

Viewmol

Version 2.3

by

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1 Purpose

VIEWMOL is a graphical front end for computational chemistry programs. It is able to graphically aid in the generation of molecular structures for computations and to visualize their results. The program's capabilities include:

- Building and editing of molecules
- Visualization of the geometry of a molecule
- Tracing of a geometry optimization or a MD trajectory
- Animation of normal vibrations or to show them as arrows
- Drawing of IR, Raman, and inelastic neutron scattering spectra
- Drawing of an MO energy level or density of states diagram
- Drawing of basis functions, molecular orbitals, and electron densities
- Display of forces acting on each atom in a certain configuration
- Display of Miller planes in crystals
- Calculation of thermodynamic properties for molecules and reactions
- Drawings generated by VIEWMOL can be saved as TIFF, HPGL, or PostScript files
- Animations of normal modes can be converted to a video file (MPEG), e. g. for inclusion into World Wide Web documents (requires additional programs available on the Internet)
- Interface to the freeware ray tracing program RAYSHADE (input file generation and use of RAYSHADE from within VIEWMOL)
- Input and output in a variety of formats, new formats can be added easily by the user

VIEWMOL includes a Python interpreter for automation.

At present VIEWMOL includes input filters for DISCOVER, DMOL, GAMESS, GAUSSIAN 9X, GULP, MOPAC, and TURBOMOLE outputs as well as for PDB files (VIEWMOL is therefore suited as a viewer for structural data on the World Wide Web). Structures can be saved as MSI car-files, MDL files, and TURBOMOLE coordinate files. VIEWMOL's file format has been added to BABEL so that BABEL can serve as an input as well as an output filter for coordinates.

VIEWMOL supports a space ball as input device.

2 Copyright

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If new input filters to read outputs of other programs or other valuable features are added or a bug is fixed the author would welcome if the additions is sent to him for inclusion into the next release of VIEWMOL.

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VIEWMOL shall be quoted as follows:

Program VIEWMOL, Version 2.3, Jörg-Rüdiger Hill, 2000.

VIEWMOL has been thoroughly tested on the computer systems mentioned below, but it cannot be excluded that there are any further bugs.

Please, report bugs to joehill@users.sourceforge.net or use
<http://sourceforge.net/projects/viewmol/>

3 Installation

VIEWMOL 2.3 was developed on a Pentium notebook running Linux using Brian Paul's OpenGL compatible library Mesa. It also compiles with OpenGLTM without any problems. VIEWMOL has been ported to Silicon Graphics computers, IBM RS/6000, DEC Alpha, Suns, and Hewlett Packard 9000/735. Since Mesa runs on any machine which has X windows it should be possible to run VIEWMOL on any machine which supports X windows. However, for best performance a native OpenGL implementation is recommended. VIEWMOL does not run on Windows PC or MacOS Macs. For use on PCs or PowerMacs Linux is required and highly recommended.

The following operating systems are presently supported:

- PC: Linux 2.2.x (tested only with Mesa)
- SGI: IRIX 6.3
- DEC Alpha: OSF1 V4.0 (tested only with Mesa; courtesy of Pablo Vitoria Garcia, qibvgap@lg.ehu.es)

On these operating systems the program was tested. Others may also work, but this is not certified. Previous versions of VIEWMOL have also been tested on the following systems:

- IBM: AIX 4.1
- HP: HP-UX 9.5 (tested only with Mesa),
- Sun: SunOS (courtesy of Keith Refson, Keith.Refson@earth.ox.ac.uk)
- PC: FreeBSD (courtesy of Pedro A. M. Vazquez, vazquez@iqm.unicamp.br)

but since the author himself does not have access to any of these systems testing of newer versions was not possible. It can, however, be assumed that VIEWMOL will at least compile on these systems.

VIEWMOL 2.3 is provided precompiled for a number of architectures. Precompiled binaries are packaged separately from source, documentation, and examples:

- x86 Linux, dynamically linked with Motif in `viewmol-2.3.bin.Linux.tgz` (requires glibc2 distribution like RedHat >= 5.0 or Debian >= 2.0)
- SGI IRIX 6.3 in `viewmol-2.3.bin.IRIX.tgz`
- IBM RS6000 AIX 4.1 in `viewmol-2.3.bin.AIX.tgz`
- DEC Alpha OSF1 in `viewmol-2.3.bin.OSF1.tgz`

VIEWMOL can be downloaded from:

- SourceForge
<http://viewmol.sourceforge.net/>
- Computational Chemistry List archive (USA)
<http://www.ccl.net/cca/software/SOURCES/C/viewmol/index.shtml>

The source code and the precompiled binaries for Linux will also be available from Metalab (formerly Sunsite, USA) or your friendly neighborhood mirror

`file://metalab.unc.edu/pub/Linux/apps/graphics/visualization`

Before installing VIEWMOL you need to make sure that the following libraries are available:

- OpenGL or Mesa (libGL.so, libGLU.so or libMesaGL.so, libMesaGLU.so)
This is probably only relevant for installations on Linux. Most Linux distributions now include Mesa.
- Motif or Lesstif (libXm.so)
This is probably only relevant for installations on Linux. Motif can be found at <http://www.motifzone.org/download/>. Lesstif can be found at <http://www.lesstif.org/>.
- Python >=1.5.2
Linux distributions usually include Python. For other system you need to obtain Python from <http://www.python.org/>

- TIFF library (libtiff.so)

Linux distributions include this library. On Silicon Graphics this library is installed if the `ifl_eoe` and `ifl_dev` packages have been installed. For other systems you need to obtain and compile this library (e. g. from sgi.com under `graphics/tiff/tiff-v3.4-tar.gz`)

Installation of the program is simple. `VIEWMOL` comes as gzipped tar file, `viewmol-2.3.src.tgz`. Unzip and untar it using `gunzip viewmol-2.3.src.tgz` and `tar -xvf viewmol-2.3.src.tar`. You get three subdirectories, `source`, `man`, and `examples`, five resource files (English, German, Russian, French, Spanish), `Xdefaults.*`, and the configuration file `viewmolrc`. Copy all files you got into an arbitrary directory. If you want to install precompiled binaries, download the appropriate file for your operating system and unpack it from the same directory you unpacked the source code. This will create a subdirectory in the source directory which holds the binaries (the name of this directory starts with the name of your operating system as you get it from `uname -s` and may contain a CPU specific ending). If you run the supported operating systems you have to set the environment variable `$VIEWMOLPATH` to point to the directory where you unpacked `VIEWMOL` (the compiled-in default for `VIEWMOLPATH` is `/usr/local/lib/viewmol`) and the installation is complete. Otherwise you have to recompile the program (cf. p. 5). The program uses dynamical memory allocation so that every size of a molecule can be handled which fits the hardware limits of your workstation.

The installation directory also contains a file `viewmolrc`. You might have a look into this file and adapt it to your needs. The format is described at page 41. In general the defaults should work fine.

`VIEWMOL` uses by default English as language, but it has been written so that other languages can easily be used¹. The distribution contains files `Xdefaults.<language>` which contain all the program messages, menus, dialog boxes etc. in other languages (currently English, German, Russian, French, and Spanish). If you want to use a different language for a system wide installation, copy the corresponding `Xdefaults.<language>` file to your applications default directory (usually `/usr/lib/X11/app-defaults`) and rename it to `Viewmol`. If you want to use a different language only for some users, instruct them to configure the language through `VIEWMOL`'s Configuration menu. `VIEWMOL` will run without any of the `Xdefaults.<language>` files installed. So if you are happy with English and want to change only a few settings you don't need to install any of the `Xdefaults.<language>` files.

`VIEWMOL` needs a few external programs for some of its functions. Once you have installed `VIEWMOL` and set `VIEWMOLPATH`, you can start `VIEWMOL`, press `Cancel` in the file selection box which will appear, and press the right mouse button in the blue `VIEWMOL` window. A popup menu will appear where the last but one option is `Configuration . . .`. Choosing this option displays a dialog where you can set path names to four external programs. These are (including their defaults)²:

Location of Web browser:	<code>netscape %s</code>
Location of Moloch:	<code>moloch</code>
Location of Rayshade:	<code>rayshade</code>
Location of display program for RLE files:	<code>xv %s</code>

If these program are installed and can be found in your path `VIEWMOL` will automatically display the correct path names in the dialog. The `%s` is a placeholder for the file name and is required for programs which use

¹The manual assumes that the English version of `VIEWMOL` is used. The shortcuts for other languages are different, but obvious in the menus.

²Moloch may be called TurboPROP if you got TURBOMOLE from Molecular Simulations Inc.

command line arguments. Once you have set these path names, choose Save from the buttons in the dialog and these settings will be stored permanently in `$HOME/.Xdefaults`.

4 Compilation

VIEWMOL 2.3 has been written in C. For compilation of VIEWMOL you need a C compiler. TIFF files are supported by the freely available TIFF library which is also necessary to compile the program. It can be found on many ftp sites, e. g. at sgi.com under `graphics/tiff/tiff-v3.4-tar.gz`. If you want to link VIEWMOL with Mesa instead of with OpenGL you will need Mesa (<http://www.mesa3d.org/>).

Linux users need Motif to compile and run the program (if the program complains about "viewmol: can't load library 'libXm.so.1'" Motif is missing). Motif is available from <http://www.motifzone.org/download/>. The Motif clone Lesstif (<http://www.lesstif.org/>) can be used with Viewmol starting with version 0.81. There are, however, some glitches with Lesstif (e. g. shortcuts don't work).

If you want to recompile the program and you are running one of the supported operating systems (this may be necessary on IBM workstations since the formats of the executables are not compatible between different releases of AIX – don't worry, IBM didn't) you may type `make` (this tries to build VIEWMOL using OpenGL on all operating systems except on Linux and FreeBSD, to build using Mesa type `make viewmol_mesa; make tm bio readgauss readmopac readgames`). The shell script `getmachine` determines the operating system you are running and sets some options for the compiler. If this does not work you should have a look into the `Makefile`. The options set are explained there. They are the following:

- **OPT**
The optimization flag for your compiler (on Linux `-O6 -mX` where X may be 486, pentium, or pentiumpro; `-O2` otherwise).
- **CFLAGS**
Additional flags for C compiler, used to handle special optimizations and some incompatibilities between different Unix versions.
- **LDFLAGS**
Additional flags needed for linker, currently only used for SGI to distinguish between different library versions.
- **INCLUDE**
The path to the include files. For OpenGL compilation this is set to point to the include files for libtiff, for Mesa compilation the path to the Mesa include files is also included (the script asks the user for these paths at the beginning, there is really no need to change anything in the file here).
- **LIBRARY**
The path to the additional libraries required. These are at least libtiff and in case of a compilation with Mesa the Mesa libraries (the script asks the user for these paths at the beginning, there is really no need to change anything in the file here).
- **LIBS**
The libraries needed to link VIEWMOL. They may differ between different operating systems and are also different for OpenGL and Mesa compilations, respectively.

The `getmachine` shell script will ask you for the path names to the TIFF and Python libraries and to the include files necessary with these libraries. If you compile with Mesa the script will also ask you for the location of the Mesa libraries and include files. You may specify these path names using environment variables if you put the name of the variable in parentheses (e. g. `$(HOME)`). These path names are assigned to the `LIBTIFF`, `TIFFINCLUDE`, `MESALIB`, `MESAINCLUDE`, `PYTHONINCLUDE`, and `LIBPYTHON` flags and stored in a file `.config.<OS>` where `<OS>` is the output of the `uname -s` command on your machine. If this file already exists, `getmachine` does not ask for these path names.

Silicon Graphics compilers on 64-bit operating systems (IRIX64 – R8000, R10000, R12000) will produce a lot of warning messages concerning casts of pointers to integers. These can be safely ignored.

The make procedure will build the program in a directory whose name depends on the operating system and type of CPU you are using. You will find all executables in this directory. After compilation follow the steps under Installation to complete the installation.

5 Synopsis

`VIEWMOL` can be called as follows:

```
viewmol [[-bio | -dmol | -gamess | -gauss | -gulp | -mopac |  
          -pdb | -tm | -tmmsi] file]
```

`VIEWMOL` has an automatic file format detection algorithm build in and should be able to identify output files of the programs supported without user intervention. `VIEWMOL` will also run Python scripts given on the command line when the Python script has the string `python` within the first 1024 characters (this is usually the case if the script starts with the common `#!/usr/local/bin/python` or similar first line). If `VIEWMOL` is called without parameters it will bring up a file selection box to select the file to be viewed.

If the option `-bio` is used the file name of a DISCOVER file has to be specified. One can use the `.car`, the `.cor`, or the `.arc` file of DISCOVER. `VIEWMOL` also looks for a file with the extension `.hessian` and tries to read the vibrational spectrum from it, if it was found.

If the option `-dmol` is used the file name of a DMOL/DSOLID/DMOL³ output file has to be specified.

If the option `-gamess` is used the file name of a GAMESS output file has to be specified.

If the option `-gauss` is used the file name of a GAUSSIAN 9X output file has to be specified.

If the option `-gulp` is used the file name of a GULP output file has to be specified.

If the option `-mopac` is used the file name of a MOPAC output file has to be specified. `VIEWMOL` also looks for a file with the extension `.gpt` and reads information about basis sets and MOs from it, if it was found.

If the option `-pdb` is used the file name of a PDB file has to be specified.

If the options `-tm` or `-tmmsi` are used the file name of a TURBOMOLE file containing at least the data group `$coord` has to be specified. `-tm` reads the original TURBOMOLE output while `-tmmsi` allows `VIEWMOL` to read TURBOMOLE outputs from the TURBOMOLE version distributed by MSI (the only difference is the ordering of the normal modes in the control file).

6 Usage and Operating Modes

6.1 Data Read From Input Files

- TURBOMOLE

The program reads the following data groups from the control file:

- \$atoms

- \$basis

- \$people

The basis functions are read from these data groups. These data will be read only if they are available.

- \$closed_shells

This data group is read to determine which molecular orbitals are occupied. The data is necessary for the calculation of electron densities. Currently, no open shell systems can be handled by VIEWMOL.

- \$symmetry

The point group of the molecule is read from this data group.

- \$coord

The Cartesian coordinates of the molecule calculated. This data group must be available.

- \$grad

The Cartesian coordinates and gradients of all previous steps of a geometry optimization. This data group will be read only if it is available.

- \$scfmo

The symmetry labels, energies, and MO coefficients are read from this data group. These data will be read if they are available and if the file contains either converged or first order molecular orbitals.

- \$symmetry

The point group of the molecule. This data group will be read only if it is available.

- \$title

The title of the calculation. This data group will be read only if it is available.

- \$vibrational_spectrum

- \$vibrational_normal_modes

The results of a force constant calculation. These two data groups will be read only if they are available.

- GAMESS

GAMESS output files are first checked for the occurrence of the string GAMESS. If it is found the necessary data are collected from this file. At the moment no vibrational spectra can be displayed for GAMESS outputs since I don't know the format of such an output.

