delete them, or simply produce a subset file with the duplicate record numbers.

Recall that the file \$TPATH/stdDB.phdb/database\_info.log contains a full accounting of all changes made to the database, including the unique identifier for each record added to the database. You may therefore use the contents of this file to construct a subset of database records for deletion. Suppose you wanted to delete the last 10 molecules from the database. To do so, simply create a subset file with the appropriate database records as follows:

We are including "\_phase.inp" at the end of the subset file name, because the Phase convention is to treat everything to the left of "\_phase.inp" as the subset name. The contents of last10 phase.inp should be as follows:

```
LIGAND_NAME = block_1/mol_127  # Title = "endo-27"  Confs = 6
LIGAND_NAME = block_1/mol_128  # Title = "endo-28"  Confs = 6
LIGAND_NAME = block_1/mol_129  # Title = "endo-29"  Confs = 18
LIGAND_NAME = block_1/mol_130  # Title = "endo-30"  Confs = 6
LIGAND_NAME = block_1/mol_131  # Title = "endo-31"  Confs = 11
LIGAND_NAME = block_1/mol_132  # Title = "endo-32"  Confs = 14
LIGAND_NAME = block_1/mol_133  # Title = "endo-33"  Confs = 4
LIGAND_NAME = block_1/mol_134  # Title = "endo-34"  Confs = 8
LIGAND_NAME = block_1/mol_135  # Title = "endo-35"  Confs = 3
LIGAND_NAME = block_1/mol_136  # Title = "endo-36"  Confs = 3
```

Everything to the right of a pound sign "#" is treated as a comment and thus ignored. Now delete these records from the database as follows:

```
Shell> $SCHRODINGER/phase database $TPATH/stdDB.phdb delete -isub last10
```

This operation should require very little time, and the tail end of the file database\_delete.log should look like:

```
c.
c.
Reading records from last10_phase.inp

Deleting 10 records . . .

Deleting structure records in block 1 . . .

Deleting summary records . . .

Updating database_master_phase.inp and database_summary.csv . . .

Total number of records stored = 126

CPU time = 0.45 sec

phase database successfully completed
```

```
Driver script for phase_database job database_delete finished Current time: Mon Dec 22 17:52:49 2014 Elapsed time = 00:00:01 phase database results for database delete are complete
```

The database now contains only 126 molecules.

## 1.8 Setting Database Permissions

When you create a database, you will naturally be the owner of all the files and directories associated with that database. As such, you will normally have read/write permissions to the files, and read/write/execute permissions to the directories. You should therefore encounter no difficulties in modifying a database that you create yourself. However, you may or may not want other users to be granted this same privilege.

If you want to be certain that you are the only person who will ever modify a particular database, then you should remove write permissions for all other users throughout the database tree. By default, the files and directories you create will normally not carry write permissions for other users (unless you change the default umask), but we demonstrate how to explicitly set those permissions here. First ensure that the TPATH environment variable is defined as described in Section 1.4, and then issue the following command:

```
Shell> chmod -R g-w,o-w $TPATH/stdDB.phdb
```

Write permissions for other users have now been removed recursively throughout the database, and you are the only person who can modify the database.

# 1.9 Database Searching: Background

In this section, material is presented that is critical to understanding how Phase 3D database searching works. Users are strongly encouraged to read this section so that they are familiar with terminology used in subsequent sections.

Before setting up and running a database search, you normally want to be in the directory where the pharmacophore hypothesis files reside. Ensure that the TPATH environment variable is defined as described in Section 1.4, then move to the directory containing the hypothesis files:

```
Shell> cd $TPATH/userFiles
```

This directory contains the following hypothesis-related files, which were created from a set of endothelin ligands:

File Name	Description
DHHRR_9.def	Pharmacophore feature definitions used to develop the hypothesis.
DHHRR_9.mae	Contains the reference ligand conformation, i.e., the full 3D chemical

	structure of the molecule that gave rise to the hypothesis.	
DHHRR_9.qsar	QSAR model.	
DHHRR_9.tab	General information about the hypothesis.	
DHHRR_9.xvol	Excluded volume definitions.	
DHHRR_9.xyz	Hypothesis site point coordinates.	

The QSAR model file and excluded volumes file will be present only if these characteristics have been defined for the hypothesis. All hypothesis files except <code>DHHRR\_9.qsar</code> contain ordinary text, so you may examine their contents, but do not modify them.

When a Phase 3D database is searched, the process is formally divided into two steps: finding matches and scoring hits.

When finding matches, the pharmacophore sites of a database molecule are searched for geometric arrangements of points that match the hypothesis with respect to feature types, intersite distances, and positional tolerances. For example, the DHHRR\_9 hypothesis contains one donor (D), two hydrophobes (H1, H2), and two aromatic rings (R1, R2). These five pharmacophore site points give rise to 10 unique intersite distances: d<sub>DH1</sub>, d<sub>DH2</sub>, d<sub>DR1</sub>, d<sub>DR2</sub>, d<sub>H1H2</sub>, d<sub>H1R1</sub>, d<sub>H1R2</sub>, d<sub>H2R1</sub>, d<sub>H2R2</sub>, and d<sub>R1R2</sub>. Accordingly, a database molecule would be scanned for occurrences of five sites of the type D, H, H, R, R, for which the 10 unique intersite distances are sufficiently close to those of the hypothesis. When such an occurrence is found, a least-squares alignment is performed to superimpose the matching site points in the database molecule onto the corresponding sites in the hypothesis, and positional tolerances are applied to determine whether pairs of aligned sites fall within a predefined radius of each other. If so, this *match* is stored in memory.

Once all matches have been recorded for a given molecule, the full 3D chemical structure of the conformer associated with each match is retrieved and aligned to the hypothesis according to the same transformation that was used to align the sites. We refer to these as *hits*, and there are various parameters and options that control which hits will survive, given a set of matches.

