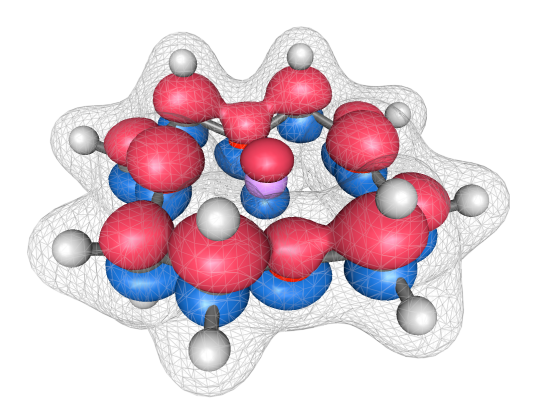
Introduction to IQmol



v
2.13 (2019) Andrew Gilbert

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1 Introduction

IQMOL is an open-source molecular editor and visualization package that runs under Windows, Mac OS X and Linux. It can read a variety of chemical file formats including xyz, cml, pdb, mol, fchk, cube data and Q-CHEM input/output. It also includes a freeform molecular builder that allows arbitrary molecular structures to be created. These structures can be optimized using molecular mechanics force fields and symmetrized to ensure the structure has the correct point group symmetry. A library of molecules and functional groups also exists, and these can be used to facilitate building more complicated molecules.

IQMOL is capable of displaying a variety of molecular properties including atomic charges, dipole moments and normal modes. Several surface types can be displayed including molecular orbitals, (spin) densities and van der Waals surfaces. These surfaces can be colored according to an arbitrary scalar field, such as the electrostatic potential. Animations are also available for vibrational frequencies, and reaction and optimization pathways.

IQMOL can operate as a stand-alone package, but has also been written to work seamlessly with the Q-CHEM (http://www.q-chem.com) computational chemistry package. A comprehensive input file generator, the Q-CHEM User Interface (QUI), provides access to most of the available options in Q-CHEM, and these options are presented in an intuitive, hierarchical fashion. The generated input files can be submitted to either local or remote servers that have the Q-CHEM software installed. In particular, a publicly accessible server is available that allows small (limited to approximately 10 mins) quantum chemistry calculations to be run without having to purchase and install Q-CHEM.

1.1 Installation

The latest version of IQMOL can be downloaded from the website:

http://iqmol.org/downloads.html

For Mac (OS X) a disk image file is provided. After downloading, Simply double-click the disk image to mount it and copy the application to the Applications directory, or any other desired location.

For Windows an installer is provided that will guide you through the installation process and will also create a shortcut to the application on your desktop.

For Linux .deb and .rpm packages are now available that can be installed using either dpkg or yum:

#> sudo dpkg -i iqmol_2.9.0.deb
#> sudo apt-get install -f

The second command is required to resolve the dependency on the Qt libraries, which you may not have installed.

#> sudo yum install iqmol-2.9.9-2.x86_64.rpm

Note that in either case you will need root permission.

1.2 Overview

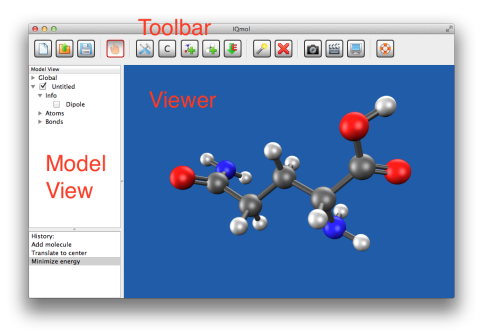


Figure 1.1: The main IQMOL window.

The main IQMOL window is shown in Fig. 1.1 and comprises the following main parts:

- The **Viewer** is the main part of the window and is where you can view and interact with your molecule.
- The **Toolbar** at the top provides access to common commands and allows selection between different viewer modes including manipulation , selection and building.
- The **Model View** panel provides a hierarchical view of the data that are available for the molecule.
- The **History** panel in the lower left shows a list of the most recent actions than can be undone, either by clicking on them or by using the Edit ► Undo menu option.

The Model View (MV) provides control over what objects are displayed in the Viewer, and also allows access to configuration options for these objects. Visibility is controlled by the associated check-box. Unchecking a check-box causes the item and all its children to become hidden. If an item does not have a check-box, for example a bond, then its visibility can only be controlled by items higher in the hierarchy.

The appearance of many objects can be configured by double-clicking the item in the MV. For example, double-clicking the molecule name brings up the Configure Molecule dialog which allows you to change the appearance of the molecular structure:

Render		
-	nd Sticks Filling	Underson Wire Frame
	: Model	Ville Flame
Flastic	woder	
Radii		
Atoms		· · · · · · · · · · · · · · · · · · ·
Bonds		0
Charges		· · · · · ·
🗹 Smalle	r Hydroger	n Atoms
	Re	set Close

Note that several molecules can be viewed concurrently in the same Viewer and the appearance of each can be configured separately.