- GAUSSIAN 9X

Gaussian output files are first checked for the occurrence of the string Entering Gaussian System. If it is found the necessary data are collected from this file. To use the wave function related topics in VIEWMOL with GAUSSIAN outputs GAUSSIAN must run with GFPRINT and

`Iop(5/33=1)`³ to print basis set and MO coefficients. Due to the vastly different outputs created by the GAUSSIAN 9X series of programs, it is not guaranteed that a particular output can be successfully read. The common types of output have been tested, but non-default routes through the program might have generated output which cannot be read.

- **DISCOVER**

The file names for DISCOVER files can be `file_name.car`, `file_name.cor`, or `file_name.arc`. The base name is used to construct the file name `file_name.hessian` (the file with frequencies and normal coordinates). All necessary data are extracted from these files.

- **DMOL/DSOLID/DMOL³**

The necessary data are collected from the `.outmol` file.

- **PDB files**

Only the Cartesian coordinates and atomic symbols are read from this file, the connectivity information is ignored and will be determined by VIEWMOL itself.

- **MOPAC**

VIEWMOL first checks for the presence of a file with the extension `.gpt` and the same basename as the MOPAC output file. This file is generated if MOPAC has been run with the keyword GRAPH. If such a file is found coordinates, basis functions, and MO coefficients are read from this file. If such a file does not exist, coordinates are read from the MOPAC output file under the header CARTESIAN COORDINATES. Finally, vibrational frequencies and normal modes are read from the MOPAC output file, if present.

6.2 The Main Window

The program displays the molecule according to the coordinates in the main window. Following manipulations are possible (cf. Figure 1):

- Holding down the left mouse button and moving the mouse horizontally
This rotates the molecule, the view point, or a light source around the y axis.
- Holding down the left mouse button and moving the mouse vertically
This rotates the molecule, the view point, or a light source around the x axis.
- Holding down the middle mouse button and moving the mouse horizontally
This rotates the molecule, the view point, or a light source around the z axis.
- Holding down one of the shift keys and the left mouse button and moving the mouse
This moves (translates) the molecule or an annotation.
- Holding down one of the shift keys and the middle button and moving the mouse
The scaling of the molecule is changed.
- Pressing the cursor keys for moving up `<↑>` or down `<↓>`
The scaling of the molecule is changed. By pressing `<↑>` the molecule will be enlarged and by pressing `<↓>` the molecule will be made smaller.

³GAUSSIAN98 seems to have a bug with respect to this option – no MO coefficients are printed anymore. Use `Iop(5/33=2)` instead which, unfortunately, also prints the density matrix.

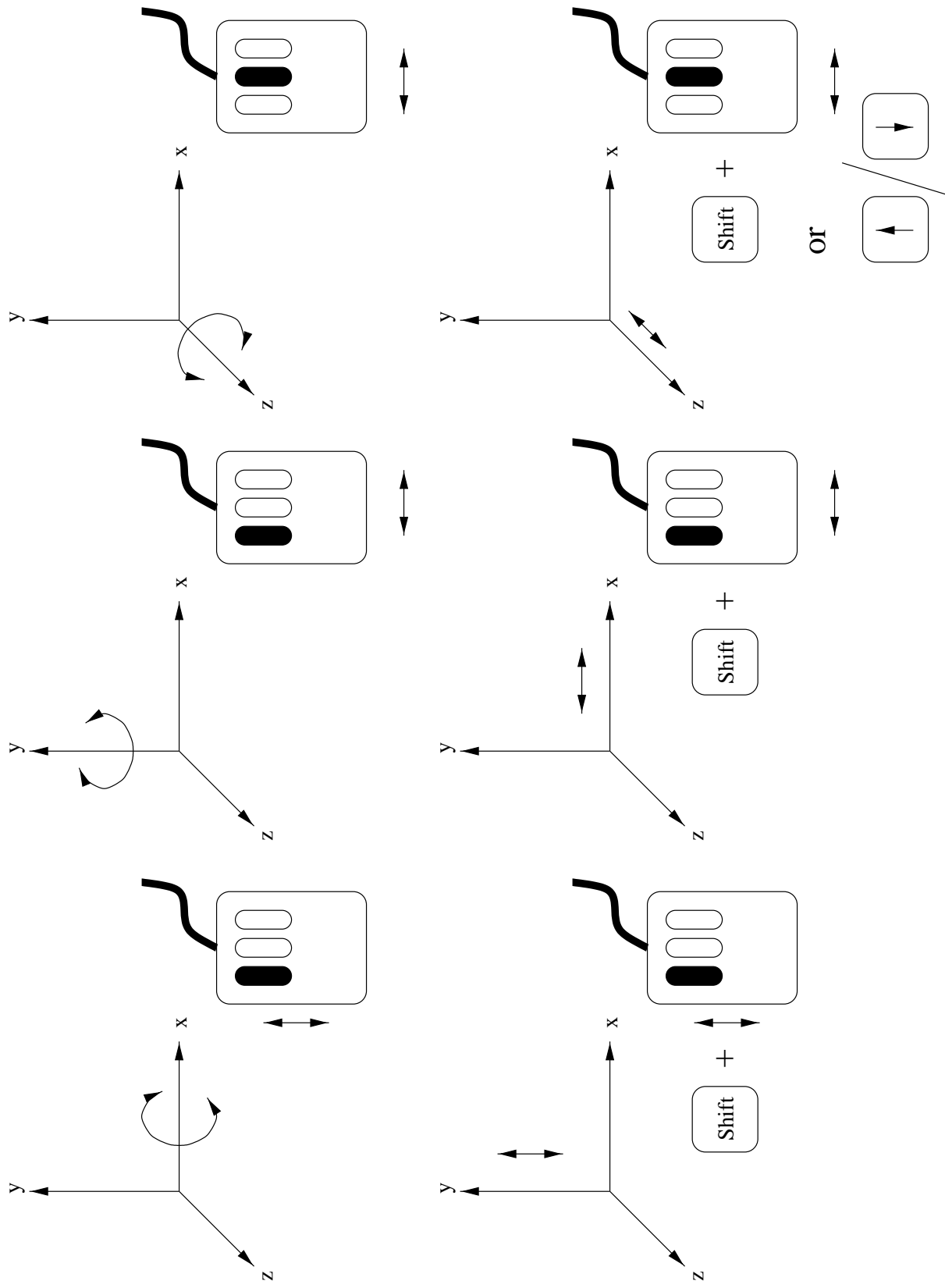


Figure 1: How the mouse can be used to manipulate a molecule or an annotation in the main window.

- Clicking on an atom with the left mouse button

This selects this atom, you will hear a beep. If you have clicked on one atom and then pressed the right mouse button, the average of all bond lengths at this atom is displayed. If you have clicked on two atoms and then pressed the right mouse button, the distance between these two atoms is displayed. If you have clicked on three atoms and then pressed the right mouse button, the angle between these three atoms is displayed. If you have clicked on four atoms and then pressed the right mouse button, the torsion angle between these four atoms is displayed. To delete the displayed values use either the Geometry menu items or repeat the steps above. Clicking with the left mouse button on an atom may be also necessary for setting or selecting some atom specific values (vide infra).

- Holding down one of the shift keys and clicking on a molecule with the left mouse button

The molecule is selected. The window title will show the name of this molecule. All subsequent translations/rotations act only on this molecule.

- Pressing the right mouse button without clicking on an atom before

A menu will appear. The menu contains the following topics (the key combination in parentheses can be used as a shortcut in the English version):

- Molecule ...

A submenu is provided with the following topics:

- * Load molecule ...

Brings up a file selection box to load a molecule.

- * Save molecule ...

Brings up a format selection dialog and a file selection box to save the currently selected molecule to file. Output formats are supported through external filters (similar to the input filters) and can be installed using options in the `viewmolrc` file (cf. p. 41). Coordinates and bond information are passed to the corresponding output filter which writes the file. Currently, the only output formats provided are MSI car-files, MDL mol-files, and TURBOMOLE.

- * Delete molecule ...

Deletes the currently selected molecule.

- * New molecule ...

Brings up the molecule editor and starts the building of a new molecule (cf. p. 27).

- * Modify molecule ...

Brings up the molecule editor to modify an existing molecule (cf. p. 27).

- Select molecule

Provides a submenu with the names of all molecules currently loaded and an item All and can be used to change the currently selected molecule. Other possibilities to select a molecule consist of clicking on the molecule (preferably while holding the shift key down) or pressing the Tab key, which cycles through all entries in the Select molecule submenu.

- Wire model (Alt+W)

The molecule will be drawn with lines. This is the default.

- Stick model (Alt+T)

The molecule will be drawn with sticks.

- Ball and stick model (Alt+A)

The molecule will be drawn with balls and sticks.

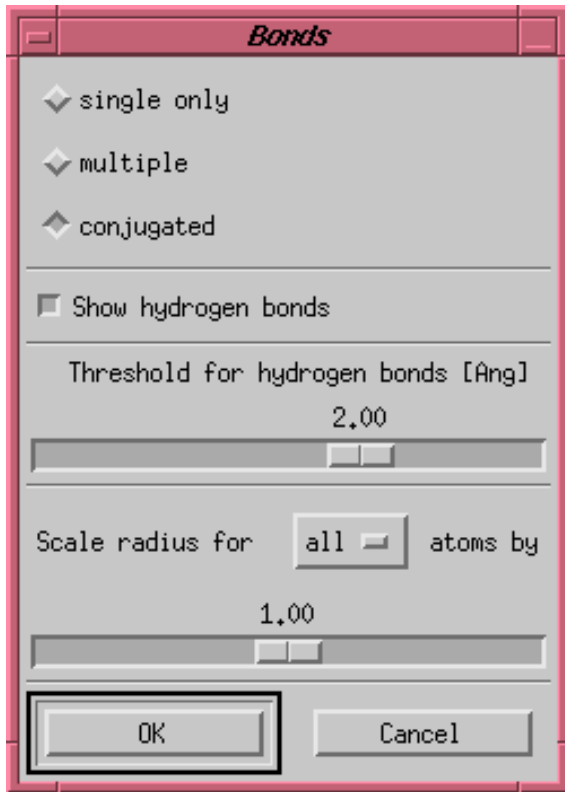


Figure 2: The dialog box for setting options for bonds.

– CPK model (Alt+C)

The molecule will be drawn with CPKs.

– Geometry ...

A submenu is provided with the following topics:

* Clear all (Ctrl+A)

If this topic is selected all labels of bond lengths, bond angles, and torsion angles are deleted.

* Clear last (Ctrl+L)

If this topic is selected the label of a bond length, bond angle, or torsion angle created last is deleted.

* Undo geometry change (Ctrl+U)

If this topic is selected, the last geometry change (cf. p. 27) is reversed. All geometry changes are buffered in the undo buffer and can be reversed one by one by repeatedly using this menu item. Geometry changes can also be reversed through the molecule editor.

– Bond types ...

Brings up the dialog box shown in Figure 2.

* single only

All bonds are drawn as single bonds.

* multiple

VIEWMOL determines the bond order for each bond considering connectivity and elements only (only the following elements are used in the determination of bond orders: H, C, N, O,

F, Si, P, S, Cl, Ge, Br, I). VIEWSMOL then draws bonds with the corresponding bond order. Bond orders can also be changed in the molecule editor.

- * conjugated

VIEWSMOL determines bond orders as for the "multiple" option. It then determines whether multiple bonds are conjugated and draws them as such. This is the default, but can be overwritten using resources (see p. 43).

- * Show hydrogen bonds

This button toggles the display of hydrogen bonds. Hydrogen bonds are determined automatically by VIEWSMOL based on a distance threshold.

- * Threshold for hydrogen bonds [Ang]

This slider can be used to set the distance threshold for the automatic determination of hydrogen bonds. A hydrogen bond is shown if the distance between a hydrogen atom and another atom is larger than the sum of their radii, but smaller than this threshold. The default is 2 Å.

- * Scale radius for all atoms by

This menu and the slider beneath it can be used to scale the Van der Waals radius for an element. Since the Van der Waals radius determines the connectivity of the atoms in the molecule, changing it will also change the connectivity. The option menu can be used to select which element to scale and the slider allows to set the scaling factor.

- Wave function ... (Alt+V)

The dialog box shown in Figure 3 is presented. This topic is currently only available with TURBOMOLE, GAUSSIAN, and MOPAC outputs, and even then only if MO coefficients and basis functions could be read in and if the calculation was closed shell. If TURBOMOLE output is used and the point group of the molecule in \$symmetry is not C1, TURBOMOLE's moloch program must be available (vide supra, p. 4) and only basis functions, occupied MO's, and electron densities can be drawn in this case. Since TURBOMOLE can handle up to g functions and GAUSSIAN can handle up to f functions the same limitations apply to VIEWSMOL.

If any wave function related drawing is displayed and the grid resolution is changed the drawing disappears and the recalculation has to be explicitly demanded by selecting this menu item again, since large molecules require significant time for the recalculation. At the top of this dialog box are five buttons which can be used to select the property which shall be shown.

- * All off

This topic disables the drawing of any wave function related topic. This is the default.

- * Basis function

This topic allows drawing of basis functions. After selecting it and closing the wave function dialog VIEWSMOL prompts for an atom in its main window. Clicking on an atom with the left mouse button will present a dialog box with all basis functions centered on this atom. After selecting one of this basis functions and pressing the OK button the corresponding basis function will be drawn.

- * Basis function in MO

This topic allows drawing of basis functions multiplied by the corresponding coefficient in a molecular orbital. This topic works similar to the previous one, except that the menu will show the MO coefficients in front of all basis functions. If no molecular orbital has been selected in the MO energy diagram window a warning message will be displayed.

- * Molecular orbital

This topic allows the drawing of a molecular orbital. If no molecular orbital has been

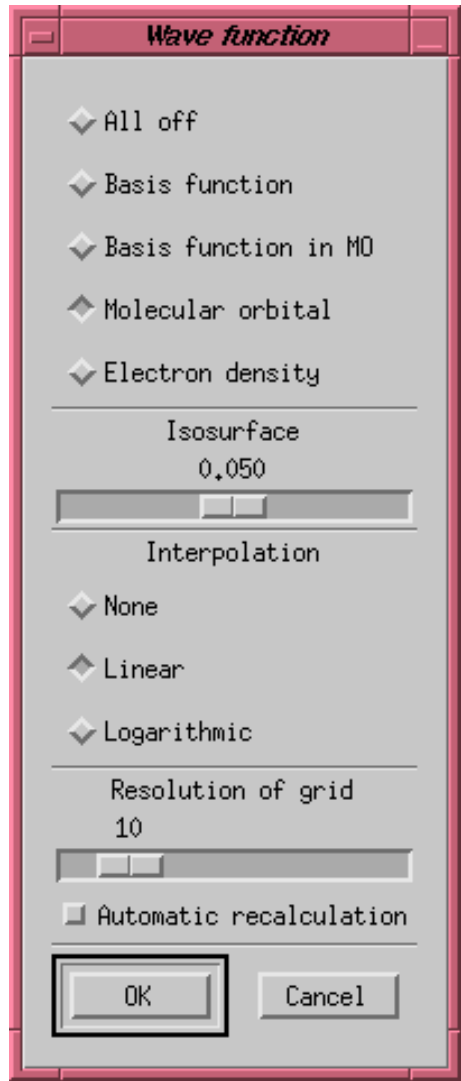


Figure 3: The dialog box for setting options for wave function related topics.

selected in the MO energy diagram window a warning message will be displayed.