First and foremost, hits are sorted in order of decreasing fitness score, a quantity that measures how well the site points in the matching conformer align to those of the hypothesis, how well the matching vector features (acceptors, donors, aromatic rings) overlay those of the hypothesis, and how well the 3D chemical structure of the hit superimposes, in an overall sense, with the reference ligand conformation. Fitness is calculated as follows:

The various fitness score terms are defined as follows:

Term	Definition
alignScore	The RMS deviation between the positions of the site points in the matching
	conformation and the site points in the hypothesis.
alignCutoff	User-adjustable parameter. Default = 1.2.

alignWeight	User-adjustable parameter. Default = 1.0.
vectorScore	Average cosine between vector features in the matching conformation and
	vector features in the reference ligand conformation.
vectorWeight	User-adjustable parameter. Default = 1.0.
volumeScore	O(A,B)/[O(A,A) + O(B,B) - O(A,B)], where $O(A,B)$ is an approximate
	overlapping volume between matching conformation A and reference
	conformation B, and $O(A,A)$ and $O(B,B)$ are approximate self-overlaps. Each
	structure is represented as a collection of heavy atom van der Waals spheres.
volumeWeight	User-adjustable parameter. Default = 1.0.

If partial matching is invoked (see <minSites>, Sections 1.10 and 1.13) and a conformer fails to match all sites in the hypothesis, a penalty is applied to alignScore. If m out of n sites are matched, alignScore is modified as follows:

alignScore 
$$\rightarrow \operatorname{sqrt}(w_{old} \cdot \operatorname{alignScore}^2 + w_{new} \cdot \operatorname{alignPenalty}^2)$$

where

$$w_{\rm old}=m/n$$
 
$$w_{\rm new}=(n-m)/n$$
 alignPenalty = User-adjustable parameter. Default = 1.2.

By adjusting parameters in the fitness function, the user can control the order in which hits are returned. For example, to emphasize alignment of vector features, the <code>vectorWeight</code> parameter should be increased. If overall molecular superposition is most important, then <code>volumeWeight</code> should be increased. Note that the volume term provides a means of forcing the shape of the hit to resemble that of the reference conformation. Since the user can limit the number of hits from each molecule and the total number of hits, the fitness function ultimately controls which hits the user sees.

Application of excluded volumes also affects the survival of hits, and the user has the option of ignoring excluded volumes if they happen to exist for a given hypothesis. Although a QSAR model does not impact which hits are obtained, it can be applied or ignored.

# 1.10 Searching Existing Conformers and Sites

In this section we demonstrate how to run a normal database searching job against the database created in Sections 1.4-1.7, so you need to complete those sections if you have not done so already.

Databases are searched using phase\_find\_matches, so type the name of that program with no arguments to see the help message:

```
Shell> $SCHRODINGER/phase find matches
```

Most options have reasonable defaults, so you need not be concerned with everything you see in the help message, although it's still a good idea to familiarize yourself with the capabilities of phase find matches.

It is now possible to search multiple databases in a single job by supplying a list of databases in a text file that ends with ".list".

While residing in the directory \$TPATH/userFiles/, issue the following command to launch a basic search against existing conformers and sites:

```
Shell> $SCHRODINGER/phase_find_matches $TPATH/stdDB.phdb DHHRR_9 basic_search \
-NOCHECKPOINT
```

As discussed before, the database need not be accessible to the machine you are logged into, so if you specify <code>-HOST</code> <code><host></code>, you could launch the job from any machine on your network that can reach <code><host></code> through Schrödinger job control.

If you are searching a large database, it is strongly recommended that you use the option -CHECKPOINT cpath> when you launch the job, so that if the job fails to complete for any reason (a disk goes offline, a cluster node goes down, a power failure occurs, etc.), you will be able to restart it by using -RESTART <path>. Note that <path> is the absolute path to a directory under which checkpoint files will be created and updated throughout the course of the search. You must have write permissions to this directory and it must be accessible to the host on which the job is run, and to all compute nodes if running the job on a cluster.

You may examine the file basic\_search.log to monitor the progress of the job. When the job has completed, the file basic\_search.okay will be created and you should see the following sort of output in basic search.log:

```
Driver script for phase_find_matches job basic_search
Current time: Mon Dec 22 18:16:35 2014
Number of CPUs requested = 1
.
.
.
.
.
Combining subjob log files . . .

*** Contents of basic_search_sub_0.log ***
.
.
.
phase_find_matches: Mon Dec 22 18:16:36 2014

Source = <TPATH>/stdDB.phdb
Hypothesis = DHHRR_9
Job name = basic_search
Options:
-NOCHECKPOINT
-subjob basic_search_sub_0
-subproc
Assigning default positional tolerances of 2.00
```

```
Applying QSAR model in DHHRR 9.qsar
Applying excluded volumes in DHHRR 9.xvol
Searching structures in database "<TPATH>/stdDB.phdb"
Pre-screening 126 database records using 2D/3D index . . .
Number of records matched = 54
Pre-screen CPU time = 0.02 sec
ID, #Confs, #Matches, #Searched
mol 1,29,25,1
mol 11,70,0,2
mol_{13,40,0,3}
mol 124,14,12,52
mol 125,6,4,53
mol 126,18,16,54
Writing 29 hits to basic search sub 0-hits.list
************************
Finished processing the current batch of 54 records
Cumulative totals:
Number of molecules searched = 54
Number of conformers searched = 2235
Number of matches found = 350
Number of molecules that produced matches = 29
Number of hits written = 29
**********************
*** End of basic search sub 0.log ***
Combining sorted hits from completed subjobs
Importing hits from basic search sub 0-hits.list . . .
Number of hits stored so far = 29
Writing 29 sorted hits to basic search-hits.maegz . . .
Hits from completed subjobs successfully combined
All subjobs successfully completed
CPU time = 0.02 sec
phase find matches successfully completed
Driver script for phase find matches job basic search finished
Current time: Mon Dec 22 18:16:37 2014
Elapsed time = 00:00:02
```

```
phase find matches results for basic search are complete
```

Observe that the pre-screen using the 2D/3D index reduced the number of records to search from 126 to 54, and that 29 of those 54 records produced hits. So while the pre-screen is not 100% effective at identifying the minimum set of records to search, it does eliminate the majority of records that cannot possibly match the hypothesis (in the present case 72 out of 97 such records were eliminated). The pre-screen is also quite fast because it requires examination of only a small amount of data per molecule.