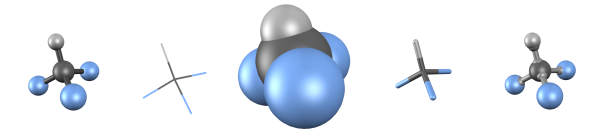


Figure 1.2: Rendering styles for molecular structures: balls and sticks, wire frame, space filling, tubes and plastic.

2 Building Molecules

2.1 Adding Atoms and Fragments

By default, IQMOL opens in build mode. This is indicated by a red border around the button in the Toolbar. The default build atom is indicated by the button and can be changed by clicking on this button. A pop-up periodic table will appear from which the desired element type can be selected.

Clicking in the empty Viewer window will create an atom of the current build element. Additional atoms can be added by clicking on an existing atom and dragging the mouse. This creates a new atom bonded to the first. To create a disconnected atom, hold down the *alt* modifier key when clicking in the Viewer window. (Note that some Linux window managers use the *alt* modifier for other purposes. This behavior can be changed using System Settings \blacktriangleright Keyboard \blacktriangleright Short-cuts menu).

Bond orders can be increased by clicking and dragging between two existing atoms. If no bond exists between the two atoms, one is created. Otherwise the bond order is increased. To decrease the bond order, the bond must first be deleted and a new bond created.



Figure 2.1: Increasing the bond order.

Functional groups can be added by clicking the \checkmark button, ensuring the Functional Group radio button is clicked and selecting the desired group from the menu. Groups are added in the same way as atoms, *i.e.* clicking and dragging from an existing atom. The empty valence is indicated by the yellow bond and shows where the group will be connected.

Entire molecules can also be added to your system by clicking the \checkmark button, ensuring the Molecules radio button is clicked, and selecting the desired molecule from the menu. Be sure to click Select or the build selection will not be updated. Unlike the other build modes, clicking anywhere in the viewer window will add the selected molecule (no mouse modifier is required). This allows several molecules of the same type to be added quickly, (e.g. for solvation), but changes the usual mouse behavior. If you accidentally add too many molecules, use the Edit \triangleright Undo menu option. When adding molecules, if you click and hold it is possible to alter the local orientation of the newly added molecule before being added to the global frame.

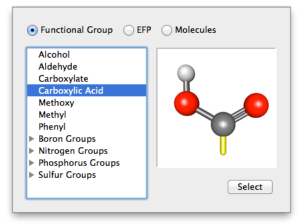


Figure 2.2: The build fragment pop-up that appears when the add fragment button is clicked.

Once the backbone of the molecule has been drawn, the Add Hydrogens button is can be clicked to automatically add hydrogen atoms to any unfilled valencies.

2.2 Optimizing Structures Using Molecular Mechanics

The builder in IQMOL is free-form, so your initial structure may look a bit wonky. To improve the geometry, click the I button. This optimizes the geometry using a molecular mechanics (MM) force field. The default force field is the Universal Force Field (UFF)[2] which has the advantage of being defined for almost all of the periodic table. However, the UFF does not perform well for systems that contain hydrogen bonds and, in these cases, it is recommended that the force field be changed using the Build \blacktriangleright Select Force Field menu option.

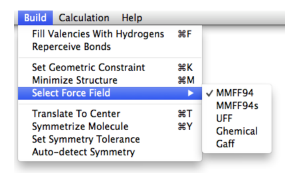


Figure 2.3: Changing the molecular mechanics force field.

2.3 Symmetrizing Structures

If your molecule has symmetry, the MM optimization is unlikely to find a structure with the desired symmetry. If this is the case, a nearly symmetric structure can be symmetrized using the Build \blacktriangleright Symmetrize Molecule menu option. Finding very high symmetry may require relaxing the tolerance using the Build \blacktriangleright Set Symmetry Tolerance menu option. Relaxing the tolerance allows the program to move the nuclear coordinates more in order to find a symmetric structure.

IQMOL uses a modified version of the SYMMOL[1] program written by Tullio Pilati and Alessandra Forni to symmetrize molecular structures.

2.4 Specifying Geometric Parameters

Specific values for geometric parameters can be set by first selecting the atoms involved and using the Build \blacktriangleright Set Geometric Constraint menu option. A dialog will appear that allows the parameter to be either set, constrained or scanned. Constrained parameters apply to any subsequent MM optimization and are also passed through to the Q-CHEM input file, if a optimization job is requested. Scan options are also passed through to Q-CHEM for scan jobs.

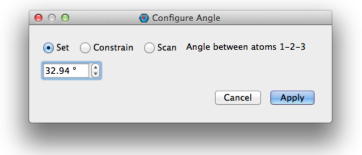


Figure 2.4: Dialog for setting geometric constraints

The type of constraint depends on the number of atoms selected:

- 1. Fixed atom position
- 2. Inter-atomic distance (or select a single bond)
- 3. Bond angle
- 4. Torsion (dihedral) angle

Active constraints are visible in the Viewer and can be deactivated by clicking the adjacent check-box in the MV.

