					Unteri			
ICM-C	hemist-Pro	User	Guide v.	3.8	•••••	1		
1 Help	Videos on	YouTu	be	•••••		3		
2 Refere	ence Guide				5			
	2.1 Graphics	Refe	rence	Guide		5		
	2.1.1 Dov	vnload	and In	stall	ICM-Brows	ser	5	
	2.1.2 Hov	v to	use the	Graphic	al Displa	ay	5	
	2.1.3 Hov	v to	make Gra	aphical	Selections		7 ogens and	
	2.1.4 Hov	v to	Convert	Proteins	, Displa	ay Hydro	egens and	
	2.1.5 Hov	na Bir	ahang Po	Ockel. Granhiae	Effects	····· / ·····	7	
	2.1.5 Hov 2.1.6 Hov	v to v to	add Labe	diapines	Annotation	ns		
	2.1.7 Hov	v to	Make Hig	h Ouali	tv Public	cation	Images8	
	2.1.8 Hov	v to	Superimpo	se Pr	otein Str	uctures.		
	2.1.9 Hov	v to	Measure	Distance	and	Angles		
	2.2 ICM-Brow	vser-Pro	Refe	rence	Guide		9	
	2.2.1 Dov	vnload	and In	stall	ICM-Brows	ser-Pro .	9	
	2.2.2 Gra	phics	~~~~~		9	)		
	2.2.3 Prot	ein	Structure	Ana	lysis		9	
	2.2.4 Surf	aces	 D		10	0 10		
	2.2.5 Sup	erimpose	PTO	Tool	••••••		)	
	2.2.0 CTy	stanograf	Δnalvei	s 10018	• • • • • • • • • • • • • • • • • • • •	10	)	
	2.2.8 Plot	ting	Tools					
	2.3 ActiveICM	1 Ret	ference	Guide	- Create	3D Molec	ular	
	Documents	for th	e Web	and Pow	verPoint .		11	
	Creating 3 2.3.1 Get	D Docur	nents Is	Straigh	tforward		11	
	2.3.1 Gett	ing	Started					
	2.3.2 Hov	v to	Create a	a Series	of Full	y-Interactive	3D	
							10	
	2.3.3 Hov 2.3.4 Hov	v lo	Display	Molecul	ar Docum	ents	PowerPoint12	,
	2.3.4 Hov 2.3.5 Hov	v to	Display	Molecul	ar Docu	ments on	the Web 13	'
	2.4 ICM-Cher	nist	Reference	Guide	2		PowerPoint12 the Web13	
	2.4.1 Hov	v to	Import,	Sketch,	and Ed	lit Chemic	als13	
	2.4.2 Hov	v to	Work wit	h Chen	nical Spre	eadsheets	13	
	2.4.3 Hov	v to	Undertake	a Ch	nemical S	earch	14	
	2.4.4 Hov	v to	Work wit	h Pharr	nacophores			
	2.4.5 Hov 2.4.6 Hov	to to	Generate	Storooi	a Cluster	ing	omers15	
	2.4.0 Hov	v to	Generate	Combi	natorial	Libraries	16	
	2.4.8 Hov	v to	Generate	Plots	and Hist	ograms .	16 16	
	2.5 ICM-Cher	nist-Pro	Refe	rence	Guide		16	
	2.5.1 Hov	v to	use the	ICM 3E	) Ligand	Editor	16	
	2.5.2 Hov		Convert			D		
	2.5.3 Liga		Energetics				10	
	2.5.4 Hov 2.5.5 Hov						10	
	2.6 Menu O		Guide	QSAK				
	2.6.1 File				1			
	2.6.2 Edit	Me	nu		2	21		
	2.6.3 Vie				2			
	2.6.4 Bioi							
	2.6.5 Too	IS N	lenu - X	ray			0	
	2.6.6 Too 2.6.7 Too	15 IV. 16 IV.	Ienu - 51	D Predici	••••••		28	
	2.6.8 Too					20	28	
	2.6.9 Too	ls N	Ienu - E	xtras				
	2.6.10 To	ols	Menu - '	Table		29		
	2.6.11 To	ols	Menu - 0	Chemical	Search	•••••	31	
	2.6.12 To							
	2.6.13 Ho							
	2.6.14 Ch 2.6.15 Do							
	2.0.15 DC	CKIIIg	wienu	••••••				

2 Reference Guide
2.6.16 MolMechanics Menu
2.6.17 Windows Menu
2.7 Tab Guide
2.7.1 Display 1ab
2.7.1 Display       Tab
2.7.4 PDB Search Tab
2.7.5 Ligedit Tab
2.7.6 Meshes Tab
3 Getting Started
<b>3 Getting Started</b>
3.1.1 The Components of the GUI
3.1.2 How to load a PDB Structure
3.1.3 How to Move a Structure in the Graphical Display36
3.1.4 How to Display Molecules using the ICM Workspace Panel
Panel
3.1.6 How to Change Protein Representation
3.1.7 How to Color
3.1.8 How to Display a Binding Pocket Surface
3.1.9 How to Save an ICM Object
3.1.10 How to Save an ICM Project File44 3.1.11 How to Drag and Drop44
3.1.11 How to Drag and Drop
3.1.13 How to Move Windows
3.1.14 How to Make a Picture
3.2 How to search and download Protein Structure, Sequences,
and Chemicals
3.2.1 Search the PDB50 How to Search the Protein Databank and Download50
3.2.2 Search Pocketome 54
3.2.3 ChEMBL Search
3.2.4 Search         SureChEMBL         57           3.2.5 BLAST         Search         57           3.2.6 Search         UniProt         58
3.2.5 BLAST Search
3.2.6 Search UniProt
3.2.8 Search Drug Bank 59
3.2.7 SearchPDB by LigandCode
3.3 Create New Objects61
3.4 Open and Read Files
3.4.1 Open with Password
3.4.2 Extract from icb file62 3.5 Saving Files
3.6 Making Selections
3.6.1 Graphical Selection Tools65
3.6.2 Quick Selection
3.6.3 How to Change the Selection Level and Mode
3.6.4 How to check what is selected
3.6.5 OrangeSelection
3.6.7 Changing a Selection
3.6.8 Filter Selection
3.6.9 Workspace Selections
3.6.10 Workspace Navigation
3.6.11 How to Select an Object72 3.6.12 How to select a Molecule
3.6.13 How to select Residues
3.6.14 Select All75
3.6.15 Selecting Neighbors
3.6.16 Selecting Neighbors: Graphical
3.6.17 Selecting Neighbors: Workspace
3.6.19 Alignment and Table Selections

3 Getting	Started
e otting	3.6.20 Making Links
	3.6.21 Tags
3.	/ Preferences
	3.7.1 Bonds Preferences
	3.7.2 DirectoriesPreferences
	3.7.4 GUI Preferences
	3.7.5 GUI Preferences
	3.7.6 Image Preferences
	3.7.6 ImagePreferences
	3.7.8 Plot Preferences
	3.7.9 RibbonPreferences
	3.7.10 Shell Preferences
	3.7.11 System Preferences
4 Protein	Structure95
	1 Convert to ICM Object
	4 1 1 Load a Protein Structure 95
	4.1.2 Converting PDB Files Into ICM Objects
4.	2 Pocket Display97
	4.2.1 Receptor Pocket Surface
	4.1.1 Dotating       a Frotering       Structure
	4.2.5 How to Display Hydrogen Bonds
4	3 Crystallographic Analysis 103
	4.3.1 Crystallographic Neighbor
	4.3.2 Crystallographic Cell
	4.2.4 Ergand       Focket       Interactions       102         3 Crystallographic       Analysis       103         4.3.1 Crystallographic       Neighbor       103         4.3.2 Crystallographic       Cell       105         4.3.3 Biomolecule       Generator       106         4.3.4 Cot       Electron       Domity       107
	4.3.4 GetElectronDensityMapMap4.3.5 Map'sOriginalCell
	4.3.5 Map's Original Cell109
	4.3.6 Contour Electron Density Map
4	4.3.7 Convert Xray Density to Grid110 4 Protein Superposition
т.	4.1 Select Proteins for Superposition
	4.4.2 Superimpose Button
	4.4.3 Superimpose by 3D113
	4.4.2 SuperimposeButton1124.4.3 Superimposeby 3D1134.4.4 SuperimposeMultipleProteins4.4.5 Appendix113
	4.4.5 Arrange as Grid114
1	4.4.6 Superimpose Sites by Atomic Property Fields
4.	5 Protein Structure Analysis
	4.5.2 Calculate RMSD
	4.5.3 Contact Areas
	4.5.4 Identify Closed Cavities
	4.5.5 Surface Area124
	4.5.6 Measure Distances
	4.5.7 Planar         Angle         125           4.5.8 Dihedral         Angle         126
	4.5.9 Ramachandran Plot Interactive
	4.5.10 Export Ramachandran Plot
	4.5.11 Protein Health
	4.5.12 Local Flexibility128
	4.5.13 Protein-Protein Interface Prediction128
	4.5.14 Identify Ligand Binding Pockets129
5 Molecula	ar Graphics133
_	1 Molecule Representation
5.	5.1.1 Wire Representation
	5.1.2 Stick and Ball (Xstick) Representation
	5.1.3 Ribbon Representation
	5.1.4 Skin Representation
	5.1.5 CPK Representation
	5.1.6 SurfaceRepresentation
	stri Disping and Ondisping Hydrogons

Table of Contents	
5 Molecular Graphics	
5.1.8 Display Hydrogen Bond140 5.1.9 Display Formal Charges142	
5.1.10 Ca (carbon alpha) Trace	
5.2 Multi-Windows 142	
5.3 Meshes - Surface - Grobs143	
5.3.1 Surfaces	
5.3.2 MolSkin	
5.3.4 Color Surface by Selection	
5.3.3 ColorSurfaceby Proximity	
5 3 6 Macroshape 147	
5.3.7 Google 3D Objects (Sketchup)	
5.3.9 Mesh Options	
5.3.10 Mesh Clipping	
5.3.10 Mesh Clipping	
5.3.12 Occlusion Shading151	
5.4 Coloring	
5.4.2 Color Background	
5.4.3 Background Image	
5.4.3 Background Image153 5.5 Lighting	
5.6 3D Stereo	
5.7 3D Printing	
5.8.1 Labeling 156	
5.8.1 Labeling	
5.8.3 Labeling Residues157	
5.8.4 Move Residue Label	
5.8.5 LabelVariables1585.8.6 Sitesand Annotation159	
5.8.7 Changing Label Colors	
5.8.7 ChangingLabelColors1635.8.8 CustomizedLabel2D or3D1635.8.9 UndisplayCustomizedLabel164	
5.8.9 Undisplay Customized Label164	
5.8.9 Undisplay       Customized       Label	
5.8.12 Displaying Tethers	
5.8.13 Displaying Distance Restraints	
5.8.14 Display Clash167	
5.8.15 Display Rainbow, Box, Ruler167 5.8.16 Display Gradient167	
5.9 Display Distances and Angles	
5.9.1 Display Distance Between Two Atoms - the quick way168	,
5.9.2 Display Planar Angle	
5.9.3 Display Dihedral Angle	
5.9.4 Delete Label	
5.10.1 Fog	
5.10.2 Shadows169	
5.10.3 Sketch Accents	
5.10.4 Elegant Ribbon & Ligand Sketch	
5.10.6 Perspective	
5.10.7 Animate View	
5.11 Graphics Shortcuts173	
5.12 Molecule Move Buttons	
5.12.1 Rotation	
5.12.2 Custom Rotation	
5.12.4 Zoom176	
5.12.5 Center	
5.12.6 Torsion Angles	
5.12.7 Connect (Move)	

5 Molecular Graphics
5.13.1 Mesh Clipping
5.14 Graphic Layers
5.15.1 High Quality Image
5.15.1 High       Quality       Image
5.15.3 Image Options
5.16.1 Making a Movie from a Set of Slides
5.16.2 Screenshot Movie
6 Slides & ActiveICM187
6.1 Making Slides
6.2 Make a Movie from a Set of Slides
6.3 How to View and Navigate Slides
6.3.2 Slide Navigation
6.4 How to Edit Slides195 6.4.1 Edit Slide195
6.4.2 Move Slide
6.5 How to Add Smooth Blending and Transition Effects
Between Slides
6.6 How to Make Molecular Documents - Link HTML Text to Slides
6.6.1 How to Add Text or Edit a Molecular Document
6.6.2 How to Make a Hyperlink Between Text and a Slide201
6.6.3 Insert         Image
6.6.5 Insert a Dialog Box204
6.6.6 How to add a check box
6.6.7 Document Navigation
6.6.8 Protect Shell Objects From Deletion
6.8 How to Embed in Microsoft PowerPoint 2003
6.9 How to Embed in MicrosoftPowerPoint20076.10 How to Embed in MicroSoftPowerPoint2010
6.11 Embed in Web Browser
6.12 How to Use ActiveICM in PowerPoint
6.13 How to Change ActiveICM Component Properties
visualisation changes214
6.14.1 PowerPoint Cache Errors217
6.15 BackgroundImages
7 Cheminformatics
7.1 Read Chemical Structures
7.1.1 Extract2D structurefrom PDB Chemical
7.1.3 Extract 3D coordinates from 3D Chemical
7.2 Save Chemical Structures and Images
7.2.2 Saving Chemical Spreadsheet as SDF
7.2.3 Export to Excel
7.2.4 Saving Chemicals in the Molecular Editor
7.2.6 Saving Chemical Images
7.2.7 IUPAC Name
7.3 WorkingwithChemicalSpreadsheets
7.3.2 Chemical Spreadsheet Template
7.3.3 Molecular Table Display
7.3.4 How to add Property Columns into a Chemical Spreadsheet
7.3.5 How to sort a column(s) in a chemical spreadsheet227

#### 7 Cheminformatics

7.3.6 How 7.3.7 Add 7.3.8 How chemical 7.3.9 How to save a chemical spreadsheet in sdf format. ....230 to export your chemical spreadsheet into 7.3.10 How Excel. 7.3.11 How 7.3.12 How ....232 7.3.13 How to use find and replace in a chemical spreadsheet. 7.3.14 How to mark and label rows in a chemical spreadsheet. to insert hyperlinks to the PDB, PubMed, and ......234 7.3.15 How Uniprot. to copy and paste 2D chemicals. ......234 to edit data inside a chemical spreadsheet. 7.3.16 How .....235 7.3.17 How to remove salts, explicit hydrogens and 7.3.18 How 7.3.19 How 7.3.20 InChi to identify duplicate chemicals in a chemical 7.3.21 How 7.3.22 How 7.3.23 How 7.3.24 Display 7.3.25 Copy 7.3.26 Edit 7.3.27 Color Table Column ......239 7.3.28 Chemical 7.3.29 Chemical 7.3.30 Zoom. table. 7.4 Molecular Click Options ......248 Dictionary ......249 7.4.3 Right 7.4.4 Chemical 7.4.5 Dictionary .....255 7.5 Chemical 7.5.1 Query Setup ......256 7.5.2 Filter 

7 Cheminformatics
Query Options
Results
Search
7.5.4 Search a Database by Text
7.5.5 Pharmacophore Drawing and Searching
7.7 Convert Chemicals to 3D
7.7.1 Converting a Chemical from the PDB 278
7.7.2 Converting a Chemical from the PDB using the ICM
7.7.1 Converting a Chemical from the PDB
7.7.3 Converting a Chemical from the PDB using the Graphical Display
Graphical Display278
7.8 Generating Chemical Fragments
7.9 Find Bioisostere
7.9.1 2D         Bioisostere
7.10 Molcart
7.10.2 Molcart Getting Started
7.10.2 Wolcart Certaing Started
7.10.3 CreatingDatabasesand ImportingData7.10.4 MolcartSearch
7.10.5 Molcart Administration
7.11 Calculate Properties
7.11.1 Number of Atoms
7.11.2 ToxScore
How the ToxScore is Calculated
How to Calculate LoxScore
7.12 StandardizeTable
7.13 Annotate By Substructure
7.14 Align/Color by 2D Scaffold297 7.15 Set Formal Charges
7.16 Enumerate Formal Charge States
7.17 Convert
7.17.1 Convert 2D to 3D298
7.17.2 3D to 2D Depiction
7.17.3 Convert Smiles or InChi to 2D
7.17.4 Convert Structure to Smiles
7.17.5 Convert to Racemic
7.19 Predict
7.20 Generate 3D Conformers 301
7.20 Generate3D Conformers3017.21 GenerateTautomers
7.22 Generate Stereoisomers
7.23 Prodrug
7.24 Ligand Energetics
7.24.1 Conformational Entropy
7.24.2 Strain
7.25 Cluster Set
7.25.2 How to select representative centers from a tree310 7.25.3 How to reorder branches and change the distance
of trees
7.25.4 How to edit the tree - labels, spacing and
coloring
7.25.5 Auto Close Tree Mode
7.25.6 Calculate Maximum Common Substructure for Nodes312
7.25.7 R Group Decomposition from Clustering Node
7.26 PCA Analysis
7.27 Visualize Chemical Space
7.27.1 MaximumCommon Substructure(MCS)Dendrogram3147.27.2 SelfOrganizedMap
7.27.2 Sen Organized Map
7.29 Merge Two Sets
7.30 Select Duplicates
-

7 Cheminformatics
7.31 MultiParameterOptimization
7.32.1 Sketch Markush Structure
7.32.2 B Groups 321
7.32.2 R-Groups
7.32.4 Enumerate by Scaffold 324
7.32.5 Enumerate by Beattion 325
7.32.4 EnumeratebyScaffold
7.33.1 R-Group Decomposition
7.33.2 SAR Table
7 33 3 Plot R-Groups 332
7.33.4 Generate SALI Table
7.33.4 GenerateSALI Table3327.33.5 MatchedPairAnalysis3337.34 SubstructureSuperposition3357.34.1 RigidSuperpositionof Compounds in a Table onto
7.34 Substructure Superposition
7.34.1 Rigid Superposition of Compounds in a Table onto
a Template in The Graphical Display
7.34.2 Rigid Substructure Superposition
7.34.3 Flexible Substructure Superposition
7.35 Alomic Property Field Superposition
7.35 Atomic       Property       Field       Superposition
7.35.2 Multiple APF Alignment of Compounds in a Table 341
7.36 APF Tools 342
7.36.1 Consensus Pharmacophore 342
7.36.1 ConsensusPharmacophore
7.36.3 APF Ligand Based Screen - Screen SD file against
template molecule(s)
template molecule(s)
8 Learn and Predict
8.1 Learn
8.1.1 About the Model
8.1.2 Model Weights
8.1.4 Save and Share Model
8.1.4 save and share woder
8.3 Fingerprint Methods 354
8.3 Fingerprint Methods
8.5 A Little Theory on Learning
9 MolScreen
9.1 Model Types
9.1.1 Kernel Chemical Classification/Regression (kcc)
models
Hybrid Eingennuinte 250
Hybrid Fingerprints
Dataset
Training
Predicted Value
9.1.2 Docking to Ligand Field Z-Score (dfz) models
APF
APF
Dataset
Training
Predicted Value
9.1.3 Docking to Ligand Field Classification/Regression

10 The	3D Fully Interactive Ligand Editor
	10.15 Side Chain Refinement
	10.16 MultipleReceptorDocking
	10.18 Dock to Atomic Property Fields
	10.19 How to Save Ligand Receptor Complex as PDB 454
	10.20 Export Docking Project
	10.21 Close and Clean Up Project455
11 Wor	king with Tables and Plots457
	11.1 Standard ICM Tables457
	11.1.1 GenerateNew Table45711.1.2 Readinga Table458
	11.1.2 Reading a Table
	11.1.5 Saving a table
	11.1.4 BasicTableNavigation45811.1.5 TableView (GridLayout)459
	11.1.6 Change Row Height459
	11.1.7 Table View Save
	11.1.10 Table         Font
	11.1.12 Mark a Row
	11.1.13 Table right click options463 11.1.14 Rename a Table463
	11.1.15 Clone a Table463 11.1.16 Delete a Table463
	11.1.17 Page Setup
	11.1.18 Print a Table
	11.1.19 Export to Excel
	11.1.20 Save a Table
	11.1.21 ChangeColumn and Row Width46411.1.22 MakingTableSelections464
	11.1.22 Making Table Selections
	11.1.24 Rename Column
	11.1.25 Inserting Columns
	11.1.26 Increment         ID         Number
	11.1.27 Column         Visibility
	11.1.29 Freeze Column
	11.1.30 Freeze Row470
	11.1.31 Column Statistics
	11.1.32 Inserting         Rows
	11.1.33 filsert finage
	11.1.35 Copy Cell
	11.1.36 Copy Selection to an ICM Table472
	11.1.37 Deleting Columns and Rows
	11.1.38 Hide and Show Columns472 11.1.39 Column Format and Custom Actions473
	11.1.40 Column Table Annotation
	11.1.40 ColumnTableAnnotation
	11.1.42 Table Filtering
	11.1.43 Append Table479
	11.1.44 Mark and Select Rows
	11.1.45 Group View by Column
	11.2 Molecular Tables
	11.3 Insert Interactive Objects into Table Cell
	11.4 Plotting Table Data
	11.4.1 Column Histogram
	11.4.2 Histogram Options
	11.4.4 Plots
	11.4.5 Multiple Curves

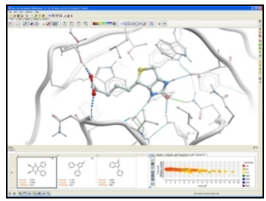
11 Working with Tables and Plots
11.4.6 Heatmap
11.4.7 Pie Chart
11.4.8 Add a title to a plot489 11.4.9 Axis Options489
11.4.9 Axis Options
11.4.11 Logarithmic Plots
11.4.12 Change Mark Shape or Size
11.4.13 Change Mark Color
11.4.14 Grid and Axis Display491 11.4.15 Least Squares Fitting492
11.4.15 Least Squales Fitting
11.4.17 Plot Selection
11.4.16 Zoom,TranslateandCenter49211.4.17 PlotSelection
11.4.19 Print Plot
11.4.20 Saving a Plot Image497 11.4.21 Table Inline Plots497
11.4.22 Annotate Plot
11.4.23 Plot R-Groups
11.4.24 Tooltips Balloons499
11.5 Principal Component Analysis
11.6 Learn and Predict501 11.7 Data Clustering
11.7 Data Clustering
11.7.2 Tree Selection
11.7.2 Tree         Selection
11.7.4 Tree View505
12 Working with Local Databases51112.1 How to make a local database.51112.2 Browse Database51112.3 Edit Database51412.4 Query Local Database516
12 ICM :- KNIME 517
13 ICM in KNIME
13 ICM in KNIME
Available Nodes
Available Nodes
Available Nodes
Available       Nodes
Available       Nodes
Available       Nodes
Available       Nodes
Available       Nodes
Available       Nodes
Available       Nodes
Available       Nodes
Available       Nodes
AvailableNodes517To useICM in KNIME52314 Tutorials52314.1 GraphicsTutorials524Introduction52414.1.1 ChangeMoleculeRepresentationand Color14.1.2 Annotation52814.1.3 Labels52914.1.4 2Dand 3D Labels14.2 CreatingFullyFullyInteractiveSlidesforPowerPointandtheWeb Tutorial535535Tutorial53514.3 ProteinStructureStructure542Tutorial542
AvailableNodes517To useICM in KNIME517 <b>14 Tutorials</b> 52314.1 GraphicsTutorials524Introduction52414.1.1 ChangeMoleculeRepresentationand Color14.1.2 Annotation52814.1.3 Labels52914.1.4 2Dand 3D Labels14.2 CreatingFullyFullyInteractiveSlides535Introduction53514.3 ProteinStructureTutorial54214.3.1 PDBSearch542542
Available       Nodes
AvailableNodes517To useICM in KNIME517 <b>14 Tutorials</b> 52314.1 GraphicsTutorials524Introduction52414.1.1 ChangeMoleculeRepresentationandColor52814.1.2 Annotation52814.1.3 Labels52914.1.4 2Dand 3D Labels14.2 CreatingFullyFullyInteractiveSlidesforPowerPointandtheWeb Tutorial53514.3 ProteinStructure535Tutorial542Introduction542Introduction54214.3.1 PDBSearch14.3.2 ConvertPDB toPDB toICM Object54314.3.4 ProteinStructureAnalysis545
AvailableNodes $517$ To useICM in KNIME $517$ <b>14 Tutorials</b> $523$ 14.1 GraphicsTutorials $524$ Introduction $524$ 14.1.1 ChangeMoleculeMoleculeRepresentationandColor $525$ 14.1.2 Annotation $528$ 14.1.3 Labels $529$ 14.1.4 2Dand 3D Labels $532$ 14.2 CreatingFullyInteractiveSlidesforPowerPointIntroduction $535$ Tutorial $535$ Introduction $542$ 14.3 ProteinStructureStructureTutorial $542$ $542$ 14.3.1 PDBSearch14.3.2 ConvertPDB toPDB toICM Object $543$ 14.3.3 ProteinSuperposition $544$ $4.3.4$ Protein $14.3.5$ ContactAreas $547$
AvailableNodes $517$ To useICM in KNIME $517$ <b>14 Tutorials</b> $523$ 14.1 GraphicsTutorials $524$ 14.1.1 GraphicsMoleculeRepresentation14.1.2 Annotation $528$ 14.1.3 Labels $529$ 14.1.4 2Dand 3D Labels $532$ 14.2 CreatingFullyIntroduction $535$ Introduction $535$ Introduction $535$ Introduction $535$ Introduction $535$ Introduction $535$ Introduction $542$ Id.3.1 PDBSearchSearch $542$ I.4.3.2 ConvertPDB to ICM ObjectI.4.3.4 ProteinStructureAnalysis $543$ I.4.3.5 ContactAreasI.4.3.6 HydrogenBondBond $548$
AvailableNodes $517$ To use ICM in KNIME $517$ <b>14 Tutorials</b> $523$ 14.1 GraphicsTutorials14.1 GraphicsTutorials14.1.1 ChangeMoleculeRepresentationand14.1.2 Annotation $524$ 14.1.3 Labels $529$ 14.1.4 2Dand 3D Labels14.2 CreatingFullyFullyInteractiveSlidesfor PowerPoint14.2 CreatingFullyIntroduction $535$ Tutorial $535$ Introduction $535$ 14.3 ProteinStructureTutorial $542$ 14.3.1 PDBSearch14.3.2 ConvertPDB toPDB toICM Object14.3.3 ProteinSuperposition14.3.4 ProteinStructureAdvance $542$ 14.3.5 ContactAreas14.3.6 HydrogenBond14.3.8 ProteinFreeparationand $548$ 14.3.8 ProteinPreparationand $548$ 14.3.8 ProteinPreparationandCrystallographicAnalysis
AvailableNodes $517$ To useICM inKNIME $517$ <b>14 Tutorials</b> $523$ 14.1 GraphicsTutorials $524$ Introduction $524$ 14.1.1 ChangeMoleculeRepresentationand14.1.2 Annotation $528$ 14.1.3 Labels $529$ 14.1.4 2Dand 3D Labels14.2 CreatingFullyInteractiveSlidesfor PowerPointandthe Web Tutorial $535$ Introduction $535$ Tutorial $542$ Introduction $542$ Introduction $542$ Introduction $542$ Introduction $542$ Introduction $542$ Id.3 ProteinStructureTutorial $542$ 14.3.1 PDBSearch14.3.2 ConvertPDB toPDB toICM Object14.3.3 ProteinStructureAdverterStructureAdverter $543$ 14.3.4 ProteinStructureAdverter $543$ 14.3.5 ContactAreas14.3.6 HydrogenBond14.3.8 ProteinPreparationand $548$ 14.3.7 PocketIdentification14.3.8 ProteinPreparationand $549$
AvailableNodes $517$ To useICM in KNIME $517$ <b>14 Tutorials</b> $517$ <b>14 Tutorials</b> $523$ 14.1 GraphicsTutorialsIntroduction $524$ 14.1.1 ChangeMoleculeRepresentationand Color14.1.2 Annotation $528$ 14.1.3 Labels $529$ 14.1.4 2Dand 3D Labels14.1.2 CreatingFullyIntroduction $535$ Introduction $535$ Introduction $535$ Introduction $542$ Id.3.1 PDBSearch14.3.2 ConvertPDB to ICM Object14.3.3 ProteinSuperposition14.3.4 ProteinSuperposition14.3.5 ContactAreas14.3.6 HydrogenBond14.3.7 PocketIdentification14.3.8 ProteinPreparationandCrystallographicAnalysisTutorial14.3.9 PDBPreparation14.3.9 PDBPreparation
AvailableNodes $517$ To useICM in KNIME $517$ <b>14 Tutorials</b> $517$ <b>14 Tutorials</b> $523$ 14.1 GraphicsTutorialsIntroduction $524$ 14.1.1 ChangeMoleculeRepresentationand Color14.1.2 Annotation $528$ 14.1.3 Labels $529$ 14.1.4 2Dand 3D Labels14.1.2 CreatingFullyIntroduction $535$ Introduction $535$ Introduction $535$ Introduction $542$ Id.3.1 PDBSearch14.3.2 ConvertPDB to ICM Object14.3.3 ProteinSuperposition14.3.4 ProteinSuperposition14.3.5 ContactAreas14.3.6 HydrogenBond14.3.7 PocketIdentification14.3.8 ProteinPreparationandCrystallographicAnalysisTutorial14.3.9 PDBPreparation14.3.9 PDBPreparation
AvailableNodes $517$ To useICM in KNIME $517$ <b>14 Tutorials</b> 14.1 GraphicsTutorials14.1.1 GraphicsTutorials14.1.2 Annotation $524$ 14.1.3 Labels $529$ 14.1.4 2Dand 3D Labels14.1.4 2Dand 3D Labels14.1.2 CreatingFullyIntroduction $535$ Introduction $535$ Introduction $535$ Introduction $535$ Introduction $535$ Introduction $542$ Tutorial $542$ Intoduction $542$ Intoduction $542$ Intoduction $542$ Intorduction $542$ Intorduction $542$ Id.3 ProteinStuperposition14.3.1 PDBSearch14.3.2 ConvertPDB to ICM Object14.3.3 ProteinSuperposition14.3.4 ProteinStuctureAnalysis $545$ 14.3.5 ContactAreas14.3.6 HydrogenBond14.3.8 ProteinPreparationand $548$ 14.3.9 PDBPreparation14.3.9 PDBPreparation24314.3.9 PDB14.3.9 PDBPreparation549

#### 14 Tutorials

14.4 Cheminformatics Tutorials554
14.4.1 Chemical Draw and Spreadsheet Tutorial554
Background
Tutorial554
14.4.2 Chemical Searching
Background563
Tutorial
14.4.3 Chemical Clustering
Background569
Tutorial
14.4.4 Working with Large Chemical Space
14.4.5 3D Pharmacophore Searching
14.4.6 2D Pharmacophore Searching579
14.4.7 How to create a Markush structure
14.4.8 How to enumerate a Markush library
14.4.9 How to decompose a library based on a Markush structure
structure
14.4.10 How to enumerate a chemical library by reaction588
14.4.11 Detecting Activity Cliffs
14.5 3D Ligand Editor Tutorial
14.6 Atomic Property Field Tutorials
14.6.1 APF Superposition606

# **ICM-Chemist-Pro User Guide v.3.8**

by Ruben Abagyan, Andrew Orry, Eugene Raush, and Maxim Totrov Copyright 2017



Sep 13 2018

Feedback.

# 1 Help Videos on YouTube

You can view webinars and how-to videos on many of the functions within ICM on YouTube.

- Link to our YouTube Channel.Link to our Webinar Archives on YouTube.

1 Help Videos on YouTube

# 2 Reference Guide

### **Chapter Contents:**

ICM-Browser Reference Guide

ICM-Browser-Pro Reference Guide

ActiveICM Reference Guide - Create 3D Molecular Documents for the Web and PowerPoint

ICM-Chemist Reference Guide

ICM-Chemist-Pro Reference Guide

#### Menu and Tab Reference Guide:

Menu Option Guide

Tab Guide

# 2.1 Graphics Reference Guide

Available in the following product(s): ICM-Browser   ICM-Browser-Pro   ICM-Chemist   ICM-Pro
For instructions on how to use ICM-Browser to make fully-interactive 3D slides and publish them in PowerPoint and the web please see the ActiveICM User Guide. ActiveICM is a free plugin for Windows PowerPoint and web browsers. Other related tutorials include:

- Graphical Display: Molecule Representation, Coloring, Labeling and Annotation
- Graphical Selections Tutorial
- Creating Fully Interactive Slides for PowerPoint and the Web Tutorial

### 2.1.1 Download and Install ICM-Browser

Getting Started: Download and Install ICM-Browser and ActiveICM.		
Download ICM-Browser Distribution.	Download	video
Install ICM-Browser Instructions.	Windows Linux Mac	
Download ActiveICM Distribution.	Download	video
Install ActiveICM.	Windows Linux Mac	

## 2.1.2 How to use the Graphical Display

```
'; winRef.document.write(str); }
```

How to use the Graphical Display		
How to search the PDB.	HTML GUI Manual	video
How to Move a Structure in the Graphical Display.	HTML GUI Manual	video
How to use the Graphics window controls.	HTML GUI Manual	
How to use the ICM Workspace Panel	HTML GUI Manual	video
How to Display a Molecule.	HTML GUI Manual	video
How to Change Protein Representation.	HTML GUI Manual	video
How to Change Ribbon Representation.	HTML GUI Manual	video
How to color wire or xstick carbon atoms.	HTML GUI Manual	video
How to Display the Residues Surrounding the Ligand Binding Pocket.	HTML GUI Manual	video
How to remove chain breaks (dotted lines).	HTML GUI Manual	video
How to Color.	HTML GUI Manual	video
How to Change the Color of Molecule Representations.	HTML GUI Manual	video
How to Change the Background Color.	HTML GUI Manual	video
How to Display a Binding Pocket Surface	HTML GUI Manual	video
How to (Un)Display Hydrogens.	HTML GUI Manual	video
How to Save an ICM Object.	HTML GUI Manual	video
How to Save an ICM Project File.	HTML GUI Manual	video
How to Drag and Drop.	HTML GUI Manual	
How to: Right Click Options.	HTML GUI Manual	

How to Move Windows.	HTML GUI Manual	video
How to Arrange Windows	HTML GUI Manual	video

# 2.1.3 How to make Graphical Selections

How to Make Selections.	HTML GUI Manual	
How to Select an Object	HTML GUI Manual	video
How to Select a Molecule	HTML GUI Manual	video
How to Select Residues	HTML GUI Manual	video
How to Select Atoms	HTML GUI Manual	video
How to Make a Spherical Selection.	HTML GUI Manual	video
How to Invert a Selection.	HTML GUI Manual	video
How to Remove a Selection.	HTML GUI Manual	video
How to Change the Selection Level and Mode.	HTML GUI Manual	video
How to Check What is Selected.	HTML GUI Manual	

# 2.1.4 How to Convert Proteins, Display Hydrogens and Ligand Binding Pocket.

Convert Protein, Display Hydrogens and Ligand Binding Pocket.		
How to Convert a PDB Structure into an ICM Object.	HTML GUI Manual	video
How to Display Ligand Binding Pocket.	HTML GUI Manual	video
How to Display Hydrogen Bonds.	HTML GUI Manual	video

# 2.1.5 How to change Graphics Effects

How to change Graphics Effects		
How to display the FOG effect.	HTML GUI Manual	video
How to display side-by-side stereo.	HTML GUI Manual	
How to display Anaglyph stereo.	HTML GUI Manual	
How to toggle full screen mode.	HTML GUI Manual	video

How to adjust perspective.	HTML GUI Manual	
How to change the lighting.	HTML GUI Manual	video
How to display sketch accents.	HTML GUI Manual	video
How to display elegant ribbon and ligand sketch.	HTML GUI Manual	video

# 2.1.6 How to add Labels and Annotations

How to add Labels and Annotations		
How to Label Residues.	HTML GUI Manual	video
How to Label Atoms.	HTML GUI Manual	video
How to Label Variables.	HTML GUI Manual	video
How to Display and Undisplay Sites.	HTML GUI Manual	
How to Make and Display Sites (Annotations).	HTML GUI Manual	
How to Make and Display 2D and 3D Labels.	HTML GUI Manual	video

# 2.1.7 How to Make High Quality Publication Images

How to Make High Quality Publication Images		
How to Toggle High Quality Display	HTML GUI Manual	video
How to Toggle Antialiasing.	HTML GUI Manual	video
How to Copy Image to ClipBoard	HTML GUI Manual	video
How to Write an Image.	HTML GUI Manual	video
How to Use the Advanced Write Image Options.	HTML GUI Manual	video
How to Add an Image to the ICM Photo Album.	HTML GUI Manual	video

# 2.1.8 How to Superimpose Protein Structures.

How to Superimpose Protein Structures.		
How to Superimpose Two or More Protein Structures.	HTML GUI Manual	video

## 2.1.9 How to Measure Distances and Angles.

How to Measure Distances and Angles.		
How to Measure Distances Between Two Atoms.	HTML GUI Manual	video
How to Measure Distances From One Atom to Many.	HTML GUI Manual	video
How to Show Corresponding Distances in Two Objects.	HTML GUI Manual	video
How to Display the Ruler Bar.	HTML GUI Manual	video

# 2.2 ICM-Browser-Pro Reference Guide

**NOTE:** ICM-Browser-Pro contains all the features in ICM-Browser. Click here for the ICM-Browser Reference Guide.

### 2.2.1 Download and Install ICM-Browser-Pro

Getting Started: Download and Install ICM-Browser-Pro	
Download ICM-Browser-Pro Distribution.	Download
Install ICM-Browser-Pro.	Windows Linux Mac

## 2.2.2 Graphics

Graphics and Movie Making	
How to generate the shadow effect.	HTML GUI Manual
How to make a screenshot movie.	HTML GUI Manual
How to make a movie from a set of slides.	HTML GUI Manual
How to move a molecule independently of the other display objects (Connect).	HTML GUI Manual

## 2.2.3 Protein Structure Analysis

Protein Structure Analysis	
How to calculate contact areas between molecules.	HTML GUI Manual
How to identify closed cavities.	HTML GUI Manual

2.1.9 How to Measure Distances and Angles.

How to calculate surface area.	HTML GUI Manual
How to generate interactive Ramachandran plots.	HTML GUI Manual

# 2.2.4 Surfaces

Surfaces	
How to generate electrostatic and binding property surfaces.	HTML GUI Manual
How to connect and rotate/translate surface (mesh).	HTML GUI Manual
How to crop a mesh/surface.	HTML GUI Manual
How to save a mesh/surface.	HTML GUI Manual

# 2.2.5 Superimpose Proteins

Superimpose Proteins	
How to superimpose proteins based on 3D by visible atoms, C-alpha, backbone or heavy atoms.	HTML GUI Manual
How to superimpose multiple proteins based on aligned residues	HTML GUI Manual

# 2.2.6 Crystallographic Tools

Crystallographic Tools	
How to contour electron density.	HTML GUI Manual
How to display crystallographic cell.	HTML GUI Manual
How to display crystallographic symmetry.	HTML GUI Manual
How to convert x-ray density to a grid.	HTML GUI Manual

# 2.2.7 Sequence Analysis

Sequence Analysis	
How to annotate an alignment - box and shade.	HTML GUI Manual
How to annotate an alignment with text.	HTML GUI Manual
How to display secondary structure in an alignment.	HTML GUI Manual

How to extract sequences from pdb files.	HTML GUI Manual
How to assign secondary structure.	HTML GUI Manual
How to link sequence, alignments, and structures.	HTML GUI Manual
How to save as image, print, and delete sequences and alignments.	HTML GUI Manual

# 2.2.8 Plotting Tools

Plotting Tools	
Make fully interactive colorful X-Y plots and histograms with up to 4 dimensions.	HTML GUI Manual
Save plot and histogram as image.	HTML GUI Manual

## 2.3 ActiveICM Reference Guide - Create 3D Molecular Documents for the Web and PowerPoint

This guide is focused on how to make fully interactive 3D documents for Windows PowerPoint and the Web. For more information on the other features in ICM-Browser please see the ICM-Browser User Guide.

# **Creating 3D Documents Is Straightforward**

Creating fully interactive 3D documents for PowerPoint, the web, and standalone browser is straightforward.