You may view the 29 hits by importing the file basic\_search-hits.maegz into Maestro. If you do so, you should see a number of properties that come from the database search:

Property	Description
Unique SMILES	Unique/canonical SMILES.
phasedb index	Database record number; same as mol_id.
Conf Index	Conformation index.
Num Sites Matched	Number of hypothesis sites matched.
	String that indicates which sites in the hit matched the hypothesis.
	The letter is the type of site that matched, and the number is the
Matched Ligand Sites	index of that site within the hit. If partial matching is invoked, a
	given site in the hypothesis may not be matched, in which case
	there would be a hyphen "-" rather than a number.
Align Score	Alignment score (see Section 1.9).
Vector Score	Vector score (see Section 1.9).
Volume Score	Volume score (see Section 1.9).
Fitness	Fitness score (see Section 1.9).
Pred Activity(1)	Predicted activity from 1-factor PLS QSAR model.
Pred Activity(2)	Predicted activity from 2-factor PLS QSAR model.
Pred Activity(3)	Predicted activity from 3-factor PLS QSAR model.

# 1.11 Creating a Flexible Database

If disk space is at a premium, a user may opt to create a database wherein only a single conformer is stored for each molecule. The necessary multi-conformer models and pharmacophore sites can be generated on-the-fly when the database is searched, without ever storing them to disk. In this section, we demonstrate how to create a database that will be searched in this manner, a so-called *flexible* database.

Ensure that the TPATH environment variable is defined described in section 1.4, and move to that directory:

Shell> cd \$TPATH

Create a new database named flexDB.phdb in the directory \$TPATH, and populate it with 100 single-conformer molecules contained in db tutorial/userFiles/singleConfs.mae.gz:

As in Section 1.4, this operation will require only a few seconds to finish. Once the import has completed, the tail end of the file create flex.log should look like:

```
LIGAND_NAME = block_1/mol_99  # Title = "862412_2" Confs = 1
LIGAND_NAME = block_1/mol_100  # Title = "862427" Confs = 1
Committing 100 records
Updating top-level database files . . .

A total of 100 records added
Total number of records stored = 100

Removing restart directory

CPU time = 1.54 sec

phase_database successfully completed

Driver script for phase_database job create_flex finished
Current time: Mon Dec 22 18:47:18 2014

Elapsed time = 00:00:03

phase database results for create flex are complete
```

Now verify that the appropriate files have been created in the database directory:

```
Shell> ls -F $TPATH/flexDB.phdb
```

You should see the following files:

```
database_dbversion database_info.log database.sqlite database_feature.ini database_ligands/ database_summary.csv database history.log database master phase.inp
```

See Section 1.4 for further details about each file. This 100-molecule database is now suitable for searching with on-the-fly generation of conformers and sites.

# 1.12 Running a Flexible Database Search

In this section we demonstrate how to setup and run a flexible database search, wherein conformers and sites are generated on-the-fly, but never written to disk. We will be using the database created in Section 1.11, so you need to complete that section if you have not already done so.

First move to the directory that holds the hypothesis files:

Then issue the following command to run the job:

The -flex flag triggers on-the-fly generation of conformers and sites. We are using default settings for conformer generation, just as we did when creating the multi-conformer database in Section 1.6.

When the job finishes, the log file flex search.log should contain the following sort of output:

```
Driver script for phase find matches job flex search
Current time: Mon Dec 22 18:50:55 2014
Number of CPUs requested = 1
Combining subjob log files . . .
*** Contents of flex search sub 0.log ***
phase find matches: Mon Dec 22 18:50:56 2014
Source = <TPATH>/flexDB.phdb
Hypothesis = DHHRR 9
Job name = flex search
Options:
-NOCHECKPOINT
-flex
-subjob flex search sub 0
-subproc
Assigning default positional tolerances of 2.00
Applying QSAR model in DHHRR 9.qsar
Applying excluded volumes in DHHRR 9.xvol
Searching structures in database "<TPATH>/flexDB.phdb"
Number of records to search = 100
ID, #Confs, #Matches, #Searched
mol 1,29,25,1
mol 3,80,0,3
mol 4, 4, 0, 4
mol_97,50,0,97
mol 98,100,0,98
mol 99,40,0,99
```

```
Writing 4 hits to flex search sub 0-hits.list
*******************
Finished processing the current batch of 100 records
Cumulative totals:
Number of molecules searched = 59
Number of conformers searched = 3763
Number of matches found = 120
Number of molecules that produced matches = 4
Number of hits written = 4
******************
*** End of flex search sub 0.log ***
Combining sorted hits from completed subjobs
Importing hits from flex search sub 0-hits.list . . .
Number of hits stored so far = 4
Writing 4 sorted hits to flex search-hits.maegz . . .
Hits from completed subjobs successfully combined
All subjobs successfully completed
CPU time = 0.00 \text{ sec}
phase find matches successfully completed
Driver script for phase find matches job flex search finished
Current time: Mon Dec 22 18:51:24 2014
Elapsed time = 00:00:29
phase find matches results for flex search are complete
```

Since the conformer generation options and the seed structures (see comments at the end of Section 1.6) in this job are identical to those used when <code>phase\_database</code> was run to create conformers for the first 100 molecules in the standard database, so we would expect the same results for the <code>flex\_search</code> and <code>basic\_search</code> jobs with respect to the first 100 molecules. Verify this by examining the files <code>flex\_search.log</code> and <code>basic\_search.log</code>. For example, you should observe identical numbers of matches and hits in these two files for <code>mol\_1</code>, <code>mol\_27</code>, <code>mol\_34</code>, and <code>mol\_43</code>, which are the four molecules that produced hits in the <code>flex\_search.job</code>.