2.5 Manipulating and Selecting Molecules

IQMOL has been designed to work best with a three-button mouse or track pad. If you are using a track pad on a Mac, it is recommended that you enable the secondary click by going to System Preferences ► Trackpad.

Manipulate mode is activated by clicking the 🛄 button in the Toolbar, and implements the following mouse functionality:

- Left click and drag: Rotate the view of the molecule.
- Middle click and drag: Zoom in and out.
- Right click and drag: Translate the view of the molecule.

It is also possible to manipulate part of the molecule independently from the rest. To do this, make a selection and press and hold the *ctrl* modifier (*command* key on Mac). The mouse movements will affect only the selected atoms as follows:

- Left click and drag: Rotate the selected atoms about their center.
- Right click and drag: Translate the selected atoms.

If only a single bond is selected, then the mouse movements have the following effects

- Left click and drag: Rotate around the axis of the bond.
- Right click and drag: Change the length of the bond.

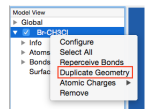
Select mode is activated by clicking the 🖉 button in the Toolbar, and implements the following mouse functionality:

- Left click: Adds atom or bond to selection.
- Click and drag: Creates a selection rectangle, all atoms and bonds within the selection rectangle are added to the selection.
- **Right click:** Removes atom or bond from selection.

It is also possible to select all, select none and invert the selection via the options under the Edit menu.

2.6 Adding Additional Conformers

For setting up jobs that use the Freezing String Method (FSM), two conformers are required corresponding to the initial and final geometries of the string. Having built the first geometry, a second geometry can be added to the molecule by right-clicking the molecule name in the MV. This brings up a context menu with the option 'Duplicate Geometry'. Selecting this menu option creates a Geometries item in the MV with two



(identical) geometries. The second of these can be modified by selecting it in the MV and manipulating selected atoms in the viewer using the *ctrl* (*command* on Mac) modifier key. If 'Freezing String' is chosen for the Calculate option in the QUI (see the next section) then both geometries will be included in the \$molecule section of the input deck, separated by ****.

3 Running Q-Chem Calculations

3.1 The QUI

IQMOL has a built-in input file generator for Q-CHEM calculations, the QUI, that can be accessed via the Calculation \blacktriangleright Q-Chem Setup menu.

		Setup	Advanced			Generated Input File:	
Job Section Calculate Method Basis Exchange	Job 1 Energy HF 6-31G HF	÷ ÷ • •	Edit Charge Multiplicity ECP Correlation		÷	\$molecule 0 0 -0.000000 H 0.000000 H 0.000000 Send \$gend \$gend \$gend	-0.1026146 0.0000000 0.4104455 -0.7920090 0.4104455 0.7920090
SCF Control							
Algorithm	DIIS	÷ (Convergence	5			
Guess	SAD	\$ N	lax Cycles	50			
Second Ba	sis None	\$	uess Mix	0%			
🗌 Dual Ba	sis Energy		Unrestricted				
							Server QChem
	n Analysis						
Wavefunctio						Reset	Cancel Subm

Figure 3.1: The Q-CHEM User Interface (QUI) dialog.

The left hand side of the QUI dialog contains controls for setting up the calculation. These are presented in a hierarchical fashion so that most commonly used options are presented in the top part of the panel, with other relevant options appearing in the lower section depending on what type of job is selected. More advanced options can be accessed via the Advanced tab. The generated input file is echoed in the panel on the right hand side of the QUI.

Clicking the Submit button will start the job running on the selected server. Once the job has completed you will be prompted to copy the results (for remote servers) before they are automatically loaded into IQMOL. The \rightleftharpoons icon appears next to the molecule name in the MV when the molecule has been updated with the results of a calculation.

	Job	Serv	er	Submit Time	Run Time	Status
26	Benzene	Quner	n	15:52:12	0:00:01	Finisnea
27	Untitled	QCher	n	16:14:12	0:00:01	Finished
28	h-atom	QCher	n	11:57:41	0:00:01	Finished
29	isoindole	QCher	n	10:01:24	0:00:36	Finished
30	cubane	QCher	n	11.22.45	0.00:01	Error
31	C3N3H3	QChe	Kill.		00:01	Finished
32	B7	QChe		iove Job	00:08	Finished
33	adamantane	QChe		ry Job	00:06	Finished
34	Raman	OChe		v Output File n Results	00-04	Finished
_	Clear List			y Results From	Server	Close

Figure 3.2: The Job Monitor dialog. Right-clicking on a selected job brings up a context menu with additional options.