- 1. Download ICM-Browser and the ActiveICM plugin. They are completely free! [video]
- 2. Open the ICM-Browser and make a series of animated fully-interactive slides showing different colored and rendered views of your molecules. [video]
- 3. Add hyperlinked HTML text to annotate and link to your slides. [video]
- Save your file in ICM-Browser and then insert into PowerPoint or the web using the ActiveICM plugin. You can also share your documents in the standalone ICM-Browser. [video - powerpoint] [video -web browser]

'; winRef.document.write(str); }

## 2.3.1 Getting Started

Getting Started: Download and Install ICM-Browser and ActiveICM.		
Download ICM-Browser Distribution.	Download	video

Install ICM-Browser Instructions.	Windows Linux Mac	
Download ActiveICM Distribution.	Download	video
Install ActiveICM.	Windows Linux Mac	

# 2.3.2 How to Create a Series of Fully-Interactive 3D Slides.

<b>Creating Slides</b> How to Create a Series of Fully-Interactive 3D Slides.	video
How to Make Fully Interactive 3D Slides	HTML GUI Manual
How to Animate Slides	HTML GUI Manual
How to View and Navigate Slides in the ICM-Browser.	HTML GUI Manual
How to Edit Slides.	HTML GUI Manual
How to Add Smooth Blending and Transition Effects Between Slides.	HTML GUI Manual

# 2.3.3 How to Create Molecular Documents

How to Create Molecular Documents: Linking Slides to HTML Text.	video
How to Create an HTML Document.	HTML GUI Manual
How to Edit an HTML Document.	HTML GUI Manual
How to Make a Hyperlink Between HTML Text and a Slide.	HTML GUI Manual

# 2.3.4 How to Display Molecular Documents in PowerPoint

How to Display Molecular Documents in PowerPoint	video
How to Embed in Microsoft PowerPoint 2003	HTML GUI Manual
How to Embed in Microsoft PowerPoint 2007	HTML GUI Manual
How to Use ActiveICM in PowerPoint	HTML GUI Manual

How to Change ActiveICM Component Properties in	HTML GUI
PowerPoint	Manual
Advanced use of ActiveICM: Macros to direct visualisation changes.	HTML GUI Manual

# 2.3.5 How to Display Molecular Documents on the Web

How to Display Molecular Documents in Web Browsers	video	
How to Display Molecular Documents in Web Browsers	HTML GUI Manual	

# 2.4 ICM-Chemist Reference Guide

## 2.4.1 How to Import, Sketch, and Edit Chemicals

How to Import, Sketch, and Edit Chemicals		
How to load the ICM Molecular Editor.	HTML GUI Manual	video
How to sketch chemicals in the ICM Molecular Editor.	HTML GUI Manual	video
How to save a 2D sketch into a chemical spreadsheet.	HTML GUI Manual	video
How to save a 2D sketch in mol format.	HTML GUI Manual	video
How to use SMILES strings to sketch a chemical.	HTML GUI Manual	video
How to load a mol, sdf or mol2 file.	HTML GUI Manual	video
How to extract a 2D sketch of a ligand in complex with a PDB structure.	HTML GUI Manual	video

# 2.4.2 How to Work with Chemical Spreadsheets

Working with Chemical Spreadsheets		
How to add columns into a chemical spreadsheet.	HTML GUI Manual	video
How to sort a column(s) in a chemical spreadsheet.	HTML GUI Manual	video
How to change the view of a chemical spreadsheet - form, table and grid.	HTML GUI Manual	video

	+	
How to copy, cut and paste columns and rows in a chemical spreadsheet.	HTML GUI Manual	video
How to show and hide columns and rows in a chemical spreadsheet.	HTML GUI Manual	video
How to save a chemical spreadsheet in sdf format.	HTML GUI Manual	video
How to export your chemical spreadsheet into Excel.	HTML GUI Manual	video
How to print a chemical spreadsheet.	HTML GUI Manual	video
How to filter columns in a chemical spreadsheet.	HTML GUI Manual	video
How to use find and replace in a chemical spreadsheet.	HTML GUI Manual	video
How to mark and label rows in a chemical spreadsheet.	HTML GUI Manual	video
How to insert hyperlinks to the PDB, PubMed, and Uniprot.	HTML GUI Manual	video
How to copy and paste 2D chemicals.	HTML GUI Manual	video
How to edit data inside a chemical spreadsheet.	HTML GUI Manual	video
How to remove salts, explicit hydrogens and standardize chemical groups.	HTML GUI Manual	video
How to calculate chemical properties in a chemical spreadsheet.	HTML GUI Manual	video
How to identify duplicate chemicals in a chemical spreadsheet.	HTML GUI Manual	video
How to compare two chemical spreadsheets.	HTML GUI Manual	video
How to merge two chemical spreadsheets.	HTML GUI Manual	video

# 2.4.3 How to Undertake a Chemical Search

How to Perform Chemical Searching		
How to setup a chemical search.	HTML GUI Manual	video
How to draw a chemical search query.	HTML GUI Manual	video

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How to add conditions to your chemical search.	HTML GUI Manual	video
How to search chemical spreadsheets, local databases and MolCart.	HTML GUI Manual	video
How to send a chemical search query.	HTML GUI Manual	video

# 2.4.4 How to Work with Pharmacophores

How to Work with Pharmacophores		
How to draw 2D pharmacophore.	HTML GUI Manual	video
How to search a 2D pharmacophore.	HTML GUI Manual	video
How to draw 3D pharmacophore.	HTML GUI Manual	video
How to extract a 3D pharmacophore from a ligand.	HTML GUI Manual	video
How to edit a 3D pharmacophore.	HTML GUI Manual	video
How to send a 3D pharmacophore search query.	HTML GUI Manual	video
How to color a 2D chemical sketch by pharmacophore feature.	HTML GUI Manual	video

# 2.4.5 How to Perform Chemical Clustering

How to Perform Chemical Clustering		
How to perform chemical clustering.	HTML GUI Manual	video
How to select representative centers from a tree.	HTML GUI Manual	video
How to reorder branches and change the distance of trees.	HTML GUI Manual	video
How to edit the tree - labels, spacing and coloring.	HTML GUI Manual	video

# 2.4.6 How to Generate Stereoisomers and Tautomers

How to Generate Stereoisomers and Tautomers		
How to generate stereoisomers.	HTML GUI Manual	video
How to generate tautomers.	HTML GUI Manual	video

## 2.4.7 How to Generate Combinatorial Libraries

How to Generate Combinatorial Libraries		
How to enumerate a Markush library.	HTML GUI Manual	video
How to decompose a library based on a Markush structure.	HTML GUI Manual	video
How to create a Markush structure.	HTML GUI Manual	video
How to enumerate a chemical library by reaction.	HTML GUI Manual	video

## 2.4.8 How to Generate Plots and Histograms

How to Generate Plots and Histograms		
How to make a histogram.	HTML GUI Manual	video
How to make an X-Y scatter plot.	HTML GUI Manual	video

# 2.5 ICM-Chemist-Pro Reference Guide

# 2.5.1 How to use the ICM 3D Ligand Editor

ICM Chemist-Pro Contains All the Tools in ICM-Chemist	ICM-Chemist Reference Guide	
ICM 3D Ligand Editor: Setup		
How to setup the ligand in the ICM 3D Ligand Editor.	HTML GUI Manual	video
How to setup the receptor in the ICM 3D Ligand Editor.	HTML GUI Manual	video
How to change the 3D Ligand Editor preferences.	HTML GUI Manual	video
ICM 3D Ligand Editor: Display		
How to display and undisplay the ligand surface representation in the ICM 3D Ligand Editor.	HTML GUI Manual	video

How to display hydrogen bonds in the ICM 3D ligand editor.	HTML GUI Manual	video
How to display energy atomic circles in the ICM 3D Ligand Editor.	HTML GUI Manual	video
How to display and undisplay hydrogen atoms in the ICM 3D Ligand Editor.	HTML GUI Manual	video
How to display unsatisfied hydrogen bonds in the ICM 3D Ligand Editor.	HTML GUI Manual	video
How to center on a ligand in the ICM 3D Ligand Editor.	HTML GUI Manual	video
ICM 3D Ligand Editor: Edit Ligand		
How to begin editing your ligand in the ICM 3D Ligand Editor.	HTML GUI Manual	video
How to undo and redo changes in the ICM 3D Ligand Editor.	HTML GUI Manual	video
How to add and sample new substituents to your ligand in the ICM 3D Ligand Editor.	HTML GUI Manual	video
How to sample more than one substituent at a time in the ICM 3D Ligand Editor.	HTML GUI Manual	video
How to edit the ligand in 2D in the ICM 3D Ligand Editor.	HTML GUI Manual	video
How to evaluate the SCORE and ligand strain.	HTML GUI Manual	video
How to add an edited ligand to a chemical spreadsheet (table).	HTML GUI Manual	video
How to change the size of the ligand binding pocket - change purple box size.	HTML GUI Manual	video
ICM 3D Ligand Editor: Docking and Minimization		
How to perform ligand minimization in the ICM 3D Ligand Editor.	HTML GUI Manual	video
How to re-dock a ligand in the ICM 3D Ligand Editor.	HTML GUI Manual	video
How to restrain (tether) atoms during docking.	HTML GUI Manual	video
How to screen databases of chemical substituents.	HTML GUI Manual	video
How to sample linkers between two chemical fragments.	HTML GUI Manual	video
	1	

## 2.5.2 How to Convert Chemicals to 3D

Convert Chemicals to 3D		
How to convert 2D chemical sketches to 3D.	HTML GUI Manual	video
How to convert 3D chemicals to 2D.	HTML GUI Manual	video
How to generate 3D ligand conformers.	HTML GUI Manual	video

## 2.5.3 Ligand Energetics

Ligand Energetics		
How to calculate conformation entropy.	HTML GUI Manual	video
How to calculate conformational strain.	HTML GUI Manual	video

## 2.5.4 How to Superimpose Chemicals

How to Superimpose Chemicals		
How to Perform Rigid Chemical Substructure Superposition.	HTML GUI Manual	video
How to Perform Flexible Chemical Substructure Superposition.	HTML GUI Manual	video
How to use Atomic Property Fields for Chemical Superposition	HTML GUI Manual	video

# 2.5.5 How to Perform QSAR

QSAR		
How to build a QSAR prediction model.	HTML GUI Manual	video
How to apply a QSAR prediction model.	HTML GUI Manual	video

# 2.6 Menu Option Guide

**Note:** Click **Next** (top right hand corner) to navigate through this chapter. Headings are listed on the left hand side (web version) or by clicking the **Contents** button on the left-hand-side of the help window in the graphical user interface.

Here we describe all the options in the drop down graphical user interface menus.

S pep Molsoft icm 3.7.2-	[NewProject *] (1 object)				
Eile Edit View Bioinfo 1	ools Homology Chemistry Docking	MolMechanics Windows	Help		
B 🛱 🖥 🔓 🤊 🐼	11 GA C : M H K & B 8	FOG S & E 🖸	A 🧕 9	1 🐔 🛛 🖬 1	t ff i
Lispius ( ingisic ( iducis ( )	inestines if search in yand if movie i				
Workspace Panel	Menu Options	8×	-		-
no selection		4			

### 2.6.1 File Menu

### 2.6.1.1 New

This option allows you to create new peptides, dna, sequences etc... and is described in the Create New Objects chapter.

### 2.6.1.2 New ICM Session

File/New ICM Session

This option allows you to generate a new clear ICM session or clone the current one.

#### 2.6.1.3 Open

File/Open

This option is described in the Open and Read chapter.

#### 2.6.1.4 Open with Password

File/Open with Password

This option is described in the Open and Read chapter.

### 2.6.1.5 Extract from ICB

File/Extract from ICB

This option is described in the Open and Read chapter.

#### 2.6.1.6 Convert to Local Database

Please see the Local Databases chapter for more information about this option.

### 2.6.1.7 Load

Options contained within the menu File/Load

PDB - read PDB from FTP, http, and local PDB

**From Multiple Object File** - A multiple object file will have a file extension \*.ob and you can select which member of the multiple object is displayed.

PFam Alignment - PFam is a collection of multiple sequence alignments enter FASTA ID

SwissProt - Download SwissProt sequence.

All Images from Dir - Read into ICM multiple image files png or jpg.

Electron Density Map - Download electron density map from Uppsala electron density server http://eds.bmc.uu.se/eds/

3D Mesh in KMZ or COLLADA Format from Google - see http://sketchup.google.com/3dwarehouse/ to download KMZ or COLLADA.

#### 2.6.1.8 Save Project

This is described in the Save File chapter.

#### 2.6.1.9 Save Project As

This is described in the Save File chapter.

#### 2.6.1.10 Save Project Compatible with ICM 3\_5

#### File/Save Project Compatible with ICM 3\_5

Use this option to save a version of your ICM project compatible with an older version of ICM. Version 3.5 or older. If you have an ICM license you can update your version of ICM by visiting our support site at www.molsft.com/support

#### 2.6.1.11 Save with Password

To save a project which is protected by a password:

- File/Save with Password
- Enter a file name or browse for a previously saved project.
- Enter a password
- Determine whether you want the file to be Fully Protected, read only or Read Only and Allow Comments .

#### 2.6.1.12 Export as ActiveICM Html

To embed in a web browser.

- 1. Download ActiveICM from here http://www.molsoft.com/getbrowser.cgi?product=activeicm&act=list (it is free!).
- Create an HTML page in ICM (File/New/Html).
   Add a series of slides.
- 4. File/Export As ActiveICM Html..

#### 2.6.1.13 Close Project

To close a project:

**File/Close Project** 

#### 2.6.1.14 Quick Image

See the High Quality Publication Image chapter.

#### 2.6.1.15 Write Image

See the High Quality Publication Image chapter.

#### 2.6.1.16 Preferences

Preferences are described in the Preferences chapter.

#### 2.6.1.17 Recent Files

Recently viewed projects and files can be easily downloaded from the "Recent Files" option. To access this:

- Select File/Recent Files.
- Select the desired project by clicking on it once.

#### 2.6.1.18 Recent PDB Codes

Quickly retrieve and display PDB structures that have recently been viewed.

- Select File/Recent PDB Codes
- Select desired PDB code by clicking on it once and it will be loaded into the graphical display.

#### 2.6.1.19 Quit

Need to close down ICM - no problem. You do one of the following:

- 1. Select File/Quit. ICM will quit without saving files.
- 2. Save and Click **X** at the upper right corner of the ICM window.
- 3. Type quit in the terminal window.

**NOTE:** You may want to save the icm session as an ICM Project file before quiting.

### 2.6.2 Edit Menu

#### 2.6.2.1 Delete

This option will delete anything that is selected.

#### 2.6.2.2 Delete All

This option will delete everything e.g. sequences, structures, tables ... Use with care!

#### 2.6.2.3 Select All

This option will select everything e.g. sequences, structures, tables...

#### 2.6.2.4 Search in Workspace

This option allow you to search for a particular text in the workspace

#### 2.6.2.5 Selection

This option allows you to make a precise selection either by neighbors or specifying a particular atom or neighbor. Click on the tabs to jump between selection levels.

#### 2.6.2.6 Invert Selection

This option will select everything that is not currently selected.

#### 2.6.2.7 Clear Selection

This option will remove all selections. For more information on selections see the Making Selections Chapter.

#### 2.6.2.8 Neighbor Selection

This option will allow you to select neighboring atoms. For more information see the Select Neighbors section in the Selections Chapter.

#### 2.6.2.9 Undo

Due to the complexities of working in an internal coordinates environment not everything can be undone or redone. Certain things like coloring and representations can be undone or redone.

#### 2.6.2.10 Redo

Due to the complexities of working in an internal coordinates environment not everything can be undone or redone. Certain things like coloring and representations can be undone or redone.

#### 2.6.2.11 Restore Recent Backup

ICM periodically makes a backup of your ICM project. If for whatever reason you lose an ICM session and you want to load the backup for the file use:

#### **Edit/Restore Recent Backup**

#### 2.6.2.12 PDB Search

See PDB Search Tab

#### 2.6.2.13 PDB Search by Field

See PDB Search Tab

#### 2.6.2.14 PDB Search by Identity

See PDB Search Tab

#### 2.6.2.15 PDB Search by Homology

See PDB Search Tab

#### 2.6.2.16 PDB Search with External Sequence

See PDB Search Tab

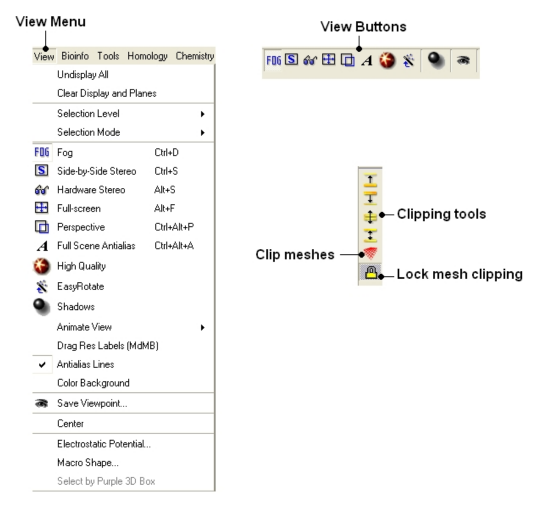
#### 2.6.2.17 Ligand Tools

See the ligand editor section of the manual.

#### 2.6.2.18 Ligand Editor Preferences

See the ligand editor section of the manual.

#### 2.6.3 View Menu



#### 2.6.3.1 Undisplay All

#### To undisplay everything currently displayed in the graphical display

• View/Undisplay All

**Note** For more details on displaying structures please see the GUI Overview chapter.

#### 2.6.3.2 Clear Display & Planes

#### To clear the display and planes

• View/Clear Display and Planes

**NOTE:** For more details on planes please see the sections on clipping tools and mesh clipping.

#### 2.6.3.3 Selection Level

There are four levels of selection - atom, residue, molecule and object. For more details on selections please see the Making Selections section.

#### 2.6.3.4 Selection Mode

There are four different ways to make selections - new, add, remove and toggle. For more details on selections please see the Making Selections section.

#### 2.6.3.5 Fog

**Fog Toggle**(Ctrl + D) : this feature creates a fog-like environment for your object, so that the part of your structure that is closer appears clear and the distant parts are faded as if they are in fog. The clipping planes control the point at which the fog begins.

• View/Fog

#### 2.6.3.6 Side-by-Side Stereo

Side-by-side stereo toggle(Ctrl + S) : this feature allows you to view your structure in 3D form without any 3D goggles.

• View/Side-by-Side Stereo

#### 2.6.3.7 Hardware Stereo

Hardware stereo toggle(Alt + S) - if you have 3D goggles and you wish to view your structure in 3D form, this feature will allow you to do so.

• View/Hardware Stereo

#### 2.6.3.8 Full Screen

**Full screen toggle**Alt\_F - this makes your graphical display fill the entire screen. If you wish to exit this mode, press escape.

View/Full Screen

#### 2.6.3.9 Perspective

**Toggle perspective Ctrl\_P** this will add perspective to your structure, enhancing depth in the graphical display.

• View/Perspective

#### 2.6.3.10 Full Scene Antialias

Anti-aliasing is the technique of minimizing the distortion artifacts known as aliasing when representing a high-resolution signal at a lower resolution. Always use this option before making high resolution images.

• View/Full Scene Antialias

#### 2.6.3.11 High Quality

**Toggle High Quality**: this option will give your ICM object better resolution and higher quality. The change in quality is most visible at a high magnification. However, if your object is very large, this feature could slow down your program.

Always use this option before making high resolution images.

• View/High Quality

#### 2.6.3.12 Easy Rotate

**Toggle easy rotation**: this feature is necessary if your structure is very large or perhaps your computer cannot quickly rotate it. It will prevent your structure from fully loading each time you rotate it, therefore speeding up the process.

• View/Easy Rotate

#### 2.6.3.13 Shadows

View/Shadows

See Graphics Effects chapter.

#### 2.6.3.14 Sketch Accents

#### To make images as shown below use:

• View/Sketch Accents

See Graphics Effects chapter.

#### 2.6.3.15 Animate View

This tool is described in more detail in the Molecular Animations and Transitions section.

#### 2.6.3.16 Drag Res Labels

#### To change the location of your residue label:

- Select View/Drag res labels.
- If your mouse has a middle mouse button, then click on handle (as shown) of the label you wish to move, and drag it to your desired area.

Click on this area *dabel* to drag your label.

> • If your mouse does not have a middle mouse button, then click on the Translation icon on the toolbar, and click on the handle (as shown) of the label you wish to move, and drag it to your desired area.

The +/- buttons on the side of the Residue and Atom buttons will shift the label. There are also other **residue label move** options available when you click and hold the residue label button. These options include **Shift to Sidechain Tips, Shift to Calphas**, and **Restore Positions** 

#### 2.6.3.17 Antialias Lines

Use this option to activate antialias lines. It is recommended to leave this option selected.

• View/Antialias Lines

#### 2.6.3.18 Color Background

#### To change the background color

- View/Color Background
- Select a color from the panel and press OK.

This option is also in the more convenient display tab.

#### 2.6.3.19 Save Viewpoint

It is possible to store a current view using the button shown below.

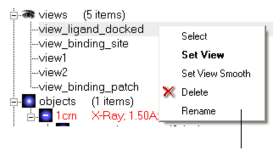


Click on the button and the current view will be stored so that you can view it later. A data entry box will be displayed asking you to name the view. All stored views can be found in the ICM workspace as shown below.

🚊 🗟 views (5 items)	
view_ligand_docked	
view_binding_site	
view1	
view2	
view_binding_patch	

• Double click on the view in the ICM Workspace to display it.

A number of view display options are available by right clicking on the view in the ICM workspace as shown below.



Store current view right click menu

The option in the right click menu called "set view smooth" returns to the view slowly showing the trajectory between the original view and the current one.

#### 2.6.3.20 Center

#### To center on an object displayed in the graphical display

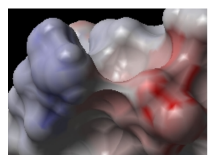
- Make a selection on the region on which you wish to center on.
- Tools/Center (or use the center button on the right hand-side of the graphical display).

#### 2.6.3.21 Electrostatic potential

This option generates the skin representation of the molecular surface colored according to the electrostatic potential calculated by the REBEL method (hydrogen atoms are ignored). REBEL is a method to solve the Poisson equation for a molecule. REBEL is a powerful implementation of the boundary element method with analytical molecular surface as dielectric boundary. This method is fast (takes seconds for a protein) and accurate. REBEL stands for Rapid Exact-Boundary ELectrostatics. The energy calculated by this method consists of the Coulomb energy and the solvation energy

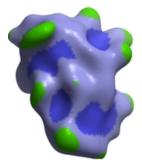
#### In order to color the skin of your molecule by electrostatic potential:

- Select View/Electrostatic potential.
- Enter the potential scale value. This is the local electrostatic potential in kcal/e.u.charge units at which the surface element is colored by extreme red or extreme blue. All higher values will have the same color. This absolute scaling is convenient to develop a feeling of electrostatic properties of molecular surfaces.
- Areas colored blue represent positive areas and red represents negative areas.



#### 2.6.3.22 Macro Shape

A macroshape allows easy viewing and manipulation of a structure. A macroshape representation is ideal for large structures which allows the user to easily identify important regions of the structure and facilitate the return to the 'standard' view of a particular molecule. The level of detail displayed in the macroshape can be controled by changing the number of harmonics, gridStep, and, contour level.



• View/Macro Shape

#### 2.6.3.23 Select by Purple 3D Box

An alternative way to make a make-selection{selection} is to use the purple 3D box. To do this:

- Select the **display** tab and the purple box button
- View/ Select by Purple 3D Box
- The atoms contained within the purple box will be selected.

## 2.6.4 Bioinfo Menu

The tools in this menu are described in the Bioinfo Menu chapter.

### 2.6.5 Tools Menu - Xray

The options in the menu are described in the Crystallographic Analysis chapter.

## 2.6.6 Tools Menu - 3D Predict

The options in this menu are described in the 3D Predict chapter.

#### 2.6.7 Tools Menu - Analysis

The options in this menu are described in the  $\ensuremath{\mathsf{Protein}}$  Structure Analysis chapter.

#### 2.6.8 Tools Menu - Superimpose

The options in this menu are described in the  $\ensuremath{\mathsf{Protein}}$  Superposition chapter.

## 2.6.9 Tools Menu - Extras

#### 2.6.9.1 Plot Function

#### To plot a function:

- Tools/Extras/Plot Function •
- Enter the Function(x) eg Sin(x)
- Enter the starting value of x (From).
- Enter the end point of x (To).
- Enter the number of points (N points).
- Click OK and your plot will be displayed next to a table of values for your function.

### 2.6.10 Tools Menu - Table

#### 2.6.10.1 Build Prediction Model

Learn and Predict tools are described here.

#### 2.6.10.2 Predict

Learn and Predict tools are described here.

#### 2.6.10.3 Cluster Set

This is described in the cluster section of the Working with Tables Chapter.

#### 2.6.10.4 Sort Table

There are a couple of ways to sort a chemical table. You can right click on the a column header and select sort or you can use the option in the menu Chemistry/Sort Table.

- Read a chemical table into ICM.
- Select the columns by which you wish to sort by as shown below. Select **Ascending** or **Descending** and for each **sort by** option and •
- ٠ then click OK

🦻 Sort Table			? 🛛
Table	drug_groups		
Sort By	mol	<ul> <li>Ascending</li> </ul>	C Descending
Then By	IDX 💌	Ascending	C Descending
Then By	MolWeight	<ul> <li>Ascending</li> </ul>	C Descending
		Ok	Cancel

#### 2.6.10.5 Merge Two Sets

#### To merge two tables:

- Read the two tables into ICM.
- Tools/Table/Merge Two Sets
- Select the first table from the drop down list (Table A) and the column you wish to use to merge the table by.
- Select merge method 1. **inner** only molecules present in BOTH A and B tables are kept; or 2. **left** ALL rows of A are kept; or 3. **right** ALL rows of B are kept.
- Select the second table from the drop down list (Table B) and the column you wish to use to mergethe table by.
- Enter a name for the output table.
- Click OK and a new table will be displayed.

🧐 Merge Two	Sets			[	? 🗙
Table A	ricinLigands2D_tauto_1	_tauto 💌	by Column	mol	•
<ul> <li>inner</li> </ul>	C left C right				
Table B	ricinLigands2D_tauto	•	by Column	mol	•
Result Name	T_join 👤				
left - ALL ro	molecules present in BOTH A and ws of A are kept ows of B are kept	B tables are ke	pt		
		Ok	Cancel	Help	

#### 2.6.10.6 Add External Columns

#### To add external columns to a table (e.g. based on a merging key):

- Read at least two tables into ICM the table you want to add to and the table you want to add the column from.
- Tools/Table/Add External Columns
- Enter the target table name and the column you wish to match each table by.
- Enter the source of the new column (Other table and column name)
- Choose to add "all the columns" from the source or "overwrite matching columns" or select the columns you want to add by selecting the "choose column" option.
- Select the columns you wish to add.

#### 2.6.10.7 Append Rows

#### To append rows from one table to another one:

- Read at least two tables into ICM the table you want to add to and the table you want to add the column from.
- Tools/Table/Áppend Rows
- Enter the name of the Target Table (where you will append).
- Enter the name of the Source Table (where you will append from).
- Select whether to Merge Columns

- Check whether you would like to Select New Rows. If you do this you can easily highlight the new rows using the mark row option. Check whether you would like to **Calculate Columns with Formulas**.

## 2.6.11 Tools Menu - Chemical Search

Chemical searching is described in the Chemistry chapter here.

### 2.6.12 Tools Menu - Molecular Editor

The molecular editor is described in the Chemistry chapter here.

## 2.6.13 Homology Menu

The options in this menu are described in the Homology Modeling Chapter.

## 2.6.14 Chemistry Menu

The tools in the **Chemistry** menu are described here.

## 2.6.15 Docking Menu

The tools in the **Docking** menu are described in the Docking chapter.

#### 2.6.16 MolMechanics Menu

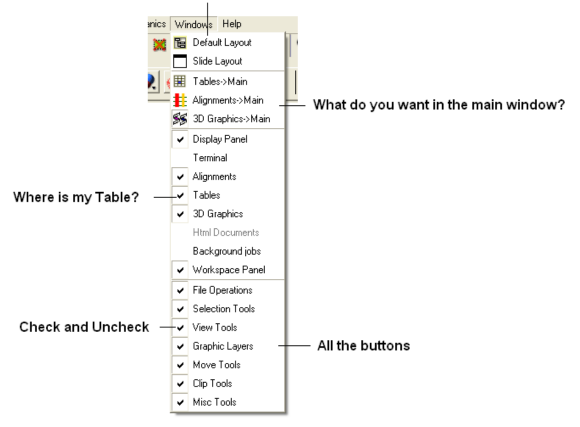
The options in this menu are described in the Molecular Mechanics chapter.

### 2.6.17 Windows Menu

This menu allows you to choose the windows you wish to display. The windows which open automatically when you first open GUI are shown in the Getting Started section. Other windows can be displayed by selecting the windows menu. For example, if you have loaded a table but cannot see it in the GUI it may be because the Tables option in the window menu hasnt been selected.

To add or remove windows from the GUI display select the 'window menu'. Other windows not included in the default display such as tables and alignments can be added.

#### I have windows open everywhere - Please bring some order.



To return to the default display option select the 'Default layout' option in the windows menu.

#### OR

Click the default layout icon.



## 2.7 Tab Guide

In this section we describe the contents of the tabs in the graphical user interface.

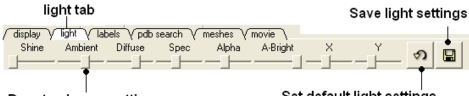
File Edit View	Bininfo Tools Hemalogy C	emistry Docking MolMechanics Windows Help
display V light V	Iabels \/ meshes \/ search \/	
Workspace Panel		8×
no selection	Tab Options	

## 2.7.1 Display Tab

The display tab contains tools for a variety of functions including structural representations, coloring, labeling and superposition. This tab is shown below.

display V light V labels V pdb search V meshes V movie	
₩. /: /:	

## 2.7.2 Light Tab

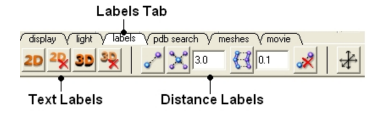


## Drag to change settings

Set default light settings

The options in this tab are described in the Lighting Section.

## 2.7.3 Labels Tab



The options in this tab are described in the labels section of this manual.

## 2.7.4 PDB Search Tab

Instructions on how to use this tab can be found in the  ${\tt Search}\ {\tt PDB}\ {\tt section}$  .

## 2.7.5 Ligedit Tab

Instructions on how to use the options in this tab can be found in the How to use the 3D Interactive Ligand Editor section.

## 2.7.6 Meshes Tab

Click on the tab button entitled **'meshes'** and three different graphical display tools are available for you to use. The three displays are surface, meshes and macroshape and are collectively referred to as meshes.

	surfaces		meshes	-	macroshape
Τ					
🔽 wi	re electrostatic 💌 🛃	[	all · 💽 🚿		N 8 step 0. V color MacroShape
/ displ	lay Y labels Y analysis Y pdb	search	) Y meshes Y mov	/ie	٦

The benefits and applications of each display are described in the section.

## **3 Getting Started**

#### **Chapter Contents:**

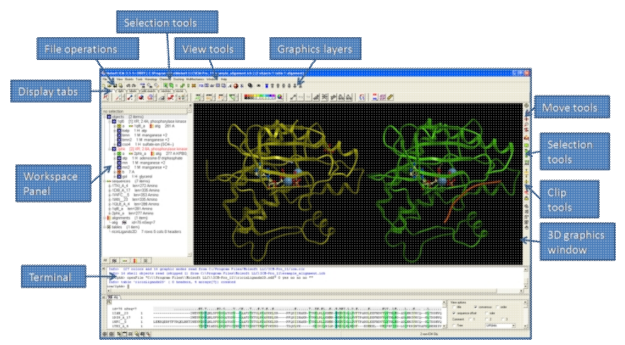
- The Basics of the Graphical User Interface
- How to search and Download Protein Structure, Sequences, and Chemicals
- How to create new Objects
- How to open files
- How to save files
- How to Make Selections
- Changing Preferences

# 3.1 The Basics of the Graphical User Interface

Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro | ICM-Chemist

## 3.1.1 The Components of the GUI

The **Graphical User Interface** (GUI) has many components. When you first use the GUI the default window layout is displayed as shown below.

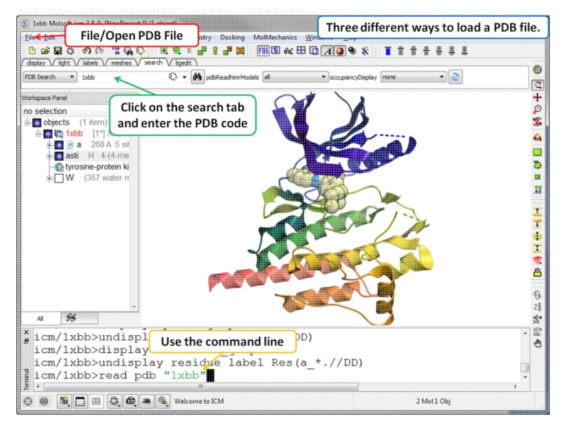


## 3.1.2 How to load a PDB Structure

There are three main ways to read in a PDB file.

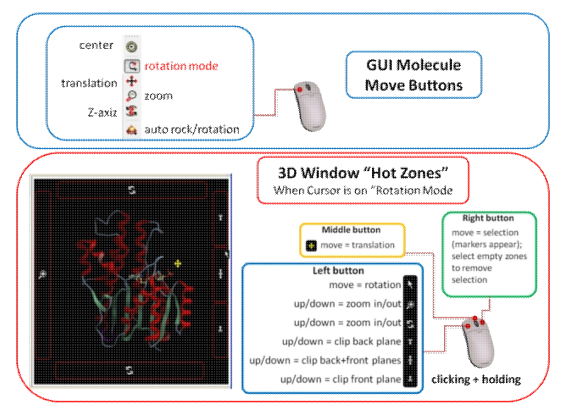
- 1. Using the command line.
- 2. Using File/Open button
- 3. Using the PDB Search tab

Other PDB search options are described in more detail in the PDB Search section of this manual.



# 3.1.3 How to Move a Structure in the Graphical Display

Available buttons and options for moving molecules around the graphical display window. This is described in more detail in the section entitled Move Buttons.



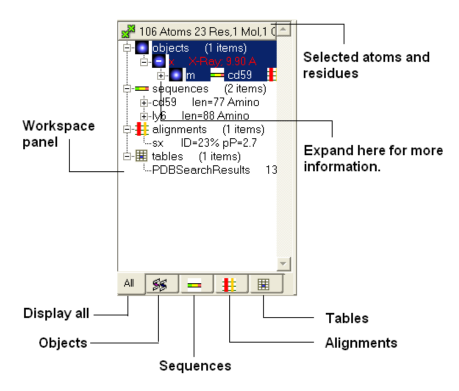
#### How to use the Graphics window controls

In the graphics window you can use various tools described elsewhere but it is helpful to know the following things:

- **Picking a tool**: the left mouse button will function according to the selected tool
- Popup menus: right click on an atom gives a pop-up menu
- Selecting in the rotation mode: the right mouse button will select atoms
- **Translating in the rotation mode**: the middle mouse button will translate the scene
- Zooming and moving clipping planes in the rotation mode: the left, top and right margins of the graphics window are reserved for other actions, zoom, z-rotation, and clipping plains, respectively. That means that even if you are picking atoms, by pressing control you can still rotate your molecule with the left-mouse-button.
- Rotating in any non-rotation mode: if you press Control in any mouse mode, e.g. zoom, pick etc., it will temporarily switch to rotation
- Escaping from the connect and continuous movement modes: pressing Escape helps to get out of certain modes, such as Full Screen, Continuous rotation or rocking, the Connect mode.
- Global rotation in the Connected mode: pressing Shift will temporarily switch to the global rotation/translation mode.

# 3.1.4 How to Display Molecules using the ICM Workspace Panel

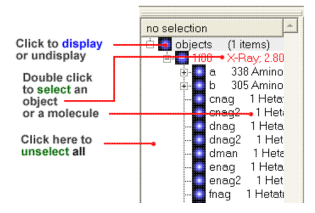
The workspace panel (located on the left hand side panel of the gui) is an important place within the graphical user interface because it displays which sequences, structures, objects, tables and alignments are currently loaded into ICM. From the ICM Workspace panel you can make graphical selections and display and undisplay molecules.



Once a structure has been loaded into ICM the individual components of that structure (i.e. amino acids, metal ions, binding sites etc) are listed in the ICM workspace.

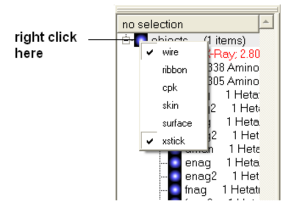
To display every component of the object except for binding sites and water atoms:

• Click on the white box next to the word object at the top of the ICM workspace. This box will be colored blue once the structure is displayed



To display the whole structure in wire, ribbon, cpk, skin, surface and xstick representations:

• Right click on the blue box next to the word object. A menu will be displayed.



• Select which representation you desire for your structure by clicking on the appropriate word. A check mark indicates the representation currently displayed. To un-display a particular representation click on the word again.

In order to clear your graphical display:

• Select View/Undisplay All

If you only wish to display part of the structure then click in the boxes further down the tree in the ICM workspace.

To display selected regions of the molecule in wire, ribbon, cpk, skin, surface and xstick representations:

• Right click on the appropriate box in the **ICM** workspace. A menu will be displayed and select the representation you would like to use (e.g. wire, ribbon etc...)

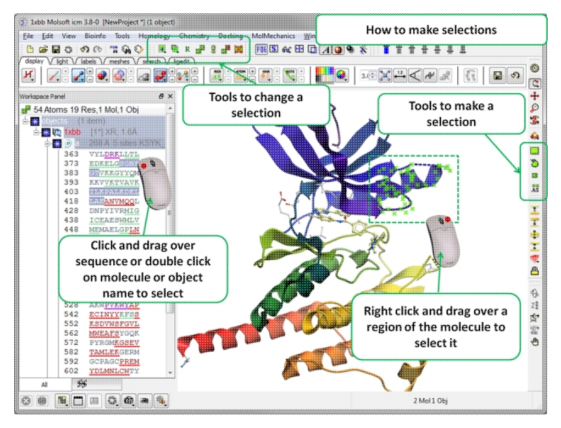
#### 3.1.5 How to Make Selections

Making selections in ICM is an important skill to master (e.g. you may want to select a binding pocket for docking or a region of a molecule for coloring). The four levels of selection are:

- 1. Atoms
- 2. Residues
- 3. Molecules
- 4. Objects (multiple molecules comprising a PDB entry)

There are several ways of making selection in ICM. The simplest is to interact directly with the graphics window - **right-click**, **hold and drag** around the area of the screen you want to select. Alternatively, in the workspace window, expand the tree of molecules and chains until the relevant protein sequences is displayed. Then left click and drag to mark residues to form a selection.

See the chapter entitled Making Selections for more information.

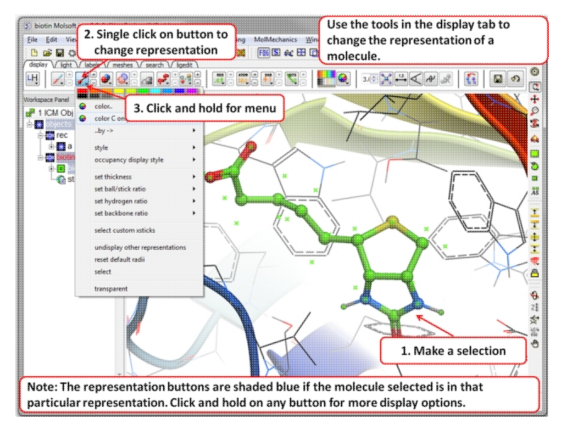


## 3.1.6 How to Change Protein Representation

To change the representation of the protein, make a selection and then use the tools in the display tab.

There are 6 main types of representation:

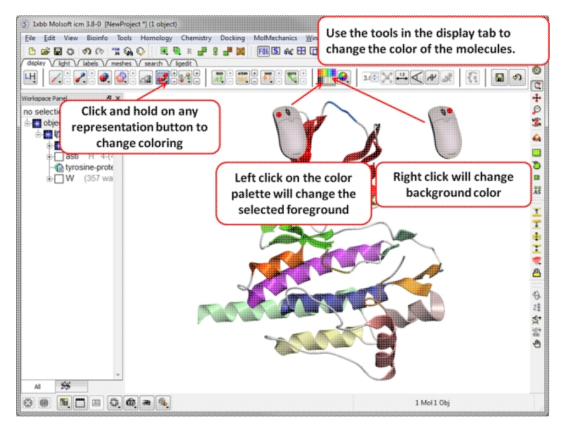
- Wire: Wires connecting covalently bound atoms of a molecule. This representation has no defined thickness as such will not make shadows. Useful for showing the chemical structure of a small molecule.
- **Xstick:** Covalent bonds are represented as cylinders whilst atoms are represented as small spheres.
- **CPK:** Atoms are represented as spheres with their respective van der Waals radius and coloured according to a standard defined by Corey, Pauling and Kultun.
- **Surface:** Solvent accessible surface. This is the center of water sphere as a water probe rolls over the molecule.
- Skin: A Connolly molecular surface over the selection. This is a smooth envelope touching the van der Waals surface of atoms as a water probe rolls over the molecule.
- **Ribbons:** Cartoon representation of protein and DNA secondary structure. Protein residues marked as alpha-helices ('H') are shown as a flat, helical ribbon, those marked as beta-sheets ('E') are shown as a flat ribbon with an arrow-head, and the rest are shown as a cylindrical "worm". If secondary elements are not defined everything will be shown as a cylindrical worm. ICM can automatically assign secondary structure: Tools/3D predict /Assign Helices and Strands



## 3.1.7 How to Color

To change the color of the representation you need to use the buttons in the display tab.

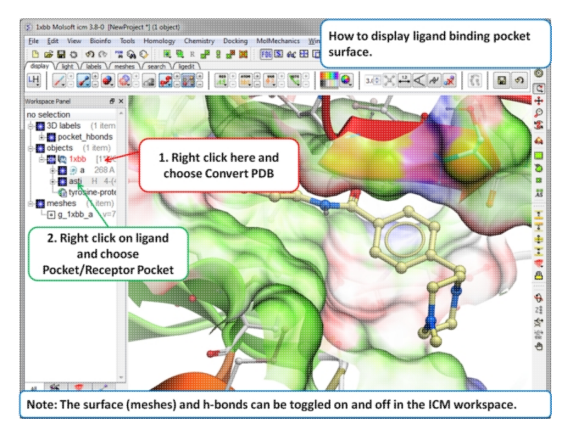
Changing the colour of a representation works in much the same way as displaying the representation itself. The selection rationale is the same followed by clicking on a colour in the palette in the display tab. It is also possible to colour different representations of the same selection independently by clicking and holding on the representation buttons in the display tab.



## 3.1.8 How to Display a Binding Pocket Surface

To display the surface of a small molecule ligand or peptide binding pocket:

- Load the PDB of interest.
- Convert PDB to ICM object. If you do not convert you will not get the properties of the pocket displayed on the surface.
- Right click on the small molecule or peptide in the ICM Workspace and select Ligand Pocket.



## 3.1.9 How to Save an ICM Object

Any ICM object such as a structure, sequence, or alignment, can be saved for use at a later time.

To save an object:

- Right click on the object name in the ICM workspace or ICM alignment editor and a menu will be displayed.
- Click on the Save As... option.
- Enter the unique name you wish to call your object in the box labeled **File name:**
- Choose which folder or directory you wish to save your object by clicking scrolling down in the box labeled **Save in:**
- Choose which file type you would like to save your object as by scrolling down in the box labeled **Save as type**. ICM structure objects should have the file ending yourfilename.ob and alignments yourfilename.ali
- Once the appropriate information has been entered click on the **Save** button in the bottom right hand section of the window.
- The object is now saved.



To save an ICM object or PDB file right click and select SaveAs..

## 3.1.10 How to Save an ICM Project File

All objects contained within an ICM session can be saved in a single file with the extension .icb. The file can then be read into ICM and the exact layout of the file will be preserved. To save a project file go to the **File** menu and select **Save Project**.

## 3.1.11 How to Drag and Drop

**NOTE:** "Drag and Drop" is a useful way of moving objects and sequences around the graphical user interface.

Sequences and objects can be moved around the graphical user interface by dragging and dropping them. All loaded sequences and objects are always displayed in the workspace panel. Select the desired object or sequence from the workspace panel by clicking and holding, move the selection to the desired location and release.

This is a useful application in the graphical user interface. For example, you may have an alignment displayed and you wish to add another sequence to the alignment. This can simply be accomplished by dragging a loaded sequence from the workspace panel into the alignment display panel. Or, you can quickly view an object by dragging and dropping it from the workspace panel into the 3D graphics window.

## 3.1.12 How to: Right Click Options

**NOTE:** If you right click on any object you will see a new menu of options related to that object.

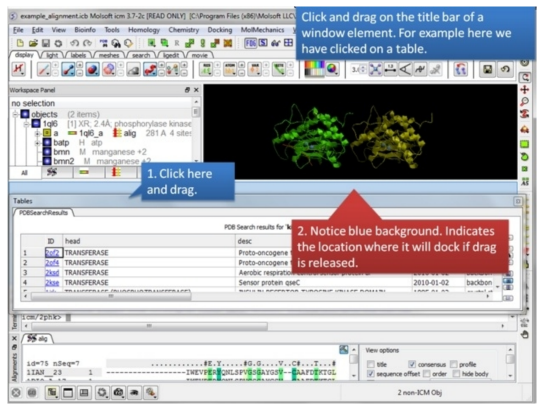
The right click mouse option can be used throughout the graphical user interface. It is a very useful means of opening up a whole new world of menus and options. Most of these options are described in this book. However, when using the graphical user interface it is always a good idea to try right clicking the mouse on an object and seeing which extra options that are available for you to use.

## 3.1.13 How to Move Windows

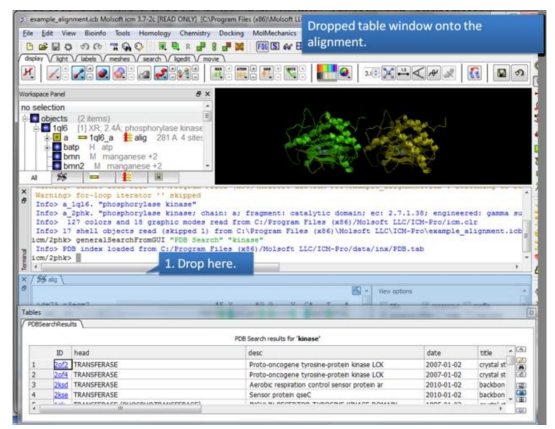
Is your graphical user interface looking a bit messy? Do you have tables, alignments, plots all over the place? Here we show you how to arrange everything to make a clearer GUI environment.

(5) example_alignment.icb Molsoft icm 3.7-2c [READ ONLY] [C\Program	m Files (x86)\Molsoft LLCV How to move wi	ndows arou	ind the ICM
File Edit View Bioinfo Tools Homology Chemistry Docl	Graphical User I	nterface	
6 2 2 5 5 6 7 9 0 2 2 2 R # 8 # 8 #		internace.	
dsplay V light V labels V meshes V search V ligedt V movie			
		<u>*</u>	<u>د</u> ا
Workspace Panel & X			+
no selection			P
biobjects (2 items) biological (2 items)			94 <b>4</b> 0 0 a
Tables			€×
PDBSearchResults			
	PDB Search results for 'kinase'		+
ID head	desc	date	title 🔺 💼 🔒
1 2012 TRANSFERASE	Proto-oncogene tyrosine-protein kinase LCK	2007-01-02	crystal st
2 2014 TRANSFERASE	Proto-oncogene tyrosine-protein kinase LCK	2007-01-02	crystal st 🖉 👳
3 2ksd TRANSFERASE	Aerobic respiration control sensor protein ar	2010-01-02	backbon - 🚍 🦲
۲ ( ا			· @
<pre>x Info&gt; 17 shell objects read (skipped 1) from icm/2phk&gt; generalSearchFromGUI "PDB Search" " Info&gt; PDB index loaded from C:/Program Files icm/2phk&gt;</pre>	kinase" (x86)/Molsoft LLC/ICM-Pro/data/inx/PD	B.tab	2≇ ★* *** • •
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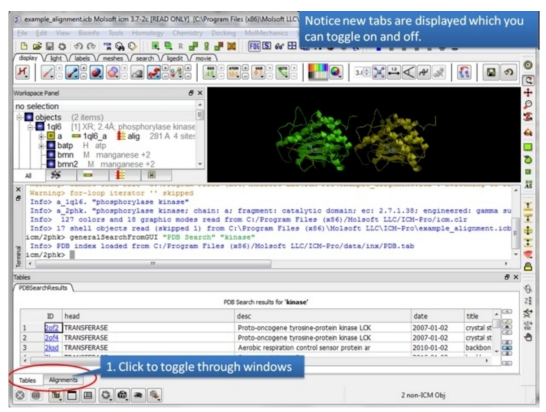
**1.** Too many windows in your GUI? You can move them around to make viewing easier.