* Electron density

This topic allows the drawing of the total electron density.

Next to these buttons there is a slider which can be used to select the value of the isosurface used to draw the property selected. Following this slider another three buttons allow the selection of the interpolation method used in drawing the property.

* None

No interpolation is done. The resulting drawing normally has a lot of edges.

* Linear

A linear interpolation is done between grid points. This gives a much smoother surface.

* Logarithmic

A logarithmic interpolation is done between grid points. This improves the quality of drawing further.

The default is linear, but this can be overwritten in the resource file (vide infra, p. 43). Following is another slider with can be used to set the resolution of the grid. As higher the number selected here as finer the grid and as smoother the resulting surface, but the calculation time goes with the third power of this number. Default is 10, but this can be overwritten in the resource file (vide infra, p. 43). At the bottom of the dialog box is a toggle button which can be used to turn automatic recalculation of MOs etc. on whenever the energy level is changed. Since these calculations can be quite time consuming, this button is off by default, but this can be overwritten in the resource file (vide infra, p. 43).

- Energy level diagram (Alt+E)

A new window will appear which shows the calculated energies of the MOs in an energy level diagram. This topic is only available using DMOL, GAMESS, GAUSSIAN 9X, MOPAC, or TURBOMOLE outputs. In TURBOMOLE outputs the data group \$scfmo must be available.

- Optimization history (Alt+O)

A diagram is plotted in a second window which shows the energies and gradient norms of the geometry optimization. With the cursor keys for moving to the left <←> and to the right <→> one can see how the geometry optimization works. Alternatively, the red cross can be dragged with the mouse.

- Show forces (Alt+F)

The calculated forces acting on the atoms are drawn as arrows. This topic is not available using DISCOVER outputs. The topic is also not available, if no forces were found for the current coordinates.

- Spectrum (Alt+S)

A new window will appear which shows the calculated spectrum for the molecule. This topic is only available if a force constants calculation has been performed.

- Thermodynamics (Alt+Y)

The dialog box shown in Figure 4 is displayed. On the top of this dialog box a number of tabs can be found. All tabs except the last one allow to select the display of thermodynamical data for one of the molecules loaded into VIEWMOL. The last tab will display thermodynamical data for one or more defined reactions among the molecules loaded.

The screen for a molecule shows the title of that molecule on top. Underneath it are displayed on the left hand side the molecular mass (in g/mol), the symmetry number, and either the rotational constants (for molecules in cm⁻¹) or the density (for solids in g/cm³). On the right hand side

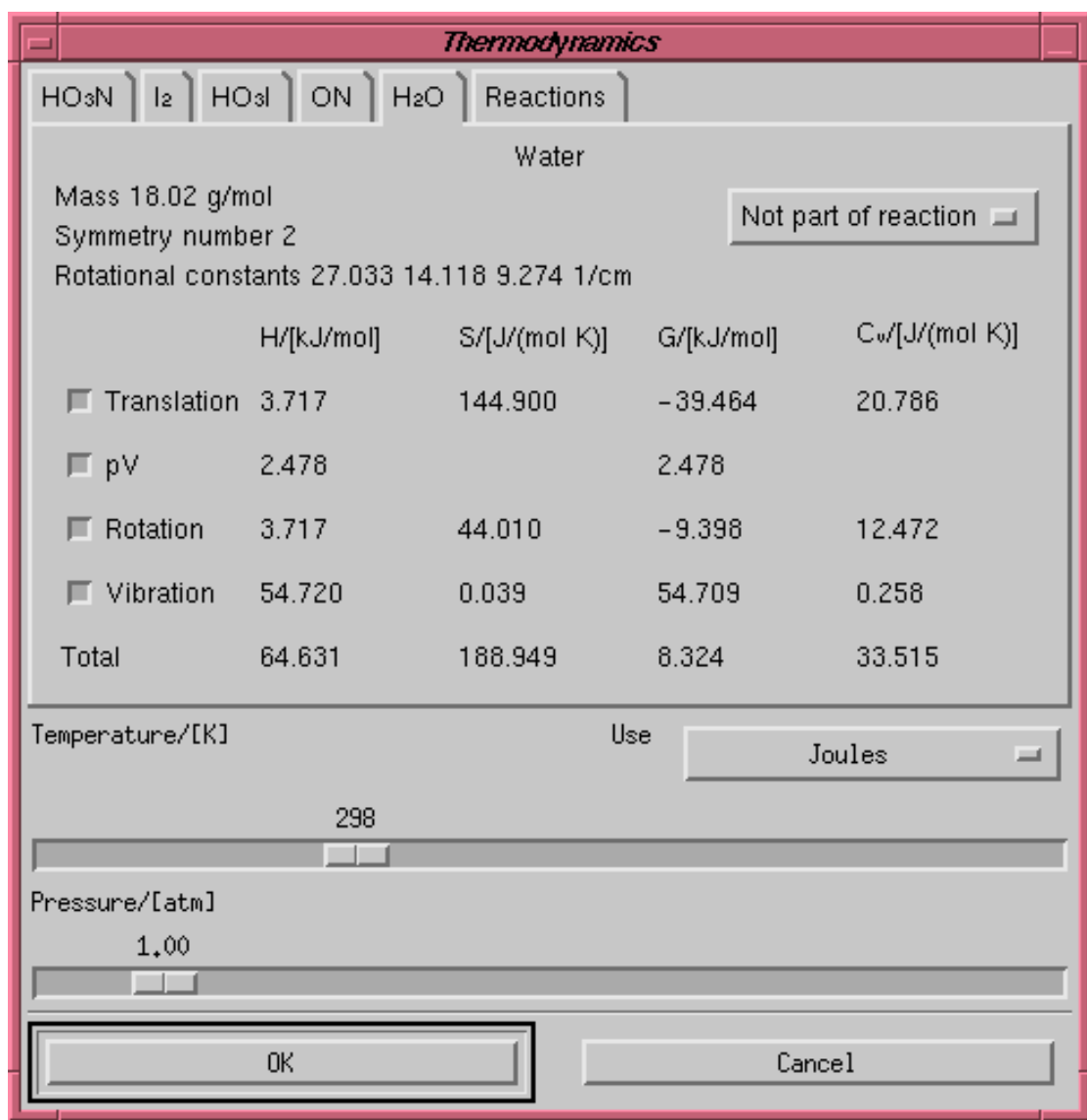


Figure 4: The dialog box for handling thermodynamics calculations.

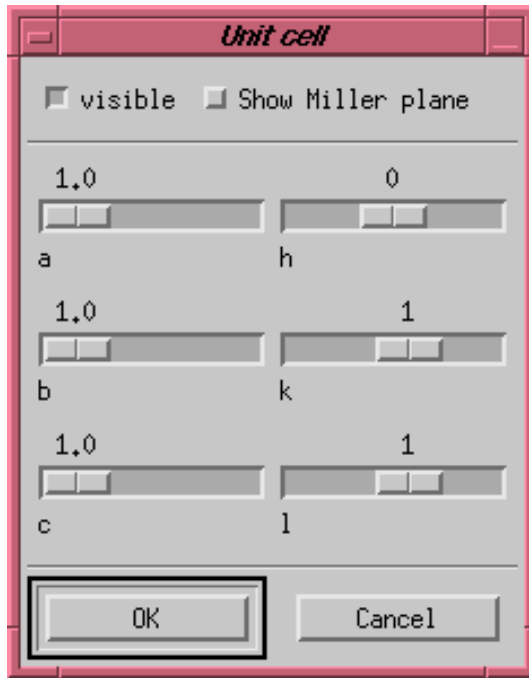


Figure 5: The dialog box for modifying the unit cell.

there is a popup menu which allows the user to select whether this molecule should be a reactant or a product in a reaction, or whether it is not involved in a reaction at all. The last item in this popup menu, All reactions, will force VIEWMOL to determine a linear independent set of possible reactions between all molecules where this item has been selected.

The remainder of the screen shows thermodynamical data for the molecule. On the left hand side there are a number of buttons which can be used to select which contributions (translation, pV, rotation, vibration) to include in the total which is used to calculate thermodynamical data for a reaction. The enthalpy, entropy, Gibbs energy, and heat capacity are listed to the right, split into contributions from translation, pV, rotation, and vibration.

The screen for reactions (not shown) shows the reaction equation on top. Underneath it are the values of reaction enthalpy, entropy, Gibbs energy, heat capacity, and the (decadic) logarithm of the equilibrium constant listed. The electronic and statistical-mechanic contributions to the reaction enthalpy are listed separately.

At the bottom all screens share a popup menu for selecting the units to be used (Joules, calories, or thermochemical calories) and two sliders. The top slider can be used to select the temperature at which the thermodynamical data are to be calculated, the bottom slider serves the same purpose for the pressure.

- Unit cell (Alt+N)

The dialog box shown in Figure 5 is displayed. The first button on the left, visible, allows to turn the display of the unit cell on or off. The three sliders on the left hand side can be used to increase or decrease the number of unit cells displayed in each crystallographic direction. By default between one and five unit cells can be selected.

The first button on the right, Show Miller plane, allows to turn the display of Miller planes on or off. The three sliders underneath this button can be used to select the Miller plane to be

displayed. By default, all combinations between -5 and 5 for the Miller indices are possible.

– Show ellipsoid of inertia (Alt+I)

The display of the ellipsoid of inertia is toggled.

– Drawing modes ... (Alt+M)

The dialog box shown in Figure 6 is provided:

* with dots

Drawing of sticks, balls, CPKs and/or molecular orbitals is done as a dot cloud.

* with lines

Drawing of sticks, balls, CPKs and/or molecular orbitals is done with meshes.

* with surface

Drawing of sticks, balls, CPKs and/or molecular orbitals is done with an opaque surface which has the properties defined in the `vewmolrc` file (these properties hold for sticks as well as for balls or CPKs).

* Lines while rotating

If this option is selected the drawing of the molecule will be done with lines during translations and rotations. This speeds up movements on low-end graphics systems. The default is on, but this can be changed in the resource file.

* Orthographic projection

The molecule is drawn using an orthographic projection.

* Perspective projection

The molecule is drawn using a perspective projection. This kind of projection resembles more closely the way the human eye perceives things.

* Move molecule

If this button is selected all translations and rotations carried out with the mouse act on the currently selected molecule.

* Move view point

If this button is selected all translations and rotations carried out with the mouse act on the viewpoint. This option is only available if perspective projection is used. Moving the view point allows the user to move into a molecule.

* Move light 1

If this button is selected all translations and rotations carried out with the mouse act on light 1.

* Move light 2

If this button is selected all translations and rotations carried out with the mouse act on light 2.

* Lights on/off, Light 1

* Lights on/off, Light 2

These two button can be used to switch lights on and off. Lights have only an effect if the drawing mode is "with surface" and either the stick, ball-and-stick, or CPK model is selected.

* Resolution of spheres

The number of polygons used for the drawing of sticks, balls and/or CPKs is changed. A higher value makes the surfaces more smoothly looking, but also decreases drawing speed. A lower value makes the surfaces rougher looking, but increases drawing speed.

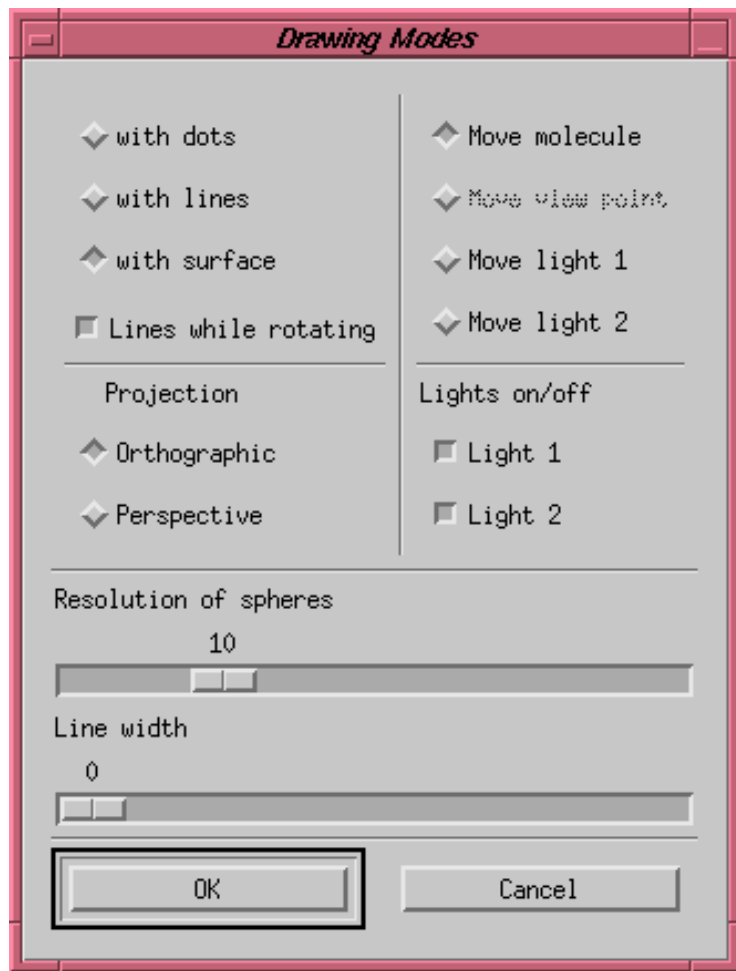


Figure 6: The dialog box for setting options for drawing style.

* Line width

The line width used for drawing the molecule as wire frame model can be selected. A value of 0 means dynamic determination of the line width based on the size of the window. This value is the default.

- Ground color (Alt+G)

The color editor appears in a separate window which can be used to change the color of the ground if perspective drawing is enabled (see description of the color editor below, p. 31).

- Background color (Alt+B)

The color editor appears in a separate window which can be used to change the background color of the window (see description of the color editor below, p. 31).

- Label atoms (Alt+L)

The atoms are labeled with atom symbols from input files. A number counting the atoms according to their order in the input is concatenated to the symbol.

- Annotate (Ctrl+N)

Using this topic annotations can be created in the main window. After selecting this topic the cursor turns into a text input cursor. Clicking at any point in the main window now allows to enter an arbitrary text string. Pressing <return> ends the annotation function. Existing annotations can be edited by simply clicking on them or deleted by deleting all characters in the string. Annotations can be moved in the same way as the molecule can be moved: hold a shift key down, click on the annotation and move the mouse. Annotations support the clipboard, i. e. annotations can be cut and pasted between applications.

- Run script (Ctrl+R)

A file selection box is displayed which allows the selection of a Python script to be run within VIEWMOL.

- Save drawing (Alt+D)

The dialog box shown in Figure 7 is provided which can be used to set file formats, file names and other options for writing the drawing to a file.

* TIFF

The current drawing of the molecule is written out as a TIFF file. The "TIFF compression" menu permits the selection of a compression algorithm. If normal modes are animated while this option is selected a series of 20 TIFF files will be written out, each contains a single frame of the animation. By using standard image manipulation tools available on the Internet it is possible to generate a video file (MPEG) from these TIFF files which can be included in multimedia documents (vide infra, p. 41).

* HPGL

The current drawing of the molecule is written out as a HPGL file for plotting on a plotter or a laser printer. This topic is not available if the drawing is done with sticks, balls, or CPKs and surfaces.