3.2 The Job Monitor

Submitted jobs can be monitored via the Calculation \blacktriangleright Job Monitor menu option. This brings up the Job Monitor dialog that provides information on the progress of jobs. Right-clicking a row in the Job Monitor brings up a context menu which allows running jobs to be killed or queried, and finished jobs to be opened.

If a job has the Error status, hovering the mouse over the status will show the error message. Alternatively, double clicking the job will open the output file, if available, and may provide additional information about why the job failed. Double-clicking a completed job will copy the results from the server (if they have not already been copied) and reload them into the main IQMOL window.

3.3 Configuring Servers

By default IQMOL is configured to submit jobs to the Q-CHEM server in Pleasanton, California. This is a publicly available server than can be used by anyone wishing to run test calculations before purchasing the Q-CHEM software. Jobs submitted to this server can access the full suite of electronic structure methods available in Q-CHEM, but are time-limited to 10 minutes.

Additional servers can be configured to access other computers that have the Q-CHEM software installed. These computers can be either local servers (*i.e.* the same machine as the one running IQMOL) or remote servers that can be connected to via SSH. To add

Connection O Lo	ocal 🔵 SSH 💿 HTTP	O HTTPS Configure
Server Name QCh	em	
Queue System We	b ‡	Configure
Remote Host		
Host Address	iqmol.q-chem.com	Port 80
Authentication	None	A T
User Name	guest	
Working Directory	(unused)	
		Test Connection

Figure 3.3: The server configuration dialog.

an additional server, go to the Calculation \blacktriangleright Edit Servers menu option and click the $\textcircled{\bullet}$ button.

The information required to configure a server depends on its type, but default options exist for each case. The following indicates the minimum that should be considered:

- Local: Set the Queue System to 'Basic' unless you know you are running queuing software on the machine. Be sure the QC and QCSCRATCH variables are set to their correct values in the Run File Template, which is accessed by clicking the Configure button.
- SSH: The host name and account on the target machine will be required in order to connect via SSH. Set the Authentication combo-box to 'Password Prompt' unless you have set up an alternative authentication protocol. Depending on how the target host has been set up, the Run File Template may require some editing to get it to work, but this will differ from machine to machine.
- **HTTP:** The default options should be suitable. Note that on Windows HTTP should be used (and is the default in v2.8) whereas for Mac and Linux HTTPS can be used and will be the default in future versions.

4 Analyzing Results

IQMOL reads in a wide variety of file types and allows the user to visualize many of the results they contain. Individual files can be opened via the File \blacktriangleright Open menu option, or alternatively by dragging and dropping the file onto the Viewer window. Directories can also be opened via the File \blacktriangleright Open Dir menu option, or by dragging and dropping the directory onto the Viewer.

Opening directories allows more than one file associated with a molecule to be loaded together, such as a Q-CHEM output file and a formatted checkpoint file. The directory name determines the base name for the molecule and IQMOL loads all files contained within the directory that have a matching base name.

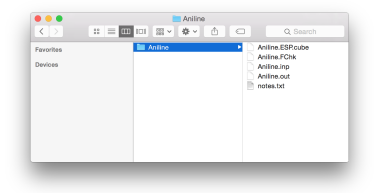


Figure 4.1: Example: Opening the Aniline directory will load the Aniline.ESP.cube, Aniline.FChk, Aniline.inp and Aniline.out files into the one molecule in the MV. The notes.txt file will be ignored as it has a different base name.

4.1 Molecular Surfaces

Several molecular surfaces can be generated without first performing a quantum chemical calculation. These include: van der Waals, pro-molecule and superposition of ionic densities (SID). The SID surface is similar to the pro-molecule surface except that it scales the densities based on the atomic charges of the atoms and may provide a more accurate representation of the density of charged systems.

To plot one of these pseudo-density surfaces, double-click the 'Surfaces' item in the MV associated with the molecule.

4.2 Plotting Molecular Orbitals From a Checkpoint File

Plotting molecular orbitals (MOs) requires a formatted checkpoint file (.fchk file extension) which is generated by default when running Q-CHEM calculations from IQMOL (GUI rem variable should be set to 2). After opening the fchk file, a 'Canonical Orbitals' item will appear in the MV under the Surfaces item. Double-clicking this brings up the Add Surface dialog:

Туре:	Alpha Orbital ᅌ						
Orbital(s):	24 (HOMO) 🗘 to 24 (HOMO) 🗘			-0.48			
			ee	-0.54	-		•
sovalue:	þ.020 🜔 🗹 Simplify Mesh		artro	-0.6			-
Quality:	Law Link		Energy/Hartree	-0.66	=		
auanty.	Low , , , , , High		nerç	-0.72			
Colors:	Negative Positive		ш	-0.78			
				-0.84	••••		
Opacity:					Alpha	I	Beta
Cancel	Add To Queue Calculate						
		Be	eta o	rbital 24:	-0.539 Eh		

Figure 4.2: The Add Surface dialog allows MOs and (spin-)densities to be plotted.