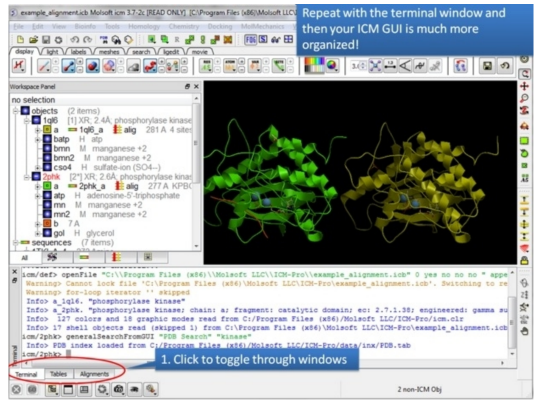
2. Click and drag on a window title bar to move it.



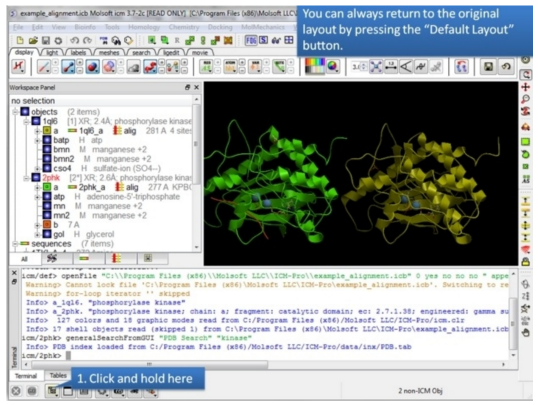
3. Drop one window on another one to dock it.



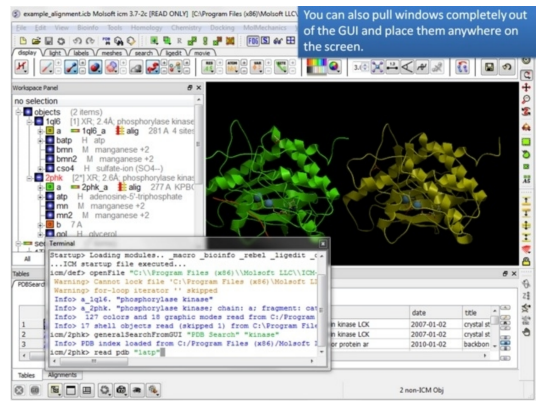
4. You can access each window by clicking on the tabs.



**5.** In this example we have the table, alignment, and terminal all in the same panel



**6.** There is a quick way to return the windows to their original location.



7. You can also move a window to any part of your screen.

**NOTE:** To return to the default display option select the 'Default layout' option in the windows menu.

#### OR

Click the default layout icon.



#### OR

Double click on the window header.

#### 3.1.13.1 How to Arrange Windows

Sometimes when using ICM you may have many items displayed such as structures, alignments and tables. As a default the graphical display is the largest and centered in the middle of the ICM graphical user interface. However if you wish to work on an alignment or table you can place the alignment or table as the main display by clicking on the buttons shown below. The larger display generally makes it easier to manipulate the alignment or table. There are ofcourse other ways to alter the layout such as tier the windows but this option is just a simple click and can sometimes come in useful.



## 3.1.14 How to Make a Picture

There are several ways of taking a picture of the contents of the 3D graphical display window see the write image section. However the easiest way is to simply click on the button in the view tools panel (see image below).

## quick high quality image

File	Edit	View	Bi	oinfo	To	pols	Н	oma
	🚔 🔚	\$	Ś	R	PDB	G,	Q	****
displ	ay \/ I	ight \/	labe	els V	mes	hes	V s	ear
PDB S	Search	•	text	or smi	les			

#### Or select /File/Quick Image

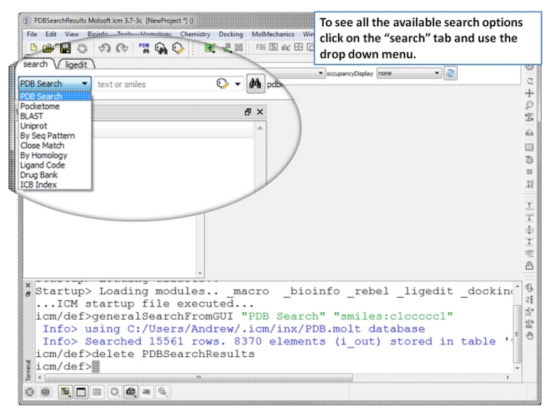
The picture will be automatically saved as a PNG file in the directory from which you loaded ICM. The default picture name is icm[n].png, where n is the number of pictures taken in one ICM session. To save in other picture formats and to change the file name see the write image section.

# **3.2 How to search and download Protein Structure, Sequences, and Chemicals**

Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro | ICM-Chemist

The **Search** tab allows you to search and download content from the following databases:

- The Protein Databank www.wwpdb.org search for protein and chemical
- The Pocketome Database http://pocketome.org/
- BLAST search the NCBI sequence database
- The UniProt Sequence database http://www.uniprot.org/
- Search the PDB by ligand code.
- Search Drug Bank http://www.drugbank.ca/
- Search PubChem https://pubchem.ncbi.nlm.nih.gov/
- Search ChEMBL https://www.ebi.ac.uk/chembl/
- Search SureChEMBL https://www.surechembl.org/search/
- Search the Crystallography Open Database http://www.crystallography.net/



Click on the search tab and then use the drop down menu to identify the database you would like to search and download from.

## 3.2.1 Search the PDB

## How to Search the Protein Databank and Download

The **PDB search tab** provides easy access to the PDB database. You can use keyword searching or type in the PDB code you are interested in. An asterisk (\*) wildcard can be used to list all the pdb files currently available in the protein databank. Different fields can be searched by using the drop down arrow as shown below. More advanced PDB search tools and how to use the PDB search result table are described in the section entititled Searching the PDB.

Once a search is complete a table of PDB files relating to your search query will be displayed. To view the PDB file in 3D in the graphical display double click on a row in the PDBSearchResults table.

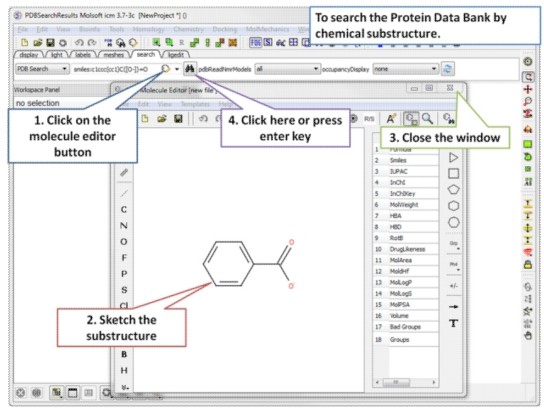
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-PDBSe	1. Enter search string		2012-01-24	Cryst CRY5	ø
	or PDB code	222 enter key	2006-01-25	Cryst	44
	of FDD code	231 2007 TRANSFERASE	2007-01-25	Cryst	₹ <sup>€</sup>
_		232 313 Transferase	2010-01-25	P13-4	
		233 2yom TRANSFERASE	2011-01-25	Cryst	1000
		234 2y60 TRANSFERASE	2011-01-25	Cryst	
		235 300W TRANSFERASE/TRANSFERASE INHIBITOR	2011-01-25	Cryst	Œ
		236 4dgg TRANSFERASE	2012-01-25		128
		237 1kg TRANSFERASE	2002-01-26	The	8.6.0
		238 1www Transferase	2005-01-26	Solut	
		239 2by TRANSFERASE	2005-01-26	Struk	
		240 2cdg TRANSFERASE	2006-01-26	Cryst	
		241 ROOX TRANSFERASE	2007-01-26	Cryst	
		242 ZODY TRANSFERASE	2007-01-26	Cryst	
		243 2202Y Transferase	2008-01-26	Cryst	
		244 3fzo TRANSFERASE	2009-01-26	Cryst	
		245 Hage TRANSFERASE 3. Double click to	012-01-26	CRY!	
		246 4aod TRANSFERASE load structure	12-01-26	CRY!	
			998-01-26	PYRI	
		248 1a3x TRANSFERASE	1998-01-26	PYRI	
		249 2tmk KINASE	1998-01-26	CRY!	
		250 1yes TRANSFERASE	2005-01-27	Cryst	
		251 1yol TRANSFERASE 252 1yom TRANSFERASE	2005-01-27	Cryst	
	-	202 IYON IKARSPERASE	2005-01-27	Cryst -	

#### To Search the PDB and Download a Structure:

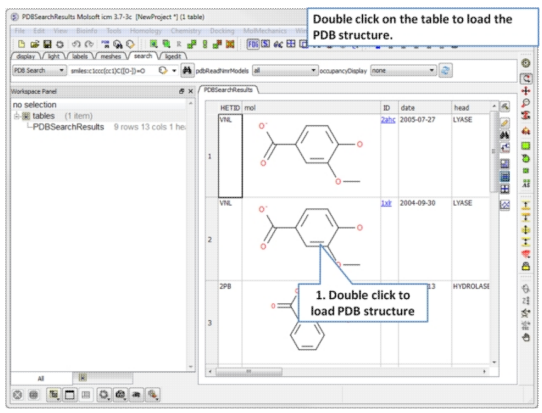
- Click on the search tab and enter a search string or PDB code.
  A table containing the results will be displayed.
  Double click on a row to load the PDB file. Read more about PDB search here.

NOTE: If you have a PDB structure already saved you can read it into ICM by going to the File Menu and selecting Open. PDB files that have been viewed previously can be loaded using File/ Recent PDB Codes.

#### 3.2.1.1 How to Search the Protein Databank by Chemical Substructure



**Step 1:** Click on the molecule editor button inside the search tab. Sketch the substructure you are interested in.



**Step 2:** A table containing the results will be displayed. Double click on a row to load the PDB file.

#### 3.2.1.2 How to Query the PDB by Sequence

There are a number of ways to search the PDB by sequences.

- 1. BLAST search the NCBI datase using the option "Sequences from PDB".
- 2. Search by **Sequence Pattern** using the option in the Edit menu.
- 3. Search by **Identity** using the option in the Edit menu.
- 4. Search by **Homology** using the option in the Edit menu.

#### 3.2.1.3 How to Query PDB by PDB Field

## To query the PDB by field (Author, Compound, PDB Header, Experiment Type, Resolution or Ligand Code

- Select Edit/PDB search by field.
- Enter the search string or value
- Click **OK** and a list of related PDB entries based on your search will be displayed in the PDBSearchResults table of the graphical user interface.

Find PDB Entries by Keywords and Fields							
Authors	i 💌	Experiment Type	×	•			
Compound	× 💌	Resolution Better Than	9.9	•			
Pdb Header	× 💌	Ligand code	×	•			
Update PDB Index							
		<u> </u>	<u>C</u> ancel	<u>H</u> elp			

#### 3.2.1.4 Load and Display NMR Structures

Use the PDB Search tab to load NMR structures from the PDB. You can use the drop down button shown below to determine how you want to display your NMR structure. You can choose to display and download the first NMR model, all models in the PDB file or all models in the PDB in a stack. If you choose the stack option the the stack will be stored in the object as described here.

display / light /	labels / meshes / search / li	igand ( movie )		
PDB Search 💌	2kd2	pdbReadNmrModels	all stack 🔍	occupancyDisplay none
			first all	
Workspace Panel		₽×	all stack	

Click for drop down m

#### 3.2.1.5 Occupancy Display

You can use the options in the PDB Search tab to control if and how the partial or zero atom occupancies are displayed. You can choose to circle or label the poor occupancy atoms.

display light labels meshes search lig	and \/ movie \		
PDB Search 🔽 1xbb	ydbReadNmrModels all stack	💙 occupancyDisplay	circle
			none
Washington Banal	E V		circle
Workspace Panel	₽×		label

>>load-pdb-hyperlinks{pdb search hyperlinks} h4-- Hyperlinks to PDB Website and UniProt {Hyperlinks to Databases}

In the PDB Search Results Table you will see blue hyperlinks that will take you directly to the PDB website or Uniprot website.

#### 3.2.1.6 Display PDB Header

To display the PDB Header for a PDB file.

- First load a PDB file into ICM (see Search PDB)
- Double click on the word header in the ICM Workspace.

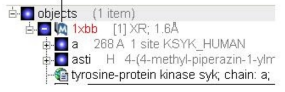
ection	ICM Workspace
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- The PDB Header information will be displayed.
- Click on the blue hyperlinked text to link to external web pages for additional information if needed.

#### 3.2.1.7 Direct link to PubMed

When you search for a PDB file and load it into ICM you will see an icon (shown below) next to your protein name in the ICM Workspace. Click the icon and you will be taken directly to the PubMed primary reference relating to the structure.

#### Direct access to PubMed

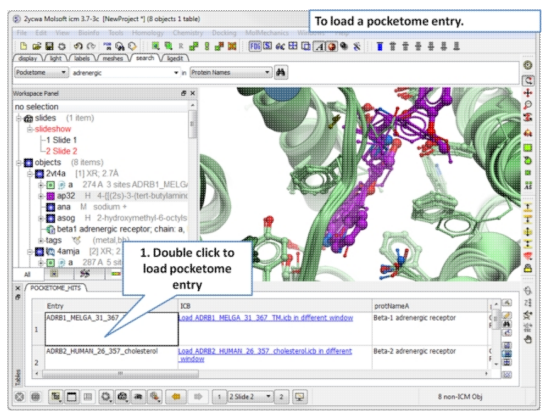


## 3.2.2 Search Pocketome

The Pocketome (www.pocketome.org) is an encyclopedia of conformational ensembles of all druggable binding sites that can be identified experimentally from co-crystal structures in the Protein Data Bank.

5 PD8SeerobRendet Malachien 37.3. Bitmiduein File Edit B & S S String	To search the Pocketome data emistry Docking MolMechanics Wir Pocketome.org	base.
C C C C C C C C C C C C C C C C C C C		<ul> <li>○中</li> <li>○+</li> <li< th=""></li<></ul>

**Step 1:** Click on the search tab and choose the Pocketome option. Check a field you would like to search e.g. Protein Name, Family, domain...



**Step 2:** A table of pocketome hits will be displayed. Browse the results and double click to load a pocketome entry.

## 3.2.3 ChEMBL Search

ChEMBL (https://www.ebi.ac.uk/chembl/) is a manually curated chemical database of bioactive molecules with drug-like properties. It is maintained by the European Bioinformatics Institute (EBI), of the European Molecular Biology Laboratory (EMBL), based at the Wellcome Trust Genome Campus, Hinxton, UK.

To search and download data from ChEMBL:

- Click on the **Search** tab.
- Choose ChEMBL from the drop down button.
- Enter a search string or search by chemical sketch by clicking on the editor button in the panel.
- Select a field to search e.g. Protein Name.
- Select whether you wish to return just activity data or all data.
- Click the search button.

A table will be displayed as shown below.

1. Search Tab/ChEMBL	table)		ChEMBL Search
	anistry Dockie 3. Ch	oose Field	111111
₩£₩8L <u>*</u> ](cox2	🔾 🗾 🏘 in Protein	Name 💌 Return Ki, Kd, IC50, EC50, XC5	50, AC50, and Potency (HT5) data only
mol 2. Enter S	tring	5. Click to search	
	CHEMBL501208	Synonyms unprot to ful PGH2_HUMAN Pro	Get Best Activity Get Ligand Summary A Click here for
6	CHEMBL525247	PGH2_HUMAN P	additional data
	CHEMBL499069	PGH2. HUMANI Pro	6
1	CHEMRI 501 208	PGH2 HIMAN Pr	

The table contains the original activity data without merging or aggregation. For instance,

Chemical 1, Protein 1, Activity 1

Chemical 1, Protein 1, Activity 2

Chemical 1, Protein 2, Activity 1

Chemical 2, Protein 1, Activity 1

Click on the button **Get Best Activity** to display the best activity for each chemical, protein pair. For instance,

Chemical 1, Protein 1, Best(Activity 1, 2, 3, ...) Chemical 1, Protein 2, Best(Activity 1, 2, 3, ...) Chemical 2, Protein 1, Best(Activity 1, 2, 3, ...)

Click on the button **Get Ligand Summary** to display a list of chemicals with all activities found in ChEMBL. For instance,

#### Chemical 1, Protein 1, Activity 1

Protein 1, Activity 2 Protein 2, Activity 1 Protein 3, Activity 1

**Chemical 2 Protein 1, Activity 1** 

Protein 2, Activity 1 Protein 2, Activity 2

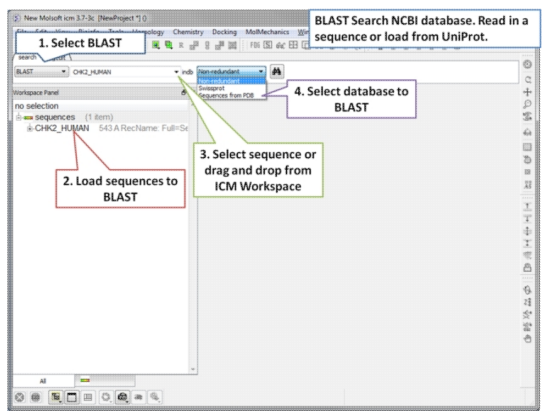
## 3.2.4 Search SureChEMBL

SureChEMBL (https://www.surechembl.org/search/) provides free access to chemical data extracted from the patent literature. To search SureChEMBL:

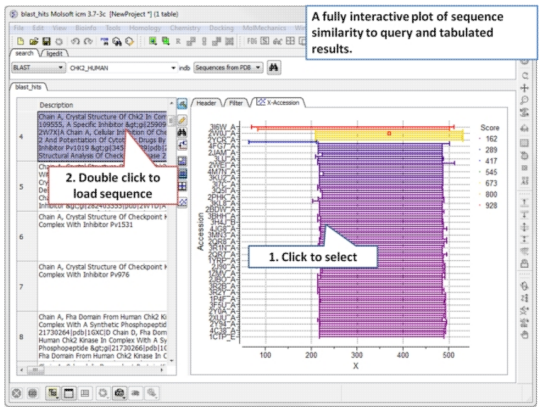
- Click on the **Search** tab.
- Choose SureChEMBL from the drop down button.
- Enter a search string or search by chemical sketch by clicking on the editor button in the panel.
- Select a field to search e.g. Protein Name.
- Select whether you wish to return just activity data or all data.
- Click the search button.

## 3.2.5 BLAST Search

To BLAST search the NCBI sequence database:

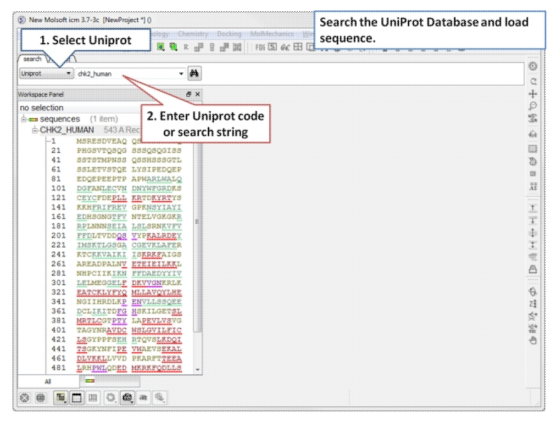


**Step 1:** Load a sequence into ICM. Select the Search tab and choose the BLAST option. Drag and drop the sequence into the search field, use the drop down menu or type the sequence name.



**Step 2:** A fully interactive table and plot of sequence conservation will be displayed. Double click to load a sequence.

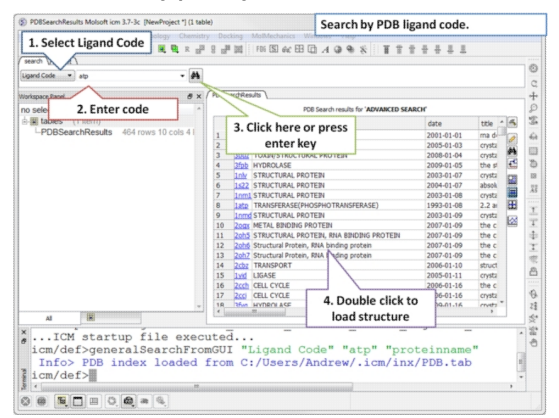
## 3.2.6 Search UniProt



**Step 1:** Click on the search tab and select **Uniprot** from the drop down menu. Enter the UniProt code. The sequence will be loaded directly into ICM and you will see it in the ICM workspace.

### 3.2.7 Search PDB by Ligand Code

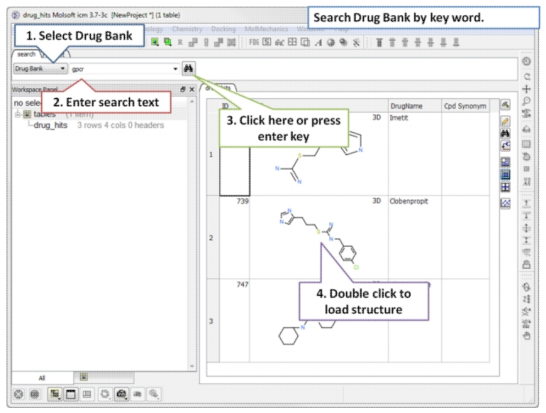
To find structures containing a particular ligand in the PDB.



**Step 1:** Click on the search tab and select **Ligand Code** from the drop down menu. Enter the Ligand code and a table of hits will be displayed.

### 3.2.8 Search Drug Bank

To search Drug Bank (http://www.drugbank.ca/)



**Step 1:** Click on the search tab and select **Drug Bank** from the drop down menu. Enter a search string and a table of results will be displayed. The table contains the original activity data without merging or aggregation. For instance,

Chemical 1, Protein 1, Activity 1 Chemical 1, Protein 1, Activity 2 Chemical 1, Protein 2, Activity 1 Chemical 2, Protein 1, Activity 1

Click on the button **Get Ligand Summary** to display a list of chemicals with all activities found in Drug Bank. The button is located in the extra panel (click on hammer - top right of table panel). For instance, **Chemical 1, Protein 1, Activity 1** 

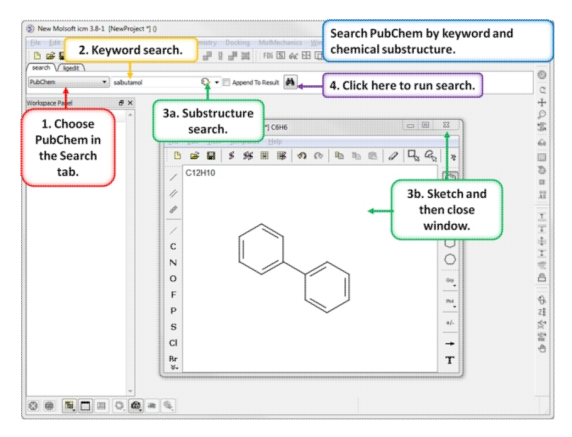
Protein 1, Activity 2 Protein 2, Activity 1 Protein 3, Activity 1

**Chemical 2 Protein 1, Activity 1** Protein 2, Activity 1

Protein 2, Activity 2

### 3.2.9 Search PubChem

To search PubChem (https://pubchem.ncbi.nlm.nih.gov/):



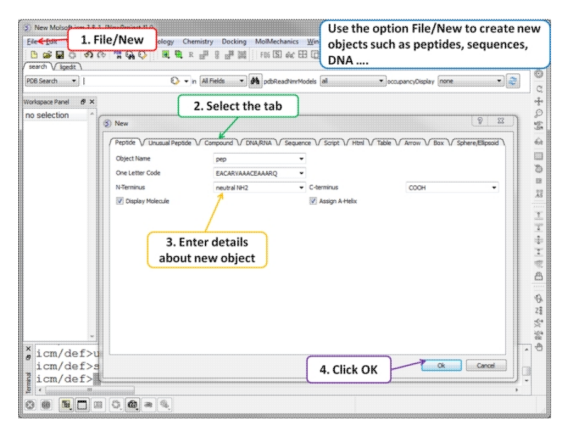
- Click on the search tab and select **PubChem** from the drop down menu.
- Enter a search string and a table of results will be displayed or • click on the chemical sketch button and sketch a substructure. When you close the chemical sketch window the smiles string will be added to the keyword search panel. Click on the "binocular" button and run the search.
- A table of hits will be displayed. •

## 3.3 Create New Objects

Available in the following product(s): ICM-Browser | ICM-Browser-Pro |ICM-Pro|ICM-Chemist

In the File menu there is an option called New. This option can be used to create new objects of the following types:

- Peptide •
- Unusual Peptide •
- Compound •
- DNA and RNA
- Sequence •
- Script
- HTML
- Table •
- Arrow • Box



# 3.4 Open and Read Files

```
Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro | ICM-Chemist
```

Any file that ICM can understand can be opened by:

• Selecting File/Open.

A file with an extension .icb is an ICM binary file and can be viewed in the GUI. A .icb file can contain many objects such as sequences, meshes, protein objects, alignments, tables etc...

### 3.4.1 Open with Password

To open a file that is password protected:

• File/Open with Password

### 3.4.2 Extract from icb file

An **icb** file is an icm project file, in some instances you may want to take objects saved in an icb file and load it in your current ICM session. This option allows you to view a tabulated list of what a icb file contains and load individual object files from it.

- File/ Extract from ICB.
- Locate the saved icb file.
- A table as shown below will be displayed
- Double-click on any of the entry to extract that object from the icb file.

examp	ble_alignment \		
	name	type	size
2	openFilePRJNAME	string	9
3	1TKI_A_4	sequence	491
4	1DI9_A_17	sequence	509
5	1WFC_5	sequence	538
6	1IAN23	sequence	415
7	1QL6_A_4	sequence	413
8	1ql6_a	sequence	907
9	2phk_a	sequence	786
10	alig	alignment	10927
11	1ql6	object	231820
12	2phk	object	235471

# 3.5 Saving Files

Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro | ICM-Chemist

Anything you have displayed in an ICM graphics session can be conveniently stored in a single file called an .icb file. To do this:

• File/Save Project

#### or

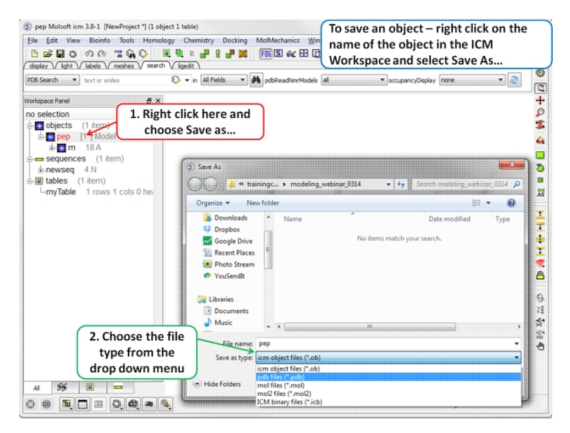
• File/Save Project As

#### A .icb file can be password protected by:

- File/Save with Password
- Enter a file name or browse for a previously saved project.
- Enter a password
- Determine whether you want the file to be Fully Protected, read only or Read Only and Allow Comments .

# Any object (e.g. protein, table, alignment...) inside the ICM GUI can be saved by:

- Right click on the name of the object in the ICM Workspace.
- Select "Save As"
- Choose the file format from the "Save as Type" drop down dialog.



# 3.6 Making Selections

Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro | ICM-Chemist

**Note:** Click **Next** (top right hand corner) to navigate through this chapter. Headings are listed on the left hand side (web version) or by clicking the **Contents** button on the left-hand-side of the help window in the graphical user interface.

There will be many occasions when you will have to make selections. For example, if you want to display a particular region or molecule contained within your protein structure or if you want to select residues around a binding pocket. If you have a molecule displayed in the graphics window, then selections will be displayed as green crosses. The selection you have made is also displayed at the top of the ICM Workspace. It is always a good idea to keep an eye on what is selected and what isnt.

There are four basic levels of selection

- 1. Object (eg a PDB structure or ICM object)
- 2. Molecule
- 3. Residue
- 4. Atom

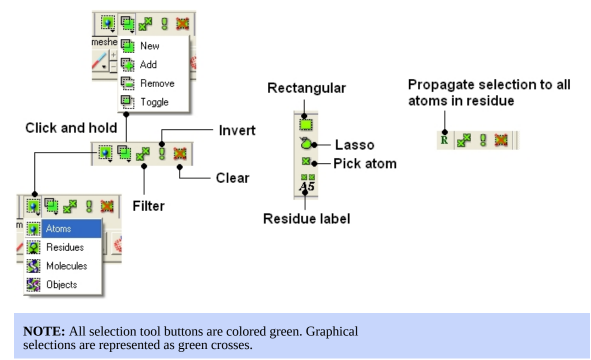
You can make selections in:

- The Graphics Display
- The ICM Workspace (Selections are highlighted in blue)
- Tables
- Sequences
- Plots

• Alignments

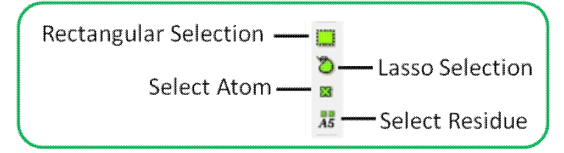
### 3.6.1 Graphical Selection Tools

The following buttons can be used to make a selection once a structure is displayed.



### 3.6.2 Quick Selection

To make a quick selection the following buttons can be used.



#### To select parts of your structure:

• Click on the **Rectangular selection icon** and click and drag around the part of the structure you wish to select.

OR

• Click on the **Lasso selection icon** and click and drag your mouse around the area of the structure you wish to select, forming a lasso around it.

#### To pick individual atoms:

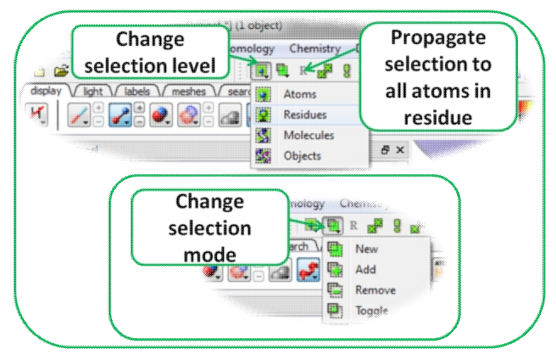
• Click on the 'pick atom' button

### 3.6.3 How to Change the Selection Level and Mode

It is possible to change the level of selection before or during the building of a selection. The selection level drop-down button can be used to do this (see image below).

For example, a C-alpha of a residue is selected but one would like to select all atoms in the residue. You can change the level to **Residues**. This selection can then be changed into all atoms of the residue by then selecting the **Atoms** level again. Or you can use the **Propagate Selection** to all Atoms button (see image below).

• Click on the **Select** objects, **Select molecules**, **Select residues**, or **Select atoms** icon, depending on which part of the structure you wish to be highlighted.

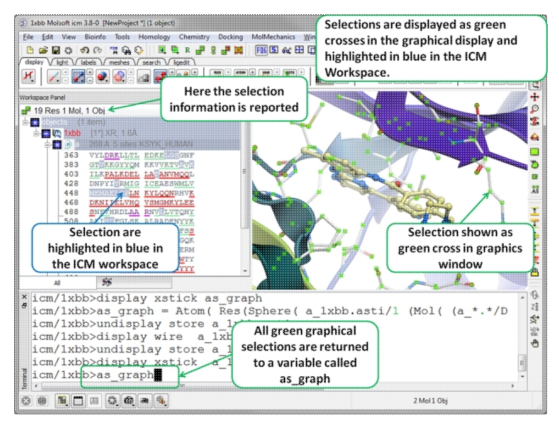


It is also important to observe the selection mode that is being used. There are four modes:

- New: new selection replaces everything selected before
- Add: new selection is added to previous selection(s), if any
- Remove: previously selection (part or whole of it), if included in the new selection will be unselected.
- Toggle: within the new selection, everything that has been selected is unselected and everything that hasni; ‰t been selected, wilbe selected

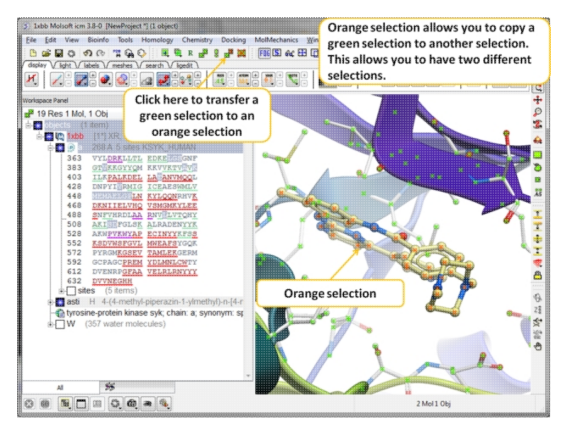
### 3.6.4 How to check what is selected.

All selections are displayed as green crosses in the graphical display and blue in the ICM Workspace. All green selections are returned to a variable called as\_graph.



### 3.6.5 Orange Selection

Sometimes it is necessary to have two different selections. The Orange selection allows you to do this it is useful for such operations as superposition and more technical procedures such as designing a protein loop. The orange selection is returned to a variable called as2\_graph.



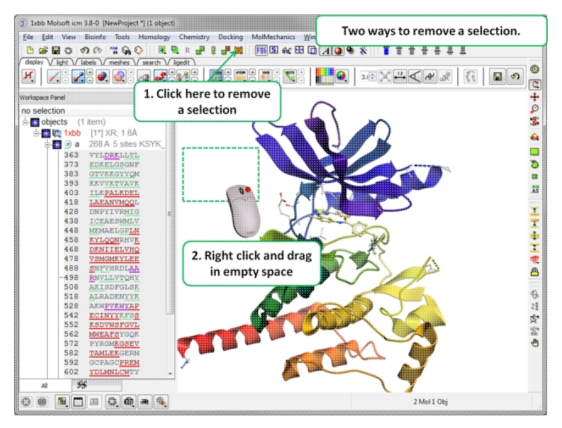
### 3.6.6 Clear Selection

To unselect everything you have previously selected:

• Simply click on the **Clear Selection** button on the selection toolbar.

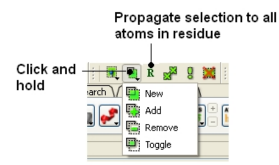
#### OR

• Right click and drag away from the displayed structure.



### 3.6.7 Changing a Selection

Once you have made a selection you may wish to add or remove parts of the selection. The buttons shown below allow you to accomplish this.



To add or remove from your current selection:

- Click on the Selection mode: add or Selection mode: remove icon on the toolbar.
- Click and drag around the part of your structure you wish to add or remove.

# You may also wish to invert your selection in a specific part of the structure.

The parts that are currently selected will become unselected, and the unselected parts will become selected.

In order to invert a selection:

• Click on the **Invert** icon on the toolbar.

If you wish to select and unselect certain regions of a selection the toggle selection button is very useful.

- Click on the Toggle selection button.
- Right click around the selections you wish to select or unselect.

**NOTE:** The selection you have made is recorded at the top of the ICM workplace. Any selection is stored in the variable as\_graph.

### 3.6.8 Filter Selection

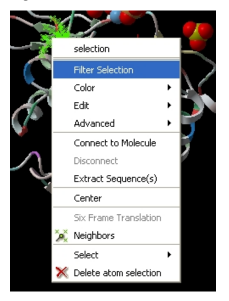
You may want to be very specific about a selection you want to make. For example you may only want to select protein backbone atoms or you might want to just select the charged residues.

The button shown below enables you to filter your selection:

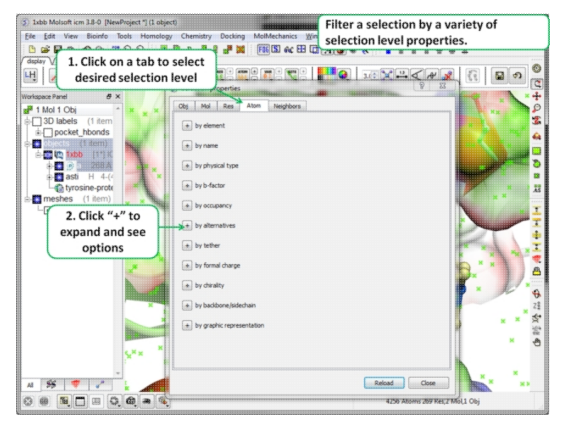


#### Or

Right click on a selection and a menu as shown below will be displayed.

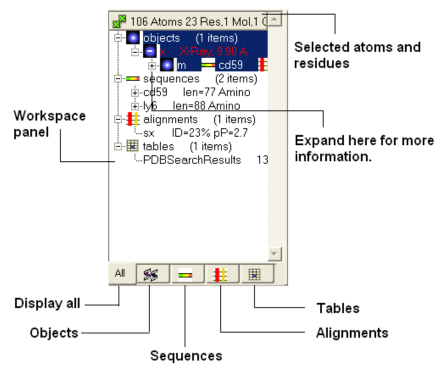


- Select the Filter Selection option.
- A dialog box will be displayed as shown below.
- Select what level of selection you want e.g. atom or residue.



### 3.6.9 Workspace Selections

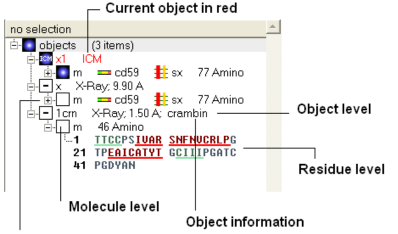
In the default GUI layout the workspace panel is located to the left of the 3D graphics display. It is a great tool for keeping track of all your sequences, pdb structures, objects, tables and alignments. As you will see in this section it also provides a way of making selections.



### 3.6.10 Workspace Navigation

Once you have mastered how to navigate the ICM workspace making a selection will become easier. Each object is divided into 3 levels:

- 1. Object Level Shown in red if it is the current object. Holds details about the structure name, X-ray, NMR, resolution etc. Importantly it will state whether the structure is an ICM object or a structure straight from the PDB. To learn how to convert a PDB into an ICM object go to the section on converting a PDB.
- 2. Molecular Level Shows the individual subunits, ligands and hetatoms of a molecule.
- 3. Residue Level Shows the sequence.



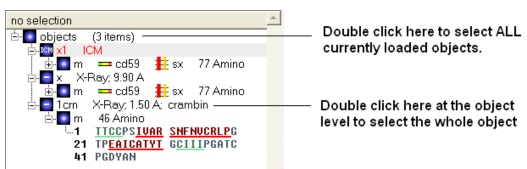
#### Click to expand tree

**NOTE:** You can expand each level of the ICM workspace by clicking the "+" button as shown above.

### 3.6.11 How to Select an Object

#### To select the whole object:

• Double click on the object level.

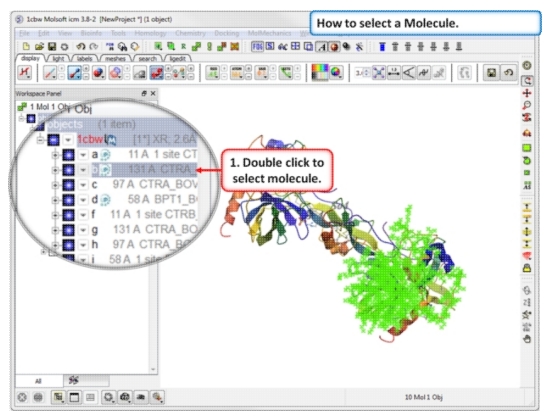


Use the CTRL button to select multiple non-contiguous objects or if they are continuous you can use double click and hold the tab button.

### 3.6.12 How to select a Molecule

#### To select a molecule(s):

• Double click on the molecule in the ICM workspace.



Use the CTRL button to select multiple non-contiguous molecules or if they are continuous you can use double click and hold the tab button.

### 3.6.13 How to select Residues

There are different options to select residues:

#### **OPTION 1:**

• Click and drag over the residues you wish to select in the ICM workspace. Selected residues will be highlighted in dark blue in the workspace and with green crosses in the graphical display.

#### Selection information is recorded here

L

	ala 184 Atoms 35 Res,1 Mol,1 Obj indiana and a state of the state of	$\bigcirc$
Click here to	E- 1188 X-Ray, 2.80 A; rhodopsin	1 7
expand tree to —	a 338 Amino OPSD_BOVIN	
show amino acid	21 RSPFEAPOYY LAEPWOFSML	
residues.	41 AAYMELLIML GEPINELTLY	The second second
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Click and drag	81 <u>VADLEMVEGG</u> <u>FTTTLYTSLH</u> 101 <u>Eventsch</u>	I Julian Mark
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you wish to	141 KPMSNFRF <u>GE</u> NHAIMGUAFT	
select.	161 <u>WUMALACAAP</u> <u>Plu</u> gwsryip 181 Egiorscgid yytpheetnn	A A A A A
	201 ESFUIYMFUU HFIIPLIUIF	
Selected	221 FCYGQLUFTU KEAAASATTQ	
residues will be	241 KAEKEUTRMU IIMUIAFLIC	L.M. X. X X X X
highlighted in	261 <u>WLPYAGUAFY</u> <u>IFTH</u> QGSDF <u>G</u> 281 <u>Rifhtipaff</u> Aktsauynpu	
the workspace	301 IYIMMNKOFR NCMUTTLCCG	
and graphics	321 KNPSTTUSKT ETSQUAPA	
window	Any selection is highlighted in the workspace as	well as in the

Any selection is highlighted in the workspace as well as in the 3D graphics window if the structure is displayed.

#### **OPTION 2:**

- Click on the **rectangular selection** icon or **lasso selection** icon on the toolbar.
- Click and drag around the residues you wish to select. Selected residues will be displayed by green crosses on the graphical display and blue in the ICM workspace.
- Use the propagate selection to residue level button.

#### **OPTION 3:**

• Right click on the a residue in the graphical display and a menu as shown here will be displayed.

	selection
	Selection Dialog
	Advanced 🕨 🕨
	Residue atoms 🔹 🕨
	Open with MolEdit
	Connect to Molecule
	Disconnect
	Extract Sequence(s)
	Center
	Annotate selection
°∕××	Neighbors
	Closed Cavities
	Select +
$\times$	Delete residue selection

- Click on **Select** and a further menu will be displayed.
- Click on Residue, Molecule or Object.



#### **OPTION 4:**

Use the select residue button.

### 3.6.14 Select All

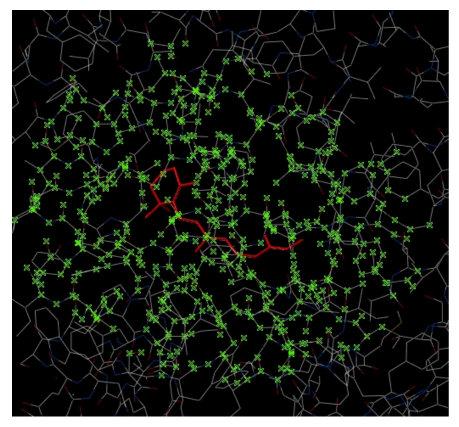
Ctrl + A will select everything in the ICM workspace, and Ctrl + Shift + A will unselect your objects.

**NOTE:** The selection you have made is always recorded at the top of the ICM workplace. If you are familiar with using the ICM terminal (See language manual) the atoms, residues, molecules or objects selected interactively in the graphics window are automatically s

### 3.6.15 Selecting Neighbors

In some instances you may only want to display or select only a subset of a structure. For example you may only wish to display the residues surrounding a ligand (as shown below (ligand red; graphical selection green crosses). The "Selecting Neighbors" option selects the residues within a shpere of a defined radius.

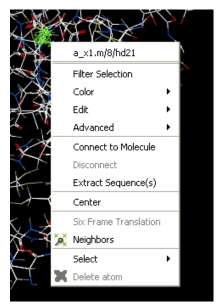
There are two ways of selecting neighbours to a particular atom or residue in ICM. Either by right clicking on the atom or residue in the graphical display or by right clicking in the ICM workspace.