* PostScript

The current drawing of the molecule is written out as a PostScript file. If this topic is selected while the drawing is done with sticks, balls, or CPKs and surfaces a resolution independent PostScript file for the drawing is written, but since PostScript does not provide a way to automatically remove hidden surfaces the drawing may contain some artifacts. The algorithm used tries to minimize these artifacts, but is not always completely successful. Sometimes changing the orientation of the molecule slightly or increasing the resolution of the spheres might help.

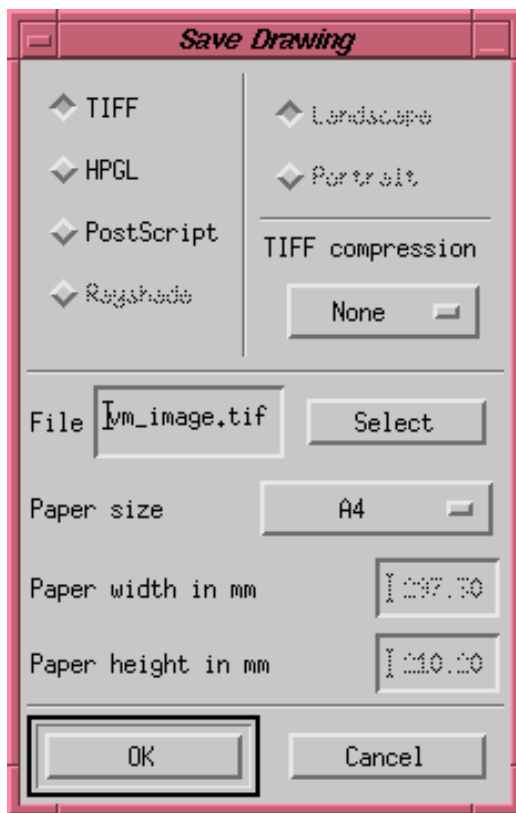


Figure 7: The dialog box for setting options to save a drawing

* Rayshade

The current drawing of the molecule is written out as an input file for RAYSHADE. This topic is only available if the drawing is done with sticks, balls, or CPKs. If molecular orbitals are drawn this topic is only available if the MO is drawn with a surface. If normal modes are animated while this option is selected a series of 20 input files for RAYSHADE will be written out, each containing a single frame of the animation.

* Landscape

* Portrait

The orientation of the drawing on the page can be chosen if the drawing is written out as either a HPGL or a Postscript file.

* TIFF compression

Can be used to select the compression mode for TIFF files. Due to a software patent on the LZW compression algorithm this compression can no longer be provided.

* File

The name for the file to be generated.

* Select

Brings up a file selection box to select the name for the file to be generated.

* Paper size

Can be used to select the paper size.

* Paper width in mm

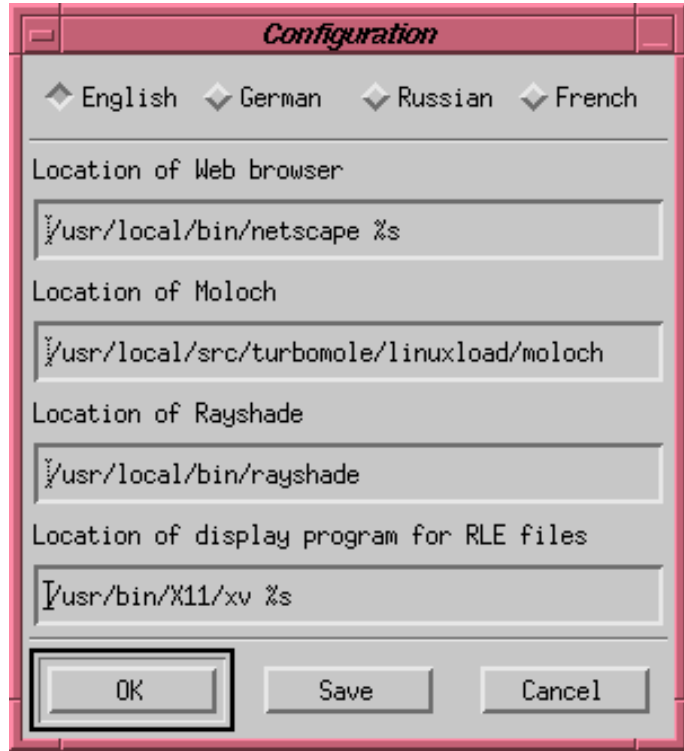


Figure 8: The dialog box for setting configuration options.

- * Paper height in mm
Input user defined paper width and height. Only available if "Paper size" is set to "User defined".
- Help/Manual (Alt+H)
This topic opens a window with the VIEWMOL manual. This topic is only available if the file viewmol.html was found in the location \$VIEWMOLPATH/man.
- Configuration ...
The dialog box shown in Figure 8 is provided. At the top of this dialog the language VIEWMOL uses in its interface can be selected. VIEWMOL loads its language specific data from files Xdefaults.<language> from the directory VIEWMOLPATH points to. The four text input fields can be used to specify the location of helper programs VIEWMOL needs for some of its operations. If the corresponding program was found in the path the dialog box will already show the correct information. If the programs specified here need file names as parameters, put %s as a place holder for the file name in the command.
At the bottom is a button Save which allows the information entered in this dialog as well as some other settings to be stored as resources in \$HOME/.Xdefaults. The following settings are saved: position and size of all open windows, window colors, selected model, selected drawing mode, selected bond type, setting of "lines while rotating", selected interpolation mode, resolution of spheres, line width, selected isosurface, selected resolution for density of states, setting of "automatic recalculation", paper size, and hydrogen bond threshold. **Note:** On Linux the setting of resources is kept across different invocations of the program. Saving the configuration and restarting VIEWMOL will therefore apparently not work. To get rid of the old resource

settings issue the command `xrdb -remove $HOME/.Xdefaults` or log out and in again.

- Quit Viewmol (Q)
This quits the program.

6.3 The Spectrum Window

Choosing Spectrum from the main window menu will result in a new window showing the calculated spectrum for the molecule. In this window the mouse acts as follows:

- Clicking with the left mouse button on a line in the spectrum
The molecule shows the corresponding normal vibration.
- Clicking with the middle mouse button in the window, holding it down and moving the mouse
This displays a rubber band box with which one can zoom into the spectrum.
- Pressing the right mouse button
A menu will appear.

The menu contains the following topics:

- Settings for spectrum ... (Alt+S)
Selecting this topic displays the dialog box shown in Figure 9.
 - * Type of spectrum
The buttons labeled All modes, IR active modes, Raman active modes, and Inelastic neutron scattering can be used to select the type of spectrum desired. IR active modes are the default.
 - * Animation
The buttons labeled Animate, Draw arrows, and Distort can be used to select whether the normal modes are to be shown animated or with arrows or whether you want to distort the molecule along a normal mode. A distorted molecule can be saved using the Save molecule option from the main menu. Animation is the default.
 - * Line shapes
The buttons Line spectrum and Gaussian spectrum can be used to select whether the spectrum is drawn as simple line spectrum or whether a Gaussian band shape [1] should be applied. Line spectrum is the default.
 - * Weights for inelastic neutron scattering
When you activate this option you can enter a value in the field to the right. After choosing the OK button all weights are drawn at the atoms in the main window and you can set an atom's weight just by clicking on it with the left mouse button. All weights are set initially to zero, so that selecting Inelastic neutron scattering as spectrum type produces nothing.
 - * Temperature
This slider can be used to set the temperature. The temperature is used for the calculation of the inelastic neutron scattering intensities and for the Gaussian shaped spectrum.
 - * Amplitude
This slider can be used to change the amplitude of the vibration. Its value is multiplied with the standard amplitude of a vibration. This slider can also be used to change the distortion of the molecule while Distort is selected.

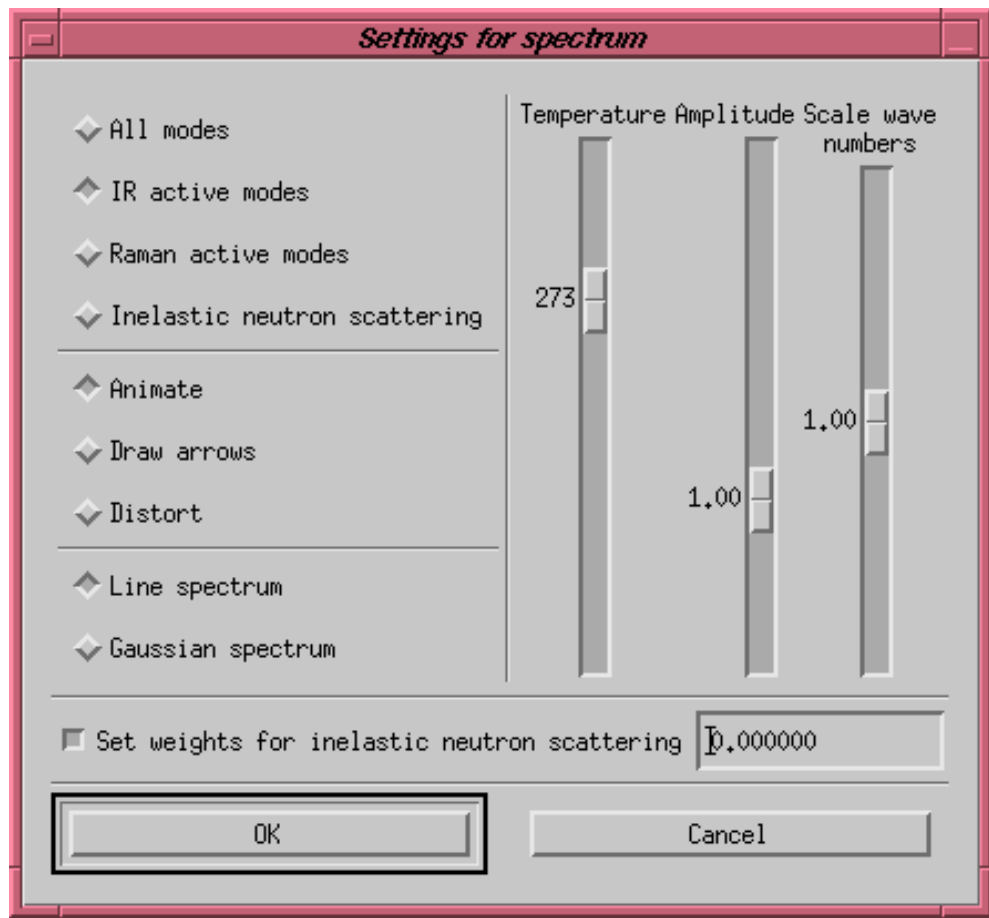


Figure 9: The dialog box for setting options for the spectrum

* Scale wave numbers

This slider can be used to scale the wave numbers.

- Select molecule

Provides a submenu with the names of all molecules currently loaded and can be used to change the molecule for the currently displayed spectrum. Another possibility to select a molecule consists of pressing the Tab key, which cycles through all entries in the **Select molecule** submenu.

- Imaginary wave numbers

If the conformation of the molecule is a saddle point you have imaginary wave numbers. The corresponding "normal modes" can be shown by selecting a imaginary wave number from this submenu.

- Read observed spectrum ... (Alt+R)

This topic can be used to read a spectrum from a file and display it along the calculated spectrum. Selecting this topic opens a file selection box to chose the file. This file has to contain a spectrum with one wave number and intensity per line. All points read are connected by a line to form a continuous spectrum (there is currently no possibility to read a line spectrum). Lines with a '#' or a letter in the first column are ignored.

- Delete observed spectrum ... (Alt+E)

This topic deletes a spectrum read with "Read observed spectrum".

- Zoom out (Alt+Z)

This topic can be used to zoom out of the spectrum after previous zoom-ins. The zoom mechanism stores all previous enlargement steps. By selecting this topic you move back one step.

- Save drawing (Alt+D)

The same dialog box as for **Save drawing** in the main menu is shown and allows you to save the spectrum as a TIFF, HPGL, or PostScript file (cf. p. 19).

- Foreground color (Alt+F)

The foreground color of the spectrum can be changed using the color editor (vide infra, p. 31).

- Background color (Alt+B)

The background color of the spectrum can be changed using the color editor (vide infra, p. 31).

- Quit spectrum (Alt+Q)

This closes the spectrum window.

• Pressing the arrow keys $<\rightarrow>$ and $<\leftarrow\rightarrow>$, respectively

The next lower and higher wave number, respectively is selected and the molecule shows the corresponding normal vibration. Normal vibrations of imaginary wave numbers can be displayed in this manner, too.

6.4 The Optimization History Window

Choosing **Optimization history** from the main window menu will result in a new window showing the energy and gradient norm in dependence of the step number of the geometry optimization. In this window the mouse acts as follows:

• Pressing the left mouse button and holding it down

The red cross showing the actual step of the geometry optimization can be moved with the mouse.



Figure 10: The dialog box for setting options for the optimization history

The main window shows the corresponding geometry. Changing the actual step can also be achieved using the cursor keys for moving to the left $<\leftarrow>$ and to the right $<\rightarrow>$, respectively.

- Pressing the right mouse button

A menu will appear.

The menu contains the following topics:

- Settings for history ... (Alt+S)

Selecting this topic displays the dialog box shown in Figure 10.

- * Energies

If this button is selected the energy curve is drawn. The default is on.

- * Gradient norms

If this button is selected the gradient norm curve is drawn. The default is on.

- * Scales

This button toggles drawing of the scales on and off. The default is on.

- Select molecule

Provides a submenu with the names of all molecules currently loaded and can be used to change the molecule for the currently displayed optimization history. Another possibility to select a molecule consists of pressing the Tab key, which cycles through all entries in the Select molecule submenu.

- Animate (Alt+A)

Selecting this topic will animate the optimization history. To stop the animation select this topic again.

- Save drawing (Alt+D)

The same dialog box as for Save drawing in the main menu is shown and allows you to save the optimization history as a TIFF, HPGL, or PostScript file (cf. p. 19).

- Color for energy (Alt+E)

The color for the energy curve can be changed using the color editor (vide infra, p. 31).

- Color for gradient norm (Alt+G)

The color for the gradient norm curve can be changed using the color editor (vide infra, p. 31).

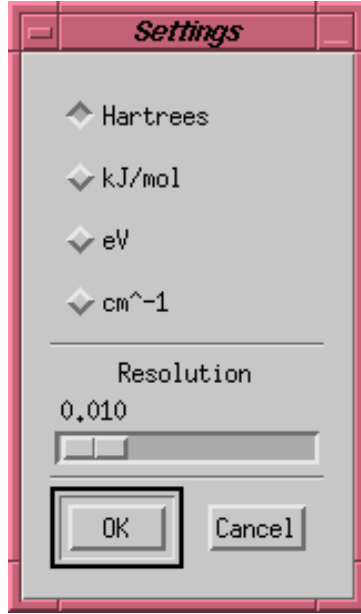


Figure 11: The dialog box for setting the energy units for the MO energy level diagram

- Background color (Alt+B)
The background color of the diagram can be changed using the color editor (vide infra, p. 31).
- Quit history (Q)
This closes the optimization history window.