The Add Surface dialog allows MOs, total densities, and spin densities to be computed. Several surfaces can be queued and computed at the same time, which is more efficient as the shell data only need to be computed once. The quality, colors and opacity of the surfaces can be be set in the dialog, and changes to these settings are saved to the preferences as default values for any subsequent surfaces generated.

Note that each tick on the quality scale corresponds to roughly four times the number of grid points as the previous one, and will therefore take roughly four times as long to compute. Also note that densities require evaluating shell-pair values and are significantly more expensive to compute that MOs, which only require shell values. For most purposes, the pseudo densities (such as the pro-molecule density) available by double-clicking the Surfaces item provide an almost identical density surface, but at a much cheaper cost.

Once computed, the individual surfaces appear as sub-items in the MV, and these can be further configured by double-clicking the item in the MV.

The Add Surface dialog also contains an interactive energy level diagram on the righthand side panel. The vertical scale can be zoomed in and out using the scroll wheel

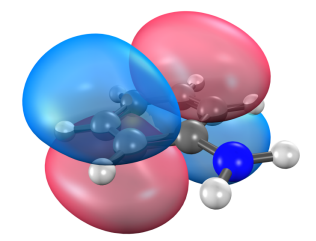


Figure 4.3: An example molecular orbital for aniline.

(or equivalent) on the mouse. The scale can also be translated by a left click-and-drag. Individual orbitals can be selected with a left-click and this will cause the energy of the selected orbital to appear below the diagram, as shown in Fig. 4.2.

4.3 Other Orbitals and Densities

Orbitals and densities other than those based on the canonical orbitals can also be visualized, if the appropriate calculation has been performed. Localized orbitals, natural transition orbitals, natural bond orbitals and attachment/detachment densities are all possible.

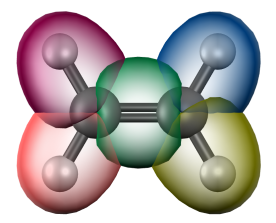


Figure 4.4: Localised σ -bonding orbitals in ethene.

Dyson orbitals can be also displayed, either from the checkpoint file (requires Q-CHEM

 \geq v5.1) or from the output file using an older version of Q-CHEM. In the latter case, the PRINT_GENERAL_BASIS option should be set to true in order for IQMOL to generate the basis function data.

```
$rem
METHOD = EOM_CCSD
BASIS = cc-pVTZ
EOM_EA_BETA = [1,1,1,0]
CC_D0_DYSON = TRUE
GUI = 2
PRINT_GENERAL_BASIS = TRUE ! Ony required for QChem < v5.1
$end</pre>
```

If available, Dyson Orbitals appear under the Surfaces item in the Model View and behave similarly to the Canonical Orbitals.

4.4 Exporting Cube File Data

Each surface requires the generation of data on a 3D grid and these data are stored internally so that subsequent calculations of the same surface (with different isovalues, for example) are much faster. To see what grid data is being stored, right-click on the MO Surfaces item in the MV to bring up the context menu. Selecting the Show Grid Info menu brings up the Grid Information dialog:

	Type	Data Size (kB)	Step Size (Å)		Bounding Box
1	Alpha 25	740.2	0.4330	(-5.97	Delete
2	Alpha 26	740.2	0.4330	(-5.97	Export Cube File Export Cube File (Switch Phase)
					Close

Figure 4.5: The Grid Information dialog shows what data is being stored and offers the option of exporting the data in a cube file format.

From this dialog it is possible to export cube files containing the grid data. Right-clicking on the desired grid brings up a context menu with the Export Cube File option. The Switch Phase option swaps the sign of the data, and may be useful when comparing different MOs where the (arbitrary) phases differ. Cube files can be saved for future plotting to avoid having to recompute the data, or for reading into another plotting package. This is described in the following section.

4.5 Visualizing Cube File Data

Cube files contain volumetric data such as electron densities, molecular orbitals or electrostatic potentials (ESP). Because the data has been pre-computed, generating surfaces using them is very quick. After opening a cube file, a Cube Data item will appear in the MV, double-clicking this item brings up the Add Surface dialog which allows an isovalue surface to be requested (Fig. 4.6). The Signed check-box causes two isosurfaces to be generated corresponding to \pm the specified isovalue, and this should be checked for data such as MOs and spin densities.

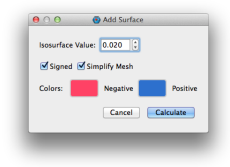


Figure 4.6: The Add Surface dialog for cube data files

Cube file data can also be used to color a surface. This will require either two cube files (one containing the surface data and the second containing the property used to color the surface) or a cube file and a checkpoint file. In either case, two files need to be loaded into the one molecule using the File \blacktriangleright Open Dir menu option, described above.

After creating a surface, double-clicking the surface item will cause the Configure Surface dialog to appear:

The Property combo-box will have the cube data as one of the options, and selecting this will cause the surface to be colored according to the data in the cube file. The gradient colors can be altered by clicking inside the gradient box.