### 3.6.16 Selecting Neighbors: Graphical

To select neighboring atoms or residues around a sphere of a certain radius:

- First select the residue(s) or atom(s) around which you wish to select neighbors. (See the Selection Toolbar Section)
- Right click on the selection and a menu as shown below will be displayed or choose Tools/Geometry/Neighbors.



• Select the Neigbors option and a data entry box as shown below will be displayed.

This option will allow you to make a spherical selection.

The window will open as displayed as below:

Select Neighb	ors	? 🗙
Select Neighbours F	or 🗙 Graphical Selecti	on (0 mol) 🔽
Radius	5. 💌	
type	visible	~
💌 exclude source		
🗹 unselect water		
Undisplay Beyond S	Selection	
Ok	Cancel	Help

- Select the molecule you wish to select neighbors around. For example you can select a ligand in the ICM Workspace and then choose the **Graphical Selection** option in the "Select Neighbors For" dialog entry box. Or alternatively you can select the object by clicking on the drop down button next to the "Select Neighbors For" dialog entry box.
- Enter the radius in Angstroms for the neighbor selection. e.g. 5.
- **Type** this option is **important.** This option relates to what is going to be selected. For example if you leave this option as **visible** and you only have ribbon representation displayed for your

receptor (e.g. when selecting neighbors for a ligand) then only backbone atoms will be selected.

Selection Type option includes:

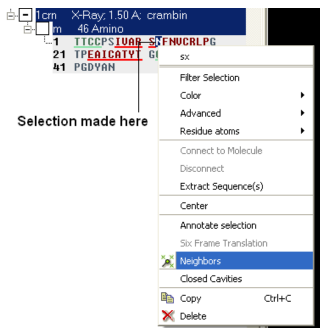
- **visible** will select all atoms displayed within the radius selected.
- **visible sidechains** will select all visible side-chains not backbone atoms.
- **same\_object\_other\_chains** will select all atoms in other chains in the same object as the original selection.
- **other objects** will select atoms in objects other than the original selection.
- **same object** will select atoms in the same object as the original selection.
- all\_objects will select atoms in all objects
- **choose\_from\_list** will allow you to select the object you wish to include in the neighbors selection.
- **exclude source** if checked will not include your original selection in the spherical selection.
- unselect water if checked will not select water molecules
- Undisplay Beyond Selection will only display the atoms selected.

**NOTE:** The selection you have made is always recorded at the top of the ICM workplace. If you are familiar with using the ICM terminal (See language manual) the atoms, residues, molecules or objects selected interactively in the graphics window are automatically saved in the variable as\_graph. Graphical selections are shown in green (crosses) or highlighted in blue in the ICM Workspace.

### 3.6.17 Selecting Neighbors: Workspace

# To select neighboring atoms or residues around a sphere of a certain radius from a residue in the ICM workspace:

- First select the residue in the ICM workspace around which you wish to select neighbors. (See the Residue Selection)
- Right click on the selection and a menu as shown below will be displayed.



- Select the Neigbors option and a data entry box as shown below will be displayed.
- Follow the instructions in the previous section.

**NOTE:** The selection you have made is always recorded at the top of the ICM workplace. If you are familiar with using the ICM terminal (See language manual) the atoms, residues, molecules or objects selected interactively in the graphics window are automatically s

### 3.6.18 Select by Residue Number

#### To select by residue number:

- Right click on the molecule you wish to make the selection in and choose Select/Res Number...
- Enter the residue numbers e.g. 3,7:12,30 Colon indicates range.

### 3.6.19 Alignment and Table Selections

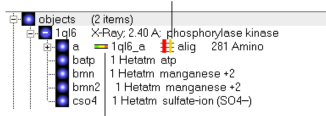
Descriptions on how to make selections in Alignments and Tables are in the sections entitled Making Selections in Alignments and Making Table Selections.

### 3.6.20 Making Links

It is sometimes necesary to make links between sequences objects and alignments. A link enables you to make selections in one environment such as an alignment and then these selections are transferred to the object such as the PDB structure displayed.

If a link is made then a symbol will be displayed next to the object in the ICM workspace. In the example shown below subunit\_a of the X-ray structure 1ql6 is linked to the sequence 1ql6\_a and the alignment called 'alig'.

#### Linked to alignment 'alig'



#### Linked to sequence 1ql6\_a

If an object is linked to an alignment a symbol as shown below will be displayed.

```
i⊡-∰talignments (1 items)
i…alig ≸≸ id=75 nSeq=7
```

Alignment is linked to an object

To link a sequence from an object - extract the sequence from the object.

- Right click on the object in the ICM workspace.
- Select extract sequence.

To link a sequence and object to an alignment.

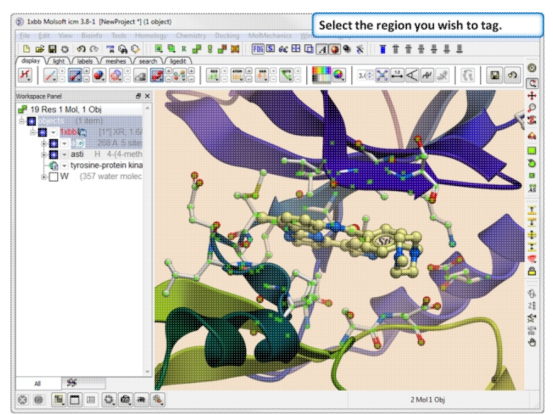
Use the extracted sequence as described above to build your alignment.

In addition a link can be made between a structure and alignment by:

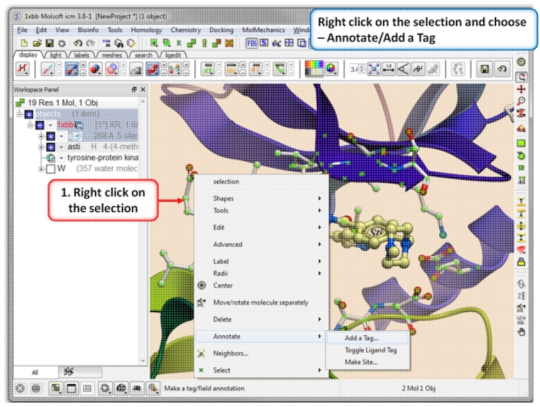
- Bioinfo/Link to Structure.
- Enter alignment name.
- OK

### 3.6.21 Tags

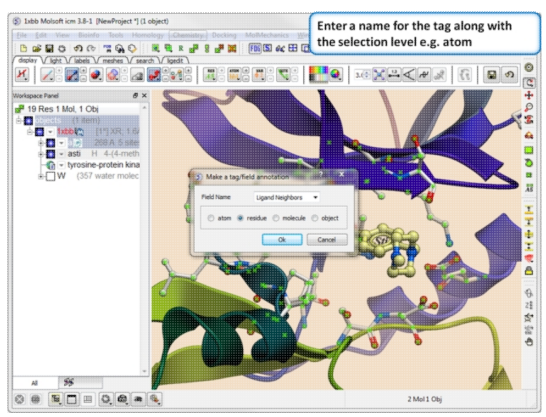
It is sometimes convenient to tag selections so you can come back and use them at a later date. To do this:



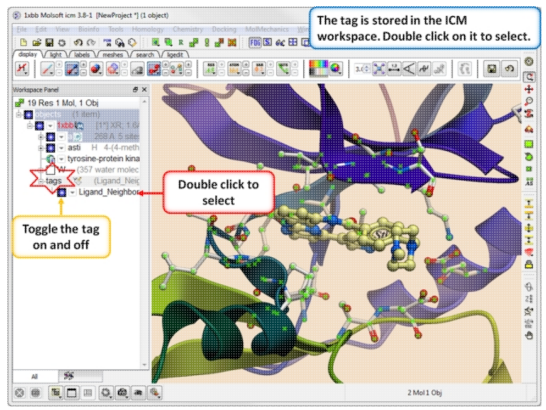
**Step 1.** Select the region you wish to tag.



Step 2. Right click on the selection and choose Annotation/Add a Tag.



**Step 3.** Give the tag a name and choose the selection level (atom, residue, molecule, or object).



**Step 4.** You will see the tag in the ICM Workspace. You can toggle the display on or off or select the tag by double clicking on it in the ICM Workspace.

# 3.7 Preferences

#### Your ICM preferences can be changed by:

• Select File/Preferences.

**NOTE:** There is a "Reset to Default" button in case you make any changes you are not happy with and also a search option.

	1. File/Preferen	
orkspace Panel	ð ×	5 23
o selection	2	
obje	System Preferences *	8 23
÷.	T Directories Graphics G	a ICMdb Image Labels Ligand Plot Ribbon Shell System Tools Search Results ()
ienew	GUILmax Sequence Length	2000 💿 🖏 GUL sequence Offset Style from residue 🔹 🛷
tabi	Blastdb Directory	C:/Program Piles (x86)/Molsoft LLC/JCM-Pro_7)/data/blast/
-my1	SEQUENCE.restore Orig Names	SEQUENCE.site Colors
	TOOLS. find Database Min Sequence	e 12 💿 🕫
11		
	Search: sequence	Defaults OK Apply Cancel
	Search: sequence	Defaults OK Acply Cancel
AI 😽	Search: sequence	

### 3.7.1 Bonds Preferences

To change Bond Preferences:

- Select File/Preferences.
- Choose the **Bonds** tab.

**GRAPHICS.ballStickRatio** - A default ratio of ball and stick radii. This ratio is applied when the styles are switched from the GUI xstick toolbar. Default (1.4)

#### **GRAPHICS.hbond Ball Period** - Default (3)

**GRAPHICS.hbondMinStrength** - parameter determines the hbond strength threshold for hbond display. The strength value is between 0. and 2. By changing 1. to 0.2 you will see more weak hydrogen bonds. Default: (1).

**GRAPHICS.hbondStyle** - determines the style in which hydrogen bonds are displayed. Here hbond-Donor, Hydrogen, and hbond-Acceptor atoms will be referred to as D, H and A, respectively,

**GRAPHICS.hetatmZoom** - The default ball and stick radii of a ligand can be different by the GRAPHICS.hetatmZoom factor. This makes a better ligand view since the ligand stands out from the surrounding protein atoms.

**GRAPHICS.stickRadius** - radius (in Angstroms) of a cylinder displayed as a part of stick or xstick graphical representation of a molecule. Individual (residue-wide) control of stick radii.

#### **GRAPHICS.xstick Backbone Ratio** - Default (1.2)

GRAPHICS.xstick Style - xstick style

**wireBondSeparation** the distance between two parallel lines representing a chemical double bond if wireStyle = "chemistry". Default (0.2 Angstroms).

GRAPHICS.distance Label Drag - enable distance label dragging

**GRAPHICS.hbondAngleSharpness** determines how the strength depends on the D-H...A(lone pair) angle. The preference can be found the general Preferences menu Default (1.7)

GRAPHICS.hbond Ball Style even, by atom size, by energy or telescopic

#### **GRAPHICS.hbond Rebuild**

GRAPHICS.hbondWidth relative width of a displayed hbond .

**GRAPHICS.hydrogenDisplay** determines the default hydrogen display mode for the display command.

```
GRAPHICS.hydrogenDisplay = "polar"
    1 = "all"  # all hydrogens are shown
    2 = "polar" <-- current choice  # polar displayed, the non-polar hidden
    3 = "none"  # no hydrogens are displayed</pre>
```

GRAPHICS.wire Width - relative width of wire Default (1)

**GRAPHICS.xstick Hydrogen Ratio** - Default (0.5)

**GRAPHICS.xstick Vw Ratio** - Default (0.6)

Wire Style - change the default wire style

### 3.7.2 Directories Preferences

#### **DIRECTORIES TAB:**

To change Directory Preferences:

- Select File/Preferences.
- Choose the **Directories** tab.

Within this tab you can select the default directories for:

**FILTER.gz, FILTER.uue, FILTER.Z, Filter.zip** allows you to read compressed files .gz, .uue, .Z, and, .zip files automatically leaving the compressed file intact.

**PDB Directory Style** - The style of your Protein Data Bank directory/directories. ICM will understand all of the listed styles, including distributions with compressed \*.gz , \*.bz2 and \*.Z fil es

**BlastDB Directory** - return directory with Blast-formatted sequence files for ICM sequence searches. You can download Blast formatted databases from here ftp://ftp.ncbi.nih.gov/blast/db/

Dock Directory - Default directory for storing docking files.

#### **CCP4 Directory**

Editor - Select a default text editor

Inx Directory - location of stored index (\*.inx) files.

**Log Directory** - when you quit an icm-session, a \_seslog.icm file is automatically stored. If the s\_logDir variable is empty, it is stored to the s\_userDir + "/log/" directory. However one can redirect it to the current working directory (".") or any other directory.

#### **Output Directory -**

**PDB Directory** - directory containing the PDB database of 3D structures. These files can also be easily downloaded directly from the PDB site if the variables are set as in the example below. PDB distributions can exist in several styles (all files in the same directory, or divided etc.).

#### **PDB Directory FTP**

#### **PDB Directory Web**

**Projects Directory** - Select the default location for storing ICM projects. Save your data in an ICM project. It is a convenient way of keeping all your structures, alignments, tables, docking results etc... in one place. A description on how to save an ICM project is described in the GUI Basics section of this manual.

**Prosite Dat** - location of the prosite.dat file a dictionary of protein sites and patterns, (Copyright by Amos Bairoch, Medical Biochemistry Department, University of Geneva, Switzerland).

Ps Viewer - Select a postscript viewer

Swissprot Dat - location of swissprot.dat file

**Temp Directory** - scratch directory for temporary files ( some montecarlo files will be saved there ).

Uniprot Dat - location of uniprot.dat file

**XPDB Directory** - Path to the ICM XPDB database of compact binary ICM objects which are annotated with the site information. The advantage of the XPDB database is the speed of reading and smaller size than PDB. XPDB entries are read about 80 times faster!

#### **TOOLS.default ChemDB**

#### **TOOLS.eds Directory**

#### **TOOLS.pdb Read Nmr Models**

```
1. = "first" : reads only one model from a multi-model (e.g. NMR) pdb file
2. = "all" : reads all models from a multi-model (e.g. NMR) pdb file and creates a separate
3. = "all stack" : creates one object and loads all other models as a stored cartesian stac
```

### 3.7.3 Graphics Preferences

To change Graphics Preferences:

- Select File/Preferences.
- Choose the **Graphics** tab.

Atom Single Style - display style of isolated atoms in the wire mode.

```
1. "tetrahedron"
2. "cross"
3. "dot"
```

GRAPHICS.clash Style - choose clash length, strain or length.

GRAPHICS.clip Grobs - enable grob clipping.

#### **GRAPHICS.clip Static -**

**GRAPHICS.grobLineWidth** - relative width of displayed lines of 3D meshes (grobs). Also affects the interatomic distance display.

**GRAPHICS.lightPosition** - X, Y and Z position of the light source in the graphics window. The X and Y coordinates are usually slightly@@ beyond the [-1. 1] range where [-1.,1.] is the size of the window, and the Z position is perpendicular to the screen and is set to 2. (do not make it negative).

**GRAPHICS.occupancyDisplay** preference controlling if and how the partical or zero atom occupancies are displayed. The abnormal occupanices are shown as circles around atoms. These following values are allowed.

1. = "none" # nothing is displayed

"circle" # a circle is displayed
 = "label" # a circle and a lable with the value (zero values are not shown)

**GRAPHICS.quality** - integer parameter controlling quality (density of graphical elements) of such representations as cpk, ball, stick, ribbon . Do not make it larger than about 20 or smaller than 1.

GRAPHICS.ruler Style - change ruler from center to side

**GRAPHICS.stereoMode** - 1. "up-and-down", 2. "line interleaved" 3. "in-a-window"

- a simple hardware stereo mode for workstations with a horizontal frame splitter.
- In the "up-and-down" mode a longer frame with two stereo images on top of each other is generated and the two halves are then superimposed with the splitter. This mode does not require anything from a graphics card, but does require a frame splitter. A frame splitter box was connected between a monitor and a graphics card output. This mode has an unpleasant side effect, the rest of the screen (beyond the OpenGl window) becomes stretched and the lower part of the screen is superimposed on the top half.
- The "line interleaved" mode can be used with a new type of frame splitter at the line level. In this case the odd lines from one stereo-image are interleaved with the even lines of another. The side-effect of this mode is that the intensity is reduced in half since at each moment one sees only one half of the lines. The splitter device for this mode can be purchased from Virex (www.virex.com). This mode produces a dark stereo image but is easily available (requires stereo goggles, e.g. from Virex).
- easily available (requires stereo goggles, e.g. from Virex).
  The "in-a-window" mode is used in SGI workstations and in a Linux workstation with an advanced graphics card supporting a quad graphics buffer. In this mode the hardware stereo regime applies only to an OpenGl window. This is the best mode but it requires an expensive graphics card (plus the stereo goggles).

**GRAPHICS.surfaceDotDensity** - Determines the number of dots per square Angstrom on the graphical solvent accessible surface.

**GRAPHICS.surfaceProbeRadius** - An increment to the van der Waals radii of atoms at thich the dotted atomic surface is calculated. It is used by the display surface command to display dotted van der Waals surface. If the GRAPHICS.surfaceProbeRadius is set to 1.4 the surface becames equivalent to the solvent accessible surface with a probe of 1.4A

**GROB.arrowRadius** - a real arrow radius in Angstoms used by the Grob( "ARROW", R\_) function. Default: 0.5. **GROB.contourSigmaIncrement** - a real increment in the sigma level used to re-contour an electron density map using the make grob m\_eds add r\_increment command. This parameter is used in the GUI when plus and minus are pressed.

**GROB.relArrow Size** - a real ratio of the arrow head radius to the arrow radius. This parameter is used by the Grob( "ARROW", R\_) function. Default: 3.0.

**shineStyle** - defines how solid surfaces of cpk , skin and grobs reflect light. Possibilities:

```
1. "white" <- default
2. "color"</pre>
```

The first option gives a more shiny and greasy look.

**GRAPHICS.center Follows Clipping** - determine the function of center button.

GRAPHICS.clashWidth - relative width of a displayed clash .

GRAPHICS.clip Skin - enable skin clipping.

**GRAPHICS.displayMapBox** - controls if the bounding box of a map is displayed

**GRAPHICS.light** - a rarray of 13 elements between 0. and 1. which controls the main properties of lighting model in GL.

**GRAPHICS.mapLineWidth** - relative width of lines and dots of a displayed map.

**GRAPHICS.occupancy Radius Ratio** - preference controlling the radius of the partical or zero atom occupancies

#### **GRAPHICS.resize Keep Scale**

**GRAPHICS.selectionStyle** - preference for the style in which the graphical selection is shown. The preference may have the following values.

**GRAPHIC.store Display** - maintains representation and coloring for an object.

**GRAPHICS.surfaceDotSize** - Determines the size of the dot on the solvent accessible graphical surface.

**GRAPHICS.transparency** - Two parameters regulating the transparency of grobs.

**GROB.atomSphereRadius** - default radius (in Angstroms) which is used to select a patch on the surface of a grob.

**GROB.relArrowHead** - a real ratio of the arrow head radius to the arrow radius.

**lineWidth** - the real width of lines used to display the wire representation of chemical bonds.

### 3.7.4 GUI Preferences

#### GUI TAB:

The options contained within the Preferences/Gui tab are described below.

**GRAPHICS.alignment Rainbow** - This option controls how alignments are colored by default.

**GRAPHICS.NtoC Rainbow** - Controls the coloring of structural representation from the N-terminal to the C-terminal

GRAPHICS.rocking - Controls default rocking motion.

GRAPHICS.rocking Speed - Controls rocking or rotation speed.

GUI.auto Save Interval - Controls auto save period (minutes)

GUI.table Row Mark Colors - Controls colors used for marking tables.

**GUI.workspaceTabStyle** - Controls the style of ICM-object tabs created in the workspace panel of ICM GUI.

**Movie.fade Nof Frames** - Controls number of frames for the fade out option in screenshot movie making.

Movie.quality - Controls the resoltuion of the movie

SEQUENCE.site Colors - Controls coloring of squence sites.

**SLIDE.ignore Fog** - Fog representations can be ignored in slide preparation if desired.

#### **GRAPHICS.discrete Rainbow -**

**GRAPHICS.rainbow Bar Style** - determines if and where the color bar will appear after a molecule is colored by an array.

GRAPHICS.rocking Range - real value of rocking range.

GUI.auto Save - auto save on or off

GUI.max Sequence Length - maximum sequence length displayed in ICM

GUI.workspace Folder Style - Workspace folder style.

**MOVIE.frame Grab Mode** - with screenshot movie making you can choose either fixed frame time or real time.

**Movie.quality Auto** - with screenshot movie making you can allow ICM to control the movie resolution.

**SLIDE.ignore Background Color** - Ignore background color when you are making a slide.

### 3.7.5 GUI Preferences

To change GUI Preferences:

- Select File/Preferences.
- Choose the **GUI** tab.

**Quality** - controls the quality (density of graphical elements) of such representations as cpk, ball, stick, ribbon . Do not make it larger than about 20 or smaller than 1. We recommend to make this parameter at least 15 if you want to make a high quality image. You can also increase the number of image resolution by making the image window 2,3,4 times larger (in the example below it is 2 times larger) than the displayed window.

Wire Style - Four different wire styles are available.

**Hydrogen Display** - Select whether you always want all hydorgens displayed or just-polar hydrogens or no hydrogens at all.

**Rainbow Scale** - determines if and where the color bar will appear after a molecule is colored by an array. Coloring by an array is one of the options of the display and color commands.

```
1. = "left" <- default choice
2. = "right"
3. = "no text"
4. = "no bar"</pre>
```

**Ball Ratio** - The ratio of ball and stick radii. This ratio is applied when the styles are switched to xstick from the GUI xstick toolbar.

**Selection Style** - Change the graphical display of your selections. Default is a green cross.

**Clash Threshold** - a clash is defined as an interatomic distance less than a sum of van der Waals radii of two atoms of interest multiplied by the clashThreshold parameter. For hydrogen bonded atoms, the distance threshold is additionally reduced by 20%. Default = 0.82

**DotSurfaceRadiusIncrement** - adius of a probe sphere used to display a dotted surface of a molecule. All van der Waals radii are expanded by this value. vwExpand=0 corresponds to the CPK surface, vwExpand=1.4 corresponds to the water-accessible surface. Be aware of the difference between the waterRadius and vwExpand parameters: waterRadius is used in

- show energy "sf"
- show [area|volume] skin
- display skin while vwExpand is used in
- show [area|volume] surface
- display surface

Default (1.4).

**H Bond Style** - How do you wish your H-Bonds to be displayed by default? Dashes, Bond Length, Bond Length and Angle.

**grobLineWidth** - relative width of displayed lines of 3D meshes (grobs). Also affects the interatomic distance display.

**general line with** - the real width of lines used to display the wire representation of chemical bonds. See also IMAGE.lineWidth parameter which controls line thickness in molecular images generated by the write postscript command, and the PLOT.lineWidth which controls the width for the plot command. Default (1.0)

single atom as - display style of isolated atoms in the wire mode.

```
    "tetrahedron"
    "cross"
    "dot"
    The size of the first two representation is controlled by the GRAPHICS.ballRadius parameter an
```

**xstickhetatomzZoom** - The default ball and stick radii of a ligand can be different. This makes a better ligand view since the ligand stands out from the surrounding protein atoms.

solid shine style - choose either white or color

**Stick Radius** - radius (in Angstroms) of a cylinder displayed as a part of stick or xstick graphical representation of a molecule. Individual (residue-wide) control of stick radii.

Stereo Mode - Select a default stereo mode

**Display Style** - A default display style can be chosen using a combination of styles.

**Water Radius** - radius of water sphere which is used to calculate an analytical molecular surface (referred to as skin) as well as the solvent-accessible surface (centers of water spheres).

clashWidth - relative width of a displayed clash.

hbondWidth - relative width of hydrogen bond display

mapLineWidth - relative width of lines and dots of a displayed map.

### 3.7.6 Image Preferences

To change Image Preferences:

- Select File/Preferences.
- Choose the **Image** tab.

IMAGE.color - logical to save color or black\_and\_white ('bw') images.

**IMAGE.gammaCorrection** - real variable to to lighten or darken the image by changing the gamma parameter. A gamma value that is greater than 1.0 will lighten the printed picture, while a gamma value that is less that 1.0 will darken it.

**IMAGE.lineWidth** - this real parameter specifies the default line width for the postscript lines.

IMAGE.orientation - image orientation.

**IMAGE.previewer** - a string parameter to specify the external filter which creates a rough binary (pixmap) postscript preview and adds it to the header of the ICM-generated high resolution bitmap or vectorized postscript files saved by the write image postscript, and write postscript , respectively .

**IMAGE.print** - unix command for printer.

**IMAGE.scale** - real variable. If non zero, controls the image scale with respect to the screen image size.

**IMAGE.stereoBase** - real variable to define the stereo base (separation between two stereo panels) in the write image postscript and write postscript command.

**IMAGE.writeScale** - an integer parameter used to increase the image resolution in the Quick Image Write tool.

**IMAGE.bondLength2D** - real length of a chemical bond (in inches) in chemical 2D drawings upon the Copy Image command.

**IMAGE.compress** - logical to toggle simple lossless compression, standard for .tif files. This compression is required to be implemented in all TIFF-reading programs.

**IMAGE.generateAlpha** - logical to toggle generation of the alpha (opacity) channel for the SGI rgb, tif and png image files to make the pixels of the background color transparent.

**IMAGE.lineWidth2D** - integer thickness of bonds in chemical 2D drawing upon the Copy Image command. This is useful for cutting and pasting from ICM to external documnents.

IMAGE.paper Size - specify paper size.

**IMAGE.previewResolution** - integer resolution of the rough bitmap preview added to the vectorized postscript file in lines per inch.

**IMAGE.printerDPI** - this integer parameter the printer resolution in Dot Per Inch (DPI). Important for the write image postscript command.

**IMAGE.stereoAngle** - real variable to define stereo angle (relative rotation of two stereo images) in the write image postscript and write postscript command.

**IMAGE.stereoText** - logical to make text labels for only one panel or both panels of the stereo diagram.

### 3.7.7 Label Preferences

To change Label Preferences:

- Select File/Preferences.
- Choose the **Labels** tab.

**atomLabelStyle** style of atom labels invoked by clicking on an the atom label button.

**GRAPHICS.displayLineLabels** - enables/disables the display of edge lengths (inter-point distances) of a grob generated with the Grob( "distance" .. ) function.

GRAPHICS.font Line Spacing - Change the spacing between lines in labels.

**GRAPHICS.resLabelDrag** - if yes, enables dragging of the displayed residue labels with the middle mouse button.

GRAPHICS. site Arrow - Highlight sites with an arrow yes or no.

**Show Res Code In Selection** - When you make a selection the icm selection language will be displayed when you right click on the selection.

**Res Label Style** - Default residue label style.

SITE.label Style - Default label sites style.

Var Label Style - Default label variable style.

**GRAPHICS.atomLabelShift** - a non-negative integer number of spaces preceding an atom label. This parameter is useful for displaying labels next to a solid representation,

GRAPHICS.fontColor - set font color

GRAPHICS.font Scale - set font size

**GRAPHICS.site Label Shift** - GRAPHICS.resLabelShift a non-negative integer number of spaces preceding a site label.

**GRAPHICS. site Label Drag** - if yes, enables dragging of the displayed site labels with the middle mouse button.

**Res Label Shift** - a non-negative integer number of spaces preceding a residue label. This parameter is useful for displaying residue labels next to a solid

**SITE.labelOffset** - (default 5. A) the real offset of the site label with respect to the residue label atom.

SITE.wrap Comment - Number of characters per comment line.

### 3.7.8 Plot Preferences

To change Plot Preferences:

- Select File/Preferences.
- Choose the **Plot** tab.

**PLOT.color** - logical to generate a color plot. Usually it does not make sense to switch it off because your b/w printer will interpret the color postscript just fine anyway.

PLOT.draw Tics logical yes or no

**PLOT.fontSize** real font size. Any reasonable number from 3. (1 mm, use a magnifying glass then) to 96.

**PLOT.lineWidth** - real line width for graphs (not the frame and tics)

**PLOT.markSize** - real mark size in points. Allowed mark types: line, cross, square, triangle, diamond, circle, star, dstar, bar, dot, SQUARE, TRIANGLE, DIAMOND, CIRCLE, STAR, DSTAR, BAR. Uppercase words indicate filled marks.

PLOT.paper Size - preference to specify plor paper size

**PLOT.rainbowStyle** - preference defining the color spectrum used by the plot area command.

**PLOT.Yratio** - real aspect ratio of the ICM plot frame. Using link option of the plot command is equivalent to setting this variable to 1.0. If PLOT.Yratio is set to 0., the ratio will be set automatically to fill out the available box optimally.

[PLOT.date ] - display date on plot

PLOT.font - preference for the title/legend font.

PLOT.labelFont - preference for the data point label font.

PLOT.logo - logical switch for the ICM-logo on the plot.

PLOT.orientation - preference for the plot orientation.

PLOT.previewer - command to local ps viewer

**PLOT.seriesLabels** - preference to indicate position of a series/color legend inside the plot frame.

### 3.7.9 Ribbon Preferences

To change Ribbon Preferences:

- Select File/Preferences.
- Choose the **Ribbon** tab.

Combo Display Style - select ribbon-cpk, atoms, ribbon-ligand, chemical

**GRAPHICS.dnaRibbonRatio** - real ratio of depth to width for the DNA ribbon .

**GRAPHICS.dnaRibbonWorm** - logical which, if yes, makes the DNA backbone ribbon round, rather than rectangular. Default: no

**GRAPHICS.dnaWormRadius** - real radius of the worm representing bases in DNA ribbon .

GRAPHICS.ribbonWidth - real width of the protein ribbon .

**GRAPHICS.wormRadius** - radius of coiled segments (i.e. those where the secondary structure is marked as "\_") of a polypeptide chain in ribbon representation. Default (0.3).

Ribbon Style - specifies default style when ribbon is displayed.

**GRAPHICS.dnaBallRadius** - DNA bases in ribbon representation are shown as balls controlled by this real parameter.

GRAPHICS.dnaRibbonWidth - real width (in Angstroms) of the DNA ribbon .

**GRAPHICS.dnaStickRadius** - real radius of the sticks representing bases in DNA ribbon .

**GRAPHICS.ribbonRatio** - real ratio of depth to width for the protein ribbon .

**GRAPHICS.ribbonWorm** - logical parameter, if yes, makes the ribbon round, rather than rectangular.

#### ribbonColorStyle -

```
- sets the ribbon coloring scheme.
1 = "type" default. colors by secondary structure type or explicit color
2 = "NtoC" colors each chain gradually blue-to-red from N- to C- (or from 5' to 3' for DN
3 = "alignment" if there is an alignment linked to a protein, color gapped backbone regions gr
4 = "reliability" 3D gaussian averaging with selectSphereRadius of alignment strength in
If ribbonColorStyle equals to 4, the conserved areas will be colored blue, while the most diver-
diverse the second strength in the second st
```

### 3.7.10 Shell Preferences

To change Shell Preferences:

- Select File/Preferences.
- Choose the **Shell** tab.

**Clash Threshold** - a clash is defined as an interatomic distance less than a sum of van der Waals radii of two atoms of interest multiplied by the clashThreshold parameter.

**Map Sigma Level** - (in Rmsd values over the mean value). Margin value used for making graphical objects contouring the 3D density map .

**Mnconf** - maximal number of conformations in the conformational stack . The stack stops growing after this number is achieved and starts replacing representative conformations with higher energy values by new conformations with superior energies, if the latter are found.

Icm Prompt - defines the ICM-prompt string.

**Select Min Grad** - default minimal gradient vector length for gradient atom selection (a\_//G). This parameter is also used by the montecarlo fast command, which requires a value of 2. to 10. for optimal performance.

**Map Atom Margin** - Margin in Angstoms around selected atoms. The margin is added to the positional boundaries to define a submap index box in the Map ( map\_source , as\_) function.

**maxColorPotential** - local electrostatic potential in kcal/e.u.charge units at which the surface element is colored by extreme red or extreme blue. All higher values will have the same color. This absolute scaling is convenient to develop a feeling of electrostatic properties of molecular surfaces. **mnSolutions** - this parameter limits the number of hits retained by the program after a search.

Real Format - format of real numbers

Water Radius - radius of water sphere which is used to calculate an analytical molecular surface

### 3.7.11 System Preferences

To change System Preferences:

- Select File/Preferences.
- Choose the **System** tab.

#### FTP.createFile -

**FTP.proxy** - string path to the proxy server for connections through firewall. Default: "" (empty string).

GUI.max Nof Recent Files - maximum number of recent files stored.

GUI.splash Screen Image - path to splash image displayed on startup

HTTP. support Cookies - http support cookies yes or no

**HTTP.user Agent** - client application used within a particular network protocol for www

Beep - warning beep yes or no

Max File Size Mb - Maximu file size in MegaBytes that can be loaded into ICM.

**USER.friends** 

#### **USER.organization**

**FTP.keep File** - (default no ). If yes, the temporary file is kept in the s\_tempDir directory. Otherwise the file is deleted.

GUI.enumberation Memory Limit - memory limit for enumeration operations.

**GUI.splash Screen Delay** 

HTTP.proxy - string for HTTP server for connection through firewall

HTTP.timeout - timeout in seconds

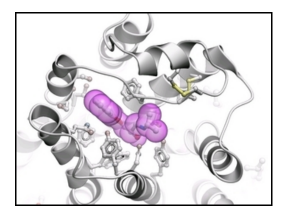
Http Read Style icm or lynx

Force Auto Bond Typing - yes/no

USER.email, USER.full Name, USER.phone

3.7.11 System Preferences

# **4 Protein Structure**



#### **Chapter Contents:**

- How to Convert Proteins and Chemicals to ICM Objects.
- Binding Site Display Tools.
- Crystallographic Analysis Tools.
- Protein Superposition.
- Protein Structure Analysis.
- Protein Structure Prediction Tools e.g. ICM Pocket Finder.

## 4.1 Convert to ICM Object

Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro | ICM-Chemist

## 4.1.1 Load a Protein Structure

There are a couple of ways to read into ICM a protein structure.

- 1. Search the PDB using the Search Tab.
- 2. Load in a PDB file that you have saved on your computer using File/Open.

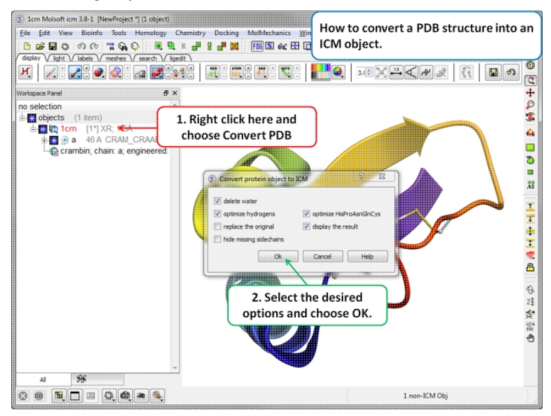
## 4.1.2 Converting PDB Files Into ICM Objects

If you are going to make any type of energy calculation in ICM (eg docking, display H-bonds, display electrostatic and binding property surfaces etc..) it is necesary to convert a protein or chemical into an ICM object. Be aware that upon conversion ICM adds missing side-chain atoms (but wont try to build missing loops) due to the nature of the internal-coordinate system. The list of residues/ atoms added is presented in the command line shell and can be reviewed. For reference, the original PDB entry is kept in the system. See the command line manual for a more complete description of what the conversion process does.

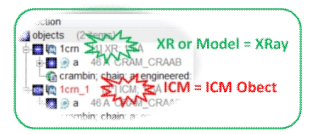
**NOTE:** Before converting a protein structure to an ICM object make sure that the chemicals contained within the structure (e.g. ligands) are correct. If an error is found you can edit the ligand as described here.

To convert a PDB structure into an ICM object follow the steps shown below:

- Right click on the name of the protein you wish to convert in the ICM Workspace.
- A dialog box will be displayed as shown below.
- If you want to delete the water molecules select **Delete Waters**
- If you want to optimize hydrogen atoms (recommended for important work) select **Optimize Hydrogens**. This option performs global optimization of hydrogens to find the best hydrogen bonding network.
- If you want to optimize the orientation of His, Pro, Asn, Gln, Cys residues then choose **optimizeHisProAsnGlnCys**. The following residues will be further optimized: His three protonation states and two rotations will be tried and the residue will be renamed according to its subtype: hie (epsilon tautomer) or hip (+). Asn and Gln (a 180 deg. flip will be tried). Cys in the vicinity of Zn, Cu, Fe and Co to cym.
- If you want to keep a copy of your PDB file uncheck the option **replace original**.
- The converted structure can be displayed immediately by checking display the result
- Uncheck the box hide missing side chains if you want ICM to build missing heavy atoms that are not reported in the PDB (due to the lack of density), they will be added according to the residue name and assigned zero occupancies. Check this box if you want residues missing heavy atoms to be hidden.



If your object is an ICM object it will display **ICM** next to the molecule in the ICM Workspace.



# 4.2 Pocket Display

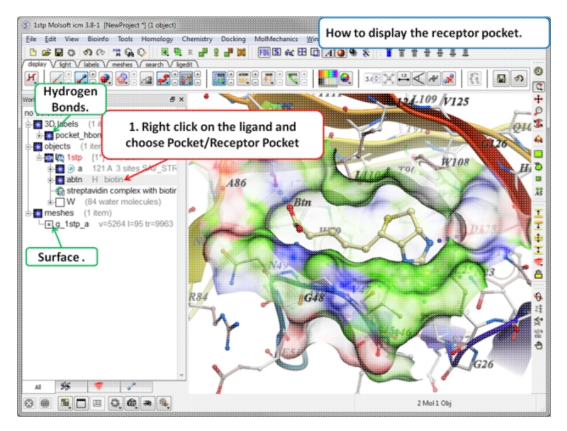
Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro

## 4.2.1 Receptor Pocket Surface.

#### To display the Receptor Pocket Surface:

- As an example we will use the PDB structure 1STP. Type 1STP in the pdb search tab and press return.
- Convert the protein to an ICM object. If you do not convert a generic surface will be displayed that is not colored by binding property.
- Right click on the ligand "abtn" and select **Pocket/Receptor Pocket**.
- Select whether you would like to display side-chain hydrogen bonds and label.
- The receptor pocket will be displayed colored by binding property

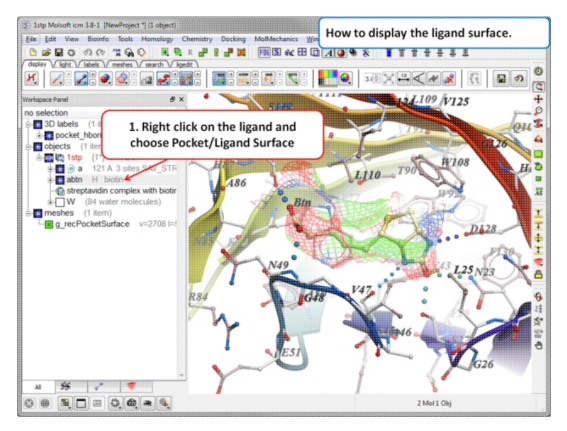
   White=aromatic lipophilic, Green=non-aromatic other (mostly aliphatic) lipophilic surface, Red=hydrogen bonding acceptor potential, Blue=hydrogen bond donor potential.
- The surface can be toggled on and off by selecting in the ICM Workspace in the **meshes** section.



## 4.2.2 Ligand Surface.

The Ligand Surface option allows you to visualize cavities that are open for ligand modifications. To display the Ligand Surface:

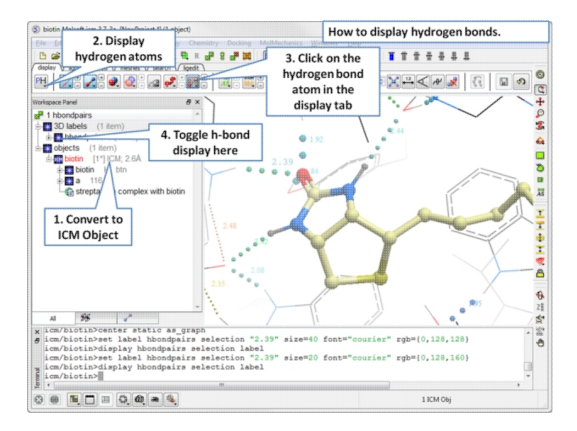
- As an example we will use the PDB structure 1STP. Type 1STP in the pdb search tab and press return.
- Convert the protein to an ICM object. If you do not convert a generic surface will be displayed that is not colored by binding property.
- Right click on the ligand "abtn" and select **Pocket/Ligand Surface**.
- The Ligand Surface will be displayed colored by binding property -White=aromatic lipophilic, Green=non-aromatic other (mostly aliphatic) lipophilic surface, Red=hydrogen bonding acceptor potential, Blue=hydrogen bond donor potential.



## 4.2.3 How to Display Hydrogen Bonds

**NOTE:** The method by which hydrogen bonds are calculated is described here in the command line manual. The GRAPHICS.hbondMinStrength parameter determines the hbond strength threshold for hbond display. The strength value is between 0. and 2. By changing 1. to 0.2 you will see more weak hydrogen bonds.

- In order to display energy related properties we need to convert the PDB file into an ICM object. Convert 1STP into an ICM object. In this example, the option "Replace the Original" was selected.
- Display the receptor in wire format and the ligand in xstick.
- Right click on the ligand and select "Neighbors" Enter 3 Angstroms and Type = Visible. Do not exclude source (the ligand) therefore remove tick from box entitled "exclude source".
- Select the display tab and then select the Display H-Bond button.

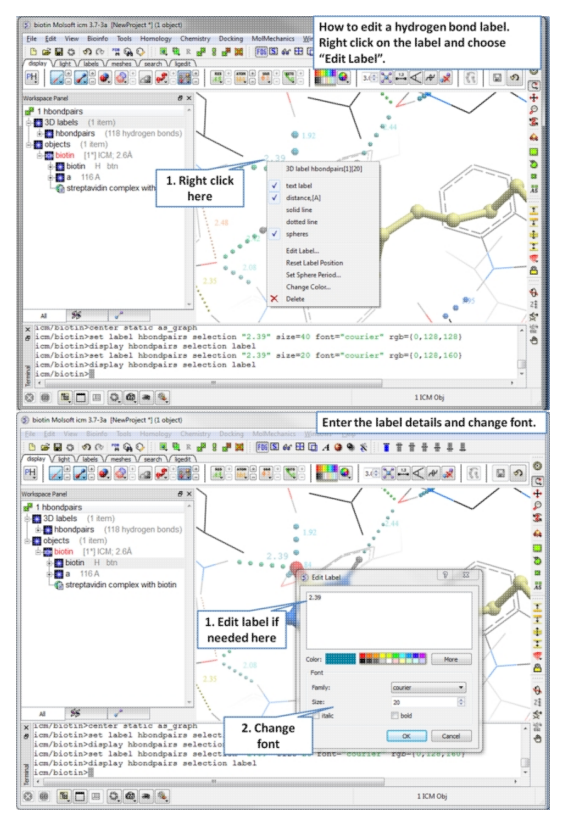


**NOTE:** Different options for displaying the H-bond can be accessed by clicking and holding on the H-bond button in the "Display" tab. The coloring of the H-bonds are red (strong - thick spheres) to blue (weak - thin spheres). Once the hydrogen bonds have been displayed they can be displayed and undisplayed in the 3D labels section of the ICM Workspace (left hand side of graphical window).

#### 4.2.3.1 Edit Hydrogen Bond Label

#### To edit a hydrogen bond label

- Right click on the label.
- Choose Edit Label.
- Make changes in the dialog box and press OK.

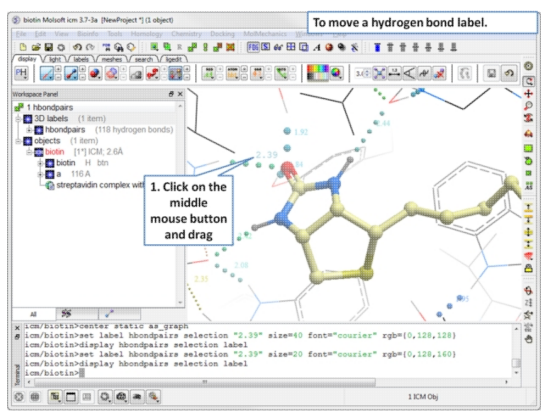


## 4.2.3.2 Move Hydrogen Bond Label

#### To move a hydrogen bond label

- Right click on the label.
- Click on the label using the middle mouse button.