6.5 The Energy Level Diagram Window

Choosing Energy level diagram from the main window menu will result in a new window showing the calculated MO energies in an energy level diagram. In this window the mouse acts as follows:

- Clicking with the left mouse button on a line
A box appears containing the symmetry and the energy for this MO. Selecting a MO can also be achieved by pressing the cursor keys for moving up $\leftarrow\uparrow\rightarrow$ and down $\leftarrow\downarrow\rightarrow$, respectively.
- Clicking in the window, pressing the middle mouse button and holding it down
With the rubber band box drawn one can zoom into the diagram.
- Pressing the right mouse button
A menu will appear.

The menu contains the following topics:

- Settings for energy level diagram ... (Alt+S)
Selecting this topic displays the dialog shown in Figure 11.
 - * Units
The four buttons can be used to select the energy unit. Available are Hartrees (default), kJ/mol, eV, and cm^{-1} .

- * Resolution for density of states

This slider can be used to change the resolution for the density of states. A smaller value results in a higher resolution. The default is 0.01. The minimum, maximum, and default values can be set in the resource file. (vide infra, p. 43).

- Select molecule

Provides a submenu with the names of all molecules currently loaded and can be used to change the molecule for the currently displayed energy level diagram. Another possibility to select a molecule consists of pressing the Tab key, which cycles through all entries in the Select molecule submenu.

- Transition (Alt+T)

This topic can be used to calculate the energy for a transition between two MO's. This topic is available only if one MO was selected by clicking on it. Choosing this menu topic followed by clicking on another MO draws a line showing the transition and a box containing the symmetry labels and the energy difference between these two MO's. Further clicks on other MO's repeat calculations of energy differences. To leave this mode click either somewhere in the window where no MO's are or select this topic from the menu again.

- Zoom out (Alt+Z)

This topic can be used to zoom out of the diagram after previous zoom-ins. The zoom mechanism stores all previous enlargement steps. By selecting this topic you move back one step.

- Save drawing (Alt+D)

The same dialog box as for Save drawing in the main menu is shown and allows you to save the energy level diagram as a TIFF, HPGL, or PostScript file (cf. p. 19).

- Draw density of states (Alt+D)

This topic can be used to toggle between the density of states and the energy level diagram. The energy level diagram is the default.

- Foreground color (Alt+F)

The foreground color of the diagram can be changed using the color editor (vide infra, p. 31).

- Background color (Alt+B)

The background color of the diagram can be changed using the color editor (vide infra, p. 31).

- Quit energy diagram (Q)

This closes the energy level diagram window.

7 Editing molecules

VIEWMOL now allows the building and editing of molecules. Bond lengths, bond angles, and torsion angles of an existing molecule can be changed, atoms can be replaced and added or deleted. New molecules can be built.

7.1 Changing bond lengths, bond angles, and torsion angles

To change a bond length, a bond angle, or a torsion angle click on the corresponding atoms and then press the right mouse button. This will display the length of the bond and the value of the bond or torsion angle, respectively (cf. p. 10). Now click on the number with the left mouse button. A cursor will appear and the

value displayed can be changed. After pressing **Return** the new value for the bond length, bond angle, or torsion angle will be set. The atom which has been clicked on first (and all atoms connected to it) will be moved. It is impossible to change bond lengths, bond angles, or torsion angles if they are part of a ring. All changes in geometry can be reversed by using **Undo geometry change** from the **Geometry** menu or the corresponding button in the molecule editor dialog box. The number of undos is unlimited.

7.2 Adding or replacing atoms

From the main window menu select the **Modify molecule** entry in the **Molecule** submenu. The dialog box shown in Figure 12 is displayed. The upper part contains the periodic table of elements and allows the selection of the element to be added or used as replacement.

In the middle there are a number of buttons for selecting different operations and certain defaults. These are the following:

- **Change geometry**

If this item has been chosen atoms can be selected with the mouse and geometry changes carried out as described in the previous section.

- **Add atom**

An atom of the element selected in the periodic table will be attached to the atom in the molecule clicked on with the left mouse button. The new bond will have the bond order selected in the editor dialog box and a bond length which is 90 % of the sum of the atomic radii (read from the `viewmolrc` file). The local geometry of the atom clicked on will be changed to reflect the current coordination of this atom (two bond partners – linear, three – trigonal planar, four – tetrahedral, five – trigonal bipyramidal, six – octahedral, seven – pentagonal bipyramidal etc.).

- **Delete atom**

An atom clicked on with the left mouse button will be deleted. If **Deleting atoms changes local geometry** is turned on the local geometry of the atom(s) connected to the deleted one will be changed in the same way as described under **Add atom**. Otherwise the local geometry remains as before.

- **Replace atom**

The atom clicked on with the left mouse button is replaced by the element selected in the periodic table. Bond lengths are adjusted to reflect the new element as long as the atom changed is not part of a ring.

- **Create bond**

A new bond will be created between the two atoms clicked on with the left mouse button. This bond will have the order selected under **Bonds are**.

- **Remove bond**

The bond between the two atoms clicked on with the left mouse button is deleted. Bonds created or removed by the user have precedence over bonds created automatically. This means that once a bond has been created by the user only the user can remove it and vice versa regardless of what happens to the molecule.

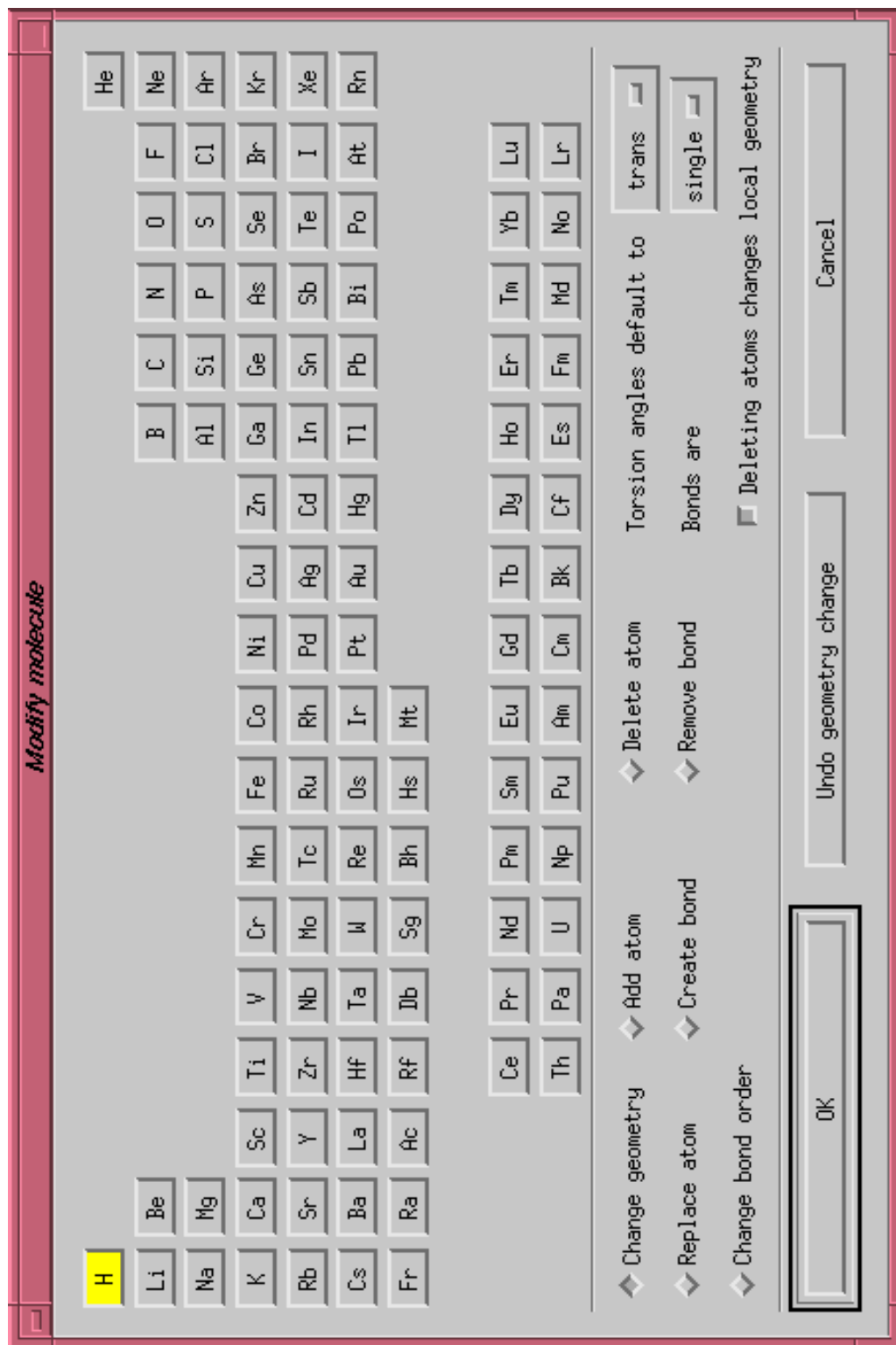


Figure 12: The molecule editor

- Change bond order

The bond between the two atoms clicked on with the left mouse button is assigned the bond border selected under Bonds are.

- Torsion angles default to

While building a molecule bond lengths of newly created bonds are set to 90 % of the sum of the atomic radii of the bonded atoms and bond angles are assigned according to the coordination. Torsion angles can be selected from this menu. Available values are trans (180°), cis (90°), gauche (60°), and $-$ gauche (-60°). This allows the construction of more complicated molecules. The torsion angle is always measured along the backbone of a molecule, i. e. while building e. g. a hydrocarbon chain the torsion angle is always measured between the carbon atoms. The backbone of a molecule is determined by counting all atoms attached to one atom and following the bonds which connect the atoms with the largest number of other atoms attached.

- Bonds are

This menu allows the selection of the bond order for bonds. Available are single, double, and triple. Bond conjugation and hydrogen bonds are determined automatically according to the respective settings in the Bond type menu.

- Deleting atoms changes local geometry

If an atom is deleted the local geometry of the atom(s) bonded to it can either be left unchanged or modified according to the new coordination. This switch can be used to select which behavior is preferred.

In the bottom row of the molecule editor dialog box is a button Undo geometry change which can be used to reverse all changes to the molecule (except changes to bonds). The number of undos is unlimited.

7.3 Example: Building cyclohexane

Cyclohexane in its chair configuration has a structure which seems quite complicated to generate, but it can easily be built using VIEWMOL's molecule builder. First, start VIEWMOL without any parameter on the command line and press Cancel in the file selection box which pops up. You should now have an empty window on your screen. Press the right mouse button within this window and select Molecule/New molecule . . . from the pop-up menu. The molecule editor will open up. We start building cyclohexane by building one of the tetrahedra:

- Select C from the periodic table.
- Click with the left mouse button in VIEWMOL's window. This will place the first carbon atom at the origin.
- Click with the left mouse button on the first carbon atom. This will attach the second carbon atom to the first.
- Click with the left mouse button on the second carbon atom. This will attach the third carbon atom to the second.
- Select H from the periodic table and click twice on the central carbon atom. The molecule should now look like the left one in Figure 13.

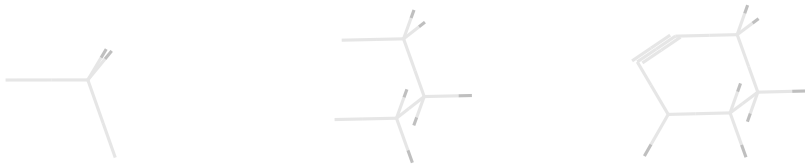
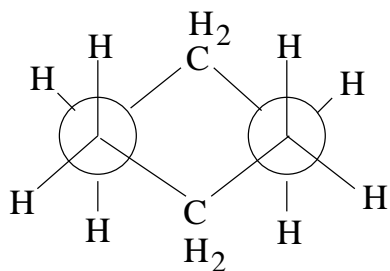


Figure 13: Building cyclohexane

To continue building cyclohexane we have to remember that the torsion angles among the carbon atoms alternate between gauche and –gauche which can be determined easily by looking at the Newman projection:



- Select C again and set the default torsion angle to gauche.
- Click on one of the end carbon atoms. Then select H and click on the same carbon atom two more times.
- Change the default torsion angle to –gauche and repeat the last two steps on the just added carbon atom. The molecule should now look like the one in the middle of Figure 13.
- Change the default torsion angle to gauche again and add a carbon atom and one hydrogen atom to the last carbon atom. The molecule will now look like the right one in Figure 13.
- Change the default torsion angle to trans and add the second hydrogen atom to the carbon atom added last.
- Now change the default torsion angle to –gauche and add one hydrogen atom to the next carbon atom in the ring, then change the default torsion angle to trans and add the second hydrogen atom.
- Finally, complete cyclohexane by adding first a hydrogen with the default set to trans and than a hydrogen with the default set to gauche to the next carbon atom in the ring.

8 The color editor

All menus contain items for changing colors. Selecting one of these items displays the color editor (cf. Figure 14). In the first row of the color editor dialog box are a number of different colored boxes. You can

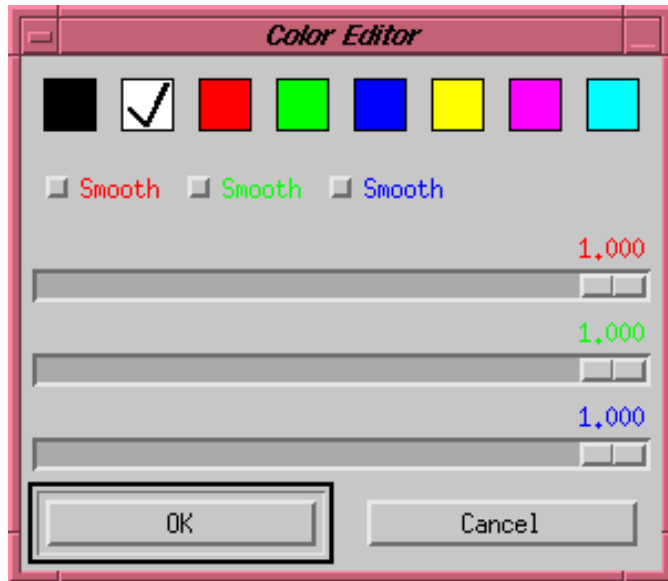


Figure 14: The color editor

select one of these colors just by clicking in the corresponding box with the left mouse button. The color you want to change in the picture changes, too, but this is not a permanent change until you click the OK button.

If none of the colors in the boxes satisfies your needs you can set up your own color by moving one of the sliders for red, green, or blue with the left mouse button. The color in the picture you want to change shows the currently selected color and by clicking the OK button you change your color permanently.

If the color editor has been brought up to edit the background color of the main window three buttons labeled Smooth are present. Activating any of these buttons will create a color ramp for the corresponding color and mix it with the other RGB components. The slider for this color then has no effect. That way interesting shading effects in the background can be generated.

9 Programming VIEWSMOL with Python

VIEWSMOL can be programmed using Python. The Python interpreter embedded in VIEWSMOL has been extended by the modules:

- atom
- element
- energylevel
- history
- label
- light

- molecule
- spectrum
- viewmol

which allow Python programs access to VIEWMOL's internal data model and functions. Each module provides a number of functions for use in Python programs.