Colors:	-0.2024	0.2028
	Swap Colors Center Gradient	
Property	Cube Data (Aniline.ESP)	
Render	• Fill Clines Dots	
Opacity		— 🗌 Clip
Surface A	rea: 172.528	
		ОК

Figure 4.7: Configure Surface dialog showing the Cube Data as an option for the property.

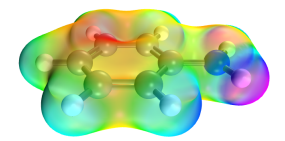


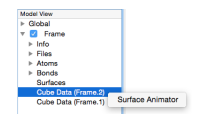
Figure 4.8: Electron density plot of aniline colored according to the electrostatic potential.

4.6 Orbital Animations

Orbital animations can be created e.g., for watching the evolution of the frontier orbitals during a reaction, or watching the relaxation of an orbital during a self-consistent field (SCF) calculation. Key frames should be saved as cube file data in a single directory. The base name of the cube files needs to match the directory name, e.g.

relaxation/relaxation.Frame.1.cube relaxation/relaxation.Frame.2.cube

Open the directory, as described in Sec. 4, and right-click on the first cube file item which appears in the MV, this will allow you to access the Surface Animator dialog:



The dialog is shown in Fig. 4.9 and can be used to adjust the key frame order and the settings for the animation, including the number of interpolation frames (higher values give smoother results, but take longer). Pressing Calculate generates each frame and individual frames can be accessed in the MV. Use the playback options to run the animation, recording it with the button, if desired.

00	Create Animation
Cube File Fr	ames
Frame.1 Frame.2	
Interpolat	ion Frames 10 🗘 🕼 😭 🜗
Surface Ren	der
Render	• Fill CLines Dots
	Simplify Mesh
Isovalue	0.020
Colors	Negative Positive
Opacity	O
Playback	
Loop	Bounce Update Bonds
Speed -	•
	Cancel Calculate

Figure 4.9: The surface animation dialog.

4.7 Potential Energy Surfaces

Several calculation types provide information about the potential energy surface (PES). These include geometry optimizations, PES scans, reaction pathways and transition state searches. On opening a Q-CHEM output file containing one of these job types a Geometries item appears in the MV and can be double-clicked to bring up the Geometries dialog.

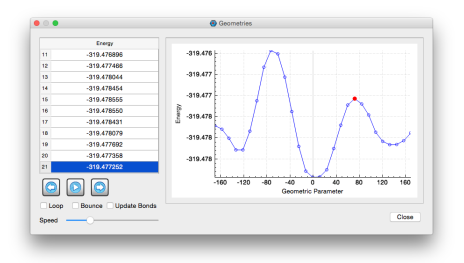


Figure 4.10: The Geometries dialog showing a selected geometry.

The pathway can be animated in the viewer using the play button, and individual frames can be selected from either the table or by selecting a dot on the energy plot. The Viewer window will be automatically updated with the corresponding structure. As with the MO energy diagrams, the energy plot can also be zoomed and scrolled.

Pathways can also be read in from an XYZ file. The format for this is simply a concatenation of regular XYZ files:

5			
-2	91.77177		
Si	0.71979	-0.08082	-0.76577
Η	0.73262	-1.27801	0.22990
Η	1.10451	0.93673	0.34825
Η	-1.32980	0.23894	-0.28240
Η	-1.22713	0.18317	0.47002
5			
-2	91.76961		
Si	0.67218	-0.07483	-0.72935
Η	0.73053	-1.29428	0.22634
Η	1.10815	0.95270	0.34714
Η	-1.29502	0.23567	-0.33016
Η	-1.19409	0.17584	0.49156

. . . .

The first line gives the number of atoms, the second (comment) line must have the corresponding energy and then the following lines have the xyz coordinates of each atom in the system. This pattern repeats for as many frames as you have available.

4.8 Vibrational Frequencies

Vibrational frequencies can be read in from a Q-CHEM output file and appear as a Frequencies item in the MV. Normal mode vectors can be visualized by selecting the associated frequency item in the MV.

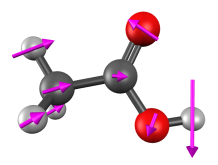


Figure 4.11: Arrows indicating a normal mode in acetic acid.

Double-clicking a frequency in the MV will cause the molecule to vibrate according to the selected mode. Double-clicking the Frequencies item in the MV will bring up the Vibrational Frequencies dialog:

This dialog contains an impulse spectrum showing the positions and relative intensities of the frequencies. Individual modes can be selected on the spectrum by clicking the hollow circles, and this will also update the Viewer window with the selected mode. The impulses can be broadened using either Gaussians or Lorentzians to give a more realistic looking spectrum, and the image can be exported by righ-clicking on the spectrum to bring up a context menu.