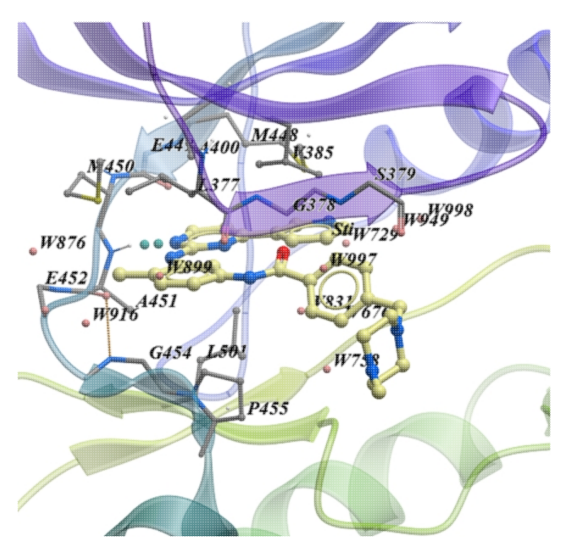
• Hold the middle mouse button down and drag.



## 4.2.4 Ligand Pocket Interactions

To view the ligand pocket interactions.

- Right click on the ligand and choose Pocket.
- Choose Ligand Pocket Interactions. Hydrogen bonds, key waters and neighboring residues will be displayed.



# 4.3 Crystallographic Analysis

Available in the following product(s): ICM-Browser-Pro | ICM-Pro

## 4.3.1 Crystallographic Neighbor

#### Theory

Molecular objects and 3D density maps may contain information about crystallographic symmetry. It consists of the following parameters:

- 1. Crystallographic group eg. P2121 that determine N (depends on a group) transformations for the atoms in the asymetric unit.
- 2. Crystallographic cell parameters A, B, C, Alpha, Beta and Gamma

To generate the coordinates within one cell one needs to apply N transformations and then to generate neigboring cells the content of one cell needs to be translated in space according to the cell position.

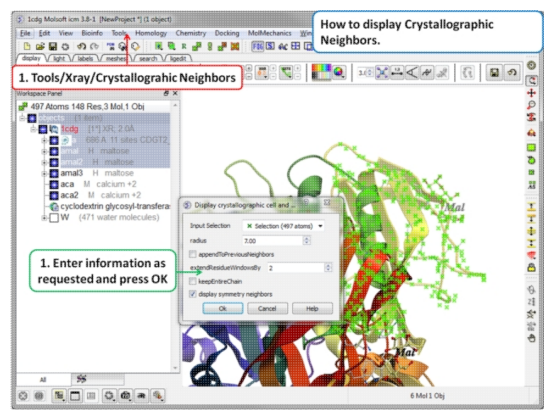
ICM has a function which generates crystallographic neighbors for the selected atoms. For large proteins it is impractical to generate neighbors for the whole molecule due to the high number of atoms in all neighboring molecules.

This information allows to generate symmetry related parts of the density or molecular objects.

#### To generate symmetry related molecules around a selection of atoms:

- Read a PDB file into ICM. For instruction see the section entitled Search PDB.
- Display the structure and select the residues around which the symmetry will be generated. For information on how to select residues see the Making Graphical Selections section.
- Select the menu Tools/Xray/Crystallograhic Neighbors.

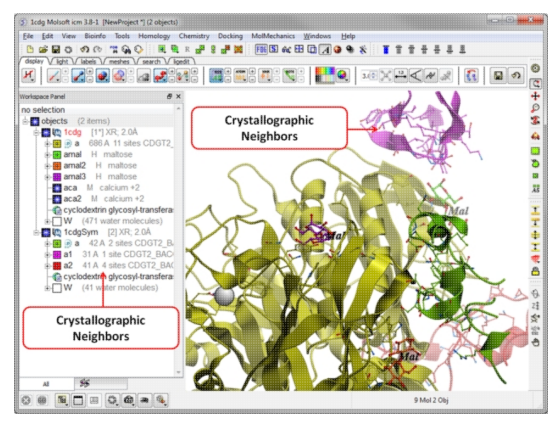
A data entry box as shown below will be displayed.



- Check that the selection is correct in the Input Selection box.
- Enter the radius around your selction from which you wish to construct the symmetry related molecules.
- If you have made symmetry related molecules previously you can select **appendToPreviousNeighbors** otherwise leave unchecked.
- The **extendResidueWindowsBy** option will allow a window of residues outside of the selection radius selected above to be displayed
- If you leave the **keepEntireChain** unchecked then a fragment of each neighbor will be created. If you check this box the full neighbor will be generated
- Check display symmetry neighbors to display them in the graphics window. The nearest neighbor residues will be displayed in xstick representation and the each neighbor colored by molecule.
- Click OK.

The crystallographic symmetry neighbors will be displayed in the Workspace. By default the object will have the object name + "Sym" and each of the neighbors will be individual molecules.

For packing analysis and display you can color each symmetry unit a different color as described in the Structural Representations Color section. This is shown in the picture below.



## 4.3.2 Crystallographic Cell

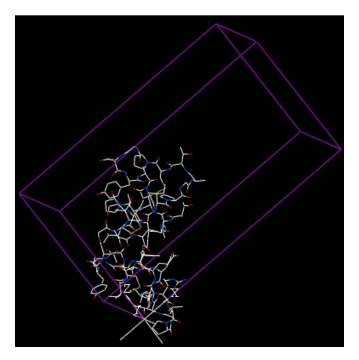
#### Theory

The crystal structure of a protein is often discussed in terms of its unit cell. The unit cell is a box containing one or more motifs, a spatial arrangement of atoms. The units cells are tiled in three-dimensional space to describe the crystal. The unit cell is given by its lattice parameters, the length of the cell edges and the angles between them, while the positions of the atoms inside the unit cell are described by the set of atomic positions measured from a lattice point.

To display the crystal cell of a PDB structure:

- Read a PDB file into ICM. For instruction see the section entitled Search PDB.
- Select the whole object. You can do this by double clicking on the name of the structure in the ICM Workspace (a selection is highlighted blue in the ICM Workspace and green crosses in the graphical display) or you can use the right-click button and drag it over the whole structure in the graphical display.
- Select the menu Tools/Xray/Crystallograhic Cell and a data entry box will be displayed.
- Click OK

The crystallographic cell will be displayed as a box as shown below.



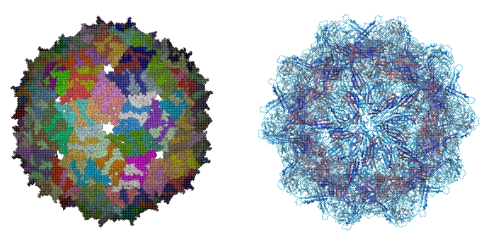
## 4.3.3 Biomolecule Generator

#### Theory

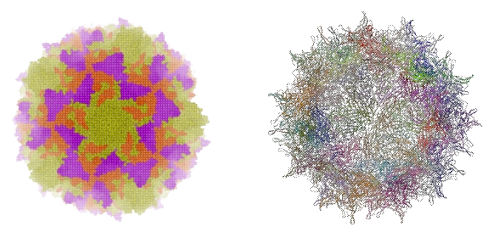
It is very useful to know how a protein from the PDB may look in a biological environment. The PDB entries solved by X-ray crystallography and deposited in the PDB contain the information about the crystal structure rather than the biologically relevant structure. For example, for a viral capsid only one instance of capsid protein complex will be deposited and only one or two molecules of haemoglobin that is a tetramer in solution maybe deposited.

In some other cases the asymetric unit may contain more than one copy of a biologically monomeric protein. ICM reads the biological unit information and has a tool to generate a biological unit. Not every PDB entry has the biological unit information.

A gallery of images created using the ICM Biomolecule generator is shown below:



**Left:** PDB: 1DWN Bacteriophage Pp7 From Pseudomonas Aeruginosa At 3.7 A Resolution **Right:** PDB: 1C8E Feline Panleukopenia Virus Empty Capsid Structure At 3.0 A Resolution



**Left:** PDB: 1AL2 P1/Mahoney Poliovirus, Single Site Mutant V1160I At 2.9 A Resolution **Right:** PDB: 1LP3 Adeno-Associated Virus (Aav-2), A Vector For Human Gene Therapy At 3.0 A Resolution

**NOTE:** Right click on a PDB structure in the ICM workspace to determine whether a structure from the PDB has biological unit information. If it does have this information then there will be an option in the menu entitled "Generate Biomolecules" if not the option will be blanked out.

#### To generate a biological unit with ICM:

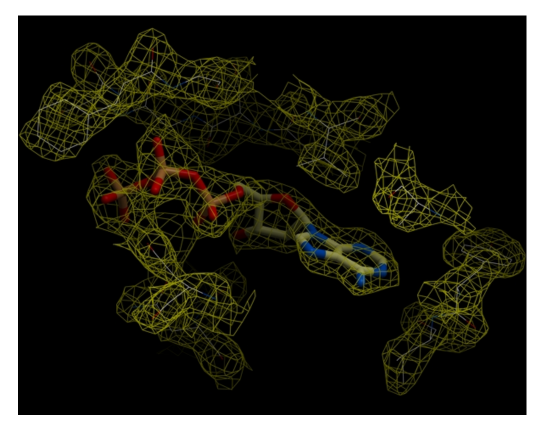
- Select the object or PDB file.
- Select the menu Tools/Xray/Biomolecule Generator.
- Tick the **makeAllBiomolecules** box.
- Click OK with very large molecules the biomolecule generation may take some time.

## 4.3.4 Get Electron Density Map

#### Theory

An electron density map is a representation of a crystal structure based on the diffraction data. The map is constructed by a summation of waves of known phase, amplitude and frequency using Fourier transform. The electron density map of a protein can be viewed along with the pdb structure. The easiest way to view the electron density map is to contour and convert it into a graphical object (mesh).

A figure showing the electron density contours surrounding the ATP molecule in pdb entry 1ATP.

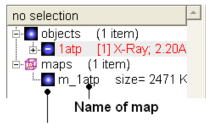


#### To load an electron density map:

- Tools/Xray/Get Electron Density Map
- Enter the PDB code of the map you would like to view.
- Click OK and the map will be downloaded from the Uppsala Electron Density Server.

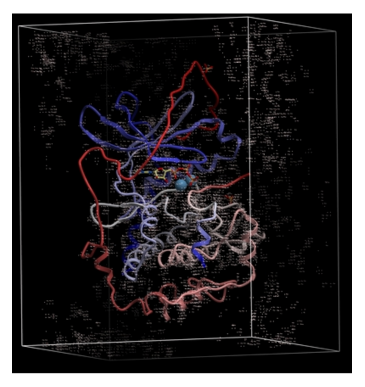
The map will be represented in the ICM Workspace as shown below.

#### ICM Workspace



Display and undisplay map here

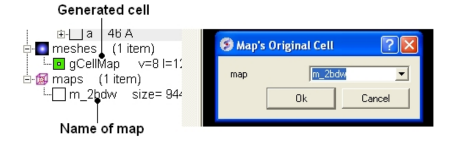
The map can be displayed as shown below however a clearer way of representing the density is to contour the map into a graphical object (mesh) as described in the following section.



## 4.3.5 Map's Original Cell

To display the original crystallographic cell of an electron density map:

- Tools/Xray/Map's Original Cell
- Enter the name of the map or use the drop-down button to locate it. If you do not know the name of the map the name can be located in the ICM Workspace.
- Click OK and the cell will be displayed. The map can be displayed and undisplayed in the **meshes** section of the ICM Workspace.

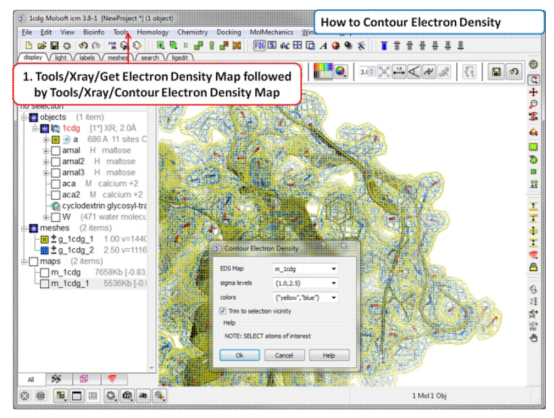


## 4.3.6 Contour Electron Density Map

#### To contour an electron density map and display as a graphical object:

- Load an electron density map as described earlier in the Load Map section.
- Read in the PDB file File/Load PDB or use the PDB search tab.
- Tools/Xray/Contour Electron Density.
- Enter the name of the map e.g. m\_lcdg the name of the map is displayed in the ICM Workspace or use the drop down arrow to locate it.

- If nothing is displayed then the whole map will be contoured. If you only want to contour a particular region of the map then you need to select the region to be contoured (e.g.the binding pocket) and then select the option **Trim to Selection Vicinity**.
- Enter a sigmaLevel value for more information see: http://www.molsoft.com/man/reals.html#mapSigmaLevel. Once the contoured map has been created the sigma level can be changed manually using the +/- buttons in the ICM workspace. Two contour levels at 1 and 2.5 sigma respectively are generated for the 2Fo -Fc map



The sigma level can be changed interactively in the ICM workspace as shown below.



Click here to increase or decrease the sigma level of the contouring

NOTE: Meshes can be cut away using the mesh clipping tools.

## 4.3.7 Convert Xray Density to Grid

For some applications, such as trying to fit a structure to a density map, you may want to extract a sub map and convert to a grid. You can do this by

- First read into ICM a map (eg File/Open or Tools/X-ray/Get Electron Density Map)
- Tools/X-ray/Convert Xray Density to Grid

- Enter the map name or use the drop down list
- Enter a grid size
- Click OK

# 4.4 Protein Superposition

Available in the following product(s): ICM-Browser-Pro | ICM-Pro



One or more proteins can be superimposed. Simply select the molecules or parts of the molecules you wish to superimpose and then use the selection of protein superimpose tools described in this section. A convenient superimpose button can be found in the Display tab (see image of button (left).

#### **Chapter Contents:**

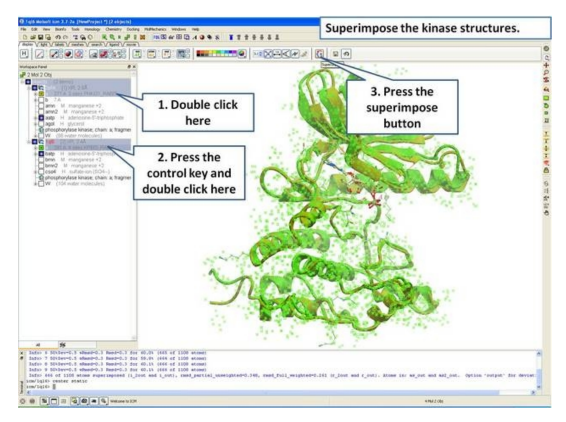
- Select Proteins for Superposition
- Superimpose Button
- Superimpose by 3D
- Superimpose Multiple Proteins
- Arrange as Grid
- Superimpose Sites by Atomic Property Fields

## 4.4.1 Select Proteins for Superposition

Before any superposition operation can be undertaken you need to select the protein structures you wish to superimpose.

One way to do this is by selecting in the ICM workspace. For other selection tools please see the Making Selections section of the manual.

• Select both receptors by double clicking on the name of the molecule in the ICM Workspace. To select two molecules use the Ctrl button or use the shift button to select a range of objects in the ICM Workspace. A receptor which is selected will be highlighted in blue in the ICM Workspace and with green crosses in the graphical display.



Once the molecules are selected you can then superimpose them using the options described in the next section of this manual.

## 4.4.2 Superimpose Button

A convenient way to superimpose two molecules is by using the superimpose button in the **display** tab, ICM will calculate the Ca-atom, backbone atom and heavy atom differences between the two structures. More advanced superimpose options can be found in the **Tools/Superimpose** menu.

To superimpose:

- First load the two structures into ICM.
- Select which parts or all of the two structure you wish to superimpose (see the chapter on Selections or the description protein-superposition-select{here}.).
- Select the **display** tab (previously called Advanced tab) at the top of the GUI.
- Select the superimpose button.



The rmsd will be displayed in the terminal window as shown below:

INTO/ 04 acoms superimposed, rmsu-1.0 icm/ly6> superimpose ( Res( as_graph ) Warning> [110] skipped 4 atom pairs w Info> 64 atoms superimposed, rmsd=1.3 icm/ly6>	& a//ca,c,n,o ) & Obj( as_graph )[1] ith zero occupancies
RMSD di	isplayed here

**NOTE:** You do not need to select the whole molecule, the superimpose button will work on small selections e.g the loop regions or domains.

## 4.4.3 Superimpose by 3D

#### To superimpose proteins by 3D:

- First display and select the proteins you wish to superimpose by 3D.
- Tools/Superimpose/Proteins by 3D
- A window as shown below will be displayed.

🧐 Automated multiple structural superposition 🛛 🔹 🔀					
Help molecules must be SELECTED and belong to different objects. To superimpose molecules in the same object clone it					
	C C alpha C Backbone C Heavy Atoms				
Static Object	a_1q6.				
sequence weight	0.5				
seed length	15 💌				
	Ok Cancel				

- Select by which atoms you wish to superimpose.
- Enter the ICM selection language description for the protein structure you wish to remain static. You can also use the drop down arrow button to select it.
- Enter the sequence weight Average local sequence alignment score.
- Enter the seed length This is the similarity window size.

## 4.4.4 Superimpose Multiple Proteins

#### To superimpoe multiple proteins:

- First display and select the proteins you wish to superimpose by 3D.
- Tools/Superimpose/Multiple Proteins
- A window as shown below will be displayed.

S Automated mul	tiple structural superposition	×			
Align Residue	es 💿 Match By Res Numbers 💿 Exact Match				
Visible Atoms	💿 Visible Atoms 💿 C alpha 💿 Backbone 💿 Heavy Atoms				
Static Object	a_glp1. 🔻				
eighted iterative superposition is single global superposition					
	Ok Cancel Help				

• Select by which method you would like to superimpose

Align Residues - Residue correspondence is established by sequence alignment using the ICM ZEGA alignment Abagyan, Batalov, 1997. Atom alignment: by atom name.

**Match by Res Numbers** - Residue alignment by residue number. Atom alignment: by atom name for pairs of identical residues or pairs of close residues (F with Y; B with D,N; D with N; E with Q or Z, Q with Z), for other residue pairs only the backbone atoms ca,c,n,o,hn,ha are aligned.

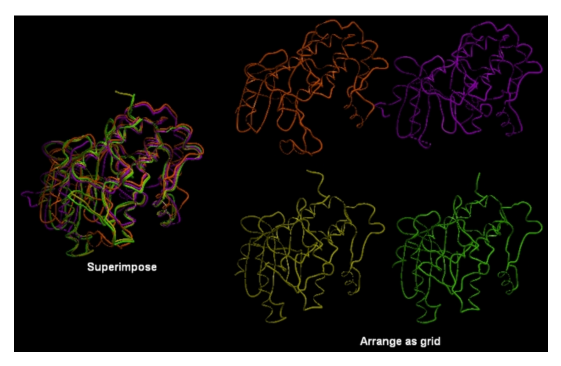
**Exact Match** - Residue alignment is by the Needleman and Wunsch method. Inside residue atoms are aligned sequentially and regardless of the name.

- Select which atoms you would like to superimpose. Visible Atoms, C alpha, Backbone, or Heavy Atoms.
- Select whether you would like to use weighted iterative superposition as described here http://www.molsoft.com/man/icm-commands.html#superimpose-minimize or non-iterative single global superposition.

## 4.4.5 Arrange as Grid

#### To separate superimposed proteins:

• Tools/Superimpose/Arrange as Grid

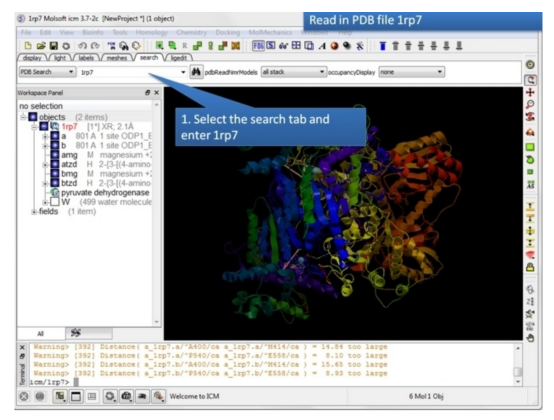


# 4.4.6 Superimpose Sites by Atomic Property Fields

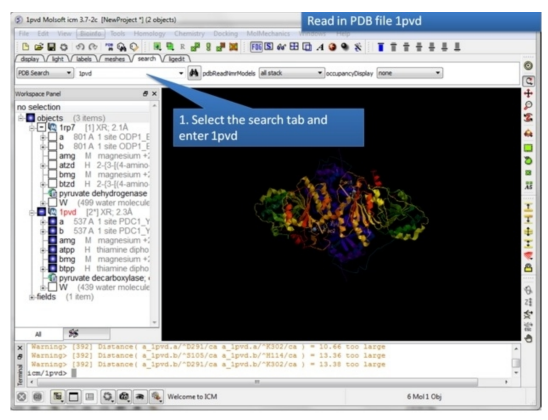
Here we describe how to superimpose and compare a ligand binding site using Atomic Property Fields. This method is described in more detail in this publication.

In this example we superimpose the ligand binding pocket of thiamine diphosphate in the binding sites of pyruvate dehydrogenase (pdb code: 1rp7) and pyruvate decarboxylase (pdb code:1pvd).Even though the sequence identity between both proteins is very low (19%) and the secondary structure surrounding the ligand undergoes considerable displacement you will see that the pockets can still be superimposed very well using the APF method.

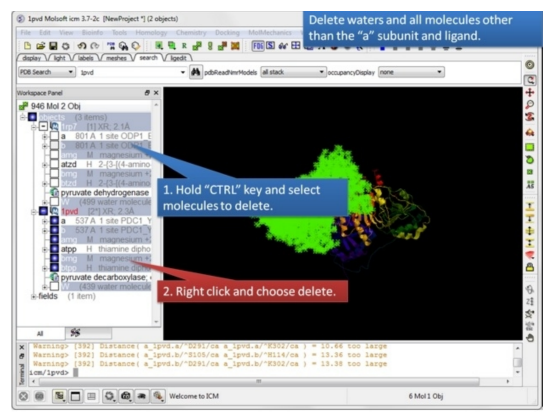
**Some Comments about Selecting the Pocket** APF pocket overlay works by superimposing a layer of receptor atoms around the pocket using APF chemical property fields of these atoms "pocket" of course is something that is not uniquely defined and is somewhat subjective. One way to define it is to use the ligand inside, in which case a layer of receptor atoms within a certain cutoff distance from the ligand is considered in superposition. This is "around selected ligand" mode. It focuses on atoms that directly interact with the ligand while largely disregarding backbone fold, which can be helpful when comparing different pockets binding similar ligands. The other mode uses input selections directly, leaving it up to the user to define the sets of atoms that comprise the two pockets. Depending on the system and degree of similarity, different selections may work better or worse. For instance in the absence of ligand one may want to detect pocket with pocket finder and then select residues around the pocket "blob". Regarding the influence of ligand on superposition, if pocket is defined via the ligand it will of course matter whether the ligands extend into certain regions or not. If external pocket definition is used, ligands will have no direct effect on superposition process.



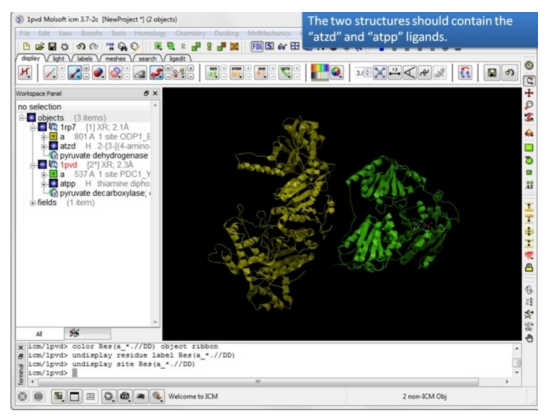
Step 1: Read in the pyruvate dehydrogenase (pdb code: 1rp7) structure.



Step 2: Read in the pyruvate decarboxylase (pdb code:1pvd).



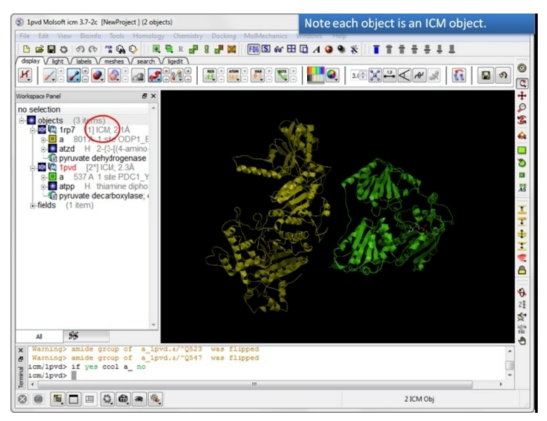
Step 3: Delete unwanted molecules in the b chain and waters.



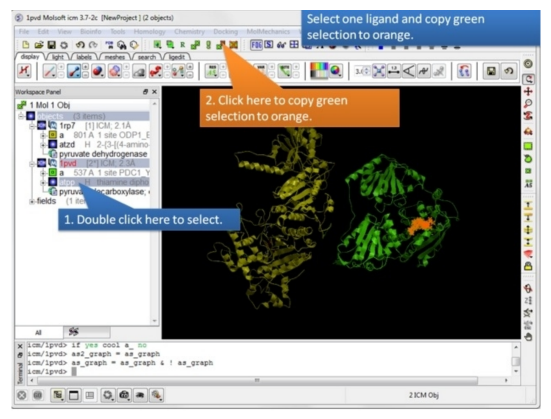
Step 4: You should have two objects containing the protein and ligand.



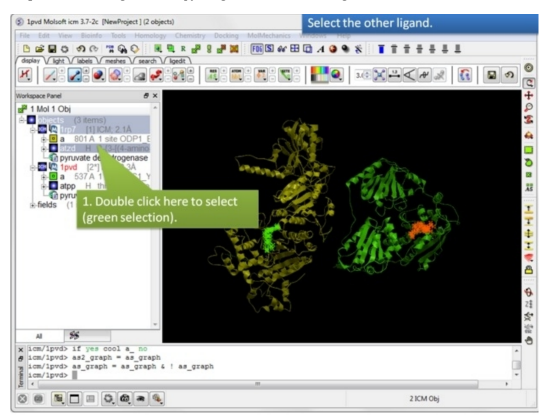
Step 5: Convert both structures to an ICM object.



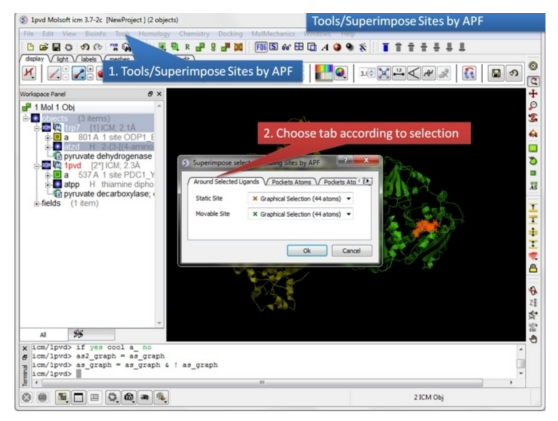
Step 6: If the pdb is converted you will see "ICM" in the workspace.



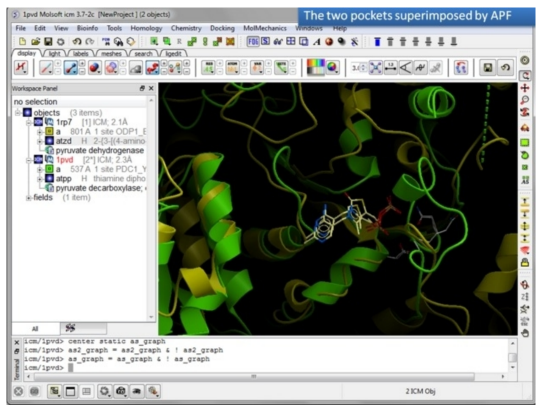
Step 7: Select one ligand and copy the green selection to orange.



Step 8: Select the other ligand (regular green selection).



**Step 9:** Select the option Tools/Superimpose/Sites by APF. Note you can also superimpose pockets - choose the appropriate tabs in the dialog box.



Step 10: Observe the superimposed site.

# 4.5 Protein Structure Analysis

Available in the following product(s): ICM-Browser-Pro | ICM-Pro

In this chapter we describe the tools available for analyzing protein structure. These tools include calculating RMSD, identifying closed cavities, calculating contact and surface area, measuring anlgles and distances, and generating Ramachandran plots.

#### **Chapter Contents:**

- Find Related Chains
- Calculate RMSD
- Contact Areas
- Identify Closed Cavities
- Surface Area
- Measure Distances
- Planar Angle
- Dihedral Angle
- Ramachandran Plot Interactive
- Export Ramachandran Plot

## 4.5.1 Find Related Chains

This option allows you to search the currently loaded PDB files or ICM objects and identify chains which are similar and/or related.

You can do this by:

- Select the objects or pdb files you want to compare.
- Tools/Analysis/Find Related Chains
- Click OK to confirm the selection you made
- A table as shown below will be displayed.

	name1	name2	len1	len2	seqid	rmsd	consensus
1	2jc6.a	2jc6.c	278	277	100	0.12	SWKKQAEDIKKIFEFKETLGTGAFSEVVLAEEKATGKLFA IENEIAVLRKIKHENIVALEDIYESPNHLYLVMQLVSGGEL
2	2jc6.a	3bhh.a	278	289	39	0.42	# # # E #G # V L #TG #A#K#I #+ # #E E# ##R #KH NIV L D # ##YLV# LV GGELF- IV + #Y E DAS #I Q#L-AV###H MG#VHRDLKPENLL##S # # PGY# PEVL #Y K#VD#W #GVI#YILL#GYPPF#DE
3	2jc6.a	3bhh.b	278	289	38	0.41	# # # E #G ## #V #TG #A#K#I #+ # #E E#
4	2jc6.a	3bhh.c	278	285	37	0.44	# ##E# # ## EA # K##+# #EE###R #KH
5	2jc6.a	3bhh.d	278	286	37	0.43	# ##E# # ## EA # K# #+ # #EE# ##R #KH
6	2jc6.c	3bhh.a	277	289	39	0.44	# # # # E #G # VL #TG #A#K#I K #E E# ##R
7	2jc6.c	3bhh.b	277	289	39	0.43	# # # # E #G ## #V #TG #A#K#I K #E E#
8	2jc6.c	3bhh.c	277	285	38	0.44	# # # # # # #V# #TG #A#K#I K #E E# ##R

**name1** = Name of query structure molecule **name2** = Name of hit **len1** = length of query **len2** = length of hit **seqid** = Sequence identity percentage **consensus** = Consensus sequence

## 4.5.2 Calculate RMSD

**NOTE:** This option is for protein structures only not for chemical compounds. You can use the command line options RMSD (http://molsoft.com/man/icm-functions.html#Rmsd) and Srmsd (http://molsoft.com/man/icm-functions.html#Srmsd) for chemicals.

#### To calculate RMSD between two protein structure:

- Read into ICM the two structures (File/Open or PDB Search or Read in Chemical) you wish to compare.
- Select one of the two molecules you wish to compare, you can do this by double clicking on the name of the structure in the ICM Workspace. Convert this selection to an Orange Selection.
- Select the second molecule, and then you should have one orange and one green selection in the graphics display.

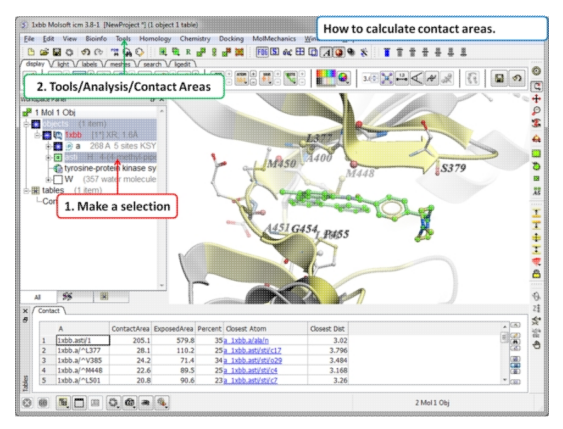
🧐 Calculate cartes	ian RMSD	×
© Superimposed	C Kept in place	
<ul> <li>No Alignment</li> </ul>	C Align Residues C Exact Match	
<ul> <li>VisibleAtoms</li> </ul>	C C alpha C Backbone C Heavy Atoms	
	Apply Close Help	

- Select whether you wish the atoms to be superimiposed onto one another or kept in place. The kept in place option would be ideal for comparing docked structures.
- Choose whether you wish to make the superposition by alignment or exactly matching the atom names.
- Select which atom types you wish to superimpose.

The **RMSD** value will be displayed in the terminal window.

#### 4.5.3 Contact Areas

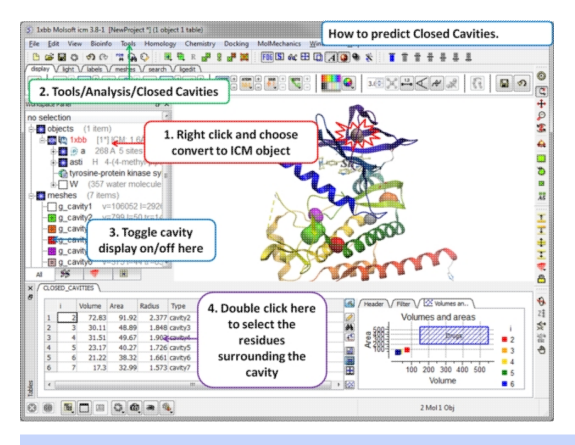
- Read in a protein structure (File/Open or PDB Search).
- Select the region you wish to analyse.
- Tools/Analysis/Contact Areas
- The xstick display in the region will be scaled according to the atom/residue contact area. For example, residues making large contacts with a ligand will be displayed in thicker xstick representation (and colored yellow) than those making less significant contacts.
- A table as shown below will be displayed. The table lists the contact area, exposed area, percentage of contact area compared to exposed, the nearest atom of a residue and the distance.



## 4.5.4 Identify Closed Cavities

This tool will identify cavities within a molecule which are completely closed. If you are looking for buried and open pockets then use icmPocketFinder.

- Read in a protein structure (File/Open or PDB Search).
- Convert the protein structure to an ICM object.
- Tools/Analysis/Closed Cavities
- Use the drop down arrow to locate the molecule you are interested in.
- Enter the minimum volume of the cavities you wish to identify.
- Click OK
- The closed cavities will be displayed in the meshes section of the ICM Workspace and a table of the cavities will be displayed. Double click on a row in the table to jump to a particular closed cavity and select the residues surrounding it.



#### What is the difference between Closed Cavity and ICMPocketFinder?

Closed pocket (cavityFinder) is purely geometrical/topological - it is a part of molecular surface that is completely disconnected from the exterior surface. It means that a probe sphere of 1.4A radius (representing a water molecule) can not pass in or out of the cavity (considering protein as completely rigid of course). icmPocketFinder identifies pockets that are likely to contain ligands (not specifically open or closed pockets). Pockets are defined based on physical interaction rather than geometric criterion. Blobs of 'pocket density' generated by icmPocketFinder represent continuous regions of space where there is significant favorable van der Waals interaction with the receptor.

## 4.5.5 Surface Area

This option calculates solvent accessible area of each selection in multiple objects and stores it in a table. If a molecule is specified in a multi-molecular object, the surface area of an isolated molecule is calculated and other molecules are ignored. The area is reported in square Anstroms and the probe radius is assumed to be the value set in the variable waterRadius.

Output: the macro creates table AREA. The empty comment field is added for user's future use. If the table exists, new rows are appended.

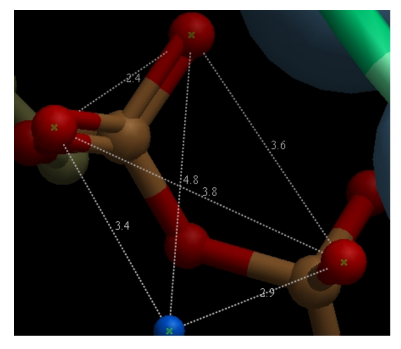
#### To calculate a surface area:

- Read in a protein structure (File/Open or PDB Search).
- Select the region you wish to analyse.
- Tools/Analysis/Surface Area
- A table will be displayed listing the residues in the selection along with the corresponding total surface area.

## 4.5.6 Measure Distances

There are two approaches to calculating and displaying distances between atoms. You can either use the options in the Labels tab or use Tools/Analysis/Distance.

#### To display **all to all** distances:



- Select the atoms between which you would like to find the distance. (See selection toolbar)
- Tools/Analysis/Distance
- Select all to all

To display intermolecular distances

- Select the atoms between which you would like to find the distance. (See selection toolbar)
- Tools/Analysis/Distance
- Select intermolecular

To display the distances between the same atoms in two objects.

- Select the atoms between which you would like to find the distance. (See selection toolbar)
- Tools/Analysis/Distance
- Select same atoms in two objects

**NOTE:** Distances can be displayed and undisplayed in the 3D labels section of the ICM Worskapce. You can change the color of a distance label by right clicking on it in the ICM Workspace. You can alse export the distance to a table.

## 4.5.7 Planar Angle

If you wish to find the planar angle between three atoms:

• Select Tools/Analysis/PlanarAngle

💈 Find planar angle between three atoms 🛛 🔹 🔀					
First atom	a_per	o.m/2/ca	•		
Second atom	a_per	o.m/6/n	•		
Third atom			•		
Help To select atoms: Right-Click, slide down to atom name and release. To see the results: look in the terminal window					
	Apply	<u>C</u> lose		<u>H</u> elp	

• Right click on the each of the three atoms which you wish to use, and select their name. The spaces next to **First atom**, **Second atom**, and **Third atom** should now contain the name of your atoms.

	a_pep.m/2/oe2	
	Selection Dialog	
	Edit	→
7	Advanced	•

• Click **Apply** to display the angle measure in the terminal window.

Angle ( a\_pep.m/6/hh21 a\_pep.m/2/oe2 a\_pep.m/3/o ) = 74.72 deg.

## 4.5.8 Dihedral Angle

In order to find the angle dihedral angle between two sets of atoms:

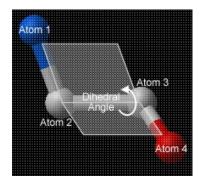
• Select Tools/Analysis/Dihedral Angles.

🂈 Find dihedral angle formed by four atoms 🛛 🔹 🔀					
First atom	a_pep.m/3/n	•	Second atom	a_pep.m/4/ca	•
Third atom	a_pep.m/6/c	-	Fourth atom		•
Help T	o select atoms: Right-(	Click, slide	down to atom nar	ne and release.	
			<u>Apply</u>	<u>C</u> lose	<u>H</u> elp

• Right click on each of the four atoms which you wish to use, and select the name of the atoms. The spaces next to **Atom 1**, **Atom 2**, **Atom 3**, and **Atom 4** should now contain the names of your atoms.

	a_pep.m/2/oe2	
	Selection Dialog	
	Edit	•
7	Advanced	•

• To find the correct angle, select your atoms according to the following diagram:



• Click **Apply** to display your dihedral angle measure in the terminal window.

## 4.5.9 Ramachandran Plot Interactive

To make an interactive ramachandran plot:

- Read in a protein structure (File/Open or PDB Search).
- Select the structure you wish to build the plot for. You can do this by double clicking on the name of the structure in the ICM Workspace (a selection is highlighted blue in the ICM Workspace and green crosses in the graphical display) or you can use the right-click button and drag it over the whole structure in the graphical display.
- Tools/Analysis/Ramachandran Plot Interactive
- The interactive ramachandran plot will be displayed in table called **RAMA**.
- You can view the **Omega**, **Phi/Psi** (Gly) or Phi/Psi angles by clicking on the tabs at the top of the plot. Each point is linked to the data in the table **RAMA** and also to the graphical display. Clicking on a point in the plot will highlight the corresponding angles in the table and also center on this region in the 3D display.

## 4.5.10 Export Ramachandran Plot

- Read in a protein structure (File/Open or PDB Search).
- Select the structure you wish to build the plot for. You can do this by double clicking on the name of the structure in the ICM Workspace (a selection is highlighted blue in the ICM Workspace and green crosses in the graphical display) or you can use the right-click button anddrag it over the whole structure in the graphical display.
- Tools/Analysis/Ramachandran Plot Export

A postscript viewer needs to be downloaded onto your machine in order to view the plot. This can be downloaded from http://www.cs.wisc.edu/~ghost/. Once this software is downloaded you need to tell ICM where it is located by typing the pathname into File/Preferences.

**NOTE:** You can always export the plot as an image directly in ICM without exporting. You can do this by right clicking on the plot and select **save as image**. Another approach could be to export the RAMA table to Excel and use the plotting tools there. You can do this by right clicking on the table name tab and selecting "Export to Excel" or save as ".csv".

## 4.5.11 Protein Health

#### Theory

The protein health option calculates the energy strain of a structure in ICM. It is generally a good idea to investigate the energy strain of any protein structure before undertaking such processes as docking. It is also essential to use this tool after making a model (see Molecular Modeling) to identify strained regions within your model and then some optimization procedure can be undertaken to rectify the problems.

The protein health option calculates the relative energy of each residue for a selection and colors the selected residues by strain. This macro uses statistics obtained in the following paper Maiorov, V.N. and Abagyan, R.A. (1998) Energy strain in three-dimensional protein structures Folding and Design, 3, 259-269.

To use the Protein Health option:

- Read in a protein structure (File/Open or PDB Search).
- Convert your PDB structure into an ICM object.
- Make a selection of the residues you wish to analyze.
- Tools/3D Predict/Protein Health and a window as shown below will be displayed.
- The scale of the coloring can be changed by altering the value within the trimEnergy data entry box.
- Click OK and the structure will be colored according to energy strain (red - high) and a table of residue energy will be displayed in a table.
- A table and plot of Normalized energies for each amino acid in the selection will be displayed. The table is ranked and colored by residues with poor normalized energies. Click on a row in the table or plot to center in on the residue.

## 4.5.12 Local Flexibility

This option systematically samples rotamers for each residue side-chain in the input selection and uses resulting conformational ensembles to evaluate energy-weighted RMSDs for every side-chain atom. These are stored in the 'field' values on atoms and can be used for example to color the structure by side-chain flexibility. Conformational entropy for each residue side-chain is also calculated and stored in a table. If 1\_entropyBfactor flag is on, the atom rmsds are normalized within the residue to reflect its total conformational entropy. If 1\_bfactor flag is set, the bfactors are reset to the same values that are placed in the atom 'field', and occupancy is set to be inversely proportional to it (O=1/(1+2\*rmsd))

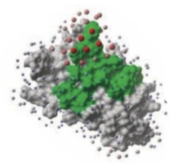
- Read pdb file (File/Open or PDB Search Tab).
- Convert to an ICM Object.
- Tools/3D Predict/Local Flexibility

## 4.5.13 Protein-Protein Interface Prediction

The ICM Optimal Docking Area method is a useful way of prediciting likely protein-protein interaction interfaces. If you do not have mutational data or other experimental data which indicates the likely protein-protein docking site this method will be useful. This procedure can save you time during the docking procedure by focusing your docking only on areas on the receptor and ligand most likely to interact.

#### Theory

**ODA** (Optimal Docking Areas) is a new method to predict protein-protein interaction sites on protein surfaces. It identifies optimal surface patches with the lowest docking desolvation energy values as calculated by atomic solvation parameters (ASP) derived from octanol/water transfer experiments and adjusted for protein-protein docking. The predictor has been benchmarked on 66 non-homologous unbound structures, and the identified interactions points (top 10 ODA hot-spots) are correctly located in 70% of the cases (80% if we disregard NMR structures). For a description of the method see Fernandez-Recio et al Proteins (2005) 127: 9632.



#### To display the optimal docking area.

- Convert the PDB file to an ICM object.
- Tools/3D Predict/Protein Interface by ODA
- If you select the **Residue Table** option the average ODA score for each residue will be displayed in a table. The lower the number the higher the chance the residue will be involved in protein-protein interactions. Regions colored red represent low ODA score and blue represents a high score.