9.1 The atom module

`getElement()`

Returns the `element` object for an atom.

`getCoordinates()`

Returns a tupel with the x, y, and z coordinates and the name of an atom.

`radius([rad])`

Sets or returns the Van der Waals radius for an atom. If `rad` is given it has to be a double. The Van der Waals radius is measured in Ångstrøms.

`radiusScaleFactor([scaleFactor])`

Sets or returns the scale factor for the radius of an atom. If `scaleFactor` is given it has to be a double. The scale factor must be greater than or equal zero.

`neutronScatteringFactor([factor])`

Sets or returns the neutron scattering factor for an atom. If `factor` is given it has to be a double. The neutron scattering factor must be greater than or equal zero.

`name([name])`

Sets or returns the name of an atom. If `name` is given it has to be a string. The maximum length of the string is eight characters.

`replace(elementSymbol)`

Replaces the atom with an atom of element `elementSymbol`. `elementSymbol` has to be a string and a valid element symbol.

`delete()`

Deletes this atom. Note: Atoms cannot be deleted using Python's `del` operator since Python works on VIEWMOL's data structures.

9.2 The element module

`darkColor([red, green, blue])`

Sets or returns the red, green, and blue values for the darkest part of an atom of an element. If `red`, `green`, and `blue` are given they have to be floats between 0.0 and 1.0. If the color is queried, it is returned as a tupel of red, green, and blue values.

`lightColor([red, green, blue])`

Sets or returns the red, green, and blue values for the lightest part of an atom of an element. If `red`, `green`, and `blue` are given they have to be floats between 0.0 and 1.0. If the color is queried, it is returned as a tupel of red, green, and blue values.

`emissionColor([red, green, blue])`

Sets or returns the red, green, and blue values of the color an atom of an element emits. If *red*, *green*, and *blue* are given they have to be floats between 0.0 and 1.0. If the color is queried, it is returned as a tuple of red, green, and blue values.

`ambientColor([red, green, blue])`

Sets or returns the red, green, and blue values of the ambient color of an atom of an element. If *red*, *green*, and *blue* are given they have to be floats between 0.0 and 1.0. If the color is queried, it is returned as a tuple of red, green, and blue values.

`specularColor([red, green, blue])`

Sets or returns the red, green, and blue values of the specular color of an atom of an element. If *red*, *green*, and *blue* are given they have to be floats between 0.0 and 1.0. If the color is queried, it is returned as a tuple of red, green, and blue values.

`shininess([shininess])`

Sets or returns the shininess of an atom of an element. If *shininess* is given it has to be a float between 0.0 and 128.0.

`transperancy([transperancy])`

Sets or returns the transperancy of an atom of an element. If *transperancy* is given it has to be a float between 0.0 and 1.0.

9.3 The energylevel module

`show()`

Displays the energy level diagram window for a molecule.

`unit([unit])`

Sets or returns the energy unit for the energy level diagram. *unit* has to be one of the integer constants HARTREE, KJ/MOL, EV, or 1/CM defined in the energylevel module.

`resolution([resolution])`

Sets or returns the resolution for the energy level diagram. *resolution* has to be a double greater than zero.

`mode([mode])`

Sets or returns the mode for the energy level diagram. *mode* has to be one of the integer constants ENERGY_LEVELS or DENSITY_OF_STATES defined in the energylevel module.

`selectMO(mo1 [, mo2])`

Selects one or two molecular orbitals. *mo1* and *mo2* have to be integers between 0 and the number of molecular orbitals.

`deselect()`

Deselects molecular orbitals which have been selected with a call to `selectMO`.

`saveDrawing(format, filename)`

Saves the energy level diagram to file. *format* has to be one of the integer constants TIFF, HPGL, or POSTSCRIPT defined in the viewmol module. *filename* has to be a string containing the name of the file the drawing is saved to.

9.4 The history module

```
show( )  
    Displays the optimization history window for a molecule.  
  
showEnergy( status )  
    Sets the displays of the energy curve to status. status has to be one of the integer constants ON or OFF defined in the viewmol module.  
  
showGradient( status )  
    Sets the displays of the gradient curve to status. status has to be one of the integer constants ON or OFF defined in the viewmol module.  
  
showScales( status )  
    Sets the displays of the scales to status. status has to be one of the integer constants ON or OFF defined in the viewmol module.  
  
animate( status )  
    Sets the animation of the optimization history to status. status has to be one of the integer constants ON or OFF defined in the viewmol module.  
  
iteration( [ iteration ] )  
    Sets or returns the iteration displayed. iteration has to be an integer between 0 and the number of optimization steps for this molecule.  
  
saveDrawing( format,filename )  
    Saves the energy level diagram to file. format has to be one of the integer constants TIFF, HPGL, or POSTSCRIPT defined in the viewmol module, filename the name of the file the drawing is saved to.
```

9.5 The label module

```
label( [ mode ] )  
    Creates a new label. mode is an optional integer denoting whether the label is editable by the user, movable by the user, or both. In the first case mode has to be set to the integer constant EDITABLE, in the second case to the integer constant MOVABLE, and in the last case to EDITABLE | MOVABLE. The integer constants are defined in the label module. The default mode is editable and movable.  
  
translate( x,y,z )  
    Sets the position of a label. x, y, and z are integers specifying the coordinates of the label in pixels. The z coordinate has no effect.  
  
text( [ text ] )  
    Sets or returns the text of a label. If present text has to be a string. The maximum number of characters for a label is limited to 255.  
  
setColor( red,green,blue,alpha )  
    Sets the red, green, blue, and alpha components of a label's color. red, green, blue, and alpha have to be floats between 0.0 and 1.0. The default color for a label is black.  
  
delete()  
    Deletes a label.
```

9.6 The light module

`rotate(x, y, z)`

Rotates a light. *x*, *y*, and *z* are the angles the light is to be rotated about the *x*, *y*, and *z* axis, respectively. These are integers and are measured in degrees.

`switch(status)`

Switches a light on or off. *status* is one of the integer constants ON or OFF defined in the `viewmol` module.

9.7 The molecule module

`molecule()`

Creates a new instance of a molecule object and returns a reference to it. Note: To obtain object references to molecules already loaded into VIEWMOL use the `getMolecules` function of the `viewmol` module.

`translate(x, y, z)`

Translates (shifts) molecule by *x*, *y*, and *z* along the *x*, *y*, and *z* axis, respectively. *x*, *y*, and *z* have to be integers and are measured in pixels of the screen.

`rotate(x, y, z)`

Rotates molecule by *x*, *y*, and *z* about *x*, *y*, and *z* axis, respectively. *x*, *y*, and *z* have to be integers and are measured in degrees.

`getSpectrum()`

Creates a new instance of a spectrum object and returns a reference to it if there is spectral information associated with this molecule.

`getEnergyLevels()`

Creates a new instance of an energy level object and returns a reference to it if there is information about energy levels associated with this molecule.

`getHistory()`

Creates a new instance of a history object and returns a reference to it if there is information about the optimization history associated with this molecule.

`showForces(status)`

Sets the display of forces for all molecules to *status*. *status* has to be one of the integer constants ON or OFF defined in the `viewmol` module.

`getAtoms()`

Returns a list containing references to all atom objects the molecule is composed off.

`getBonds()`

Returns a list of tupels describing all bonds in the molecule. The tupels consists of three integers (`atom1, atom2, order`) where `atom1` and `atom2` are the indices of the two atoms forming the bond and `order` is the bond order. The bond order can be -1, -2, 1, 2, or 3 for hydrogen/Van der Waals, conjugated, single, double, and triple bonds, respectively.

`getWavenumbers()`

Returns a list of tupels describing all wave numbers of the molecule. The tupel consists of four floats and one string (`waveNumber, IRIIntensity, RamanIntensity, INSIntensity`)

`ty, symmetry)` where `waveNumber` is the wave number in cm^{-1} , `IRIntensity`, `RamanIntensity`, and `INSIntensity` are the IR, Raman, and inelastic neutron scattering intensities in per cent, and `symmetry` is a label describing the symmetry of the mode.

`title([title])`

Sets or returns the title of a molecule. `title` has to be a string. The maximum length of the title is limited to 255 characters.

`bondAverage(atom)`

Returns the average of the lengths of all bonds involving atom `atom` in Ångstrøms. `atom` is an atom object.

`bondLength(atom1, atom2 [, length, unit])`

Returns or sets the length of the bond between atoms `atom1` and `atom2`. `atom1` and `atom2` have to be atom objects, the bond length is returned in Ångstrøms. If `length` and `unit` are given, the bond length is set. `length` is a double, `unit` a string containing either `Ang`, `au` or `bohr`, or `pm` for Ångstrøms, atomic units, or picometers. Everything else is interpreted as Ångstrøms.

`bondAngle(atom1, atom2, atom3 [, angle])`

Returns or sets the bond angle `atom1–atom2–atom3`. `atom1`, `atom2`, `atom3` are atom objects. If `angle` is given the bond angle is set. `angle` has to be a double and is measured in degrees.

`torsionAngle(atom1, atoms2, atom3, atom4 [, torsionAngle])`

Returns or sets the torsion angle `atom1–atom2–atom3–atom4`. `atom1`, `atom2`, `atom3`, `atom4` are atom objects. If `torsionAngle` is given the torsion angle is set. `torsionAngle` has to be a double and is measured in degrees.

`getThermodynamics(property, type)`

Returns a thermodynamical property of the molecule. `property` and `type` are integers. `property` can be one of the integer constants `ENTHALPY`, `ENTROPY`, `GIBBS_ENERGY`, or `HEAT_CAPACITY` defined in the `molecule` module. `type` can be one of the integer constants `TRANSLATION`, `PV`, `ROTATION`, `VIBRATION`, or `TOTAL` also defined in the `molecule` module. The returned thermodynamic property will be in SI units.

`reaction([side])`

Sets or returns whether the molecule is a reactant or a product in a reaction. `side` is an integer and can be set to one of the integer constants `REACTANT`, `PRODUCT`, or `ALLREACTIONS` defined in the `molecule` module.

`showElectrons(type [, gridResolution, interpolation])`

Displays wave function related properties of the molecule. `type` is an integer and can be set to one of the integer constants `BASIS_FUNCTION`, `BASIS_IN_MO`, `MOLECULAR_ORBITAL`, or `DENSITY` defined in the `molecule` module. `gridResolution` is a double specifying the resolution of the grid used to calculate the isosurface. Larger values for `gridResolution` result in smoother displays. `interpolation` is an optional integer and can be one of the integer constants `IP_NONE` (no interpolation), `IP_LINEAR` (linear interpolation), or `IP_LOG` (logarithmic interpolation) defined in the `molecule` module.

`selectBasisfunction(atom, name, count)`

Selects a basis function for display. `atom` is an atom object specifying which atom the basis function belongs to. `name` is a string specifying what kind of basis function (s, p, d, etc.) to select. `count` is an integer specifying which of the s, p, d, etc. functions to select. Assume atom 1 is a carbon atom

in a calculation using a DZVP basis set. It therefore has three s functions. `selectBasisfunction(1, "s", 1)` would select the 1s function, `selectBasisfunction(1, "s", 2)` the first 2s function, and `selectBasisfunction(1, "s", 3)` the second 2s function.

`unitCell(visible [, afac, bfac, cfac])`

Sets visibility and number of replicas of unit cell. *visible* has to be one of the integer constants ON or OFF defined in the `vewmol` module to turn display of the unit cell on or off. *afac*, *bfac*, and *cfac* are doubles specifying the number of replicas of the unit cell to be displayed along the a, b, and c axis, respectively. Fractions are allowed for *afac*, *bfac*, and *cfac*.

`millerPlane(visible [, h, k, l])`

Sets visibility and orientation of Miller plane. *visible* has to be one of the integer constants ON or OFF defined in the `vewmol` module to turn display of a Miller plane on or off. *h*, *k*, and *l* are integers specifying the Miller indices of the plane to display.

`addAtom(symbol [, attach])`

Adds an atom to the molecule. *symbol* is a string containing the element symbol of the atom to add. *attach* is an atom object specifying the atom the newly added atom should be attached to. *attach* can be omitted, but this is only useful for adding the first atom to a molecule.

9.8 The spectrum module

`show()`

Displays the spectrum window for a molecule.

`mode([mode])`

Displays the mode specified or returns the mode displayed. *mode* has to be an integer between 0 and the maximum number of vibrational modes for this molecule.

`deselect()`

Deselects modes previously selected with a call to `mode`.

`type([type])`

Sets or returns the type of the spectrum displayed. *type* has to be one of the integer constants `SPECTRUM_ALL`, `SPECTRUM_IR`, `SPECTRUM_RAMAN`, or `SPECTRUM_INS` defined in the `spectrum` module to set the display to all modes, IR active modes, Raman active modes, or inelastic neutron scattering display.

`display([type])`

Sets or returns the way normal modes are displayed. *type* has to be one of the integer constants `ANIMATE`, `ARROWS`, or `DISTORT` defined in the `spectrum` module to display normal modes animated, with arrows, or as distortion.

`style([style])`

Sets or returns the display style of the spectrum. *style* has to be one of the integer constants `LINES` or `GAUSSIANS` defined in the `spectrum` module to set the display style to line spectrum or gaussian spectrum.

`amplitude([amplitude])`

Sets or returns the amplitude of a normal mode. *amplitude* has to be a double.

`scaleFactor([factor])`

Sets or returns the scale factor for the wave numbers in a spectrum. *factor* has to be a double.

`zoom(x1, y1, x2, y2)`

Sets the zoom of the spectrum. *x1*, *y1*, *x2*, and *y2* are doubles specifying the minimum and maximum values of wave numbers and intensities, respectively, to be displayed. *x1* and *x2* as well as *y1* and *y2* cannot be equal.

`saveDrawing(format, filename)`

Saves the spectrum to file. *format* has to be one of the constants TIFF, HPGL, or POSTSCRIPT defined in the `vewmol` module, *filename* the name of the file the drawing is to be saved to.

The spectrum window can be closed by deleting the spectrum object.

9.9 The `vewmol` module

`load(filename)`

Loads a molecule into VIEWMOL. *filename* has to be a string containing the name (and path if necessary) of the file to load.

`save(molecule, filename, format)`

Saves molecule *molecule* in the format *format* to file *filename*. *molecule* has to be a molecule object, *filename* a string giving the name of the file (including path, if appropriate) the molecule is to be saved to, and *format* a string describing the format in which the molecule is to be saved. *format* can be any of the strings given after the `output` keyword in `vewmolrc` (currently car, arc, mol, or tm).

`delete(molecule)`

Deletes molecule *molecule*. *molecule* has to be a molecule object. Note: Molecules cannot be deleted using Python's `del` operator since Python works on VIEWMOL's data structures.

`getMolecules()`

Returns a list of the molecules loaded into VIEWMOL.

`getLights()`

Returns a list of the available lights.

`getLabels()`

Returns a list of all labels known to VIEWMOL.

`model([model])`

Sets or returns the model used to display molecules. *model* has to be one of the integer constants WIREMODEL, STICKMODEL, BALLMODEL, or CPKMODEL defined in the `vewmol` module to set the model to wire model, stick model, ball-and-stick model, and CPK model, respectively.