## 1.13 Using Site Masks

When the phase\_find\_matches <minSites> parameter is smaller than the number of sites in the hypothesis, partial matching is invoked. So in the case of a 5-point hypothesis, one could set <minSites> equal to 4 to indicate that the search should consider 4-point matches if no 5-point

matches/hits are found. This does not mean that every single 4-point match will always be identified; rather, it means that if a given molecule fails to produce any 5-point matches or if all hits from 5-point matches get filtered out by excluded volumes or hard scoring cutoffs (see phase\_find\_matches -hard option), the search will move on to consider 4-point matches. However, if a molecule produces at least one 5-point match, and the associated hit survives all filters, then there will be no subsequent search for 4-point matches. Note that this is the default partial matching behavior. One can force phase\_find\_matches to exhaustively consider all 5-point and all 4-point matches by supplying the -ex flag.

Continuing with the 5-point hypothesis example, any 4 points of the correct type which also satisfy the geometric matching criteria would constitute a partial match. However, suppose you know that a ligand cannot bind to a particular receptor unless it contains a positive site and an aromatic ring. In that case, you would be interested in 4-point matches only if they contain those pharmacophoric elements.

To enable this sort of searching you must create a *site mask* file. Consider the hypothesis we've been using, for which the site types are D, H, H, R, and R. Suppose we wanted to require that every partial match contain the donor site and the first aromatic site. The corresponding site mask file should then contain the following 5 lines:

```
4 D 1
5 H 0
6 H 0
9 R 1
11 R 0
```

The first two columns are identical to those in <code>DHHRR\_9.xyz</code>, whereas the third column contains the mask value. A value of "1" indicates that the site must be matched, and "0" means that matching is optional. With this site mask and with <code><minSites>= 4</code>, matches involving all 5 sites would first be sought. If no 5-point matches/hits are found, the search would move onto 4-point matches, and the site mask would be applied.

In order to be recognized, the site mask file must be named <hypoID>.mask, where <hypoID> is the prefix used for all the other hypothesis-related files. Accordingly, while in the userFiles directory, use a text editor to create the file DHHRR\_9.mask, type the above 5 lines into the file, using spaces (not tabs) to separate values, and save your changes.

While still in the userFiles directory, launch the database searching job as follows:

```
Shell> $SCHRODINGER/phase_find_matches $TPATH/stdDB.phdb DHHRR_9 site_mask \
-match 4 -NOCHECKPOINT
```

When the job finishes, examine the file <code>site\_mask.log</code>. You should find that a total of 3308 matches were identified, and a total of 68 hits were written to <code>site\_mask-hits.maegz</code>. In the original <code>basic\_search</code> job, where partial matching wasn't used, there were only 350 total matches and 29 hits. So by requiring only 4 sites to match, many additional matches and hits were found.

To verify that every hit in site\_mask-hits.maegz matches both the donor site and the first aromatic site, issue the following command:

```
Shell> gunzip -c site mask-hits.maegz | grep "D("
```

This should produce the output:

```
"D(5) H(-) H(6) R(10) R(9)"
"D(4) H(9) H(10) R(13) R(14)"
"D(5) H(8) H(9) R(10) R(-)"

"D(2) H(6) H(-) R(9) R(10)"
"D(9) H(11) H(12) R(15) R(17)"
"D(7) H(-) H(11) R(15) R(16)"
```

As discussed in Section 1.10, these Matched Ligand Sites strings indicate how the hit matched the hypothesis. For example, "D(5) H(-) H(6) R(10) R(9)" indicates that the  $5^{th}$  site in the hit, a donor, was matched to the first site in the hypothesis, the  $6^{th}$  site in the hit, a hydrophobe, was matched to the  $3^{rd}$  site in the hypotheses, etc. When a site in the hypothesis is not matched by the hit, a hyphen appears, e.g., H(-). Note that you will see no hyphens for the donor site or the first aromatic site because the site mask required that both of these be matched.

# 1.14 Using Feature-Matching Rules

Suppose that you wish to allow a hydrophobic site in the hypothesis to match either a hydrophobic or aromatic site in the database you are searching. Suppose further that if the hydrophobic site is not matched to one of these two types of sites, you never want it to inadvertently match an ionizable site. These sorts of conditions may be imposed by defining *feature-matching rules*.

A feature-matching rule for a given site in the hypothesis consists of a string of *permitted features*, and an optional string of *prohibited features*. As the names suggest, permitted features define what a given site is allowed to match, while prohibited features indicate what that site is *not allowed* to match. A rule must be created for each site in the hypothesis, following the same order as in <hypoid>.xyz.

Feature-matching rules are best explained by way of example, so consider the following set of rules for the 5-point hypothesis we have been using:

```
4 D
5 HR NP
6 H
9 R
11 RH
```

The first rule "4 D" is just the default way of matching, i.e., donor site 4 is permitted to match only another donor site. However, the second rule "5 HR NP" indicates that the hydrophobic site 5 is permitted to match either a hydrophobic or an aromatic site, but if it does not match either of these, then it *cannot* match a negative ionizable or positive ionizable site.

Note that the first character in each permitted feature string must be the actual site type that appears in <code>DHHRR\_9.xyz</code>, (so, e.g., the last rule is "11 RH" rather than "11 HR") and none of the permitted features may appear in the prohibited feature string. Also note that prohibited features are only relevant in the case of a partial match.

When a given site in the hypothesis is allowed to match more than one type of site in the database (e.g., the  $2^{nd}$  and  $5^{th}$  rules above), this condition is described as using *mixed permitted* features. An important consequence of doing this is that all vector scoring will be turned off, so that matches between vector and non-vector features (e.g.,  $H \leftrightarrow R$ ) aren't automatically scored lower than matches between two vector features.