## 4.5.14 Identify Ligand Binding Pockets

#### Theory

The ICM Pocket Finder method (1-2) uses only the protein structure for the prediction of cavities and clefts. No prior knowledge of the substrate is required. The position and size of the ligand-binding pocket are determined based on a transformation of the Lennard-Jones potential by convolution with a Gaussian kernel of a certain size, a grid map of a binding potential and construction of equipotential surfaces along the maps. The pockets are displayed graphically as a surface and the dimensions of each pocket are presented in an interactive table and plot.

Factors that can influence ligand binding to a pocket include the pocket volume and area, buriedness, hydrophobicity, and how compact the pocket is. All these properties are calculated using ICMPocketFinder and tabulated. Scientists at Merck used MolSoft.s ICMPocketFinder algorithm to define a way for quantifying "drugability" of a protein target (3). The metric they used is called Drug-Like-Density (DLID), and this score is also provided in the results table. A good example of the use of the icmPocketFinder method is in the database called Pocketome. The Pocketome (www.pocketome.org) is an encyclopedia of conformational ensembles of all druggable binding sites that can be identified experimentally from co-crystal structures in the Protein Data Bank (4).

1. An, J., Totrov, M. & Abagyan, R. Pocketome via comprehensive identification and classification of ligand binding envelopes. Mol. Cell. Proteomics 4, 752 (2005).

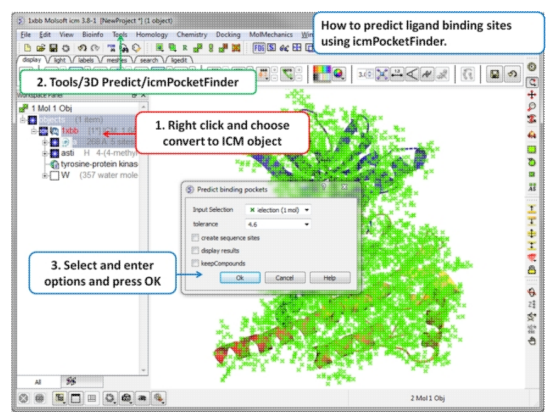
2. Abagyan, R. & Kufareva, I. The flexible pocketome engine for structural chemogenomics. Methods Mol. Biol. Clifton NJ 575, 249.279

#### (2009).

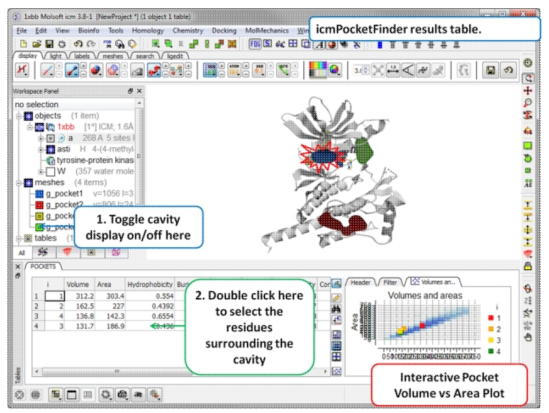
3. Sheridan, R. P., Maiorov, V. N., Holloway, M. K., Cornell, W. D. & Gao, Y.-D. Drug-like density: a method of quantifying the .bindability. of a protein target based on a very large set of pockets and drug-like ligands from the Protein Data Bank. J. Chem. Inf. Model. 50, 2029.2040 (2010).

4. Kufareva, I., Ilatovskiy, A. V. & Abagyan, R. Pocketome: an encyclopedia of small-molecule binding sites in 4D. Nucleic Acids Res. 40, D535.540 (2012).

#### To predict pockets:



- Read in a protein structure (File/Open or PDB Search).
- Convert the protein structure to an ICM object.
- Tools/3D Predict/icmPocketFinder
- Enter a tolerance level (4.6 is the default value and we recommended you to use this). The lower the tolerance value the more pockets predicted and the higher the tolerance the less pockets predicted.
- Check the box **create sequence sites** if you wish the site to be labeled.
- Check the box **display results** to see the predicted pockets as grobs in the display panel.
- Check the box **keep compounds** if you wish the compounds (ligands) in the receptor to be included in the prediction. If you dont check this box the pockets will be calculated based on the receptor without ligands.
- Click OK to run icmPocketFinder and the results will be displayed in a table.



#### Results

- Click on a row of the results table to select the residues surrounding the pocket.
- The pocket view can be toggled on or off in the meshes section of the ICM Workspace. Right click on the name of the mesh in the ICM workspace to change properties such as color.
- A fully interactive plot (Årea vs Volume) is also provided, pockets that fall within the blue shaded region have a "drug-like" volume and size.

#### About the icmPocketFinder Results Table

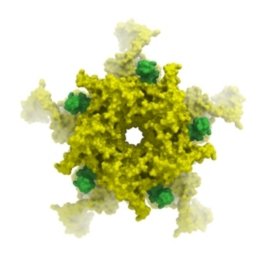
Factors that can influence ligand binding to a pocket include the pocket volume and area, buriedness, hydrophobicity, and how compact the pocket is. These values are reported in the results table:

- Volume of the pocket in <sup>^</sup>.
- Area of the pocket in <sup>^</sup>.
- **Hyrdophobicity** represents the percentage of the pocket surface s in contact with hydrophobic protein residues (values can range from 0-1)
- **Buriedness** The buriedness parameter is calculated as follows: One measures the solvent accessible surface area of the pocket (probe radius, 1.4) in isolation. Then one measures the solvent accessible surface area of the pocket covered by its shell. The ratio of the second number to the first is the fraction buried. The lowest possible value is 0.5; i.e. the pocket is completely open and the surface flat. The highest is 1.0, i.e., completely buried.
- **DLID** Merck's Drug-like density score (see Sheridan et al JCIM 2010). >0.5 is considered "druggable".
- **Radius** gives an indication of how sprherical a pocket is. (3/4 Volume / Pi )^1/3, i.e. radius of an ideal spherical cavity of the same volume as a pocket blob.
- Nonsphericity give an indication of how spherical the pocket is. Area / (area of ideal spherical cavity). It is 1.0 if the cavity is spehrical

- **Conservation** Average conservation (%identity) of residues in contact with the pocket. Calculation is performed if multiple alignment is present and linked to the protein chain being analyzed.
- **RelCons** Above conservation, relative to the average over the entire chain.
- **Type** returns the ICM selection for the residues forming the pocket.

## **5 Molecular Graphics**

In this chapter we describe how to make beautiful graphical representations of molecules and manipulate them in the 3D graphics window. This includes how to change color, light, representations, clipping planes, and how to use built in graphics effects. We also teach how to label and annotate molecules displayed in the graphical user interface.



#### **Chapter Contents:**

- Molecule Representation
- Meshes-Surface-Grobs
- Coloring
- Lighting
- Labeling and Annotation
- Display Distances and Angles
- Graphics Effects
- Graphics Shortcuts
- Molecule Move Buttons
- Clipping Tools
- Graphic Layers
- High Quality Publication Images
- Movies

## **5.1 Molecule Representation**

Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro

To change the molecule display representation:

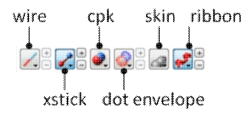
- Select the atoms, residues, molecules, or objects you wish to change in the graphical display or in the ICM Workspace.
- Then use the molecule representation (e.g. wire, ribbon) options in the Display Tab.

The display tab contains tools for a variety of functions including structural representations, coloring, labeling and superposition.

There are six main types of structural representation in ICM. They are wire, ball and stick (Xstick), ribbon, skin, CPK and dot envelope (surface).

#### To display one of these representations:

• Click on the representation button you desire in the **display** tab.



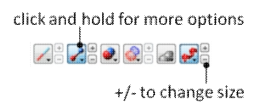
# To remove a displayed representation or to toggle between display and undisplay:

• Click on the corresponding representation button in the **display** tab.

**Identify which representations are displayed:** The button display will change appearance (shaded blue) when selected. This makes it easier to identify which representations are currently being displayed.



**Change the display and size of the representations** Many characteristics of the graphical representation such as color can be changed by clicking and **holding** on the button. The size can be changed by clicking the plus(+) and minus(-) buttons next to them.



## 5.1.1 Wire Representation

Click and hold on the **wire representation** button. A menu will be displayed as shown below.

Wire — Display	analysis	V pdb search V meshes V movie
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	1 12	normal
	'n	thick 💢
	1 12	undisplay other representations
	۹۲	reset to default
	2	

#### To change the wire style:

• Click and hold on the **wire representation** button and then click on **wire, chemistry** or **tree.** 

#### To change the size of the wire representation:

• Click and hold on the **wire representation** button and then click on **thin**, **normal** or **thick**.

**NOTE:** Clicking on the +/- next to the **wire representation** button also changes the thickness of the wire representation.

#### To undisplay representations other than wire:

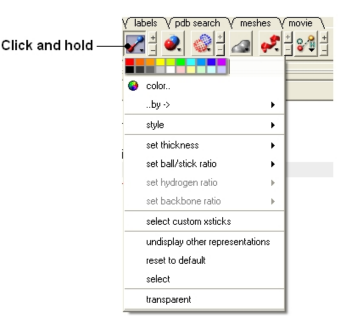
• Click and hold on the **wire representation** button and then click on undisplay other representations.

If you make a mistake or you are not happy with the way your structure is displayed with the wire representation:

• Click and hold on the **wire representation** button and then click on reset to default.

## 5.1.2 Stick and Ball (Xstick) Representation

Click and hold on the **stick and ball representation** button. A menu will be displayed as shown below.



#### To change the style of the Xstick representation:

• Click and hold on the **stick and ball representation** button and then click on style and choose either **plain**, **chemistry**, **Chem Plus**, or **Chem Plus Aromatic**.

#### To change the size of the Xstick representation:

• Click and hold on the **stick and ball representation** button and then click on **set thickness**, **set ball/stick ratio**, **set hydrogen ratio**, and **set backbone ratio**.

**NOTE:** Clicking on the +/- next to the **xstick representation** button also changes the thickness of the xstick representation.

#### In order to make some parts of your picture clearer, the xstick representation can be set to transparent:

• Click and hold on the **stick and ball representation** button and then click on transparent.

#### To undisplay representations other than xstick:

• Click and hold on the **stick and ball representation button** and then click on undisplay other representations.

#### If you make a mistake or you are not happy with the way your structure is displayed with the xstick representation:

• Click and hold on the **stick and ball representation** button and then click on reset to default.

## 5.1.3 Ribbon Representation

Click and hold on the ribbon representation button and a menu will be displayed.

#### To change the style of the Ribbon representation:

• Click and hold on the **ribbon representation button** and then click on a style option: **smooth**, **wide**, **wide smooth**, **cylinders**, **protein worm**.

#### To assign secondary structure:

To accurately represent the secondary structure of the molecule in ribbon representation you may wish to assign secondary structure:

• Click and hold on the **ribbon representation** button and then click on **assign sec. structure**.

## To make some parts of your picture clearer, the ribbon representation can be set to transparent:

• Click and hold on the **ribbon representation button** and then click on **transparent**.

#### To undisplay representations other than ribbon:

• Click and hold on the **ribbon representation** button and then click on undisplay other representations.

## If you make a mistake or you are not happy with the way your structure is displayed with the ribbon representation:

• Click and hold on the **ribbon representation button** and then click on **reset to default**.

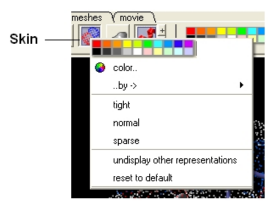
**NOTE:** Always use the **ICM** assign sec.\*\* structure tool in the ribbon right click menu to get accurate secondary structure assignment. This is particularly important when studying helices which may have non-cannonical elements within them such as 3/10 or pi. To view non-cannonical helix segments use the segment option in the ribbon right click menu.

#### To change the display of chain breaks (dotted lines):

- Click and hold on the ribbon representaion button.
- Select the options Display Chain Breaks or Display Chain Break label.

## 5.1.4 Skin Representation

Click and hold on the **skin representation button**. A menu will be displayed as shown below.



## To make some parts of your picture clearer, the skin representation can be set to tight, normal or sparse:

• Click and hold on the **skin representation button** and then click on either tight, normal or sparse.

#### To undisplay representations other than skin:

#### 5.1.3 Ribbon Representation

• Click and hold on the **skin representation button** and then click on **undisplay other representations**.

If you make a mistake or you are not happy with the way your structure is displayed with the skin representation:

• Click and hold on the **skin representation button** and then click on \*\* reset to default\*\*.

**NOTE:** Sometimes due to singularity problems holes may appear within the skin surface. To cure this infliction select atoms nearby and right click select Advanced->RandomizeAtoms

## 5.1.5 CPK Representation

Click and hold on the **CPK representation button**. A menu will be displayed as shown below.



To undisplay representations other than CPK:

• Click and hold on the **CPK representation button** and then click on **undisplay other representations**.

If you make a mistake or you are not happy with the way your structure is displayed with the cpk representation.

• Click and hold on the **CPK representation button** and then click on **reset to default**.

## 5.1.6 Surface Representation

Click and hold on the surface representation button. A menu will be displayed as shown below.

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	tight		1
	normal		
	sparse		
	set dot density	۲	
	set probe radius	۲	
	set dot size	۲	
	undisplay other representation	s	
	reset to default		
	select		

#### To change the style of the surface representation:

• Click and hold on the **surface representation button** and then click on **tight**, **normal**, or **surface**.

#### To undisplay representations other than surface:

• Click and hold on the **surface representation button** and then click on undisplay other representations.

## If you make a mistake or you are not happy with the way your structure is displayed with the surface representation:

• Click and hold on the **surface representation button** and then click on **reset to default**.

## 5.1.7 Display and Undisplay Hydrogens

To display and undisplay hydrogens. Click and hold on the **"Change Hydrogen Display"** button shown below. Multiple single clicks will toggle through the hyrogen display options.

- Display Tab
- Click and hold on the "Change Hydrogen Display" button shown below.

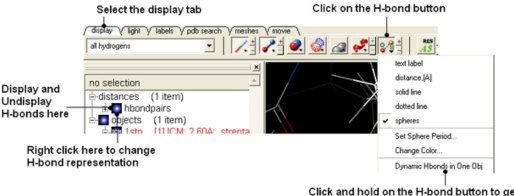
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н	No Hydrogens				
PH	Polar Hydrogens				₽×
н	All Hydrogens				<u></u>
LH	Ligand All, Rec Polar				
	Display Formal Charges				
	Ribbon+CPK				
	Ligand+Ribbon				
1	Atoms				
	Chemical				
1	Undisplay Beyond Selection				
1	Elegant Ribbon+Ligand Sketch				
1	Green Wireframe				

## 5.1.8 Display Hydrogen Bond

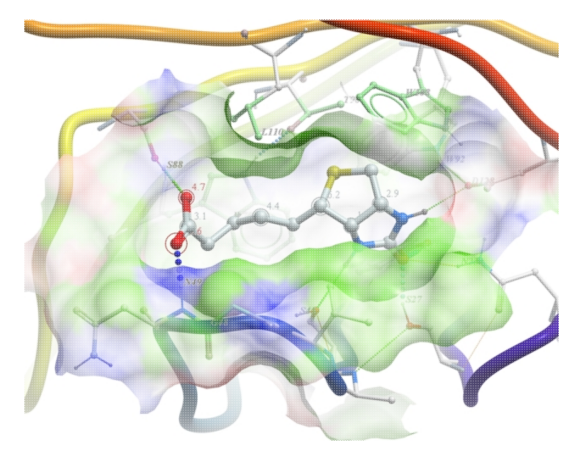
**NOTE:** The method by which hydrogen bonds are calculated is described here in the command line manual. The GRAPHICS.hbondMinStrength parameter determines the hbond strength threshold for hbond display. The strength value is between 0. and 2. By changing 1. to 0.2 you will see more weak hydrogen bonds.

In order to display potential hydrogen bonds in your structure:

- Convert to an ICM Object
- Make a selection if you are trying to display the H-bonds between a ligand and the receptor make sure the ligand is part of the selection.
- Click the Display Tab.
- Click on the **Toggle H-bonds** icon in the **display** tab.

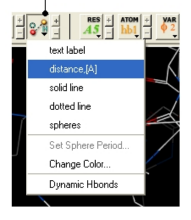


Click and hold on the H-bond button to get access to other display options



- Click the +/- on the right of the H-Bond button to change thickness of H-bond representation.
- Click and hold the button to change representation or use the **hbondpairs** option in the ICM Workspace.

## Click and hold



#### What do the default coloring of the H-bond represent?

Longer and shorter H-X distances in the hydrogen bond are color-coded, from red to blue, respectively.

**NOTE** Dynamic hydrogen bonds can be set by clicking and holding on the **H-bond toggle** button in the **Display** tab. Hydrogen bonds will then respond to any changes made to the ligand.

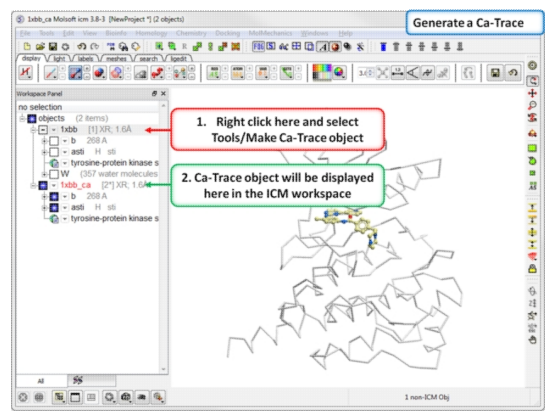
## 5.1.9 Display Formal Charges

You can display formal charges by clicking and holding on the "Change Hydrogen Display" button in the Display tab.

## 5.1.10 Ca (carbon alpha) Trace

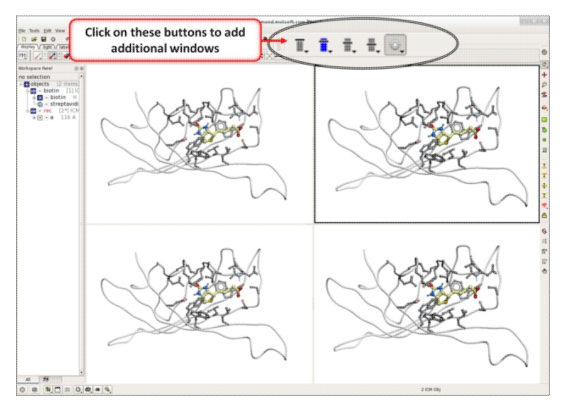
To generate a carbon alpha (Ca) trace:

- Right click on the object in the ICM workspace and select Tools/Make Ca-Trace object.
- The Ca-Trace object will be displayed in the ICM workspace.

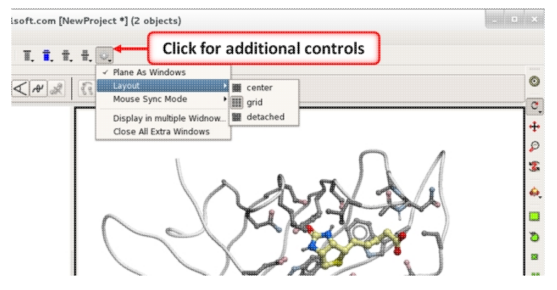


## 5.2 Multi-Windows

To display more than one window click on the buttons shown below.



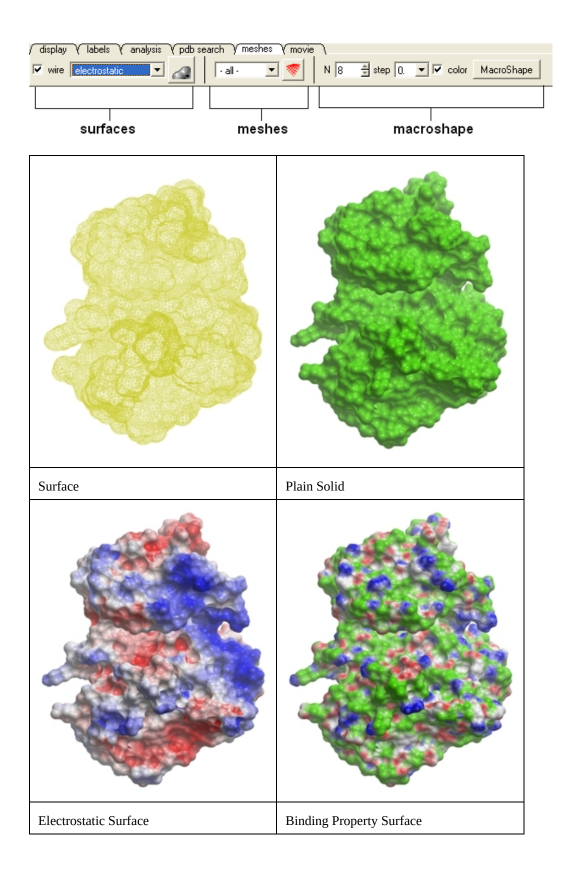
To control the layout and mouse sync mode click on the button shown below.

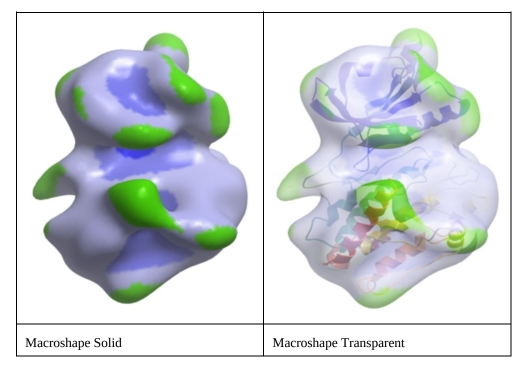


5.3 Meshes - Surface - Grobs

Available in the following product(s): ICM-Browser-Pro | ICM-Pro

Click on the tab button entitled **'meshes'** and more graphics tools for surfaces are available. In ICM surfaces are sometimes referred to as meshes or graphical objects (Grobs).

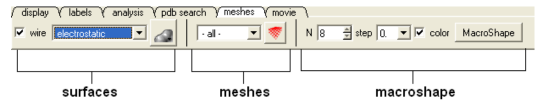




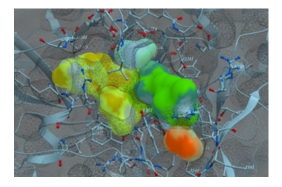
## 5.3.1 Surfaces

The surface of your structure can be displayed and colored by **electrostatics** or **binding** properties. To do this:

- Load a structure into ICM File/Open or tab-pdb{PDB Search}
- Convert the structure into an ICM object.
- Select the 'meshes' tab button.
- Click on the drop down arrow menu shown below and select which surface you wish to generate.
- Click on the generate surface button next to the drop down arrow.



## 5.3.2 MolSkin



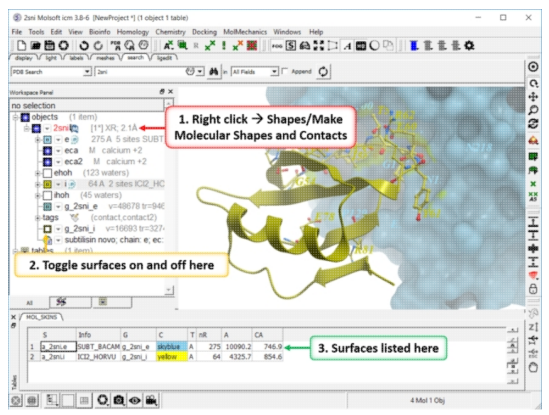
MolSkin is a new macro that instantly makes publication quality graphics and it gives each molecule a color and colors contact patches. It

undertakes the following:

- Molecular surfaces are generated with those assigned colors and occlusion shaded.
- Carbons of each molecule are colored by its own consistent color.
- Labels, ribbons, are colored accordingly.
- Surfaces are gradually cross-colored by each other to mark the contact patches.
- The labels are brought to the surfaces.

To use MolSkin:

- Right click on the object in the ICM workspace and choose Shapes/Make/Molecular Shapes and Contacts.
- Once the surfaces are made they will be listed in the ICM workspace where they can be toggled on or off.



## 5.3.3 Color Surface by Proximity

To color a surface by proximity to neighboring molecules:

- Make a surface.
- Right click on the molecule you want to color by proximity and choose Shapes/Make/Color by Proximity.

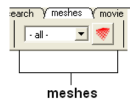
## 5.3.4 Color Surface by Selection

To color a surface by a selection:

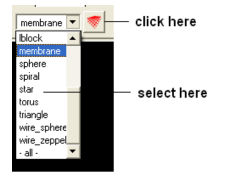
- Make a surface.
- Make a selection.
- Right click on the molecule you want to color by selection and choose Shapes/Make/Color by Selection.

## 5.3.5 Meshes

A variety of shapes can be constructed automatically using ICM. These shapes are referred to as meshes. The types of shapes you can build are shown in the drop down option in the meshesh tab

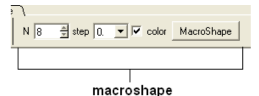


To make a shape select it from the menu by clicking on the down arrow and then click the button next to the menu. The shape will then be displayed in the 3D graphics window.



## 5.3.6 Macroshape

A macroshape can be constructed and allows easy viewing and manipulation of the structural representation. A macroshape representation is ideal for large structures which allows the user to easily identify important regions of the structure and facilitate the return to the 'standard' view of a particular molecule. All the buttons needed to display a macroshape structure are shown below in the 'meshes' tab.



#### To construct a macroshape:

- Load a molecule into ICM File/Open or tab-pdb{PDB Search}
- Select the amount of detail required in the shape by increasing the values in 'N' or 'step' data entry box (note the default values are usually sufficient).
- Check the 'color' if you wish your molecule to be colored.
- Click the button labeled 'MacroShape'.

Macroshape can also be used from the View menu: View/Macro Shape

## 5.3.7 Google 3D Objects (Sketchup)

#### To read in a 3D Mesh from Google in KMZ or COLLADA format:

- File/Load/ 3D Mesh in KMZ or COLLADA Format from Google
- Search for the object you would like to view and download it.
- To read the file go to File/Open

## 5.3.8 Display or Undisplay Meshes or Surfaces

To display or undisplay the surface click in the box in the ICM workspace as shown below:

#### ICM Workspace

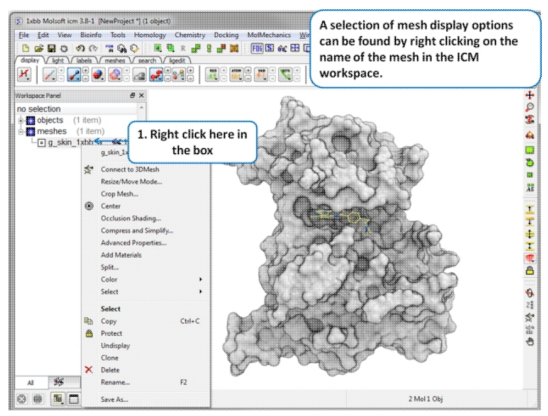


or undisplay surface

**NOTE:** All surfaces, meshes and macroshapes come under the one heading of **meshes** in the workspace panel.

## 5.3.9 Mesh Options.

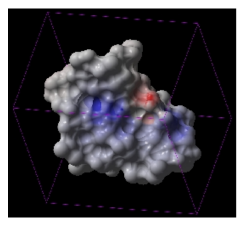
A number of options relating to meshes can be found by right clicking on the mesh in the ICM Workspace. This section describes some of these options.



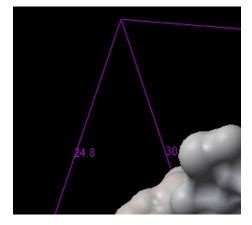
### 5.3.9.1 Resize Mesh

Once a mesh has been created you can move it and resize it. To do this:

- Locate the mesh in the ICM Workspace and right click on it
- Select the **Resize/Move Mode** in the menu.
- A purple box as shown below will surround the molecule.



To resize the mesh click on one of the corners of the box and drag to the required size. The number displayed on the edges of the box represent the dimensions.



#### 5.3.9.2 Move Mesh

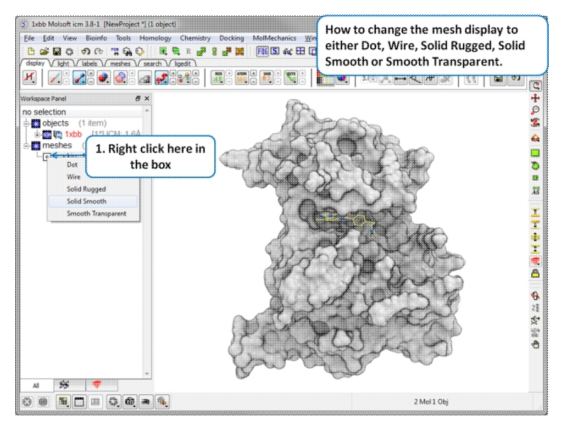
To move a mesh:

- Locate the mesh in the ICM Workspace and right click on it
- Select the **Resize/Move Mode** in the menu.
- Click on the mesh in the graphical display with the **middle** mouse button.

#### 5.3.9.3 Mesh Representation

There are five different display modes:

- Dot
- Wire
- Solid Rugged
- Solid Smooth
- Smooth Transparent.



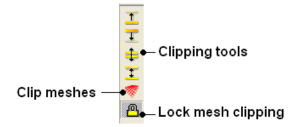
### 5.3.9.4 Mesh Color and Lighting

There are a number of options to color and change the display of the mesh. These options can be accessed simply by right clicking on the mesh name in the ICM Workspace as shown below.

## 5.3.10 Mesh Clipping

Clipping tools can be used to adjust the frames of the mesh independently of other objects.

The buttons shown below can be used for this purpose.



The buttons used for clipping are described in the section entitled Clipping Tools.

To clip the skin independently of the object as shown in the image above:

- Display object and mesh.
- Click and hold on clip meshes button(red button) and select Reset Clipping to Default and check the option Clipping enabled
- Click on the front clipping plane button.

• (Optional) To lock the mesh clipping click on the yellow lock mesh clipping button.

## 5.3.11 Save Mesh

You can save a mesh as a wavefront object by right clicking on the mesh in the ICM Workspace and selecting **SaveAs**.

## 5.3.12 Occlusion Shading

The occulusion shading option provides better representation of depth within a cavity. The color of each surface element of a grob (mesh) is changed by mixing its own color with the background depending on the burial of the surface element.

#### To add occlusion shading:

- Right click on the mesh in the ICM Workspace and select **Occlusion Shading**. The occlusion shading value can also be changed before generating the mesh in the meshes tab.
- Enter a depth value default is 0.8. Higher values will generate a more dramatic shading.

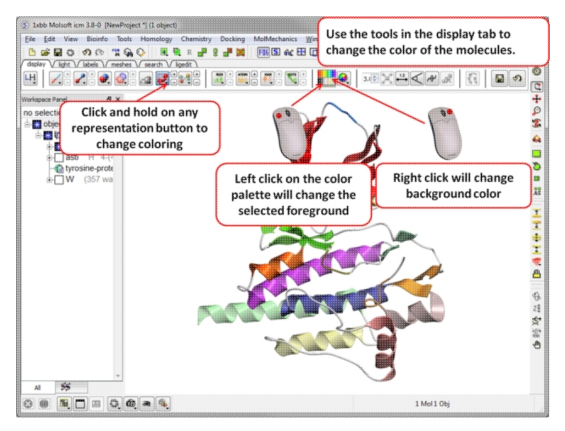
See example here.

## 5.4 Coloring

Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro

To change the coloring of molecules:

- Select the atoms, residues, molecules, or objects you wish to color in the graphical display or in the ICM Workspace.
- Then use the color options in the Display Tab.



## 5.4.1 Coloring

To change the color of a structural representation such as CPK, Xstick, wire or ribbon.

- Select the atoms, residues, molecules, or objects you wish to color.
- Click and hold on the structural representation button for the representation you wish to color (e.g. wire, ribbon etc...) in the **Display** tab.
- Select a color by clicking color.

## To color by a particular parameter such as atom type, b-factor, secondary structure etc...

- Click and hold on the structural representation button for the representation you wish to color (e.g. wire, ribbon etc...) in the **Display** tab.
- Select .. by-> option

#### To change the color of everything displayed :

Click on the color palette in the Display Tab.

## 5.4.2 Color Background

#### To change the color of the background:

• Select View/Color background.

🏅 Color bad	kground		? 🛛
Color	black	<b>.</b>	
	<u>0</u> k	<u>C</u> ancel	<u>H</u> elp

• Click on the square of your desired color. If you are not satisfied with the color palate, click on the arrow next to the colors to customize a color.

#### OR

• Right click on a color in the colors panel in the display tab.



Right click here in the display panel to set the background color.

## 5.4.3 Background Image

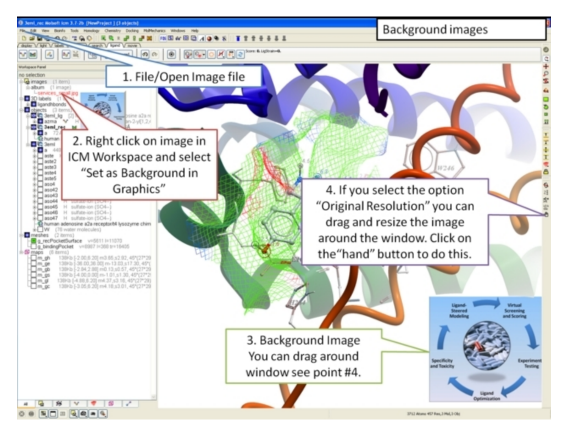
A background image can be added to the graphical display. This can be useful for making cool images or for comparing structures (e.g.compare displayed object with background image of object).

#### To add a background image from an image file (png or jpeg):

- File/Open Image
- Right click on the image in the ICM Workspace and select "Set as Background in Graphics."
- Choose one of the following: (1) Original Resoution, (2) Original Resolution Centered, or (3) Scale to Fit
- Enter the Scale (%)
- Enter whether you want to keep existing background.

To move and resize a background image: Version 3.7-2b and higher.

- File/Open Image
- Right click on the image in the ICM Workspace and select "Set as Background in Graphics."
- Choose the option **Original Resoution**.
- Click on the drag atom button (looks like a hand).
- Click and drag image.
- Remember to deselect drag atom button.

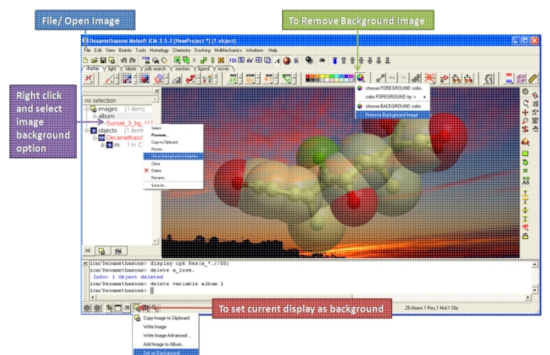


#### To set current display as background image:

• Click and hold on the "Copy Image to Clipboard" button at the bottom of the gui and select the "Set as Background" option.

#### To remove a background image:

• Select the **display** tab and then click and hold on the color sphere button and select "Remove Background Image".



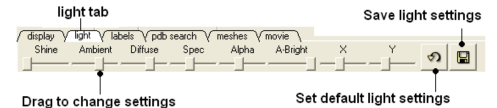
- Right click on the image in the ICM Workspace and select "Set as Background in Graphics."
- Choose the option **Remove From Background**

## 5.5 Lighting

OR

Available in the following product(s):  $\texttt{ICM-Browser} \mid \texttt{ICM-Browser-Pro} \mid \texttt{ICM-Pro}$ 

These options are in the light tab



CLick and drag the sliders to change the lighting. You can also save your preferred lighting settings and return to default.

Shine - shininess property of the solid material

Ambient - ambient light intensity of RGB for ambient light

Diffuse - diffuse light intensity of RGB for diffuse light

Spec - specular light intensity of RGB for specular light

Alpha - transparency setting for grob

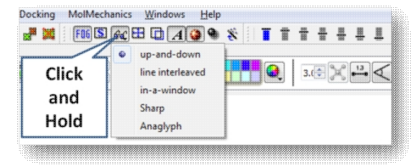
A-Bright - light intensity shinning on grob

 $\mathbf{X}$  and  $\mathbf{Y}$  - Change the position of the light source in the graphics window

## 5.6 3D Stereo

```
Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro
```

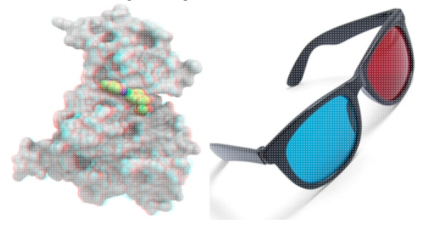
Click and hold on the stereo hardware button (see image below).



There are 5 different options available which are described here http://molsoft.com/man/icm-table.html#GRAPHICS.stereoMode.

- 1. up-and-down
- 2. line interleaved
- 3. in-a-window
- 4. Sharp
- 5. Anaglyph

The Anaglyph option is the easiest to used with inexpensive 3D glasses and and without any expensive 3D compatible hardware or monitors. The 3D effect is better with a lighter background.



## 5.7 3D Printing

You can save surfaces in the STL format for 3D printing. To do this:

- Make a mesh e.g. go to the meshes tab plain solid.
- The mesh (surface) will then be displayed in the ICM workspace (left hand side).
- Right click on the mesh and choose Save as.. and choose stl from the format option list

## **5.8 Labeling and Annotation**

Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro

To add labels or display or undisplay pre-defined annotation:

- Select the atoms, residues, molecules, or objects you wish to label in the graphical display or in the ICM Workspace.
- Then use the label options in the Display Tab.

To add new user-defined annotation:

- Select the atoms, residues, molecules, or objects you wish to label in the graphical display or in the ICM Workspace.
- Right click on the selection and choose "Annotate Selection".

## 5.8.1 Labeling

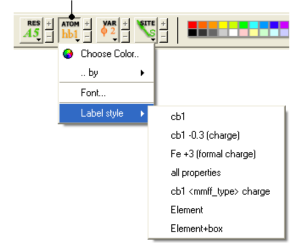
Labeling options are contained within the Labels or Display Tab. In many cases clicking and holding a label button will allow you to view more options.

## 5.8.2 Labeling Atoms

Select the atoms you wish to label (see display structure or selection toolbar).

- Select the **display** tab.
- Click the label ATOM button.

### Click and hold to change label options



#### To change the level of label detail:

• Click and hold the **label ATOM** button and select the desired level of label detail, color or style.

## 5.8.3 Labeling Residues

#### To label residues:

- Select the **display** tab.
- Select the residues you wish to label (see display structure or selection toolbar).
- Click the label RES button.

## Click and hold for more options

A5         Hb1         \$	
by->	J
Label Style 🔹 🕨	A5
Drag Labels	Ala 5
Font	ALA 5
Shift to Sidechain Tips	Ala
Shift to Calphas	ALA
Restore Positions	Alanine 5
	5
	A
	A
	<molname></molname>
	<objname></objname>

#### To change the level of label detail:

• Click and hold the **label RES** button and select the desired level of label detail or style.

## 5.8.4 Move Residue Label

#### To change the location of your residue label:

- Select View/Drag res labels.
- If your mouse has a middle mouse button, then click on handle (as shown) of the label you wish to move, and drag it to your desired area.

Click on	abel
this area	-Dubei
to drag	
your	
label.	

• If your mouse has no middle mouse button, then click on the Translation icon on the toolbar, and click on the handle (as shown) of the label you wish to move, and drag it to your desired area.

The +/- buttons on the side of the Residue and Atom buttons will shift the label. There are also other **residue label move** options available when you click and hold the residue label button. These options include **Shift to Sidechain Tips, Shift to Calphas**, and **Restore Positions** 

## 5.8.5 Label Variables

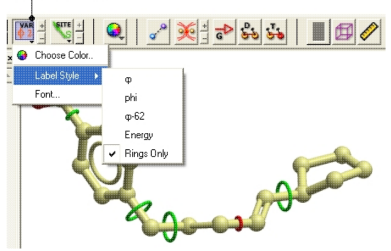
## To label variable angles (dihedral-torsion, planar and phase angle) the molecule needs to be converted into an ICM object.

- Convert the molecle to an ICM object.
- Select the atoms for which you would like to display the variables (see display structure or selection toolbar).

- Click on the **toggle variable label** button shown above located in the **display** tab.
- Change the font size by using the +/- buttons.
- Change the font and color by clicking and holding on the variable atom label button.

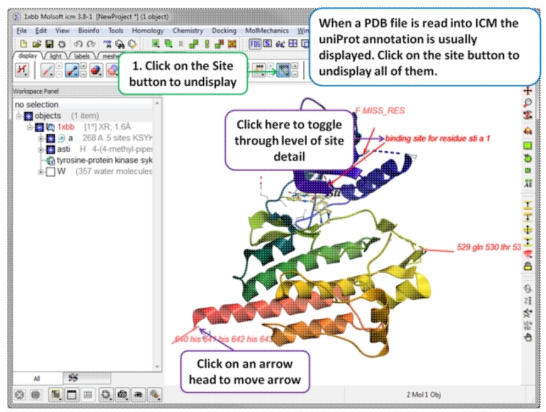
To change the variable label style click and hold the variable atom label button as shown below.

### Click and hold

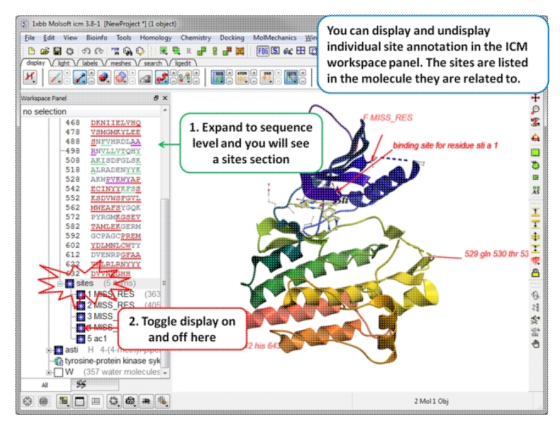


Rings of varying diameter and color are superimiposed on rotatable bonds. Green rings with large diameter are considered less constrained than rings with small green rings. Red rings are highly constrained. When the **Label Style/Energy** option is selected the first number displayed represents the bond angle, the second the energy and the third the worst energy that could be achieved by rotating the bond.

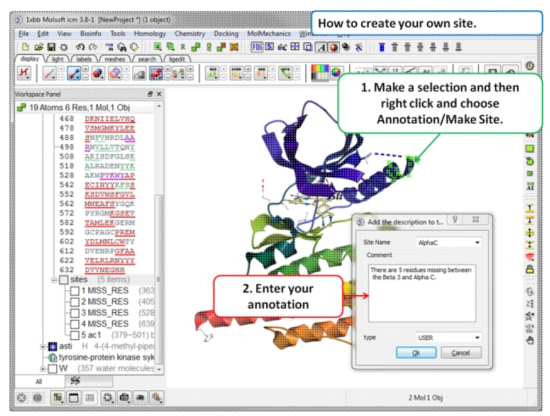
## 5.8.6 Sites and Annotation



When a PDB file is read into ICM the sequence functional/mutation sites listed in Uniprot are automatically displayed. To undisplay all displayed sites click on the **Sites** button in the display tab. Click in the bottom left hand corner of the annotated site to change the level of detail or click on an arrow head to move the location.

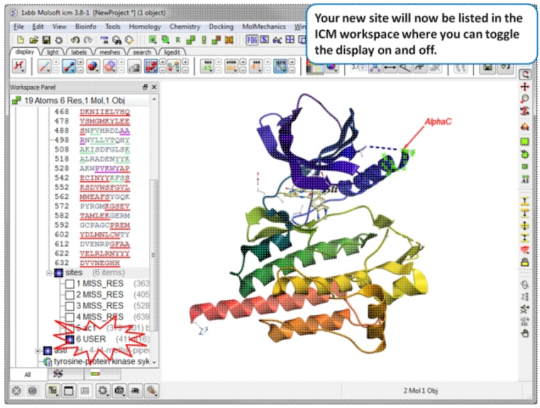


Individual sites can be displayed or undisplayed in the ICM workspace.