`drawingMode([mode])`

Sets or returns the drawing mode for molecules. *mode* has to be one of the integer constants DOT, LINE, or SURFACE defined in the `vewmol` module to set the drawing mode correspondingly.

`projection([projection])`

Sets or returns the projection. *projection* has to be one of the integer constants ORTHO or PERSPECTIVE defined in the `vewmol` module.

`sphereResolution([resolution])`

Sets or returns the resolution for spheres and cylinders. *resolution* has to be an integer. Higher resolutions result in smoother looking spheres and cylinders.

`lineWidth([width])`

Sets or returns the line width for wire model displays. *width* has to be an integer. If *width* is set to zero the line width is calculated based on the size of the window.

`groundColor([red, green, blue])`

Sets or returns the color of the ground displayed if the projection is set to PERSPECTIVE. *red*, *green*, and *blue* are floats specifying the red, green, and blue components of the ground color. They have to be between 0.0 and 1.0. If the ground color is retrieved, a tuple with the red, green, and blue values is returned.

`backgroundColor([red, green, blue])`

Sets or returns the color of the background. *red*, *green*, and *blue* are floats specifying the red, green, and blue components of the background color. They have to be between 0.0 and 1.0. If the background color is retrieved, a tuple with the red, green, and blue values is returned.

`labelAtoms(status)`

Specifies whether atoms should be labeled. *status* is an integer set to one of the constants ON or OFF defined in the `vewmol` module.

`saveDrawing(format, filename)`

Saves the drawing to file. *format* has to be one of the constants TIFF, HPGL, POSTSCRIPT, or RAYSHADE defined in the `vewmol` module, *filename* the name of the file the drawing is to be saved to.

`isosurface([level])`

Sets or returns which isosurface to display for wave function related drawings. *level* has to be a double.

`showThermodynamics([select])`

Displays the thermodynamics dialog. *select* is an integer specifying which tab to display. This integer has to be either one of the integer constants REACTION or OFF defined in the `vewmol` module or an integer between 1 and the number of molecule loaded. In the first case the reaction page is displayed, in the second case the thermodynamics dialog is closed, and in all other cases the page for the corresponding molecule is shown.

`redraw()`

Redraws the main window of VIEWMOL. Redraws are necessary to make changes visible performed using other methods of the `vewmol` module.

`getFramesPerSecond()`

Returns the drawing speed of the last redraw of VIEWMOL's main window in frames per second.

`quit()`

Quits VIEWMOL.

10 Adapting the Program to a Different Language

VIEWMOL has been written to take full advantage of the language independence of X windows. All program messages, menus, dialog box texts etc. are stored outside of the program in resource files. Therefore it is possible that different users can run the same VIEWMOL executable in different languages on the same

computer. Currently, five languages are supported: English, French⁴, German, Spanish⁵, and Russian⁶. To adapt VIEWMOL to another language only the `Xdefaults.<language>` file has to be translated. For a native speaker of the language this will take between 45 and 60 minutes.

The `Xdefaults.<language>` files provided with VIEWMOL have three sections. The first section is related to the installation of external support program, the second section contains default settings (see p. 43). The third section contains all language dependent messages and this is the only section which needs translation. To translate, translate everything right of the colon. The strings '%s', '%d', or '%f' mark the position of names or numbers which are filled in by the program and must remain in the translated version at the appropriate position. After translation install the new `Xdefaults.<language>` file as described in the Installation section (cf. p. 2) and VIEWMOL will talk in your language. The author would appreciate to get a copy of the translated resource file for inclusion in the next public release.

11 The making of multimedia files

If normal modes are animated and the user selects `Save drawing/TIFF` from the main window menu a series of TIFF files is written out, one for each frame of the animation (currently 20 frames which cannot be changed by the user). These TIFF files can easily be converted to a video file (MPEG) showing the animation using standard image manipulation tools from the Internet. One possible MPEG encoder is `mpeg_encode` which is available from `ftp://mm-ftp.CS.Berkeley.EDU` via anonymous ftp. This encoder expects its input files either in PPM, PNM, or YUV format. To convert the TIFF files written by VIEWMOL you can use the `PBMPLUS` or `NETPBM` libraries which have a filter `tiff2pnm` (you also need `pnmflip`, since `tiff2pnm` changes the orientation of the picture). The following shell script will do the conversion (for sh and ksh users) if the default files from VIEWMOL have been used:

```
for i in vm_image*.tiff
do
  j=`basename $i tiff`pnm
  tiff2pnm $i | pnmflip -topbottom > $j
done
```

The resulting PNM files can then be processed by `mpeg_encode` to produce a MPEG file which can, e. g., be included into a World Wide Web document.

Selecting `Save drawing/Rayshade` from the main window menu with an animation running will write a series of 20 input files for RAYSHADE. These files can also be processed by RAYSHADE and used to generate a movie of the vibration. This process can, however, be very time consuming.

12 Data files

VIEWMOL uses a data file named `viewmolrc` for getting informations about atoms and available input and output filters. There may be three of these files. VIEWMOL looks at first in the current directory for

⁴Many thanks to Ludovic Douillard (douillard@DRECAM.cea.fr).

⁵Many thanks to Jose R. Valverde (jrvvalverde@cnb.uam.es).

⁶My Russian is a little bit rusty. Apologies for any grammar mistakes and/or unrecognizable meanings. I would appreciate a check by a native speaker.

this file, then in the users HOME directory for a file `.viewmolrc` and finally in the directory where the environment variable `VIEWMOLPATH` points to. In one of these three locations such a file must be found. The file should contain the following data:

- Lines of the format:

```
option <name of option> <name of input filter> [<command line options for input filter>]  
" <characteristic string>"
```

These lines define the input filter for VIEWMOL and the command line options connected with them (i. e. you can change the command line option if you want). `<option>` is the command line option VIEWMOL expects on its command line or the word `default`. The input filter connected with `default` is used if no command line option is passed to VIEWMOL. If no default input filter is specified VIEWMOL displays a file selection box in this case. `<name of input filter>` is the path to and name of the input filter executable. If the input filter requires command line option (e. g. a file name) they can be specified after the name of the input filter. `%s` is used as a placeholder for file names. The path or name of the input filter can contain environment variables or the string `$OSNAME`. The latter is replaced by the subdirectory name for the machine VIEWMOL is currently running on. `" <characteristic string>"` is a string which is used to identify the type of a particular file. The first 1024 characters of an input file passed to VIEWMOL are scanned for this string and the input filter connected with the string is then used to read the file. Therefore these strings have to be unique for each input filter and have to be in every file of a certain type within the first 1024 characters. Since most programs write their names out at the beginning these restrictions seem to be no problem.

- Lines of the format:

```
output <reference to resources> <name of output filter> [<command line options for output filter>]  
%s
```

These lines define the output filter for VIEWMOL. `<reference to resources>` is an arbitrary string which must refer to a resource in the `Xdefaults` file. This string is used to provide the label for the output filter in the output filter selection box. `<name of output filter>` is the path to and name of the output filter executable. All output filters should at least accept the name of the output file from their command lines. If additional parameters are required they can also be specified after the name of the output filter. `%s` is used as a placeholder for the output file name. Environment variables or the string `$OSNAME` can be used in the same way as for input filters.

- Lines of the format:

```
<symbol> <rad> <rd> <gd> <bd> <rl> <gl> <bl> <surface>
```

`<symbol>` is an atomic symbol, `<rad>` is the Van der Waals radius of this atom in Ångstrøms, `<rd>`, `<gd>` and `<bd>` are the red, green and blue color for the darkest part of this atom and `<rl>`, `<gl>` and `<bl>` are the red, green and blue color for the lightest part of the atom. There are four reserved strings for `<symbol>`. If `<symbol>` is `bd` the following description describes a hydrogen bond. The `<rad>` field is also interpreted as the radius of all bond sticks. All other fields are only applied to hydrogen bonds. `uc` specifies a unit cell corner. Radius and color given here affect the appearance of the unit cell. `ps` and `ms` specify the surface properties for the positive and negative isosurface, respectively, used to draw wave function related topics. In these cases the radius is not used. `<surface>` is an optional specification for the surface used when stick, ball, or CPK drawing with surfaces is activated. `<surface>` is a list of one or more of the following options

- `emission <r> <g> `

The emission color of the surface. Using this option causes the surface to emit light. `<r>`, `<g>`, and `` are the red, green and blue components for the light color.

- ambient <r> <g>
The ambient light which is reflected by the surface. <r>, <g>, and are the red, green and blue components for the light color.
- specular <r> <g>
The specular light which is reflected by the surface. <r>, <g>, and are the red, green and blue components for the light color.
- shininess <n>
A parameter which determines the kind of reflection. <n> can be in the range 0 ... 128.
- alpha <n>
This parameter determines the transparency of the surface.

All color specifications can be between 0.0 and 1.0. The total length of a line specifying an atom is restricted to 132 characters. The keywords for the surface specifications can be abbreviated with the first two letters.

Any line starting with '#' is treated as a comment.

VIEWMOL makes extensive use of X Windows resources. All standard search algorithms for the location of the resources apply (see e. g. O'Reilly books on X Windows). VIEWMOL has English resources compiled in. Resources for other languages are provided in files `Xdefaults.<language>` and might be installed as described in the installation section of this manual (p. 2).

The following resources are used to specify the defaults. They can be overwritten in the user's `$HOME/.Xdefaults` file or, in part, by the configuration options available in the program. Defaults configurable from within the program are marked with an asterisk (*).

<code>Viewmol*geometry:</code>	500x500+50+50 (*)
<code>Viewmol.history.geometry:</code>	500x250+50+590 (*)
<code>Viewmol.spectrum.geometry:</code>	500x250+50+590 (*)
<code>Viewmol.MODiagram.geometry:</code>	250x500+565+50 (*)
<code>Viewmol.Bell:</code>	<no default>
<code>Viewmol.model:</code>	wire (*)
<code>Viewmol.drawingMode:</code>	surface (*)
<code>Viewmol.bondType:</code>	conjugated (*)
<code>Viewmol.sphereResolution:</code>	10 (*)
<code>Viewmol.simplifyWhileRotating:</code>	True (*)
<code>Viewmol.interpolation:</code>	linear (*)
<code>Viewmol.bondLength:</code>	%7.4f Ang
<code>Viewmol.bondAngle:</code>	%7.2f deg
<code>Viewmol.torsionAngle:</code>	%7.2f deg
<code>Viewmol.wavenumbers:</code>	0:5000
<code>Viewmol.isosurface:</code>	0.05 (*)
<code>Viewmol.reservedColors:</code>	0
<code>Viewmol*spectrumForm*amplitudeSlider.decimalPoints:</code>	2
<code>Viewmol*spectrumForm*amplitudeSlider.minimum:</code>	-250
<code>Viewmol*spectrumForm*amplitudeSlider.maximum:</code>	250
<code>Viewmol*spectrumForm*scaleSlider.decimalPoints:</code>	2
<code>Viewmol*spectrumForm*scaleSlider.minimum:</code>	50

Viewmol*spectrumForm*scaleSlider.maximum:	150
Viewmol*wavefunctionForm*level.minimum:	1
Viewmol*wavefunctionForm*level.maximum:	100
Viewmol*wavefunctionForm*grid.minimum:	4
Viewmol*wavefunctionForm*grid.maximum:	20
Viewmol*wavefunctionForm*grid.value:	10
Viewmol*MODiagramForm*resolution.minimum:	1
Viewmol*MODiagramForm*resolution.maximum:	1000
Viewmol*MODiagramForm*resolution.decimalPoints:	3
Viewmol*MODiagramForm*resolution.value:	10
Viewmol.paperSize:	A4 (*)
Viewmol.viewer*font:	variable
Viewmol.spectrum*font:	variable
Viewmol.history*font:	variable
Viewmol.MODiagram*font:	variable
Viewmol.viewer.background:	white (*)
Viewmol.viewer.foreground:	gray75 (*)
Viewmol*spectrum.spectrum.background:	white (*)
Viewmol*spectrum.spectrum.foreground:	black (*)
Viewmol*history.history.background:	white (*)
Viewmol*history.history.foreground:	blue (*)
Viewmol*MODiagram.MODiagram.background:	white (*)
Viewmol*MODiagram.MODiagram.foreground:	black (*)
Viewmol*foreground:	black (*)

The Viewmol.Bell resource is the only resource which does not have a default. As long as this resource is not set the standard keyboard bell is rung as soon as a selection in one of the windows is made by mouse click. This resource can be set to the name (and command line parameters) of any program which shall be run instead, preferably one which produces a nicer sound effect.

The Viewmol.model resource can be set to wire, stick, ball, or cpk. The Viewmol.drawingMode resource can be set to dot, line, or surface. The Viewmol.bondType resource can be set to single, multiple, or conjugated. The Viewmol.interpolation resource can be set to none, linear, or logarithmic. The resources for specifying formats for bond lengths, bond angles, and torsion angles have to contain a valid C format string for printing a floating point number. The resource for the bond lengths recognizes Ang, pm, bohr, and au in the format string as units and converts the bond lengths accordingly. The Viewmol.reserved-Colors resource can be used to limited the number of colors allocated by VIEWMOL if it runs in colormap mode. VIEWMOL tries to allocate as much colors as it can. This might interfere with others program. In this case Viewmol.reserved-Colors can be used to tell VIEWMOL to leave the specified number of colors unallocated. In case of the specifications for the sliders the values given for minimum and maximum have to be multiplied by $10^{decimalPoints}$. I. e. if the number of decimals is to be changed also minimum and maximum have to be changed. Paper sizes currently recognized are A5, A4, A3, Letter, Legal, and <width> x <height> where <width> and <height> are in millimeters.

The Viewmol.viewer.foreground resource is used for the color of the ground if perspective drawing is enabled.

13 Programming Your Own Input Filter

VIEWMOL can be easily adapted to read outputs of other programs or other file formats. All you have to do is to write a new input filter which extracts the data from the corresponding file. These input filter are stand-alone programs and can be written in every programming language you want. Examples in C and awk are included.

The input filter has to read the following data from the output file and write them to its standard output in the format described below. This format follows the file format of TURBOMOLE very closely. A few sections had to be extended to allow data which are currently not supported by TURBOMOLE (e. g. unit cells).

- the Cartesian coordinates and atom symbols (required)

Write to standard output in the following format:

```
$coord factor
  x1    y1    z1    symbol1    xyz
  x2    y2    z2    symbol2    xyz
  ...
  ...
```

factor is the conversion factor the coordinates have to be multiplied with to convert them to Ångstrøms. Any combination of x, y, and z at the end of the line (optional) indicates that the corresponding atom has been kept fixed in that direction during a geometry optimization. Consequently, VIEWMOL will not draw the forces acting on this atom in the fixed direction.

- the title (optional)

Write to standard output in the following format:

```
$title
title
```

- the wave numbers and intensities (optional)

Write to standard output in the following format:

```
$vibrational spectrum
  symmetry1  wavenumber1  IR-intensity1  Raman-intensity1
  symmetry2  wavenumber2  IR-intensity2  Raman-intensity2
  ...
  ...
```

symmetry is the symmetry label for the vibrational mode, wavenumber is its wave number and IR-intensity and Raman-intensity are its IR and Raman intensity, respectively. If the symmetry labels for the vibrational modes are unknown they should be set to a default (e. g. A1).