The horizontal scale of the spectrum can be zoomed (using the scroll wheel on the mouse) and translated (left-click and drag) to give greater detail.

4.8.1 Isotopic Substitution

IQMOL makes it easy to set up isotopic substitution loops for a Q-CHEM vibrational frequency calculation. First select the atoms you wish to substitute in the Viewer and

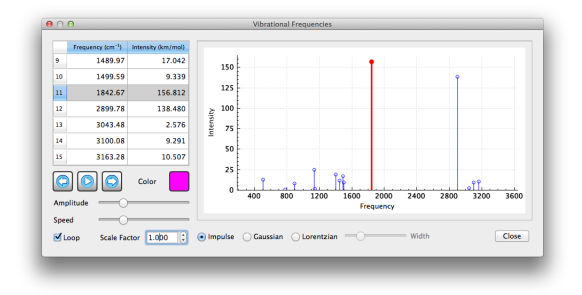


Figure 4.12: The Vibrational Frequencies dialog.

then select the Build \blacktriangleright Set Isotopes menu option. A dialog will appear that allows you to change the isotope for the selected atoms, along with the ability to change the pressure and temperature used for calculating the thermochemical data. Once the masses have

Temperatur	e 298.18 K	C Pressure 1.00 atm
Element	Isotopic Mass	Indicies
Hydrogen	H(2) 🗘	3, 4, 5
Carbon	C(std) 🗘	1
		Cancel OK

Figure 4.13: Isotope substitution dialog allowing non-default masses to be specified for the selected atoms. Note that only one non-default mass is allowed for each element type.

been selected, click OK and an Isotopes item appears in the MV containing the substitution information. This can be edited or checked by double-clicking on the appropriate Substitution item in the MV. Different substitutions can be set up by selecting different atoms in the Viewer and/or different masses and these will be looped over in the Q-CHEM frequency calculation. Each substitution loop only requires a re-weighting of the Hessian before diagonalization and so multiple loops can be obtained very quickly once the Hessian has been calculated.

4.9 NMR Spectra

NMR shieldings and chemical shifts can be visualized using IQMOL. A Q-CHEM calculation (JOB_TYPE = NMR) must first be run and the output file loaded into IQMOL. Shielding constants can be displayed as atom labels in the viewer by selecting the Display ▶ Atom Labels ▶ NMR option.

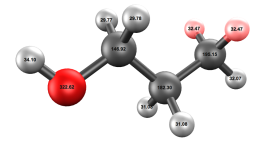


Figure 4.14: Propanol showing the NMR shieldings

An NMR item will appear on in the MV and double-clicking this brings up the NMR Spectrum dialog.

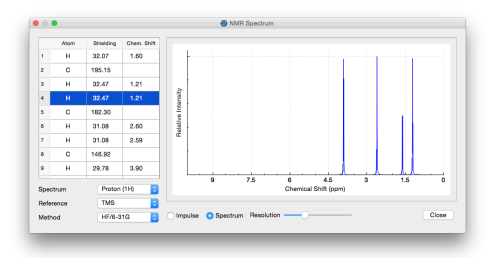


Figure 4.15: The NMR Spectrum dialog

Various spectra can be displayed (¹H, ¹³C etc.) and if a reference is available (*e.g.* TMS) for the given level of theory, chemical shifts can also be displayed. Selecting rows in the table will also highlight the corresponding atoms in the viewer for easy identification. If impulses

are plotted, then these can also be selected to identify which atoms are responsible for the selected signal.

As with the vibrational frequencies spectra, NMR spectra can be exported by rightclicking on the image to bring up the context menu.

5 Appearance

5.1 Viewer Camera

When viewing a molecule, the coordinates of the atoms and surfaces are fixed in the world reference frame. When manipulating the molecule using the mouse (see § 2.5) it is actually the camera that changes position and orientation. If more precise control of the camera is required (*e.g.* for reproducing a particular image), then the camera can be configured via the Display \blacktriangleright Camera menu option. By default the camera uses a perspective projection to give better depth perception, but an orthographic projection can also be selected, if required.

Orthogra	aphic 💿 F	Perspectiv	e FoV	45 🗘
Position r	10.25	0 90	0 4	330 🗘
		Rese	t 🛛	ОК

Figure 5.1: The Configure Camera dialog allows precise control over position and orientation of the Viewer camera.

5.2 Clipping Planes

A Global clipping plane is available and can be viewed by enabling the Clipping Plane item under Global in the MV. Clipping planes can be useful for showing surface information without obscuring the underlying molecular structure. While selected, the clipping plane can be arbitrarily translated and rotated using the manipulate selection mode described in § 2.5. Alternatively, a precise orientation of the clipping plane can be set by doubleclicking the Clipping Plane item in the MV to open the configuration dialog.

Individual surfaces can be clipped, by selecting the Clip check-box in in the surface configurator dialog (see Fig. 4.7) for each surface. Once a surface has been clipped, the clipping plane remains active even if hidden by unchecking the Clipping Plane item check-box in the MV.