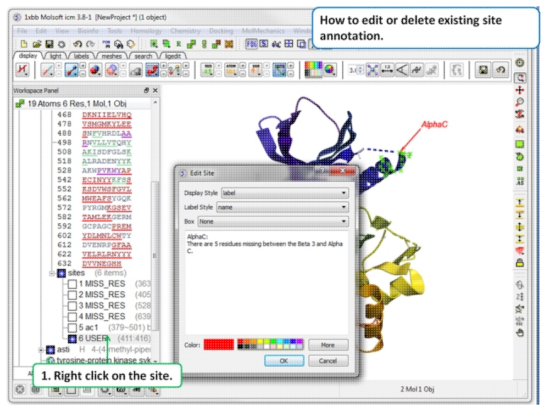


To create your own annotated site:

- Select the region you wish to annotate as a Site.
- Right click on the selection.
- Choose the option Annotate/Make Site
- Enter the annotation into the text box and select ok



Your new site will now be listed in the ICM Workspace.



To edit or delete a site:

• Right click on the site in the ICM workspace.

## 5.8.7 Changing Label Colors

#### To change the color of any label:

• Click and hold down the required label button and a menu as shown below will be displayed.



• Select color.

## 5.8.8 Customized Label 2D or 3D

To generate a customized a label:

- Select the labels tab.
- Select either 2D or 3D button.
- Enter your label and select the desired color, font and size.

2	Select 2D or 3D label	Undisplay labe		
- L	S New Label	ubel here	•	_ Add custom text
	Color: Color: Font	times	More	
	Size: italic Help:	20	Cancel	

To **edit** or **delete** a label - right click on the label in the graphical display as shown below.

### Right click here to Edit or Delete label

Crystal Structure of 2CI label 1: C Edit X Delete	rysta S Edit Label		
	Color: Font Family: Size: italic	CRN Courier 20 bold OK	More Cancel

To move a custom 3D label you need to right click on it in the ICM workspace (under meshes) and choose Connect/Move independently. Press the Esc key to disconnect when you have finished moving it.

## 5.8.9 Undisplay Customized Label

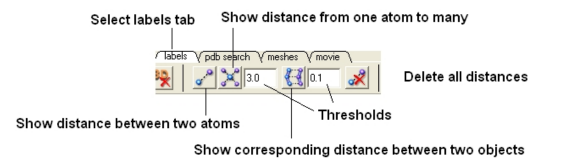
**Undisplay Residue, Atom, and Variable Label** Any label that is displayed can be undisplayed by selecting the region of the molecule related to the label and clicking on the corresponding label button in the labels tab. For example if you wish to undisplay an atom label - click the atom label button. If a label is displayed the coresponding button in the **display** tab will be shaded blue. When you delete the button will return to grey. 2D and 3D labels have an undisplay button (red cross on the button see customized label section).

Undisplay 2D or 3D label Click on the undisplay label button in labels tab.

**NOTE:** A label can also be deleted by right clicking on the label in the graphical display and selecting **delete**.

## 5.8.10 Labeling Distances

Within the **labels** tab there are tools for calculating and displaying distances. These tools can also be found in the Tools/Analysis menu.



#### To display distance between two atoms:

- Click on the labels tab (previously called advanced tab).
- Select the atoms between which you would like to find the distance. (See selection toolbar)
- Click on the 'Show Distances Between Two Atoms' Button
- The distance will be displayed in angstroms, in green.



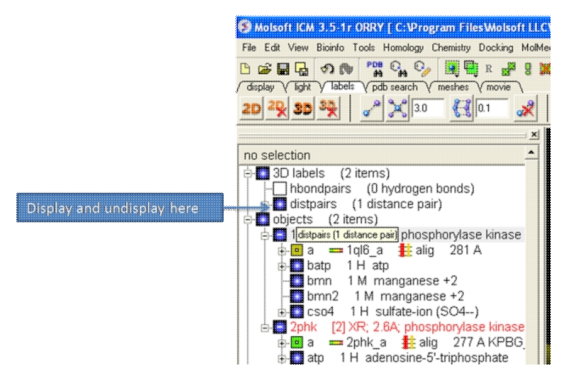
#### To find the distance from one atom to many:

- Click on the labels tab (previously called advanced tab).
- Select the atom from which you wish to measure the distance from (See selection toolbar)
- Click on the 'Show Distances From One Atom To Many' button.
- The distances will be displayed in green.

The maximal and minimal distances can be selected by entering values in the boxes shown here (below) in the labels tab (previously called Advanced tab).



**NOTE:** Distances can be displayed and undisplayed in the 3D labels section of the ICM Workspace. See image below.



#### To change the color of the distance label

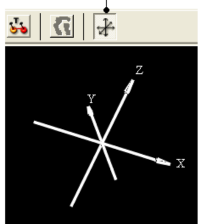
• Right click on the **distpairs** under the **3D labels** section of the ICM workspace and select **Change Color**.

## 5.8.11 (Un)display Origin

To display and undisplay the axis of the coordinate frame (origin):

• Select the **labels** tab and select the **toggle origin** button.

Display or undisplay origin button - located in the labels tab



## 5.8.12 Displaying Tethers

#### Theory

A tether is a harmonic restraint pulling an atom in the current object to a static point in space. This point is represented by an atom in another object. Typically, it is used to relate the geometry of an ICM molecular object with that of, say, an X-ray structure whose geometry is considered as a target. Tethers can be imposed between atoms of an ICM-object and atoms belonging to another object, which is static and may be a non-ICM-object. You cannot create tethers in ICM-Browser, however, if the project that you have loaded contains tethers between two objects, then they can be displayed:

- Click on the display tab (previously called advanced tab).
- Click on the 'Toggle Tethers' button.

### 5.8.13 Displaying Distance Restraints

#### Theory

A distance restraint imposes a penalty function on the distance between two atoms in the same object. You cannot create distance restraints in ICM-Browser, however, if the project that you have loaded contains distance restraints, then they can be displayed:

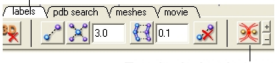
- Click on the **display tab** (previously called advanced tab).
- Click on the 'Toggle distance restraints' button.

## 5.8.14 Display Clash

To display a clash the file needs to be an ICM Object.

- Select the region around which you would like to identify clashes.
- Select the display tabs and the "toggle clashes" button shown below.

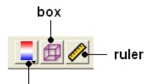
#### Labels tab



Toggle clashes button

# 5.8.15 Display Rainbow, Box, Ruler

To (un)display a rainbow scale, box or ruler use the buttons shown below located in the Labels Tab.



rainbow (click and hold to change colors)

# 5.8.16 Display Gradient

This button is located in the **display** tab.

#### Toggle energy gradient button



This option is described in detail in the language manual http://www.molsoft.com/man/icm-commands.html#display-gradient

# **5.9 Display Distances and Angles**

Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro

# 5.9.1 Display Distance Between Two Atoms - the quick way

- Click on the **Display** tab
- Click on the **Distance between two atoms** button shown below.
- Click on the atoms you wish to measure.
- Distance will be displayed in the graphical display. You can turn this on and off in the ICM Workspace panel under the heading **3D labels**.



Display distance

# 5.9.2 Display Planar Angle

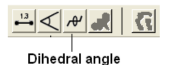
- Select the **display** tab.
- Select three atoms.
- Select the button shown below.



Display planar angle

# 5.9.3 Display Dihedral Angle

- Select the **display** tab.
- Select four atoms.
- Select the button shown below.



# 5.9.4 Delete Label

To delete distance or angle labels

- Select the **display** tab.
- Select the delete distance or angle label button shown below.

#### Delete distance or angle labels button in display tab



# 5.10 Graphics Effects

Available in the following product(s):  $\texttt{ICM-Browser} \mid \texttt{ICM-Browser-Pro} \mid \texttt{ICM-Pro}$ 

All the visual effects tools can be accesed by the View Menu or click on the corresponding button in the View Tools panel shown below.



# 5.10.1 Fog

**Fog Toggle**(Ctrl + D) : this feature creates a fog-like environment for your object, so that the part of your structure that is closer appears clear and the distant parts are faded as if they are in fog. The clipping planes control the point at which the fog begins.

• View/Fog

### 5.10.2 Shadows

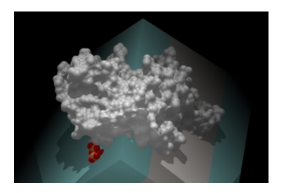
• View/Shadows

#### OR

select the shadow button shown below.

# Toggle shadow

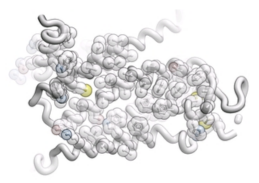




## 5.10.3 Sketch Accents

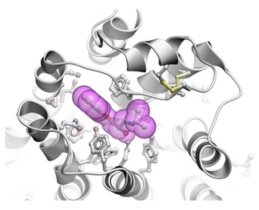
#### To make images as shown below use:

• View/Sketch Accents



# 5.10.4 Elegant Ribbon & Ligand Sketch

- Display Tab
- Click and hold Hydrogen button
  Select Elegant Ribbon+Sketch



# 5.10.5 Occlusion Shading

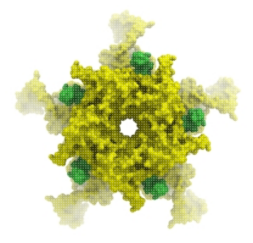
Occlusion shading is for surfaces and meshes to give a perception of depth.

#### To add occlusion shading:

Right click on the mesh in the ICM Workspace and select Occlusion Shading. The occlusion shading value can also be changed before •

generating the mesh in the meshes tab.

• Enter a depth value - default is 0.8. Higher values will generate a more dramatic shading.



# 5.10.6 Perspective

**Toggle perspective Ctrl\_P** this will add perspective to your structure, enhancing depth in the graphical display.

• View/Perspective

# 5.10.7 Animate View

Learn how to build fully interactive and interruptable animations.

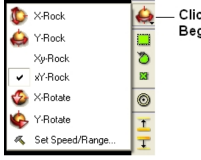


Smooth Animated Transitions

### 5.10.7.1 Make Animation

To quickly produce an ICM Molecular Animation:

- Click and hold down the "Begin rocking/rotation" button shown in the picture below.
- Choose from the following options X-Rock, Y-Rock, Xy-Rock, XY-Rock, XY-Rock, X-Rotate, and Y-Rotate.



#### Click and HOLD DOWN the Begin rocking/rotation button

#### 5.10.7.2 Change Speed, Range and Cycle Length of Animation

To change the speed, range and cycle length of the animation:

- Click and hold down the "Begin rocking/rotation" button shown in the picture above.
- Choose the set speed range option and change the speed and range using the drag bars. Any change will appear in the graphical display behind this box.
- If desired you can change the number of cycles of the animation. This is an ideal tool for screen-shot movie making.

S Rocking Preferences
Speed         1.00           Range         1.00
Cycles
C Endless movement     Number of cycles     4
OK Cancel Set Defaults

**NOTE:** There is a return to default button in the Rocking Preferences dialog box shown above and defaut values can be changed in File/Preferences/Gui.

**NOTE:** Default rocking speed can be changed in the File/Preferences/Gui menu.

#### 5.10.7.3 Interrupt Animation

An ICM Animation or Transition is fully interactive and is interrupted by a single click of the mouse.

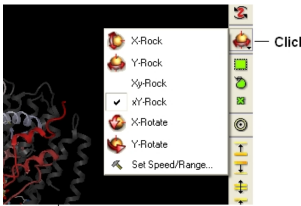
To stop or change an animation or transition:

• Click the "Begin rocking/rotation" button shown in the picture below.

To temporarily halt an animation or transition:

• Click in the graphical display. Once you release the mouse button the animation will start again.

**NOTE:** If you click on the graphical display during an animation the animation will be interrupted. Whilst clicking and holding the mouse button other operations can be performed such as zooming and selections.



Click in the graphical display window to temporarily interrupt an animation

### 5.10.7.4 Saving an Animation

An animation can be saved in an ICM project:

File/Save Project

Or

as a slide.

# 5.11 Graphics Shortcuts

Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro

The left mouse button can be mapped onto different graphics tools which can be selected from the right hand tool bar.

Note: (1) You can access many non-rotation modes directly from the rotation mode by using Middle and Right-mouse buttons, as well as by using the right, top and left margins of the graphics window. (2) You can access the rotation mode from non-rotation modes by pressing Ctrl.

- rotation ( **the default** , press Ctrl if you in the non-rotation modes )
- translation ( the middle mouse button in the rotation mode)
- zooming in and out by dragging the mouse up and down (the left margin in the rotation mode, or use the mouse **wheel**)
- Z-rotation ( the top margin in the rotation mode)
- selecting by box (the right mouse click in the rotation mode)
- selecting by lasso ( Ctrl-draw lasso in the rotation mode )
- picking out atoms ( a toggle )
- picking out and labeling residues (a toggle)
- moving the front clipping plane ( the top section of the right margin in the rotation mode)
- moving the rear clipping plane ( the bottom section of the right margin in the rotation mode)
- moving the slab ( the middle section of right margin in the rotation mode)
- unclipping ( Ctrl-U )
- rotating torsions (Ctrl-left-mouse-click in the rotation mode)

Click here to stop an animation

• connect and unconnect separate molecules to movement controls

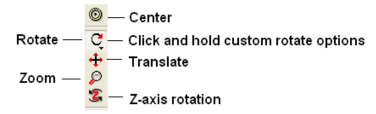
Many useful graphics tips are summarized here.

```
NOTE: Key mouse controls are summarized in the command line manual here http://www.molsoft.com/man/graphics-controls.html
```

# 5.12 Molecule Move Buttons

```
Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro
```

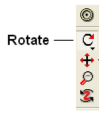
To move your structure it must first be displayed in the graphics window (for instructions on how to display a structure see the Display Tab). All of the following options are displayed in the Move Tools toolbar (shown below).



# 5.12.1 Rotation

In order to achieve the best pose for a picture or to enable the study of a certain region of your structure in more detail you may need to rotate the structure:

• Click on the **rotation** icon on the toolbar.



• Click and drag on your structure in the display window until it is in the desired position.

# 5.12.2 Custom Rotation

An option is provided to customize the rotation of the molecule. This allows exact rotation by a specified number of degrees.

Rotate — Click and hold custom rotate options

- Click and hold down the rotation button and a data entry box as shown below will be displayed.
- Enter the number of degrees of rotation you require and in which X, Y or Z coordinate.

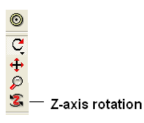
📀 Custom Scen	e Rotation		? 🔀
Rotate around: -	CΥ	СZ	
Rotate by: 180.			degrees
45.	90.	180.	270.
	Apply	OK	Cancel

#### To continuously rotate the picture:

- Click on the **continuous rotation** icon on the toolbar.
- Click, hold, and slightly move your mouse anywhere on the graphical display window. The point at which you hold your mouse, is the direction to which the object will turn.
- Positioning the mouse towards the center of the display will move the object slower than if the mouse is positioned towards the edge of the graphical display.

In order to rotate your picture around the Z-axis:

• Click on the **Z-axis rotation** icon on the toolbar.

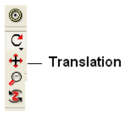


• Click and drag your object around the Z-axis until it is in the desired position.

# 5.12.3 Translation

To translate your structure up, down, left, or right:

• Click on the **translation** icon on the toolbar.



• Click and drag on your structure in the display window until it is in the desired position.

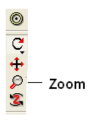
When you are displaying more than one object and you wish to translate one object in relation to the other on the Z-axis:

- Right click on the name of the object you wish to move in the ICM workspace and select connect to object. This object is now independent from the other object and can now be manipulated separately.
- Click on the **Z translate** icon on the toolbar.
- Click and drag your structure along the Z-axis, moving it closer or further from your unconnected structure.
- Once you are finished, right click on the name of the object which is connected, and click on disconnect.

## 5.12.4 Zoom

#### To zoom in or out of your structure:

• Click on the **zoom** icon on the toolbar.



• Click and drag your mouse up to zoom in and down to zoom out.

You can also zoom in and out directly with the right-mouse-button *without* explicitly switching to the zoom tool, if you use the **left 5%-margin** of the graphics window.

## 5.12.5 Center

# To restore your picture to the center of the graphical display window or to center on a selection:

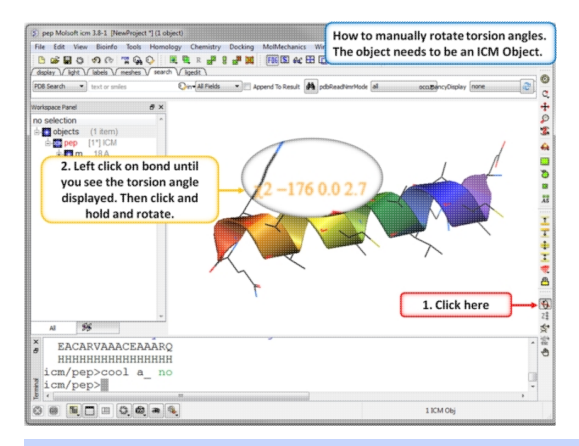
- Make a selection of the region you wish to zoom into if no selection is made the whole structure will be centered.
- Click on the **center** icon on the toolbar.



# 5.12.6 Torsion Angles

# To alter the torsion angle of certain residues of your structure manually:

- Convert your pdb structure into an ICM object.
- Click on the **change torsion angles** icon on the toolbar (see image below).
- Click on the bond until you see the torsion angle displayed in yellow (see image below). Click and hold on the bond around which you wish to rotate a residue. The changing torsion angle will be displayed in orange.



**NOTE:** This option can be used more effectively in conjunction with the variable label option.

#### To alter torsions by entering specific angle values:

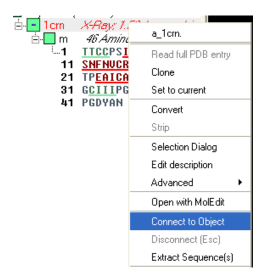
• You can edit torsions by specifically defining the exact angle as described here.

# 5.12.7 Connect (Move)

When there is more than one object displayed in the graphical display window the objects are connected to one another. If you wish to move or manipulate one object independently from the others you need to **connect** to it

#### To do this from the ICM Workspace:

• Right click on the name of the object you wish to move in the ICM workspace and select **Connect to Object**. The object will now be colored yellow.

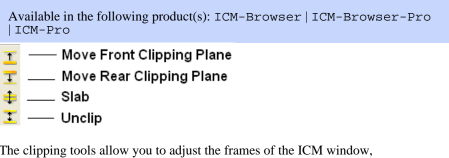


- The object is now controlled separately from the rest of your objects by your mouse.
- Disconnect your object by once again right clicking on the name of the object in the ICM Workspace and selecting disconnect in the drop down menu or **Press the ESCAPE key**.

Note: you can temporarily switch to the global rotation in the connected state if you press Shift

Note: use the Escape button to disconnect

# 5.13 Clipping Tools



The clipping tools allow you to adjust the frames of the ICM window, changing the clipping planes.

Clipping planes can also be moved *without* switching to the clipping tool, if you click the right hand margin of the graphics window:

- The top section of the right 5% margin of the graphics window: moves the back clipping plane
- The middle section of the right 5% margin of the graphics window: moves the slab (both clipping planes)
- The bottom section of the right 5% margin of the graphics window: moves the front clipping plane

In order to move the front or rear clipping planes of your screen:

- Click on the **Move front clipping plane** or **Move rear clipping plane** icons on the toolbar.
- Click and drag the respective plane frontward or backward, depending on how you wish to clip it.

You can also move the **slab** of viewing window, keeping the distance between the front and back clipping planes. In order to adjust the area

of the structure where your viewing window is located:

- Click on the **Slab** icon on the toolbar.
- Click and drag the slab frontward or backward, depending on the desired area of the structure you wish to see.

If you have made changes to the clipping planes which you do not wish to keep or you wish to automatically fit your entire structure within the clipping planes:

• Click on the **Unclip** icon on the toolbar. This will automatically set the clipping planes to fit your object.

## 5.13.1 Mesh Clipping

Clipping tools can be used to adjust the frames of the mesh independently of other objects. This is described here.

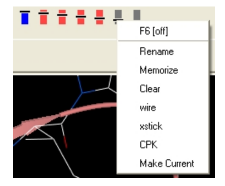
# 5.14 Graphic Layers

Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro

To display and undisplay layers of a structure you can use the buttons shown below. Seven layers can be created and within each layer different structural representations can be displayed.



Right click on one of the layer buttons and a number of options can be chosen as shown below.



To change the display in one of the layers:

- Right click on one of the layer buttons.
- Select a representation wire, xstick or CPK.
- You can do this for each of the seven layer buttons.
- Click on the layer button to display and undisplay. If the layer button is shaded red then the layer is not displayed. If the layer button is shaded light blue then it is displayed. You can switch between layers by clicking on the button or using the. You can use the **memorize** button to store a particular representation and **clear** to remove a memorized representation.

# 5.15 High Quality Publication Images

```
Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro
```

# 5.15.1 High Quality Image

#### To make high quality publication images:

The first step is to improve the quality of the image using the **High Quality Image** and **Antialias** buttons shown below.



high quality image

#### To save and write an image:

• Select **File/Write Image** and the following window will be displayed:

Image File Nam	e icm	Browse
🖲 png 🔘	tif 🔘 rgb 🔘 targa 🔘 eps	

Choose one of the tabs depending on the image type you require:

- Antialiasing allows you to make png, tif, targa, eps or rgb file types.
- High Resolution PNG Allows you to specify a resolution increase and transparent backgroun.
- **High Resolution** Allows you to specify a resolution increase for tif, targa and rgb files.
- Vectorized Postscript This creates a vectorized postscript model of the screen image. Instead of the bitmap snapshot this command generates lines, solid triangles and text strings corresponding to the displayed objects. Since the postscript language is directly interpreted by high-end printers, the printed image may be even higher quality than the displayed image. The final resolution is limited only by the printer since the original image is not pixelized. Warning: there may be inevitable side-effects for some types of solid images at the intersection lines of solid surfaces (i.e. large scale cpk representation, hint: use display skin instead).

\*\*NOTE: With some graphics drivers the high resolution image may be distorted or multiple representations of the image in one frame. Please see the FAQs for a solution to this.

### 5.15.2 Quick Image

A quick image can be saved using this option. The image will be saved as icm1.png in the current directory in which you are working. Each subsequent image produced will be incrementally numbered.

This option is also available via a button as shown below:

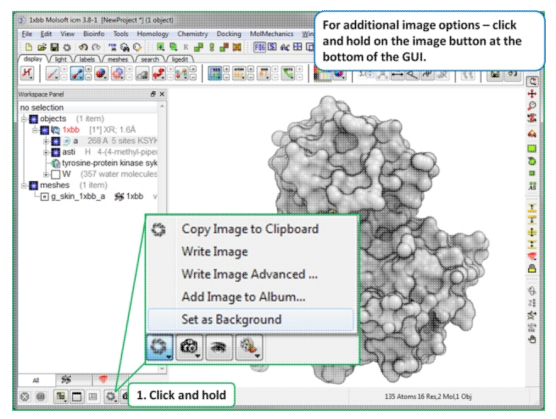


Quick Image Button

## 5.15.3 Image Options

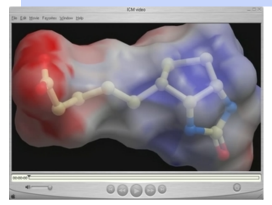
Click and hold on the image button at the bottom of the GUI to:

- Copy image to clipboard (single click on the button also does this).
- Resize image (Write Image Advanced)
- Transparent Background Write Image Advanced)
- Store an image in ICM (Write Image Advanced)
- If you are making an ICM document you may want to store images inside ICM (Add Image to Album).



# 5.16 Movie Making

Available in the following product(s): ICM-Browser-Pro | ICM-Pro



A movie is an excellent way of communicating 3D structural data. The resulting movie can easily be transfered into other applications such as Microsoft Powerpoint.

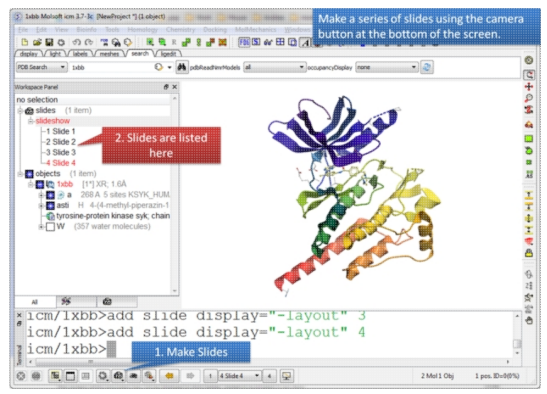
There are two ways to make a movie:

- Make a movie directly from a series of slides.
- Make a movie directly from screenshots.

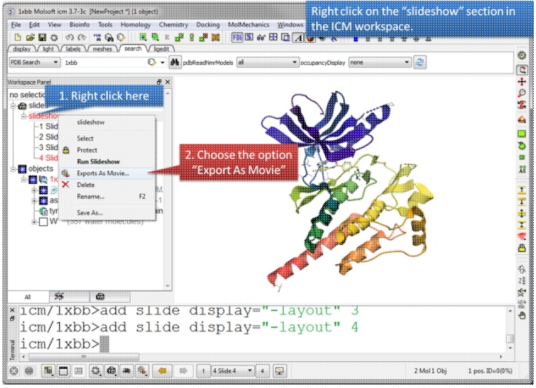
An alternative to a movie is a fully interactive 3D document embedded in Windows PowerPoint or a Web Browser, this can be undertaken using slides and the ActiveICM plugin.

# 5.16.1 Making a Movie from a Set of Slides

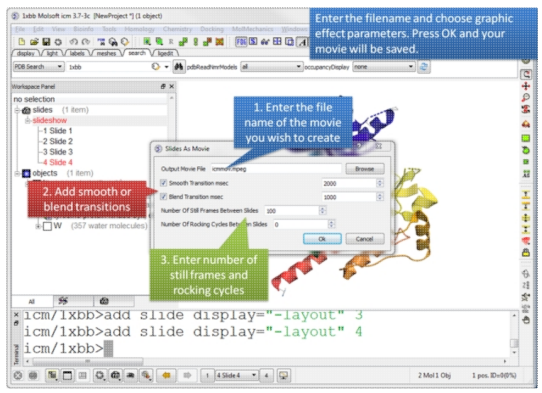
To make a movie from a set of slides:



Make a series of slides using the camera button.

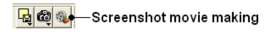


Right click on the slideshow in the ICM Workspace and choose the option "Export as Movie".



Enter the filename for your movie and set the movie effect parameters. Press OK and the movie will be written to a file.

# 5.16.2 Screenshot Movie



To make a Screen-grabbing Movie follow these steps:

- Resize the graphical display to the screen size/resolution you need. You may also want to select the high quality image button and antialiasing to improve the quality of the movie or add visual effects such as shadows.
- To begin making a movie click on the movie making button at the bottom of the graphical user interface (as shown below).
- Enter a file name for your movie and select the movie format (.mov, .avi, mpeg).

	Enter filename		
File name: Save as type: COL A_ NO	icmmov QuickTime files (*.mov *.qt) QuickTime files (*.mov *.qt) Windows video files (*.avi) DMPEG video files (*.mpg *.mpeg)	•	-Choose movie format

#### Screenshot movie making button

**NOTE:** If you want to make a movie to include in a PowerPoint presentation you need to save the movie in AVI format.

• To begin recording the screenshot movie click on the red **Record video** button. Anything displayed in the graphical display will be recorded, for example you can record animations and transitions. Specifying the number of cycles in the animation (rocking, rotation) is an ideal tool for screen-shot movie making. If you have a fast computer you can use **Realtime screen grabbing** which can be selected by clicking and holding the **Record video** button. The real time option can also be set in File/Preferences/Gui menu.



#### Record video button

• The length of the movie in minutes, seconds and milliseconds is displayed in the top right hand corner of the graphical display.



• You can pause the movie and fade out by clicking on the button shown below. The number of frames for the fading out option can be controled using the option in File/Preferences/Gui

#### Pause and fade out



#### Pause recording

• You can record a smooth transition from a previous frame by clicking on the button shown below.

#### Smooth transition from previous frame



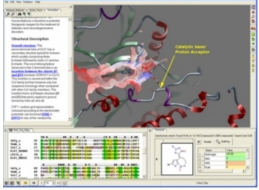
**NOTE:** Anything you do in the graphical display will be recorded in the movie. For example you can change representations, lighting, add new molecules etc. This can be achieved in a more controled manner using the **pause** and **record smooth transition** button.

- Once you have paused the recording the viewpoint and representation of the molecules can be changed and a smooth transition from the previous frame can be generated by selecting the **Record smooth transition from previous frame** button.
- To stop recording a video press the button shown below.

#### Record smooth transition from previous frame



# 6 Slides & ActiveICM



In this chapter you will find a description of the tools available to create files (.icb) containing fully-interactive three-dimensional (3D) molecules and two-dimensional (2D) data. These files can contain multiple interactive views and animations of molecular structures and objects in conjunction with related hyperlinked text, chemical, biological sequence, alignment and data views. The files are small and easily transferable and downloadable. The files can be used

for Molecular Presentation and Documents inside the ICM browser or displayed on the web and in PowerPoint using the ActiveICM plugin. For examples of ICM Molecular Documents please see MolSoft's ActiveICM product page at www.molsoft.com/activeicm.html

#### **Chapter Contents:**

- Making Molecular Slides
- How to View and Navigate Slides
- How to Edit Slides
- How to Add Smooth Blending and Transition Effects Between Slides
- How to Make Molecular Documents Link HTML Text to Slides
- ActiveICM ActiveICM enables you to view and display ICM graphical slides and animations interactively inside Windows Microsoft PowerPoint and web browsers such as Internet Exporter and Mozilla Firefox.
- IcmJs is a JavaScript/HTML5 version of the ActiveICM plugin.

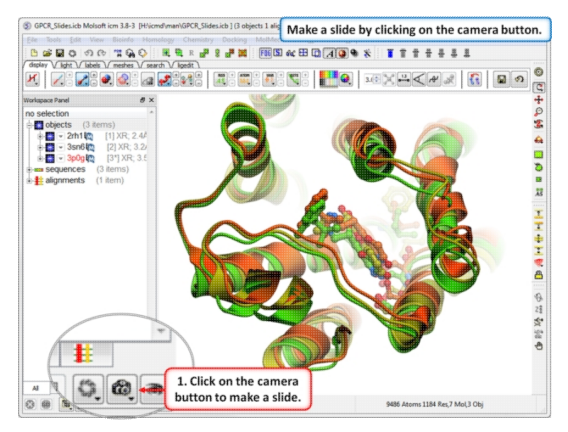
# 6.1 Making Slides

Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro

A slide enables you you to store a large number of different 3D visualization along with text and window layout. The following information can be stored in a slide:

- Viewpoint (e.g. rotation, translation, zoom, lighting and depth effects)
- A set of atom-specific graphical representations such as surfaces, which can be represented as smooth, transparent or wireframe, ball-and-stick models, CPK (space-filling) models.
- A set of atom, residue or distance labels on any of the atomic items
- A set of arbitrary 3D textual annotations assigned to a point in space
- A set of arbitrary 2D annotations assigned to specific 2D coordinates on a screen
- Parameters of the parametric animation
- Window layout for when the slides are viewed in the browser.
- Current table(s)
- Sequences and Alignments
- HTML text with hyperlinks
- 2D images
- For each grob (mesh): representation and colors.

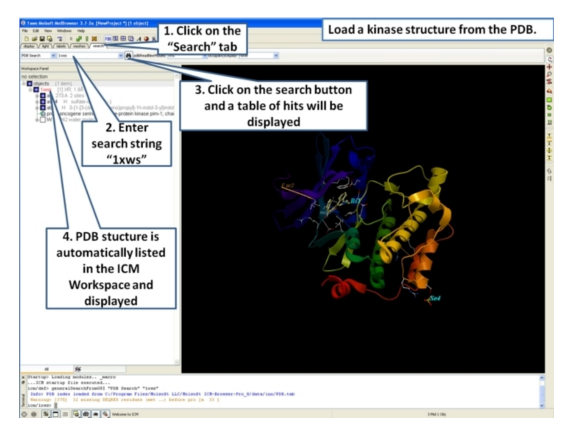
A slide can be made by clicking on the Camera button at the bottom of the GUI.



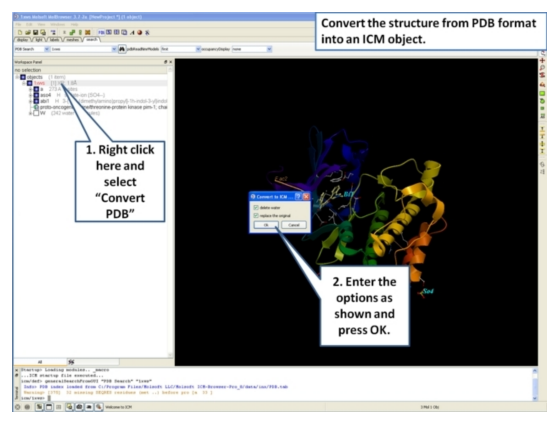
Slides can be viewed in the GUI, can be edited and graphical effects can be added to them. Slides can also be embedded into the web, or PowerPoint using ActiveICM or viewed in ICM-Browser (or ICM-Pro).

This tutorial takes you through the steps to create a series of fully interactive 3D slides. Another tutorial can be found here. To begin making ICM Molecular Slides:

• First load the structure or structures you wish to display in your first slide. Additional structures, labels etc and text can be added at any point during the slide making process. In this example we will load the PDB file 1XWS a PIM1 kinase.

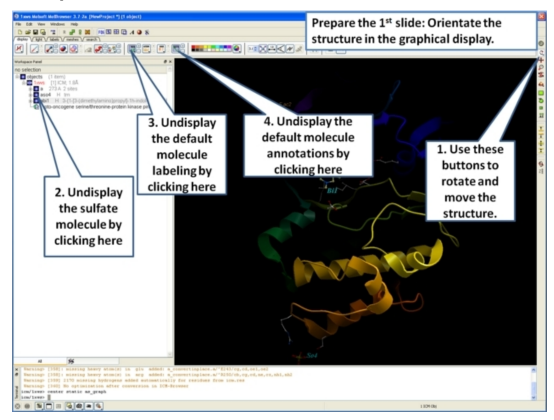


• Next, we will convert the PDB file to an ICM object so we can make slides of the ligand-receptor hydrogen bonds and binding pocket surface.

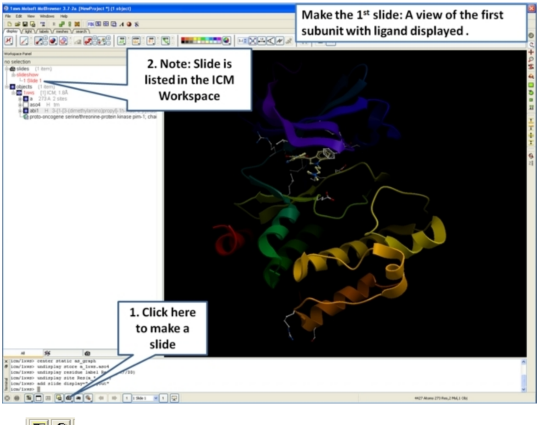


• Now we are going to prepare the first slide by rotating the protein structure to an orientation which allows the viewer to see

the key features of the kinase. For example the bulge in the hinge region (between the N- and C- lobes) which is unique to PIM proteins.



• Next, make the first slide by clicking on the camera button at the bottom of the graphical user interface.





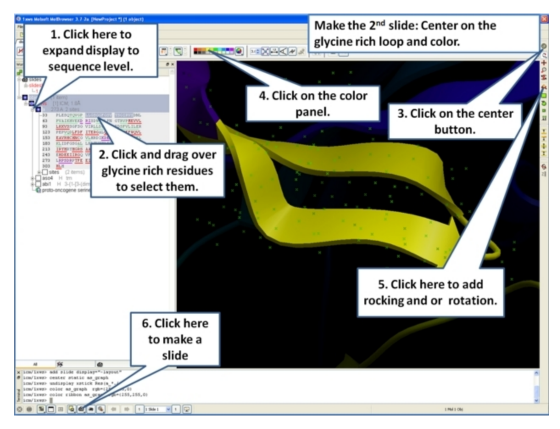
## Click to add slide

• Once you have clicked on the camera button you will see that the first slide has been generated. The first slide is shown in the <u>ICM Workspace window as shown below</u>.

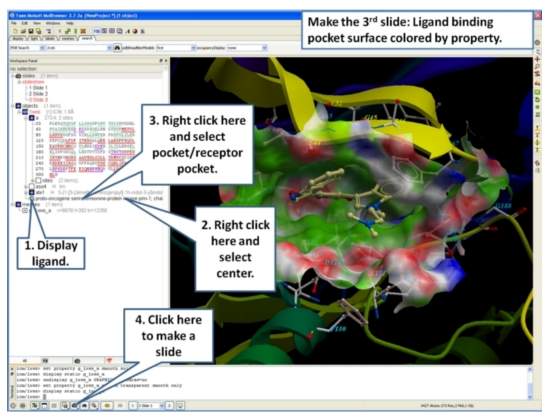


The number and name of the first slide is displayed in the ICM Workspace

• Slides can consist of Static views or Transitions and Animations. Here we will zoom into the flexible glycine rich region of the kinase which lays across the roof of the ATP-binding pocket. Click on the camera button and make the second slide



• Next, we will make a slide of the surface of the ligand binding pocket colored by binding property.



• Now save the document as an icb file. Go to File/Save as...

# 6.2 Make a Movie from a Set of Slides

Available in the following product(s): ICM-Browser-Pro | ICM-Pro

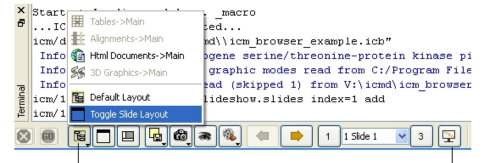
Please see the movie chapter on how to make a movie from a set of slides.

# 6.3 How to View and Navigate Slides

Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro

## 6.3.1 View Slide Show

To view a slide show select the buttons shown below:



#### Click and hold and select Toggle Slide Layout

Run Slideshow

**NOTE:** Slides are associated with the objects currently loaded into ICM. Therefore if you delete an object then the slides will not work. However if you delete an object and then re-read the same object with the same name and structure the slides will be ok.

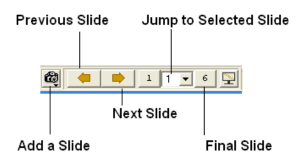
#### To save a slide show

• File/Save Project

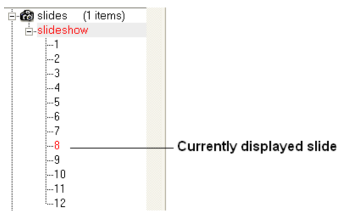
## 6.3.2 Slide Navigation

You can make as many slides as you wish as described in the Making Molecular Slides section.

To navigate through the slides you can use the buttons shown below, the cursor keys for some operations or the right click options in the ICM Workspace.

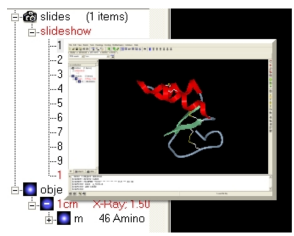


The slide currently displayed is highlighted in red in the ICM Workspace.



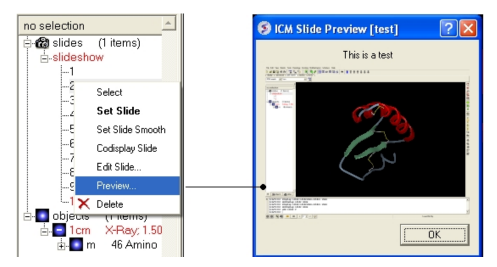
To jump to another slide right click and select "Set Slide".

All slides are displayed in the ICM Workspace. You can hover the mouse over a slide name in the ICM Workspace and a thumbnail sketch of the slide is displayed as shown below. This can be used for slide navigation purposes.



Hover mouse over slide name in the ICM Workspace and a thumbnail sketch of that slide will be displayed.

Or you can right click on the name of the slide in the ICM Workspace and select the option "Preview".



# 6.4 How to Edit Slides

```
Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro
```

You can jump to the slide you wish to edit by following the slide navigation instructions.

# 6.4.1 Edit Slide

**Edit slide contents:** To edit the content of a slide the procedure is to add a new slide and then delete the old one or use the "overwrite current slide" option as shown below:

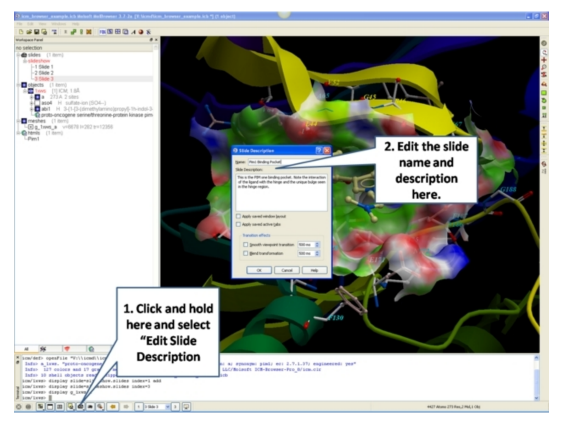
• Click and hold down on the camera button.



### Click and hold

#### To edit a slide description.

• Click and hold down on the camera button and select the option "Edit Slide Description".



- Enter the name of the slide
- Enter a description of the slide.
- If you wish to keep the current window layout or active tabs check the boxes provided

#### To delete a slide:

• Right click on the name of the slide in the ICM Workspace and select Delete.

#### To change the name of a slide

• Right click on the name of the slide in the ICM Workspace and select Edit Slide.

## 6.4.2 Move Slide

To change the slide's position in the slideshow use the Move Current Slide option and select the new position from the list.

- Click and hold on the "make slide button".
- Select Move Current Slide.



### Click and hold

• Select the position in the slide show where you want to move the slide to.



#### Co-display more than one slide

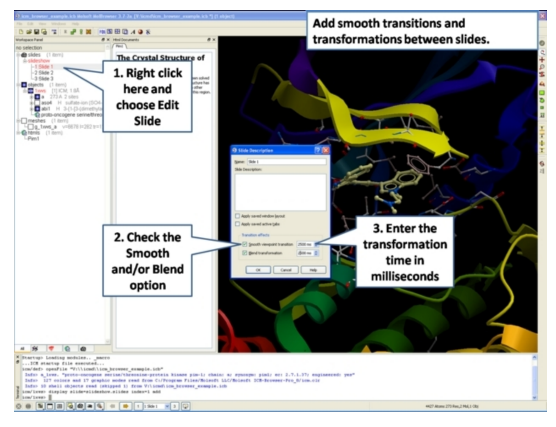
- Right click on the name of the slide in the ICM Workspace you wish to co-display with the curently displayed slide.
- Select the option co-display slide.
- Select the option co-display slide.

# 6.5 How to Add Smooth Blending and Transition Effects Between Slides

Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro

#### How to add smooth and blend transitions to a slide.

- Right click on the name of the slide in the ICM Workspace.
- Select Edit Slide.
- Select the desired transition effect **smooth** or **blend** as shown below.
- Select the length of the transition in milli seconds.



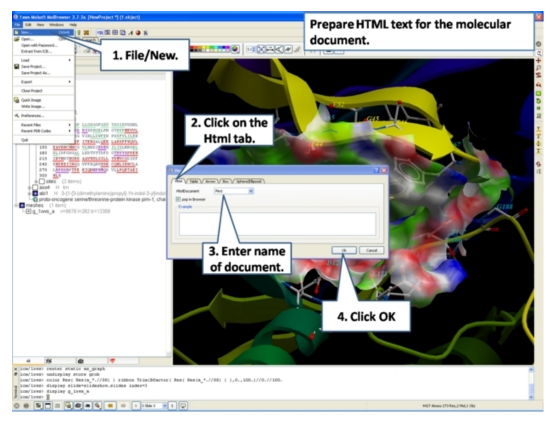
# 6.6 How to Make Molecular Documents -Link HTML Text to Slides

Available in the following product(s): ICM-Browser | ICM-Browser-Pro | ICM-Pro

An ICM Molecular Document contains text and images which can be hyperlinked to the graphical display. Click on the hyperlinked text and then a fully-interactive 3D slide will be displayed. The hyperlinks are usually linked to a set of slides but can also be linked to a series of commands in a script, a web page, a table or alignment. Once a molecular document has been made you can view it in the ICM-Browser (File/Save Project .icb file) or download ActiveICM and view it in a web page or Powerpoint.