- normal coordinates (optional)

Write to standard output in the following format:

```
$vibrational normal modes
  i1  i2  nm(1,1)  nm(2,1)  nm(3,1)  nm(4,1)  nm(5,1)
  i1  i2  nm(6,1)  ...       nm(3*natom,1)
  i1  i2  nm(1,2)  nm(2,2)  nm(3,2)  nm(4,2)  nm(5,2)
```

```

i1  i2  nm(6,2) ...      nm(3*natom,2)
...
i1  i2  nm(1,nmodes) ...      nm(5,nmodes)
i1  i2  nm(6,nmodes) ...  nm(3*natom,nmodes)

```

i1 and *i2* are integers which are skipped during reading. *nm(i, j)* are the normal mode coefficients. They have to be provided ordered by Cartesian coordinates (all x components of the first atom first, then all y components of the first atom etc.).

- optimization history or MD trajectory (optional)

Write to standard output in the following format:

```

$grad          cartesian gradients
cycle = nc SCF energy = E_nc |dE/dxyz| = gradnorm_nc
[unitcell a b c alpha beta gamma]
x1  y1  z1  symbol1
x2  y2  z2  symbol2
...
xn  yn  zn  symboln
gx1 gy1 gz1
gx2 gy2 gz2
...
gxn gyn gzn
cycle = nc+1 SCF energy = E_nc+1 |dE/dxyz| = gradnorm_nc+1
...

```

nc is a counter for the cycle, *E_nc* is the energy for the configuration of cycle *nc*, and *gradnorm_nc* is the gradient norm of cycle *nc*. The line starting with *unitcell* is optional and can be used to specify the current unit cell e. g. during a constant pressure MD run. The *x*, *y*, and *z* are the Cartesian coordinates for each atom, *symbol* is the atomic symbol. The *gx*, *gy*, and *gz* are the gradients for each atom. This structure can be repeated for as many cycles as necessary.

- MO energies and coefficients (optional)

Write to standard output in the following format:

```

$scfmo [symmetrized] [gaussian]
n symmetry_label_n eigenvalue=MO_E_n nsaos=norb
moc(n,1) moc(n,2) moc(n,3) moc(n,4)
moc(n,5) ...      moc(n,norb)
n+1 symmetry_label_n+1 eigenvalue=MO_E_n+1 nsaos=norb
...

```

The string *symmetrized* after *\$scfmo* is optional and can be used to notify *VIEWMOL* of the fact that the MO coefficients are with respect to symmetrized AOs rather than with respect to AOs. *VIEWMOL* needs *moloch* from the *TURBOMOLE* package to handle symmetrized AOs. If *moloch* is not installed and symmetrized AOs are input, MOs and electron densities cannot be drawn. The string *gaussian* is also optional and notifies *VIEWMOL* that the MO coefficients are normalized and ordered *GAUSSIAN* style. *n* is a counter counting the MOs, *symmetry_label_n* is the symmetry label for MO *n*, *MO_E_n* is the MO energy for MO *n*, and *norb* is the total number of orbitals. The *moc(n,i)* are the MO coefficients for MO *n*.

- basis functions and occupation numbers (optional)

Write to standard output in the following format:

```
$atoms
atom_symbol1 list_of_indices1 \
  basis=basis_set_name1
atom_symbol2 list_of_indices2 \
  basis=basis_set_name2
...
$basis
*
basis_set_name1
*
  number_of_primitives  angular_momentum
  exponent1  coefficient1
  exponent2  coefficient2
  ...
  exponentn  coefficientn
  number_of_primitives  angular_momentum
  ...
*
basis_set_name2
*
  ...
*
$closed_shells
  symmetry_label      list_of_indices    (2)
$pople   [6d/10f/15g]
```

`atom_symbol` is the atom symbol of an element and `list_of_indices` contains the indices of all atoms of the particular element according to the list of coordinates read in under `$coord`. The list can be either comma separated and/or contain hyphens for indicating ranges (e. g. c 1,3,7-10 is a valid descriptor). `Basis_set_name` can be an arbitrary string describing a particular basis set. It is only used to find the corresponding basis set in the list read under `basis`. This list simply states the name for a basis set and then lists the primitive functions which make up a contracted Gaussians starting with the number of primitives in that particular contracted Gaussian and its angular momentum (s, p, d, f, ...). Than the exponents and contraction coefficients are listed line by line. This is repeated for all contracted Gaussians of that particular basis set. `$closed_shells` is used to tell VIEWSMOL which MOs are occupied and which are not. `symmetry_label` is the symmetry label for a number of MOs and `list_of_indices` is a list of integers stating which of the MOs of that particular symmetry are occupied. This list can be either comma-separated or contain hyphens to indicate ranges of MOs. **Note:** `$closed_shells` has to appear after `$scfmo` in the output written by the input filter. `$pople` is used to indicate that d, f, or g functions have 6, 10, or 15 components instead of 5, 7, or 9. **Note:** This data group has to appear after the `$coord` or `$grad` in the output. Otherwise VIEWSMOL will fail.

- the unit cell (optional)

Write to standard output in the following form:

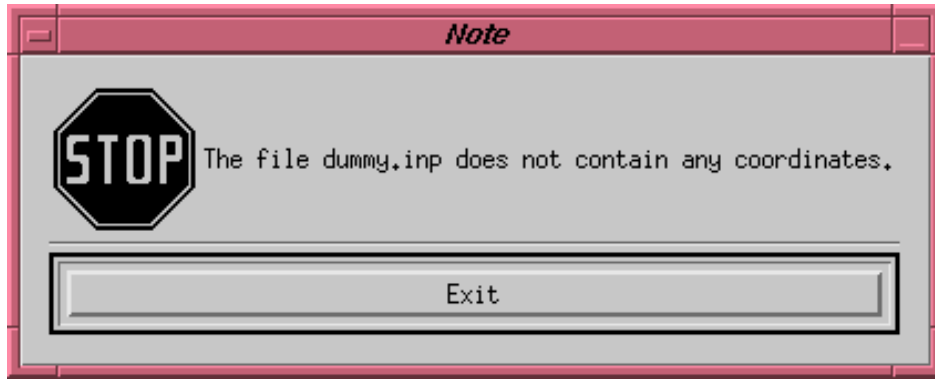


Figure 15: The error dialog produced by the sample error message

```
$unitcell      a      b      c      alpha     beta     gamma
```

- errors occurring during file processing (optional)
Write to standard output in the following form:

```
$error errorLabel severity additionalInformation
```

`errorLabel` is an arbitrary one word label which refers to an error message in the resources. `severity` is a label for the severity of the error. Set it to 0 if the program can continue despite this error. Set it to 1 if the program must stop. `additionalInformation` is any additional information you want to be displayed in the error message (e. g. the name of a file which was not found). Currently, the following `errorLabels` are in use: `noFile`, `notConverged`, `unsupportedVersion`, `wrongFiletype`, `noCoordinates`, `noEnergy`, and `unknownErrorMessage`. If your input filter wants to return an error because it is missing coordinates in the input file "dummy.inp" you can have it writing the following line to standard output:

```
$error missingCoordinates 1 dummy.inp
```

Then you have to specify a resource for the error message in `$HOME/.Xdefaults`:

```
Viewmol.missingCoordinates: The file %s does not  
contain any coordinates.
```

With these two lines in place any encounter of no coordinates in an input file will lead to the display of the error dialog in Figure 15. There is no need to recompile VIEWMOL to achieve this.

The last line of the data written to standard output by the input filter must be `$end`.

The input filter can be installed by adding a line to the `viewmolrc` file.

14 Programming Your Own Output Filter

VIEWMOL can be easily adapted to write files in any format. All you have to do is to write a new output filter which formats the data provided by VIEWMOL. These output filters are stand-alone programs and can be written in every programming language you want. Examples in awk are included.

The output filter has to accept the following data from its standard input and write them to a file whose name is given as a command line parameter to the filter. VIEWMOL passes the following data groups to the output filter:

- the unit cell (if present) is sent in the following format (a, b, and c in atomic units, the angle in degrees)

```
$unitcell a b c alpha beta gamma
```

- the Cartesian coordinates are sent in the following format (in atomic units)

```
$coord
  x1    y1    z1    symbol1
  x2    y2    z2    symbol2
  ...
  ...
```

- the bond information is sent in the following format

```
$bonds
  atom1 atom2 bond_order
  ...
  ...
```

where `atom1` and `atom2` are the numbers of the atoms according to the list in `$coord` which form the bond. `bond_order` is the actual order of the bond, `-2` if the bond is part of a conjugated system, or `-1` if it is a hydrogen bond.

`$end` is passed to the output filter as last line.

15 Limitations

VIEWMOL currently cannot handle GAUSSIAN outputs which contain Cartesian f functions (10f).

If VIEWMOL runs in color map (that should only happen on IBM RS6000 with Sabine graphics adapters, in this case the background of the title screen has a constant color instead of the usual dark top and lighter bottom) shadows are not drawn if the drawing mode is "with surface".

If TIFF files are saved make sure no other window (including dialog boxes) overlaps with the window to be saved. The information is read from the screen and overlapping windows will show up in the saved file.

HPGL outputs for drawings which are labeled with non-latin characters will not contain any labels. HPGL output has only support for German umlauts, Postscript provides all ISO-8859-1 and KOI-8 characters.

If VIEWMOL has been linked with a Mesa version < 2.1 and is running on Linux the "Lines while rotating" button should always be set to on (the default). If it is set to off, a model is drawn with surfaces, and tried to rotate, VIEWMOL will crash with a floating point exception. This is a bug in early Mesa versions.

VIEWMOL is now usable with Lesstif (> 0.81). There are, however, some glitches, e. g. shortcuts don't work.

16 Frequently asked questions

1. Is there a home page for VIEWMOL ?

There is a home page at <http://viewmol.sourceforge.net/> and a project page at <http://sourceforge.net/projects/viewmol/>.

2. VIEWMOL on Linux reports on start up:

`viewmol: can't load library 'libMesaGLU.so.3'`
or another library.

VIEWMOL does not find a dynamical linked library it needs. The reason for this is that either the library is not installed, the wrong version is installed, or the dynamic linker is not set up to find this library. VIEWMOL needs the following dynamic libraries:

```
libtiff.so.3
libGLU.so.1
libGL.so.1
libXm.so.2
libXp.so.6
libXi.so.6
libXext.so.6
libXt.so.6
libX11.so.6
libpthread.so.0
libdb.so.3
libutil.so.1
libdl.so.2
libm.so.6
libc.so.6
libjpeg.so.62
libz.so.1
libSM.so.6
libICE.so.6
libXmu.so.6
/lib/ld-linux.so.2
```

These libraries can normally be found in `/lib`, `/usr/lib`, and `/usr/X11R6/lib`. The dynamic linker checks the major version number and will refuse any library where the major version number does not match. The minor version number does not matter. The dynamic linker has to be set up to search the directories which contain these libraries. This is done in the file `/etc/ld.so.conf`. After modifying this file run `ldconfig -v` as root. Alternatively, the environment variable `LD_LIBRARY_PATH` can be set to point to these directories.

3. After the splash screen appears on Linux VIEWMOL stops with a segmentation fault
Segmentation faults occur reproducibly for certain operations on Linux

There is a compatibility problem with a library, usually libc. Unfortunately, there are so many different library versions in use among the different distributions, that there is a good chance to run into problems. If it is impossible to use the exact same distribution VIEWMOL was compiled on (see README file), recompilation usually helps.

4. VIEWMOL on Linux reports on start up:

```
viewmol: Symbol 'jpeg_resync_to_restart' is not defined.
```

There are two different versions of `libtiff.so` distributed with different Linux distributions. One contains jpeg code (Debian) the other doesn't (RedHat). VIEWMOL has now been linked with the version which does not contain jpeg code so that this error will probably not occur anymore. If this error occurs only a recompilation will help. Please notify the maker of your Linux distribution so that they can make their distribution compatible.

5. When I try to recompile VIEWMOL I get a lot of error messages:

```
cc -c -Wall -DLINUX -I/usr/compat/linux/usr/include/GL/
-I/usr/compat/linux/usr/include/gr/ -O6 -m486
-fomit-frame-pointer
./annotate.c
./annotate.c:19: X11/StringDefs.h: No such file or directory
./annotate.c:20: X11/cursorfont.h: No such file or directory
./annotate.c:21: Xm/Xm.h: No such file or directory
./annotate.c:22: Xm/Text.h: No such file or directory
In file included from ./annotate.c:24:
./viewmol.h:20: X11/Intrinsic.h: No such file or directory
./viewmol.h:21: GL/gl.h: No such file or directory
./viewmol.h:22: GL/glx.h: No such file or directory
*** Error code 1
```

To recompile VIEWMOL you need to install X windows and OpenGL development environments which, in most Linux distributions, are separate packages. In this case you are missing all X windows and OpenGL header files. You also need the development environment of Lesstif or Motif for all header files in `/usr/include/Xm`.

6. While trying to recompile VIEWMOL the link step fails with:

```
ld:
Unresolved: __eprintf

*** Error code 1 (bu21)
```

or a similar message referring to `__eprintf` (encountered on SGIs and IBMs so far only). There is a problem with the TIFF library you are linking with. If you have built the library yourself make sure it was built on the same machine as where you try to link VIEWMOL. If you have been trying to use a vendor supplied version of the TIFF library try to download and compile the library yourself.

17 History, Authors, and Contributors

VIEWMOL started its life somewhere in 1991 as a tool to draw IR and Raman spectra from Turbomole outputs. Since drawing only spectra soon turned out to be insufficient for writing a PhD thesis, capabilities were added to draw the molecule and animate normal modes. Since other people in the Arbeitsgruppe Quantenchemie an der Humboldt-Universität zu Berlin, Max-Planck-Gesellschaft got interested in the program and wanted extensions for other program's output Andreas Bünger (a then 16 year old high school student on a practical course in the group) and Andreas Bleiber started to write an input filter for GAUSSIAN 9X. Arne Dummer wrote a filter for DMOL. In the course of the research performed in the group other capabilities were asked for and added by the original authors and by Mariann Krossner (calculation of inelastic neutron scattering intensities) and Andries de Man (extension of GAUSSIAN 9X input filter to read density functional outputs). Version 1.2 was presented at the German/Austrian Academic Software Award competition in 1993 and honored as outstanding achievement (cf. <http://www.ask.uni-karlsruhe.de/asksam/sampages/htmltxt/viewmol.html>). With the advent of Linux it was recognized that the original Fortran/IrisGL version would be difficult to port to more affordable hardware. Version 2.0 of the program was a complete rewrite in C/OpenGL by Jörg-Rüdiger Hill now mainly done on a Linux system. Development of the program continues on Linux using Brian Paul's terrific implementation of OpenGL, Mesa.

Contributions, mainly in form of bug reports, code snippets, and enhancement requests have come from a number of people. In no particular order (and hopefully without forgetting somebody) I want to thank:

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- French: Ludovic Douillard (douillard@DRECAM.cea.fr)
- Spanish: Jose R. Valverde (jrvalverde@cnb.uam.es)

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