Feature-matching rules must be stored in a file named <hypoID>.rules, so while residing in the userFiles directory, use a text editor to create DHHRR\_9.rules, type the above rules into that file, using spaces to separate values, and save the changes.

Launch the database searching job as follows:

```
Shell> $SCHRODINGER/phase_find_matches $TPATH/stdDB.phdb DHHRR_9 feature_rules \
-match 4 -NOCHECKPOINT
```

Be aware that the site mask created in the previous section is also being applied, so the donor site and the first aromatic site must be matched. There are no logical inconsistencies in combining this site mask and the current feature rules. However, it would be inconsistent if the site mask required the first hydrophobic site to be matched, because we have defined prohibited features for that site, which are applied only if the site is not matched to a permitted feature.

When the job finishes, examine the file feature\_rules.log. You will see that, compared to the previous site\_mask search, the total number of matches has increased from 3308 to 5959, and the number of hits has increased from 68 to 84. This is because the use of mixed permitted features leads to a greater number of ways to match the hypothesis to a given molecule.

To examine the various mappings in the hit file, issue the following command:

```
Shell> gunzip -c feature rules-hits.maegz | grep "D("
```

This should produce the output:

```
"D(5) H(-) H(6) R(10) R(9)"
"D(6) H(8) H(9) R(10) R(12)"
"D(5) H(6) H(7) R(10) R(12)"
```

```
"D(3) R(8) H(6) R(7) R(9)"
"D(8) R(15) H(-) R(12) R(14)"
"D(8) H(15) H(16) R(18) H(13)"
```

Observe that among the 84 hits, there are many instances of the first hydrophobic site in the hypothesis being matched to an aromatic ring, e.g., "D(6) R(15) H(-) R(13) R(14) ", and the second aromatic ring in the hypothesis being matched to a hydrophobic site, e.g., "D(3) R(11) H(-) R(12) H(9) ".

Verifying that the prohibited rules have been enforced is not so straightforward, but it's possible to do so by visual inspection of the hits that fail to match the first hydrophobic site.

In order to avoid inadvertent application of feature-matching rules in subsequent sections, remove DHHRR 9.rules before proceeding:

```
Shell> rm DHHRR 9.rules
```

## 1.15 Using Feature-Specific Positional Tolerances

As noted in Section 1.9, database matches are found by comparing intersite distances from the hypothesis with those of the database conformers. A single tolerance <code><deltaDist></code> is applied to all intersite distances, and a single positional tolerance is applied to all matching sites after alignment, which means that different types of pharmacophore features are treated equivalently. Because certain types of ligand-receptor interactions are stronger and more specific, it often makes sense to define different positional tolerances for matching different types of features. To enable this, you must create a file containing *feature-specific positional tolerances*.

Note that <code><deltaDist></code> is still applied to identify all the initial matches, so the value of this parameter may be adjusted independently of the positional tolerances. While the mathematical relationship between intersite distance matches and positional matches is quite complex for hypotheses with several sites, choosing a value of <code><deltaDist></code> that is equal to the sum of the two largest positional tolerances will ensure that <code><deltaDist></code> is no more stringent than the positional tolerances. Be aware that if you supply a set of positional tolerances and you do not specify a value for <code><deltaDist></code>, this rule is applied to assign a default value. Because this often leads to a fairly large <code><deltaDist></code> (e.g., 3 or 4 Å), the number of intersite distance matches may be excessive, and the computational cost of searching may increase significantly. In most cases, using such a large <code><deltaDist></code> is not worth the extra cost because the vast majority of the additional matches will provide poor overall fits to the hypothesis.

To be recognized, the feature-specific tolerances file must be named <hypoid>.tol. While in the userFiles directory, use a text editor to create the file DHHRR\_9.tol and type in the following 9 lines of text, using spaces to separate the letters from the numbers:

<sup>&</sup>lt;sup>3</sup> If the hypothesis contains two or more sites of a given type, and the positional tolerance assigned to that type is the largest, one would simply double that tolerance to obtain a <deltaDist> that is no more stringent than the positional tolerances.

```
A 1.50
D 1.50
H 2.00
N 0.75
P 0.75
R 2.00
X 1.50
Y 1.50
Z 1.50
```

Note that we are defining a tolerance in Å for every possible feature type (including custom features X, Y, Z). Strictly speaking, we really need only define tolerances for D, H and R in order to search with the current hypothesis, but you should get into the habit of considering all possible features because you may want to use a single set of tolerances for all of your database searching. Be aware that if you accidentally omit a feature type that's contained in your hypothesis, a default tolerance of 1.0 Å will be used.

It should be apparent that this mechanism does not allow you to assign different tolerances to different instances of the same feature type. So with our current hypothesis, the same tolerance would be applied to both hydrophobic sites, and the same tolerance would be applied to both aromatic sites. Section 1.16 describes how to define site-specific tolerances that allow you to overcome this restriction.

Save the file DHHRR\_9.tol and launch the job as follows:

```
Shell> $SCHRODINGER/phase_find_matches $TPATH/stdDB.phdb DHHRR_9 feature_tol \
-d 2.0 -NOCHECKPOINT
```

We have not used the option -match <minSites>, so all 5 sites are required to match and the existing site mask will have no effect. We are overriding the default <deltaDist> value, which would have been 4.0 since the two largest positional tolerances are 2.0.

When the job finishes, examine the file feature\_tol.log. You will see that a total of 228 matches were found, and a total of 29 hits were written to feature\_tol-hits.maegz. This is a reduction from the 350 matches found when positional tolerances were not used. The number of hits has remained the same, however.