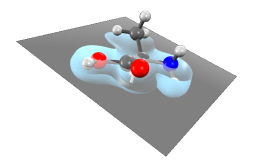


Figure 5.2: Image showing a clipped density surface allowing the underlying molecular structure to be more easily seen.

5.3 Shaders

IQMOL uses GLSL shaders to enhance the appearance of the Viewer. Shaders can be made active by going to the Display \blacktriangleright Appearance menu option, and selecting the Shader panel.

	Shader POV-Ray
Shader Pho	ong 🗘
Ambient	
Diffuse	
Highlights	O
Noise Intensi	ty 🔘
Saturation	O
Shininess	
Options	
	e Edges 🗹 Enhance Transparency here Lighting
Lights	
✓ Front ✓ Lower	☑ Highlight 🛛 Left
Save As Defa	oK

Figure 5.3: Configuring shader options

Changes to the various shader options will take immediate effect and can be saved as the default by clicking Save As Default. Some shaders have several lights associated with them, and these can be turned on or off to give different lighting effects.

5.4 Exporting POV-Ray Files

IQMOL can generate scene files (.pov) for use with an external POV-Ray package. POV-Ray uses ray tracing algorithms to generate images of very high quality and resolution. POV-Ray is open source, but several official and unofficial packages exist with pre-compiled binaries, in particular testing was done using the Mac version of MegaPOV. Binaries can be downloaded from the following sites:

- Mac: http://megapov.inetart.net/povrayunofficial_mac
- Windows: http://www.povray.org/download

To export a scene file use the Edit \blacktriangleright Generate POV-Ray Input menu option. Scene files are plain text with instructions for the POV-Ray program, and can be edited by the user for fine tuning before processing. IQMOL can apply several effects (*e.g.* surface textures) to the molecule, and these can be configured by opening the Appearance dialog (Display \blacktriangleright Appearance menu option).

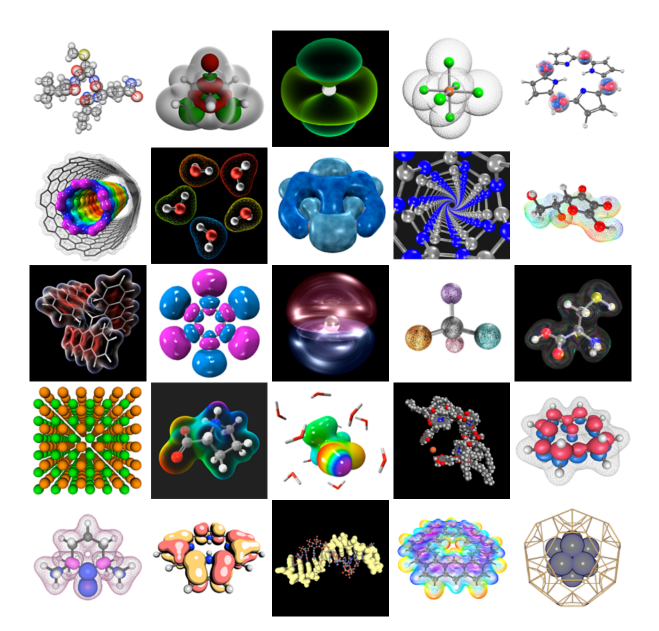
	Shader	POV-Ra		
Phong sha Global	der setting	s are use	d for ligh	iting
Height	480 🗘	Width	640	0
Backgro	Background Black			
Gamma	3.0 🗘			
Textures				
Atom	Rough		0	
Bond	Crumple	d	\$	
Surface	Bubble		0	
Save As De	fault			ок

Figure 5.4: Appearance dialog showing Options for POV-Ray scene files.

Note that IQMOL attempts to generate a scene file that matches as closely as possible what is shown in the Viewer window. However, there may be slight variations in lighting, color and camera angle, and some trial and error may be required to obtain a pleasing image. In particular, the Gamma value may need experimenting with, and changes to the ambient and diffuse lighting (via the shader options) may be required depending on if a light or dark background is selected for the scene.

6 Sample Images

Here are some examples of the kinds of images that can be generated using IQMOL's shader and rendering options. Most of these are simply screen shots, however, tiles 3, 4 and 5 in the middle row were obtained by running POV-Ray on the IQMOL generated scene file.



References

- [1] PILATI, T., AND FORNI, A. Symmol: a program to find the maximum symmetry group in an atom cluster given a prefixed tolerance. J. Appl. Cryst. 31 (1998), 503.
- [2] RAPPE, A. K., CASEWIT, C. J., COLWELL, K. S., GODDARD III, W. A., AND SKIFF, W. M. Uff, a full periodic table force field for molecular mechanics and molecular dynamics simulations. J. Am. Chem. Soc. 114 (1992), 10024.