In order to avoid inadvertent application of these positional tolerances in subsequent sections, remove <code>DHHRR 9.tol</code> before proceeding:

```
Shell> rm DHHRR 9.tol
```

## 1.16 Using Site-Specific Positional Tolerances

When using feature-specific tolerances, as described in Section 1.15, it is not possible to distinguish between different instances of the same feature type. So if your hypothesis contains

two hydrophobic sites, both would be matched with the same tolerance. However, there may be cases where you will want to use different tolerances, and we describe how to do that here.

Site-specific tolerances must be stored in a file named <hypoID>.dxyz, so use a text editor to create the file DHHRR\_9.dxyz and type in the following 5 lines of text, using spaces to separate the values on each line:

```
4 D 1.50
5 H 2.00
6 H 1.00
9 R 2.00
11 R 1.00
```

Observe that we are placing tighter tolerances on matching the second hydrophobic site and the second aromatic site. After saving the file, launch the job as follows:

```
Shell> $SCHRODINGER/phase_find_matches $TPATH/stdDB.phdb DHHRR_9 site_tol \ -d 2.0 -NOCHECKPOINT
```

When the job finishes, examine the file site\_tol.log. You will see that a total of 126 matches were found and a total of 24 hits were written to site\_tol-hits.maegz. This is a reduction from the 228 matches and 29 hits found when positional tolerances of 2.0 were applied to both hydrophobic sites and both aromatic sites.

In order to avoid inadvertent application of site-specific matching tolerances in subsequent sections, remove <code>DHHRR\_9.dxyz</code> file before proceeding:

```
Shell> rm DHHRR 9.dxyz
```

## 1.17 Working with Database Subsets

There may be instances where a user wishes to perform operations on only a subset of a Phase database. For example, suppose you wanted to restrict a database search to the set of molecules that produced hits in a previous search. Or perhaps you have added new molecules to a database that already contains conformers and sites, and you wish to generate conformers and sites for only the new molecules. Or maybe you want to create a database subset by performing a logical operation (AND, OR, NOT) on two existing subsets. In this section, we demonstrate how to accomplish these sorts of tasks using phase\_database.

To see the available options, type the following:

```
Shell> $SCHRODINGER/phase_database -help_subset
```

Observe that you may create subsets from a hit file, from conformer and site queries, from a file containing a list of titles, and from a logical combination of existing subsets.

We shall first demonstrate how to create a subset from a hit file and then show how to restrict a database search to that subset. Ensure that you are in the userFiles directory and that the file basic\_search-hits.maegz is present. With the TPATH environment variable defined as described in Section 1.4, issue the following command to create a subset from the aforementioned hit file:

We are using the <code>-NOJOBID</code> flag so that the task runs as a foreground process rather than as a Schrödinger job. When the prompt returns, the current directory should contain the subset file <code>myHits\_phase.inp</code>, which holds <code>LIGAND\_NAME</code> records for the 29 molecules in <code>basic\_search-hits.maegz</code>:

Note that Phase subset files always have a name of the form <subset>\_phase.inp, and that they normally contain a LIGAND\_DIR record in addition to the LIGAND\_NAME records, although the LIGAND\_DIR record is optional.

Now suppose you wanted to repeat the search that identified those 29 hits, but you wanted to use a tighter intersite distance matching tolerance. In that case, there is no need to search the entire database, because molecules that did not match the hypothesis in the initial search cannot possibly match it if the tolerance is reduced. Therefore, you can restrict the search to the 29 molecules in the subset you just created. Accordingly, launch the following database searching job:

```
Shell> $SCHRODINGER/phase_find_matches $TPATH/stdDB.phdb DHHRR_9 tight_match \
-isub myHits -d 1.3 -NOCHECKPOINT
```

When the job finishes, examine the file tight\_match.log. You should find that only 17 of the 29 molecules satisfied the stricter distance matching tolerance.

Suppose you wanted to determine which of the 29 molecules *failed* to satisfy the new distance-matching tolerance. First, create a subset from the 17-molecule hit file as follows:

```
Shell> $SCHRODINGER/phase_database $TPATH/stdDB.phdb subset \
-hits tight match-hits.maegz -osub tightHits -NOJOBID
```

Now use the logical NOT operator to subtract the 17-member subset from the original 29-member subset:

```
Shell> $SCHRODINGER/phase_database $TPATH/stdDB.phdb subset \
-logic NOT -isub1 myHits -isub2 tightHits -osub missedHits -NOJOBID
```

The file missedHits\_phase.inp should now contain LIGAND\_NAME records for only the molecules that failed to satisfy the 1.3 Å tolerance, i.e.,

```
LIGAND_DIR = database_ligands
LIGAND_NAME = block_1/mol_1
LIGAND_NAME = block_1/mol_27
LIGAND_NAME = block_1/mol_34
LIGAND_NAME = block_1/mol_43
LIGAND_NAME = block_1/mol_118
LIGAND_NAME = block_1/mol_119
LIGAND_NAME = block_1/mol_120
LIGAND_NAME = block_1/mol_121
LIGAND_NAME = block_1/mol_121
LIGAND_NAME = block_1/mol_123
LIGAND_NAME = block_1/mol_124
LIGAND_NAME = block_1/mol_125
LIGAND_NAME = block_1/mol_125
LIGAND_NAME = block_1/mol_126
```

As a final example, we illustrate how subsets are used to restrict conformer and site creation to a portion of a database. Issue the following command to add the single-conformer molecules in \$TPATH/userFiles/newMol.mae.gz to the database you created in Sections 1.4-1.7:

This operation will require about one second to finish. The tail end of the file database import.log should look like:

At this point, the database contains 126 molecules with multiple conformers and pharmacophore sites, and 25 molecules with single conformers and no sites. To bring this database up-to-date, we need to generate conformers and sites for only the last 25 molecules. Observe that when you import structures, phase\_database always creates a subset file, <jobName>\_new\_phase.inp, with the newly created record numbers. We used the default job name for the last import, so the records we need are contained in the file database\_import\_new\_phase.inp.

To create sites and conformers for only the 25 newly added molecules, launch a phase database revise job as follows:

```
Shell> $SCHRODINGER/phase_database $TPATH/stdDB.phdb revise new_mol \
-confs all -isub database import new
```

When the job finishes, you should see the following sort of output at the end of the file new mol.log:

All 151 molecules in the database now have multiple conformers and pharmacophore sites.

## 1.18 Working with Database Properties

The Maestro and SD files that you import to create a Phase database usually contain any number of properties for each molecule, and these properties are stored within the block database files under the database\_ligands/ directory. You may also generate Canvas properties when you run a revise job with the -props flag, and these are also stored in the block database files. The phase\_database "extract" task may be used to extract all such properties to a separate SQLite database that can be searched using SQL queries.

We will be extracting properties from the database \$TPATH/stdDB.phdb, so you need to complete Sections 1.4-1.7 and Section 1.17 in order to obtain the results presented in this section.

To see the available options for the extract task, type the following:

```
Shell> $SCHRODINGER/phase database -help extract
```

Although we will not be using the -append flag in this tutorial, it's important to be aware of it. If you create a large database, extract properties, then import a few new structures to the database, you can extract properties for just the newly added records by using -append. This is much faster than extracting properties for the entire database. Note, however, that once the extraction has been done for the new records, use of the -append flag in a subsequent extraction will not re-extract properties for the newly added records. For this reason, if you intent to run a revise job to create conformers and sites for the newly added records, you should extract properties after the revise job. This will guarantee that any changes which take place as a result of the revise job (e.g., num\_confs, has\_sites), will be reflected in the extracted properties. If you inadvertently do the extraction before running the revise job, you will need to re-extract properties for the entire database in order to get up-to-date properties.

While in the userFiles/ directory, issue the following command to launch the property extraction job:

```
Shell> $SCHRODINGER/phase database $TPATH/stdDB.phdb extract -map -stats
```

The job should require only a second or two to finish, after which you will find the following new files in the directory \$TPATH/stdDB.phdb:

File Name	Description
database.sqlite	SQLite database with all extracted properties.
database_props.csv	Comma-separated file with a copy of all the properties in
	database.sqlite.
database_props_map.csv	Contains mappings between the SQLite property names and the
	source property names in the block database files. Created only
	if the -map flag is used.
database_props_stats.csv	Property statistics: count, min, max, avg, where count is the
	number of records for which the property exists. Created only if
	the -stats flag is used.

The file database\_props\_map.csv is particularly useful because it provides a vertical list of all the properties that can be used in SQL queries. In cases where SQLiteProp and SourceProp differ, the former should be used when performing a query. Be aware that for imported properties with a conformational dependence (e.g., r\_mmod\_Potential\_Energy-MMFF94s), only the value for the very first conformer is stored in the block database files and in the SQLite property database.

Now that <code>\$TPATH/stdDB.phdb/database.sqlite</code> has been created, we can pose a SQL query to find records that satisfy some property filter. This is done by running a <code>phase\_database</code> "query" job. To see the available options, type the following

```
Shell> $SCHRODINGER/phase database -help query
```

Observe that the query may involve records that satisfy a particular condition, records that contain specific values of a given property, or records that match a SMARTS string. In the case of SMARTS, it's not actually necessary to extract properties before performing the query.

Suppose you wanted to identify molecules with a value of r\_user\_Activity greater than 7.5. To accomplish this, issue the following command:

```
Shell> $SCHRODINGER/phase_database $TPATH/stdDB.phdb query active \
    -where "r user Activity > 7.5"
```

The query must be enclosed in single or double quotes so that it is interpreted as a single argument by the shell. When the job finishes, the files active\_matches.csv and active\_matches.phase.inp will be created in the current directory. Use a text editor or spreadsheet program to examine active\_matches.csv and verify that there are 13 matching records, and r\_user\_Activity is greater than 7.5 in each case. Verify that the subset file active\_matches\_phase.inp contains the same 13 records.

The following are examples of other valid -where queries:

```
-where "num_confs = 1"
-where "title <> 'endo-1'"
-where "r_user_Activity > 7.5 OR num_confs > 10"
-where "r_user_Activity BETWEEN 7.0 AND 8.0"
-where "title LIKE 'serot%'"
```

Note that in some cases single quotes must be used within the double quotes to ensure that the query is correctly interpreted by the SQLite engine.

## 1.19 Converting Databases

Databases created with previous versions of Phase using <code>phasedb\_manage/phasedb\_confsites</code> (or using the Phase GUI in 2011 or earlier) are not compatible with the tools described in this tutorial, so you will need to convert those databases to the new format in order to search them or to make modifications to them.

A database may be converted to the new format, or merged into an existing database, by running a phase database "convert" job. To see the available options, type the following:

```
Shell> $SCHRODINGER/phase database -help convert
```

The destination database is the usual cpath/name> argument we have been specifying with
every phase\_database command, whereas the source database is the one that is being
converted/merged, and it may be stored in the old or new format.

Ensure that the TPATH environment variable is defined as described in Section 1.4, and move to that directory:

```
Shell> cd $TPATH
```

The tutorial comes with a compressed tar archive that contains a small database created using Phase 3.3, which was the last release before the official transition to the current database format. Extract that archive as follows:

```
Shell> gunzip -c userFiles/Phase 3.3 DB.tar.gz | tar xf -
```

Now list the contents of the extracted database directory:

```
Shell> 1s -F Phase 3.3 DB
```

You should see the following files:

```
oldDB_dbInfo.log oldDB_feature.ini oldDB_master_phase.inp
oldDB dbversion oldDB ligands/ oldDB phasedb
```

Submit the conversion job as follows:

old to new.log:

```
Shell> $SCHRODINGER/phase_database $TPATH/Phase_4.2_DB.phdb convert old_to_new \
-source $TPATH/Phase 3.3 DB/oldDB -new
```

As with other phase\_database jobs, it is possible to launch a conversion job from a local host that does not have physical access to the source or destination databases, but the remote host must have access to both databases via the paths supplied in the launch command.

Since only 36 records are being converted, this job should finish in less than a minute. When the conversion is complete, you should see the following sort of output at the end of the file

```
phase_database results for old_to_new_sub_0 are complete

*** End of old_to_new_sub_0.log ***

All subjobs successfully completed. Updating top-level database files . . .

Conversion of 36 out of 36 records completed

Average number of conformers per converted record = 28.81

Total number of records stored = 36

Removing restart directory

CPU time = 0.35 sec

phase_database successfully completed

Driver script for phase_database job old_to_new finished

Current time: Mon Dec 22 20:00:45 2014

Elapsed time = 00:00:11

phase database results for old to new are complete
```

Now list the contents of the destination database directory:

```
Shell> ls -F Phase_4.2_DB.phdb
```

You should see the following files:

Verify that there are 36 LIGAND\_NAME records in the file database\_master\_phase.inp and that the directory database\_ligands/block1 contains the files block\_struct\_1.sqlite.

To compare the sizes of the source and destination databases, issue the following commands:

```
Shell> du -k Phase_3.3_DB
Shell> du -k Phase 4.2 DB.phdb
```

You should see that the source database consumes 1136 blocks (1 kB per block), whereas the destination database consumes only 408 blocks, which is a reduction of 64%. In most cases, databases stored in the new format are only 30-45% as large as those stored in the old format.

At this point you may safely delete the source database as follows:

```
Shell> rm -rf Phase 3.3 DB
```

# 1.20 Exporting Structures from a Database

<sup>&</sup>lt;sup>4</sup> The database sizes you observe may differ slightly from what is reported here.

At some point you may wish to export structures from a Phase database and write those structures to Maestro or SD files. In this section we demonstrate how this may be done by running a phase database "export" job.

We will be exporting structures from the database \$TPATH/stdDB.phdb, so you need to complete Sections 1.4-1.7 and Section 1.17 in order to obtain the results presented in this section.

To view the available options type the following:

```
Shell> $SCHRODINGER/phase database help export
```

Observe that if running under Schrödinger job control, the *<base>* parameter must contain an absolute path that exists on the host where the job will run.

For large databases, the exported structures may require a great deal of disk space, so phase\_database has an option to impose a quota on the total disk space consumed by the files it creates. The program will abort as soon as that quota is reached. So if there are, say, 10 Gbytes of available disk space on the filesystem that you will be exporting to, it's a good idea to set the quota to something smaller than 10 Gbytes, particularly if the filesystem is shared with other users.

Ensure that the TPATH environment variable is defined as described in Section 1.4, and move to the directory \$TPATH/userFiles:

```
Shell> cd $TPATH/userFiles
```

Issue the following command to export the first conformer only for each molecule in \$TPATH/stdDB.phdb, and write those structures to a series of SD files, with a maximum of 100 structures per file:

The export should only take a second or two, after which you should find the files stdDB\_export\_1.sdfgz and stdDB\_export\_2.sdfgz in the current directory, with 100 and 51 structures, respectively.

# 1.21 Running on Multiple Processors

phase\_database "revise" and "convert" tasks and phase\_find\_matches may be run as multiple-CPU jobs on an appropriately configured cluster, a multiprocessor workstation, or across a series of networked hosts. Because a Phase database is held in a fixed, central location, (i.e., not copied between hosts at runtime), these jobs are not intended to be run in a grid computing environment.

It is beyond the scope of this tutorial to provide step-by-step instructions for setting up an environment to allow multiple-CPU jobs, so it is assumed that the user is already able to run other Schrödinger software on multiple CPUs. For further details, please consult the Product Installation Guide.

Aside from configuration issues, the first requirement is that the database be located in a directory that is uniformly accessible to all machines that will need access to the database. This means the absolute path to the database must be identical on all the applicable CPUs. Further, phase\_database jobs must operate on multiple blocks in order to use multiple CPUs. For example, if you wished to split a phase\_database revise job across 4 CPUs, you would have to be creating sites/conformers/properties for at least 4 database blocks. If you wished to split a phase\_database convert job across 4 CPUs, the converted records would have to be spread over at least 4 blocks in the destination database. This restriction exists because different subjobs cannot be allowed to perform write operations on the same block database at the same time. Doing so would result in a damaged block database. No such restriction exists for phase find matches jobs because they require read access only.

Assuming these requirements are met, a phase\_database or phase\_find\_matches job may be split across multiple processors by using the -HOST option to indicate how many CPUs should be used.

Suppose the file \$schrodinger.hosts contains a host entry named hostx with16 processors, and you wish to split a phase\_database revise job across 4 of those processors. Accordingly, you could issue the following sort of command to launch the job:

Similarly, if you wanted to run a phase\_database convert job or a phase\_find\_matches job on 4 processors, you could use commands like:

```
Shell> SCHRODINGER/phase_database <path/name> convert multi_cpu \
    -source <source> -new -HOST hostX:4

Shell> $SCHRODINGER/phase_find_matches <path/name> <hypoID> multi_cpu \
    -HOST hostX:4 -NOCHECKPOINT
```

If you do not currently have access to a machine with multiple processors, you can still observe how jobs are split up by mimicking a request for 4 processors on the *localhost*. For example:

This command will result in the launching of 4 subjobs that will run as concurrent processes on the machine you are logged into. The behavior will be much the same as if you were actually running on 4 independent CPUs, except that it won't be any faster than a regular single-processor job.

If you wish to split a job across more than one host on the network, then you must enclose the list of hosts in double quotes and separate them with spaces. For example:

```
Shell> $SCHRODINGER/phase_find_matches <path/name> <hypoID> multi_host \
    -HOST "hostX:4 hostY:2 hostZ:8" -NOCHECKPOINT
```

In this example, a total of 14 subjobs would be launched.