#### To begin creating an ICM document

- File/New/ and click on the HTML tab.
- Enter some text. E.g the Name of the HTML document. Formatting can be changed as described in the edit section below.
- Click OK



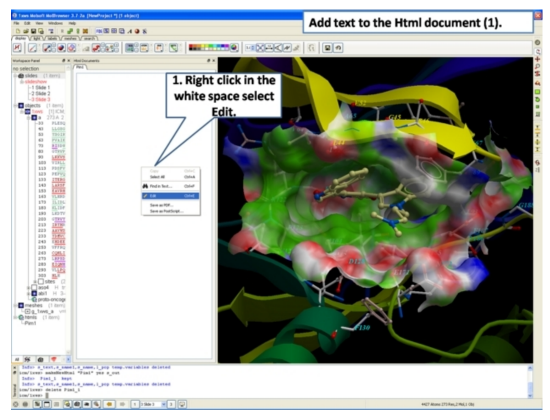
• A HTML text panel will be displayed in the graphical user interface.

**NOTE:** You can add multiple documents into a single file. The documents will be accessible via tabs at the top of the HTML panel.

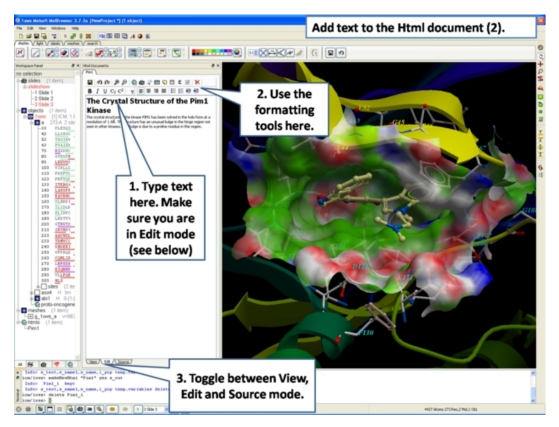
# 6.6.1 How to Add Text or Edit a Molecular Document

#### To edit the HTML text in the graphical display

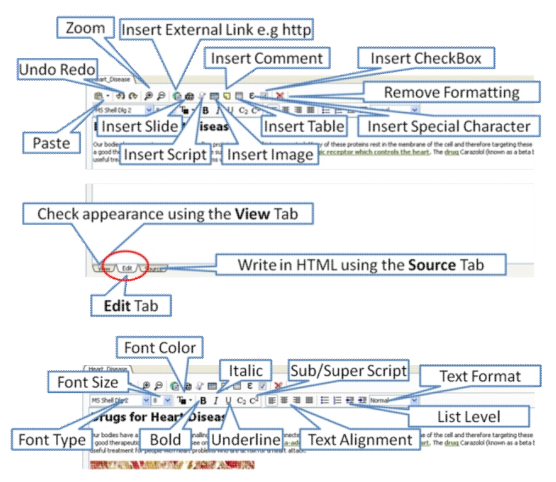
- First create an HTML document and the text panel will be displayed in the graphical user interface.
- Right click in the body of the text display panel and select Edit.



• Enter text and use the formatting tools provided in the panel above the text editor. Make sure you have selected the **Edit** tab in the HTML editor. You can see your page in the **View** tab or write directly in HTML in the **Source** tab.



The key formatting tools in the HTML editor are shown below.

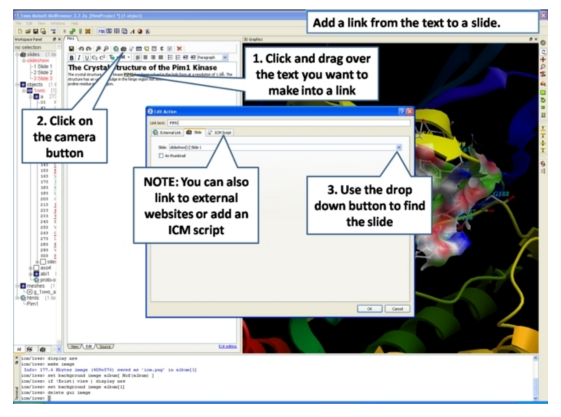


# 6.6.2 How to Make a Hyperlink Between Text and a Slide

#### To make a hyperlink between the text and the graphical display (slide)

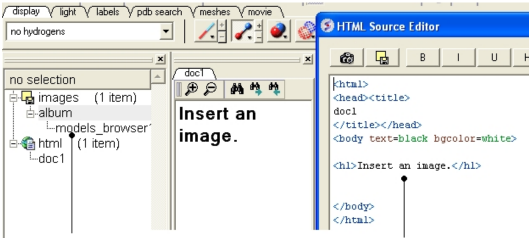
Make a slide or set of slides of the graphical display you wish to link to. See Making Molecular Slides for help on this. Once slides have been created:

- File/New/Html
- Right click in the body of the text display panel.
- Select Edit.
- Highlight the text you wish to link to a graphical display you can do this by left clicking and dragging over the text (selected text will be highlighted in blue).
- Click on the "Camera button" in the HTML editor formatting tool panel.
- Select the Slide tab.
- Select which number slide you wish the text to be linked to from the drop down menu.
- There is an option to display the slide as a thumbnail image in the text document panel. Check if appropriate.



## 6.6.3 Insert Image

**NOTE:** The easiest way to add images (PNG or JPEG) into an ICM Document is to use drag and drop. You can drag and drop the image into the ICM Workspace or go to File/Open. Once the image is in the album in the ICM Workspace you can then drag it from the ICM workspace into the HTML editor.



#### Drag and Drop from the ICM Workspace to the HTML source editor

#### Another way to insert a picture into the HTML text panel

• First read the image into the ICM photo album File/Open OR Drag and Drop from directory into the ICM Workspace.

The image name and preview will then be displayed in the ICM Workspace.



- Create HTML text File/New/HTML. Add text.
- Right click in the HTML window and select 'Edit Source'.
- Right click on the position in the ICM Script Editor where you would like to insert the image.
- Select 'Insert Image'

❤ Edit HTML Image	O X			
Image source: 2 gui2.png				
Image Size				
Width: 160	Reset <u>O</u> riginal			
Height: 120				
✓ Keep original aspect ratio				
OK Cancel				

- Select the image name source.
- Choose the desired Width and Height.

- Click OK.
- Click Save in the ICM Script Editor.

## 6.6.4 Insert Script

#### How to insert a script to the text panel

There are 3 ways to add a script - described in more detail below

1. Drag and drop script from ICM Workspace

2. In the HTML Source Editor - right click and select Insert Slide or Action

3. Create an "inline" script

These methods are described below:

#### **Drag and Drop Method**

- Create a script File/New/Script
- The script will be displayed in the ICM Workspace.
- Right click in the HTML Text Panel (for instructions on how to create this panel see create molecular document) and select edit source and the HTML Source Editor will be displayed.
- Click-Drag and Drop the script into the HTML Source Editor

A line as shown below will be added.

<a href="#icm/script/script1">text placed here will be displayed as a link in the document</a>

#### Another way to add a script to the document is to Insert Action:

- Right click in the body of the text display panel.
- Select Edit Source
- Highlight the text you wish to link to a graphical display you can do this by left clicking and dragging over the text (selected text will be highlighted in blue).
- Right click and select 'Insert Slide or Action' or select the button in the HTML Source Editor and a window as shown below will be displayed.

🦻 Create/Edit a	Slide or a built-in ICM Script	<b>?</b> ×
Internal Link Name	Linking to a script	
Highlighted text	Link text to a script	_
	, Help: start script from #dialog{"name" to generate a dialog}	
C Display Slide:	slideshow[1] 🗹 🔽 As Thumbnail 🔽 Smooth Transition: 2000 🚖	msec
	nd) ICM commands	11000
	script here	_
Add your .		
	OK Cano	el

- Select the option [Arguments and] ICM commands
- Add script in the editor provided
- Select ok

#### **Inline Script**

A script can be added to the HTML text in the following way

- Right click in the body of the text display panel.
- Select Edit Source
- Enter script in the format as shown below.

```
<!--icmscript name="script2"
#dialog{"Test"}
# i_number1 (2)
# i_number2 (3)
print $1 + $2
--><a name="script2" href="#_">script2</a>
```

## 6.6.5 Insert a Dialog Box

Dialog boxes are provided to enable a viewer to interact with a presentation or document file. The dialog box will be a gui data entry box. For an example here is a script to prompt the user of the file to enter a pdb code:

```
#dialog{"Read PDB File"}
# s_pdbcode (1crn)
read pdb $1
ds a_1.
```

💙 Read PDB File			ť	3	ĸ
Please Enter PDB (	Code	1 crn		•	
	<u>c</u>	<u>)</u> k	Cance	91	

The code above can be saved as a script or inside the html text. To do this:

- 1. Right click on the HTML text display and select "Edit Source".
- 2. Highlight the text you wish to link to a dialog box and then select the right click and select 'Insert Slide or Action' or select the button in the HTML Source Editor and a window as shown below will be displayed.

Create/Edit a Slide or	a built-in ICM Script	OX
Internal Link Name	Examplescript2	
Highlighted text	Example Script 1	
Hel	p: start script from #dialog{"name"} to generate a dialog.	
C Display Slide :	slideshow[1] 🔽 🗖 Smooth Transition: 2000 🊔 ms	
• [Arguments and a statements and a statement of the s	d] ICM commands	
#dialog{"Read # s_pdbcode ( read pdb \$1 ds a_1.		
	OK Cance	»

OR.

- 1. Right click on the HTML text display and select "Edit Source".
- 2. Add a link to a script as shown below.

<a href="#icm/script/script1">Example Script 2</a>

### 6.6.6 How to add a check box.

To add a check box:

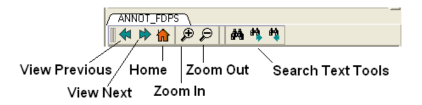
- Click on the check box button in the HTML editor tools panel (see below).
- You can choose the check box to toggle an object or molecule on or off. Or you can be more specific about the toggle actions using the ICM command line. You can choose different actions for On and

Changes	Kout       Arial       Image: Comparison of the compa
7	insert Checkbox ? ×
	Object Type     Objects     Image: a_1xbb.       Image: Customize Toggle Actions       User %1 to refer to the object name above. For example: 'display %1'       OnCheck     OnUncheck
	ds cpk a_lkbb.asti
	Ok Cancel

## 6.6.7 Document Navigation

Off.

The following buttons shown below aid document navigation. Also remember that more than one document can be stored and the header of each document file will be displayed in multiple tabs in the text panel window.

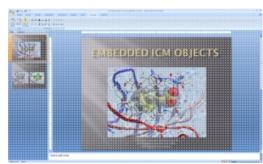


## 6.6.8 Protect Shell Objects From Deletion

When making a molecular document you can protect objects from deletion by the person who reads your document by:

- Right click on the object in the ICM Workspace.
- Select the **Protect** option.

## 6.7 ActiveICM



ActiveICM enables you to view and

display ICM graphical slides and animations interactively inside Windows Microsoft PowerPoint and web browsers such as Internet Exporter and Mozilla Firefox.

#### **Chapter Contents:**

- How to Embed in Microsoft PowerPoint 2003
- How to Embed in Microsoft PowerPoint 2007
- How to Embed in Microsoft PowerPoint 2010
- How to Embed in a Web Browser
- How to Use ActiveICM in PowerPoint
- How to Change ActiveICM Component Properties
- Advanced use of activeICM: Macros to direct visualisation changes
- Background Images

# 6.8 How to Embed in Microsoft PowerPoint 2003

#### Setup

- Download ActiveICM from www.molsoft.com/support
- Save an ICM file (.icb) containing slides. Click here to see how to make slides.

#### Embed icb file

- Open the Insert menu from the top bar of PowerPoint and select **Object**
- This opens up the Object dialogue. Select ActiveIcmCtlClass:

• Create <u>n</u> ew	Object type: ActiveIcmCtl Class Adobe Acrobat 7.0 Document		ОК
Create from <u>fi</u> le	Bitmap Image Calendar Control 11.0 DatePicker Control GroupWise Secure Mime Control GWComposeCtl Class Microsoft Equation 3.0	v	Cancel
	rts a new ActiveIcmCtl Class object into your entation.		

- Click on OK. A file dialogue will then be opened. Open the ICB file you wish to use via this dialogue. IMPORTANT: To avoid later problems, make sure the ICB file is in the same folder as the PowerPoint file.
- A low-resolution snapshot of the first slide in the ICB file will be shown in the activeICM control you created. You can change the shape of the control by dragging the corners of the control with

the mouse, once selected.

Right-click on the activeICM control and select the Properties menu item

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Side 1 of 1		efault Design	English (U.K.)	63	, fe

• Save the PowerPoint presentation

## 6.9 How to Embed in Microsoft PowerPoint 2007

#### Setup

- Download ActiveICM from www.molsoft.com/support ٠
- Save an ICM file (.icb) containing slides. Click here to see how • to make slides.

NOTE: Here are the instructions for ActiveICM in Microsoft Office 2007, for older versions of PowerPoint see here.

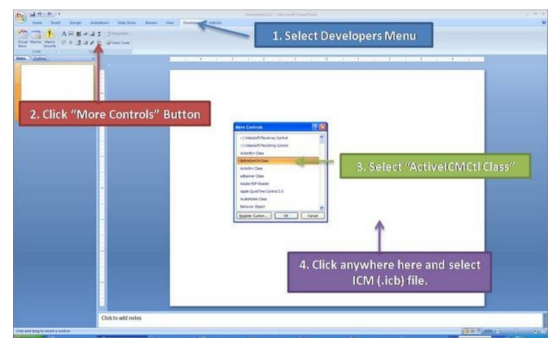
#### **Enable the Developer Menu:**

- Click the Microsoft Office Button (button top left), and then click PowerPoint Options.
- In the **PowerPoint Options** dialog box, click Popular. Under **Top options for working with PowerPoint**, selet the **Show** • Developer tab in the Ribbon check box, and then click OK.

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Tata Lafi - "Office Teamer"	Click to add note	_	 		(Station c)

#### Insert ActiveICM into PowerPoint:

- Select the **Developer** menu.
- Select the More Controls button in the Controls field.
- Select ActiveICMCtl Class from the list of controls and click OK.
- Click the mouse anywhere in the white PowerPoint space and a dialog box will be displayed asking you to select your ICM (.icb) file.
- Click and drag at the corners of the image to resize the normal way you would resize an object in PowerPoint.



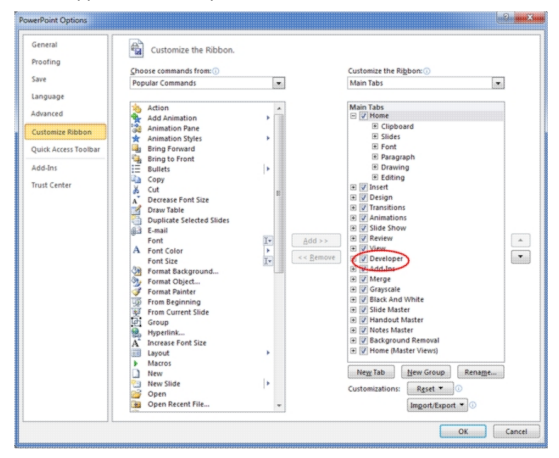
## 6.10 How to Embed in MicroSoft PowerPoint 2010

Setup

- Download ActiveICM from www.molsoft.com/support . Please always check you are using the latest version.
- Save an ICM file (.icb) containing slides. Click here to see how to make slides.

#### **Enable the Developer Menu:**

- Click on the File tab, and then click Options menu.
- Select the **Customize Ribbon** option and then check the **Developer** option in the right hand side panel (see image below).
- Click on the **Developer Menu**.
- **IMPORTANT!** Click OK and then you will see the Developer Menu in the Ribbon at the top of PowerPoint.
- Select the Developer menu.
- Select the More Controls button in the Controls field.
- Select ActiveICMCtl Class from the list of controls and click OK.
- Click the mouse anywhere in the white PowerPoint space and a dialog box will be displayed asking you to select your ICM (.icb) file.
- Click and drag at the corners of the image to resize the normal way you would resize an object in PowerPoint.



## 6.11 Embed in Web Browser

**NOTE:** There is now a plugin-free version using JavaScript/HTML5 - click here for more information.

To embed in a web browser.

 Download ActiveICM from here http://www.molsoft.com/getbrowser.cgi?product=activeicm&act=list (it is free!).

- 2. Create an HTML page in ICM (File/New/Html).
- 3. Add a series of slides.
- 4. File/Export As ActiveICM Html..

**NOTE.** There is an issue with FireFox21 because it disables the ActiveICM plugin. The workaround at the moment is:

- type about:config in the location (address) bar and press the "Enter" key to open the about:config page, just like you open a website by typing the URL in the location bar.
- if you see a warning message then you can confirm that you want to access the about:config page.
- in the Search bar at the top of the about:config page type 'load\_appdir\_plugins'
- double click on the preference to set value to true
- restart FF

## 6.12 How to Use ActiveICM in PowerPoint

#### **\*\*IMPORTANT** There are two ways to open a presentation:

- Double click on the ppt file in windows folder. (in this case PowerPoint will set the current directory to the one which contains the file and there should be no problems with both relative and absolute paths)
- Open ppt through the "File-Open" or recent files. (in this case PowerPoint DOES NOT SET the current directory to the one which contains the file -> relative path might not work and user will be prompted to locate the ICB file unless file is found in absolute location)

#### To view the slides you must be in Slide Show mode

• Press the **F5** button to start the **Slide Show**. In edit mode (i.e. not presentation mode), the control is shown as a static image  $i_{i,i}^{\infty}$  it is not possible to interact with the ICB file. Therefore, to prepare the presentation so that the control shows the correct initial visualisations it is necessary to run the PowerPoint slide(s) in presentation mode

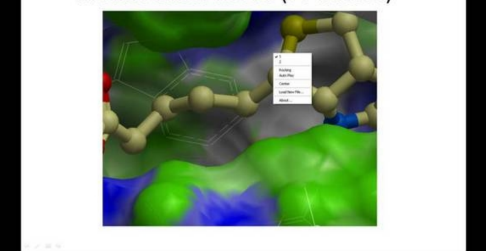
#### **Change Slides**

• Use the left and right cursor keys to change slides.

## A number of other options can be accessed by right clicking on the slide. These options include:

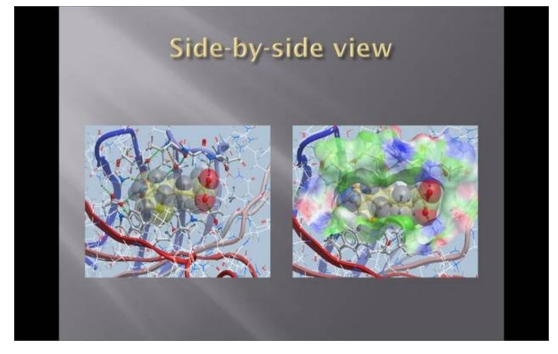
- Select Slide
- Auto Play
- Set on/off rocking
- Center
- Load a new ICM File

## ICM Embedded Into PowerPoint in Slide Show Mode (F5 button)



### You can also add multiple ActiveICM 3D displays in one slide:

• To display multiple ActiveICM 3D displays in one slide just copy the original display or repeat the steps described above. All powerpoint slides should point to the same ICM file (.icb) but they can point to different slides.



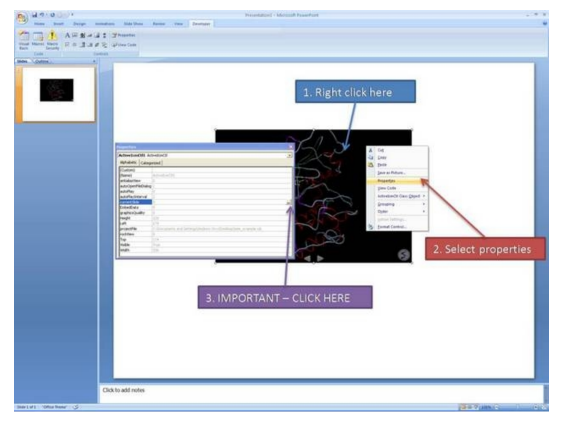
## 6.13 How to Change ActiveICM Component Properties

A number of properties of ActiveICM can be changed once embedded in powerpoint. The options include:

- Select the first slide to be displayed. •
- Set slide auto play. •
- Set auto play of a script. •
- Embed the powerpoint file and the icb file all into one file.

To change these options:

- Right click on your embedded activeICM in Powerpoint.
  Select **Properties** and click on the button shown below.



• A Property Pages window will then be displayed as shown below.

ActiveIcmCtII ActiveIcmCtI Aphabetic Categorized  Aphabetic Categorized  (Custom) (Name) ActiveIcmCtI1  ActiveIcmCtI  ActiveIcmCtI1  ActiveIcmCtI1  ActiveIcmCtI1  ActiveIc	
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6.13 How to Change ActiveICM ComponentProperties

To change the file name of the icb file linked to activeICM: Simply type in the path to the file or use the browse option.

**To change the current ICM slide:** Use the drop down button next to **Current ICM Slide** to select the slide you wish to display first in your presentation.

**To auto play slides:** Check the **Auto Play Slides** box and select the interval between slides option. A range of slides can be played by entering the number of the slides separated by a comma.

To auto play a script: Select whether you want the script to run On Click or On Slide then select the script from the script to play drop down button. You should first save your script in the icb file.

**To embed the icb file in the ppt file** Click the **Embed File into Control** option. **Important** - Please save your PowerPoint file in the t 1997-2003 ppt format not pptx.

# 6.14 Advanced use of activeICM: Macros to direct visualisation changes

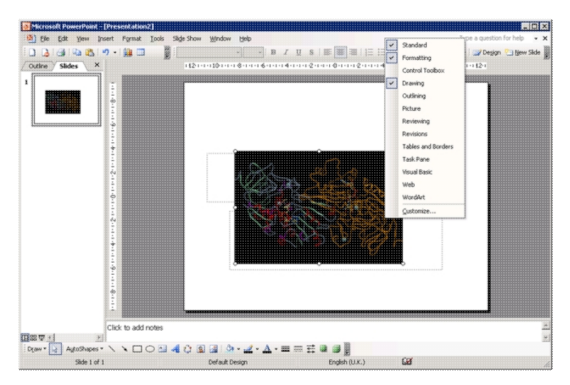
Documentation kindly provided by Dr. Brian Marsden (SGC Oxford http://www.sgc.ox.ac.uk/people/brian/)

It is possible to write simple VisualBasic scripts to avoid having to use the right-click menu approach to changing activeICM control slides within the control itself. This allows one to place buttons outside of the activeICM control, but in the same PowerPoint slide, which controls the control's behaviour. Below are a couple of useful examples of this approach.

#### Creating a button to set the control's active slide:

#### Insert a button Office 2003

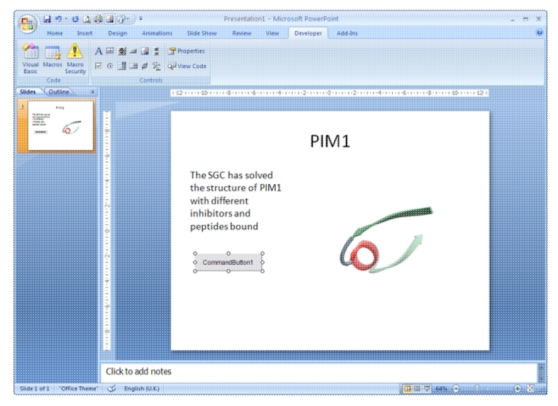
• In edit mode, make sure the control toolbox toolbar is shown by right-clicking the blank area at the top of the top bar and ensuring **Control Toolbox** is ticked.



• Click on an icon in the Control Toolbox which corresponds to the sort of button you wish to use. Then click and drag in the PowerPoint slide to generate the button.

#### Insert a button Office 2007:

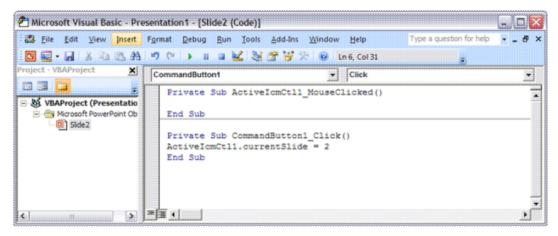
• In edit mode, click on an icon in the Developer menu or ribbon which corresponds to the sort of button you wish to use. Then click and drag in the PowerPoint slide to generate the button.



• Double-click on the new button to open the VisualBasic editor with two empty functions pre-defined. The first one pertains to the control itself and can be ignored in this context \*For the second function (which is for the newly-created button), copy the following into the editor, between the two lines of function code:

ActiveIcmCtll.currentSlide = 2

- This sets the current activeICM control's slide to be number 3 **note** that the value placed in this code needs to be 1 less than the actual slide number (confusing, no?). Obviously, use a value here that makes sense in the context of your ICB file.
- This should leave the editor looking like this:



- Close the Visual Basic editor
- To change the physical properties of the button e.g. text, colour e.t.c.right-click on the button and select the Properties menu option. This opens up a dialogue as below, where many properties of the button can be changed:

ommandButton1 CommandButton							
Alphabetic Categ	gorized						
(Name)	CommandButton 1	1					
Accelerator							
AutoSize	False						
BackColor	8H8000000F&						
BackStyle	1 - fm8ackStyleOpaque						
Caption	CommandButton 1						
Enabled	True						
Font	Arial						
ForeColor	&H80000012&						
Height	39.75						
Left	53.875						
Locked	False						
MouseIcon	(None)						
MousePointer	0 - fmMousePointerDefault						
Picture	(None)						
PicturePosition	7 - fmPicturePositionAboveCenter						
TakeFocusOnClick	True						
Тор	321						
Visible	True	No.					
Width	153	~					

• Using this dialogue, it should be possible to disguise the button to look like normal text (for example) which can be clicked on during the presentation to change the visualisation of the

control, apparently magically. Note that the button will only work in presentation mode.

• IMPORTANT: In Office 2007, remember to save the PowerPoint presentation now as a pptm file that is, a macro-enabled PowerPoint file otherwise the macros will not work next time you load the presentation.

**Other code examples:** Just copy and paste the example of interest inside the function for the button in the Visual Basic editor. Code that enables a button to cycle through the ICB files slides in order (including wrap-around)

```
currentSlide = ActiveIcmCtll.currentSlide
numSlides = ActiveIcmCtll.nofSlides
If currentSlide = numSlides - 1 Then
ActiveIcmCtll.currentSlide = 0
Else
ActiveIcmCtll.currentSlide = currentSlide + 1
End If
```

## 6.14.1 PowerPoint Cache Errors

PowerPoint caches some information about active controls. Sometimes after an ActiveICM upgrade you may get an error when trying to access some property or method: "Wrong number of arguments or invalid property assignment" or something similar.

In this case you need to close PowerPoint and remove all files from the location below:

C:\Documents and Settings\seva\Local Settings\Temp\PPT11.0

## 6.15 Background Images

In ActiveICM version 1.1-5 and higher you can add background images to your icb file and display in PowerPoint and the web using ActiveICM. The documentation on how to insert background images can be found here.

## 6.16 ICM JavaScript (IcmJS)

IcmJS (formerly known as ActiveIcmJS) is a JavaScript/HTML5 version of the ActiveICM plugin. You can read more about IcmJS here.

To make a fully interactive 3D molecular web page from ICM and embed using IcmJS:

- Use ICM vesion 3.8-5 or higher. If you do not have access to this please contact MolSoft to request it.
- Make a set of slides.
- Go to File/Export as ActiveICM Html and then select the IcmJS check box.
- The IcmJS file will then be saved to your computer which you can open in any modern browser e.g. Chrome, FireFox or Safari.

6.16 ICM JavaScript (IcmJS)

## 7 Cheminformatics

#### **Chapter Contents:**

- Read Chemical Format Files
- Save Chemical Structures and Images
- Chemical Spreadsheets (Chemical Tables)
- Sketch Chemicals Molecular Editor
- Chemical Search
- Convert Chemicals to 3D
- ToxScore
- Generate Chemical Fragments
- Find Bioisostere
- MolCart Chemical Cartridge
- Calculate Properties
- Standardize Table
- Annotate Chemical by Substructure
- Align and Color by 2D Scaffold
- Set Formal Charges
- Enumerate Formal Charge States
- Convert
- Build Prediction Model
- Predict using Model
- Generate 3D ConformersGenerate Tautomers
- Generate Stereoisomers
- Calculate Ligand Energetics
- Chemical Clustering
- PCA Analysis
- Visualize Chemical Space
- Compare Two Sets (tables)
- Merge Two Sets (tables)
- Select Duplicates
- Multi Parameter Optimzation
- Combinatorial Chemistry
- SAR Analysis
- Chemical Superposition
- APF Superposition
- APF Tools

## 7.1 Read Chemical Structures

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro |ICM-VLS

#### Video

Chemical structures can be read into ICM from MOL/MOL2, SMILES, and SDF files or you can construct your own structures by drawing them in the ICM Molecular Editor. Chemicals can also be searched and downloaded via the Search Tab using the PDB Chemical Substructure Search, search by PDB Ligand Code, or by searching DrugBank, ChEMBL, SureChEMBL or PubChem.

#### To read a chemical format file:

- Select File/Open.
- Select the chemical structure file you wish to open: MOL, MOL2 or • SDF.
- Once selected the file will be displayed as a chemical table (See ICM molecular tables section).

#### If you know the chemical smiles string for the compound you can build the 2D molecule by:

• Select File/New.

- Click the **Compound** tab at the top of the window.
- Enter a name for the compound.
- Type in the **Smiles String** in the Smiles String data entry box. Remember to delete the previous string.
- Check the boxes Display Molecule Delete Other Objects according to your preference.
- Click the OK button.

#### Smiles can be read from a text file into a chemical table by:

- File/Open and select Files of type: Smiles format
- Then you can convert SMILES to 2D sketch using Chemistry/Convert.

#### Smiles can be read directly into the ICM Molecular Editor:

- Open the ICM Molecular Editor window.
- Select Edit/Add Smiles

#### Multi mol2 3D files can be read by:

• File/Open and select Files of type: mol2 format.

## Chemicals can be read directly from ChEMBL, SureCHEMBL, PubChem, and DrugBank

• Click on the search tab and find the database you wish to search in the drop down dialog box.

### 7.1.1 Extract 2D structure from PDB Chemical

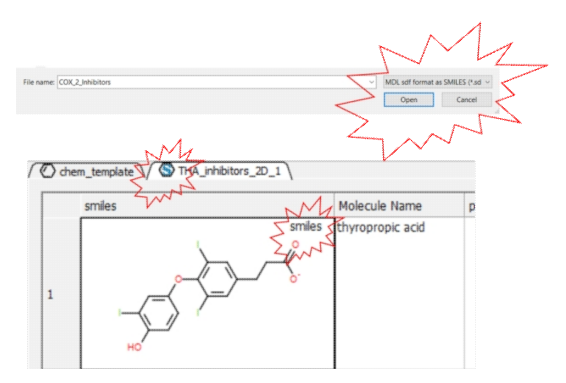
## To extract a 2D chemical representation from a PDB file into a chemical spreadsheet.

- Read the PDB file.
- Right click on the ligand and choose **Extract Ligand**.
- Choose whether you want to extract the 3D coordinates.
- Choose whether you want to append to an existing table (chemical spreadsheet).

### 7.1.2 Read Large SDF Files

The ability to read large SDF files (e.g. >100K chemicals) depends on the memory of your computer. A work around for this is available in ICM.

- File/Open and in the Filter Section choose "MDL sdf format as SMILES"
- The table header will contain an "S" symbol representing that the table contains a SMILES column rather than mol. Each cell will also be labeled "Smiles".



### 7.1.3 Extract 3D coordinates from 3D Chemical.

To extract the 3D coordinates of a chemical into a chemical spreadsheet e.g for 3D QSAR or after chemical superposition.

- Select the 3D chemicals in the ICM workspace. One or more can be selected by clicking holding the CTRL key and clicking. A range of chemicals can be selected by holding the keyboard tab and clicking.
- Right click on any of the selected chemicals in the ICM workspace and choose **Extract Ligand**.
- Select "keep 3D coordinates.
- Append to an existing table if needed.

## 7.2 Save Chemical Structures and Images

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

There are a variety of ways to save chemical structures. Chemicals can be saved in mol, sdf and smiles format from a chemical table (spreadsheet), molecular edior or from the ICM-Workspace. An image of the 2D sketch can be saved as an image from a chemical table.

### 7.2.1 Save Chemical Spreadsheet Workspace

If you wish to save your chemical spreadsheet and plots and setup.

• File/Save As - ICM Binary Format (.icb)

### 7.2.2 Saving Chemical Spreadsheet as SDF

#### To save all the chemicals in a table as an SDF file:

• Right click on the chemical table header.

#### 7.1.2 Read Large SDF Files

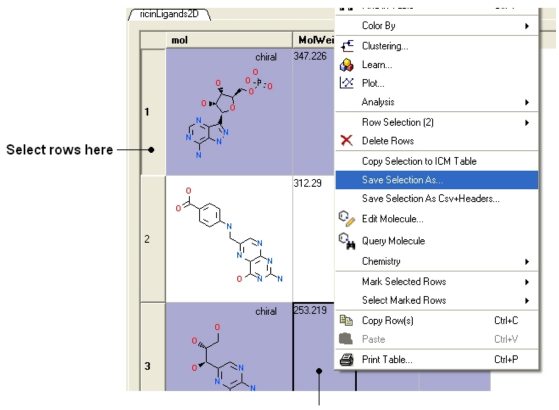
- Select Save as.. •
- Select Save as type: (SD file, Mol file or comma separated value -• csv)

_							
1	ricinLig	jands2D	ricinLigands2[	)			
		mol	Select		loWeight 7.226	MolPSA 157.205	MolLogS -1.44554
			Clone		7.220	197.209	-1.44004
			🗙 Delete				
	1	0	Rename				
		N.	📕 Save				
		"Ý	Save As	_			
		o		31	12.29	114.926	-4.43001
	2		N L				
	2						
			o Ka Ka				

### **Right click here**

#### To save selected chemicals in a table as an SDF file:

- Select the row(s) of the chemicals you wish to save in SDF format. Row selections in tables is described in the Tables chapter.
  Right click on any of the selected rows and select Save Selection
- As....



### **Right click here**

### 7.2.3 Export to Excel

To export to Excel:

• Right click on the table header and select "Export to Excel".

	ang MolMechanics Windows Help R x <sup>×</sup> I x <sup>×</sup> III       roc S ⇔ ** □ A □ O ⊡     I I II
	All Fields T Append
1. Right click	Xdict Xdict
	Export + O
	69 Copy Ctri+C Clone Clone
	Rename
	70 Page Setup  Print Table Ctri+P
	Export To Excel
	Save Save As
	71 H <sub>2</sub> N

## 7.2.4 Saving Chemicals in the Molecular Editor

Chemicals drawn in the ICM Molecular Editor can be saved:

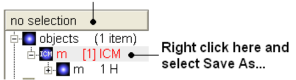
• File/Save or File/Save As... or Edit/Copy as SMILES

### 7.2.5 Saving Chemicals 3D Chemicals

If you have converted a chemical sketch into 3D, the 3D structure will be displayed in the ICM Workspace. To save this structure in mol format:

- Right click on the name of the chemical in the ICM Workspace.
- Select Save As..

#### ICM Workspace

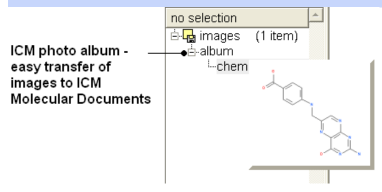


## 7.2.6 Saving Chemical Images

#### To save an image of a chemical sketch:

- The chemical needs to be displayed in a chemical table.
- Right click on the 2D image
- Select Save Molecule As Image

**NOTE** Using the right click options you can also save the image to the **clipboard** or copy the molecule to the ICM Image Album for use in Molecular Documents.



## 7.2.7 IUPAC Name

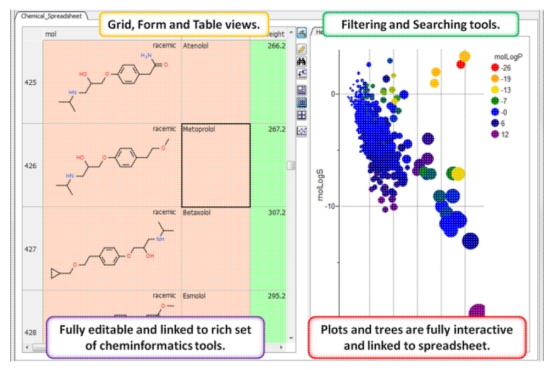
The IUPAC nomenclature for a compound can be generated on the fly for a chemical. You can view the IUPAC name in the Molecule Editor or you can insert a column into a chemical spreadsheet with the IUPAC name.

## 7.3 Working with Chemical Spreadsheets.

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

When an sdf file is read into ICM it is displayed as a chemical spreadsheet. Many of the operations you can perform on chemical spreadsheets (Molecular Tables) are described in the table section of this manual. Some useful chemical-only options are described in this section.

#### An example of an ICM molecular table:



## 7.3.1 Read and Display Chemical Spreadsheet

When an sdf file is read (File/Open SDF) into ICM it is displayed as a chemical spreadsheet.

## 7.3.2 Chemical Spreadsheet Template

#### |Video|

To create a chemical spreadsheet table template:

- Chemistry/Create Empty Chemical Table
- Enter a name for the table.
- Choose whether you wish to add an ICM column and the prefix.
- Enter the number of empty roads you would like to insert.
- Click Next to select property columns.
- You can add plots and histograms to the template.

You can apply the template to new data by:

• Cut and paste or append the new data into the template or double click in a cell and edit the chemical. The chemical properties will be automatically calculated and the plots/histograms populated with data.

You can save a template by:

- Right click on the template listed in the ICM workspace (left hand side).
- Select Save As.. and choose Save as type ICM binary .icb

You can open a template by:

• File/Open file type ICM Binary .icb.

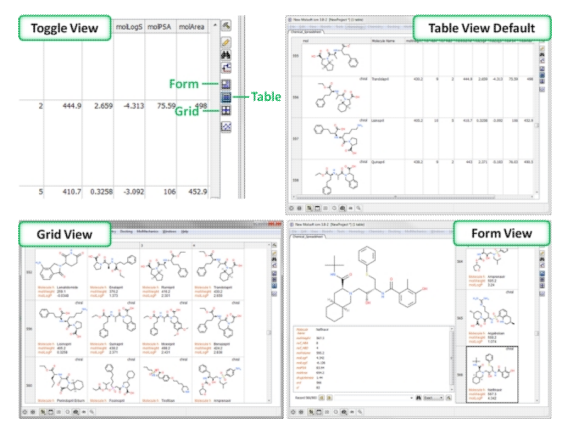
## 7.3.3 Molecular Table Display

|Video|

#### There are three built-in table views.

- Table View (Default)
- Grid View select the columns you wish to display in a grid.
- Form View

You can toggle between each view using the buttons on the right hand side of the table shown below.



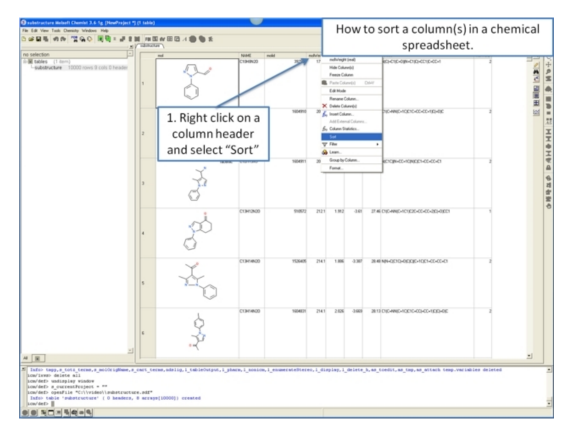
# 7.3.4 How to add Property Columns into a Chemical Spreadsheet.

|Video|

To add chemical properties (e.g. MW, LogP, PSA etc...):

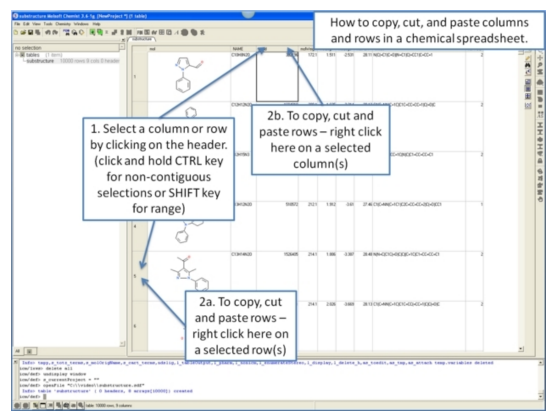
• Chemistry/Calculate Properties

# 7.3.5 How to sort a column(s) in a chemical spreadsheet.



## 7.3.6 How to copy, cut and paste columns and rows in a chemical spreadsheet.

For large data it is recommended to use Tools/Table/Append rows.



7.3.5 How to sort a column(s) in a chemical spread sheet.

## 7.3.7 Add New Data to Chemical Spreadsheet

To add new data to a chemical spreadsheet:

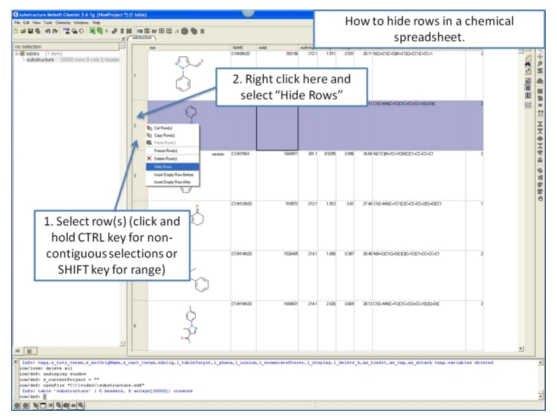
• Use Tools/Table/Append rows.

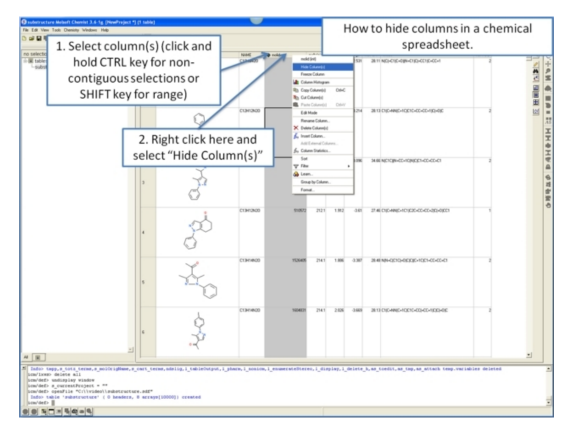
### 7.3.7.1 Highlight New Data

To highlight new data added to chemical spreadsheet.

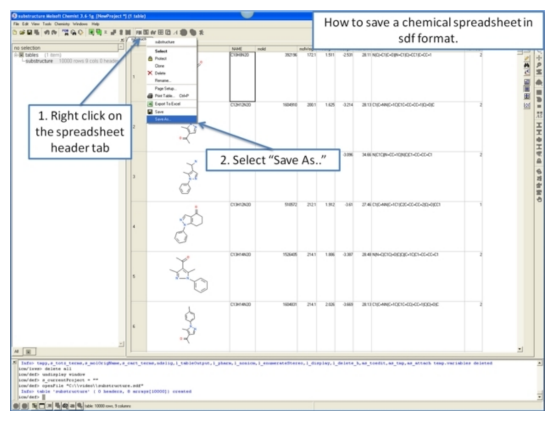
- Use Tools/Table/Append rows.Choose the option to Select New Rows.
- Mark the selected rows. •

### 7.3.8 How to show and hide columns and rows in a chemical spreadsheet.

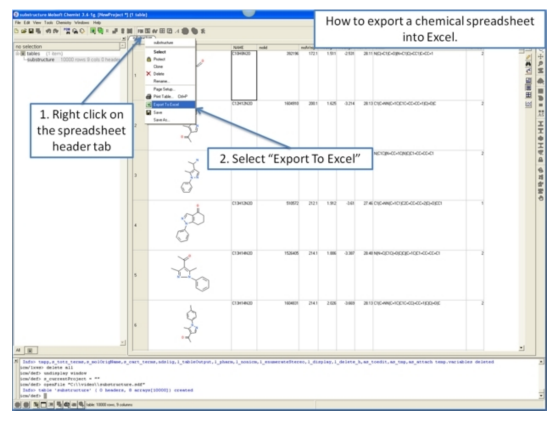




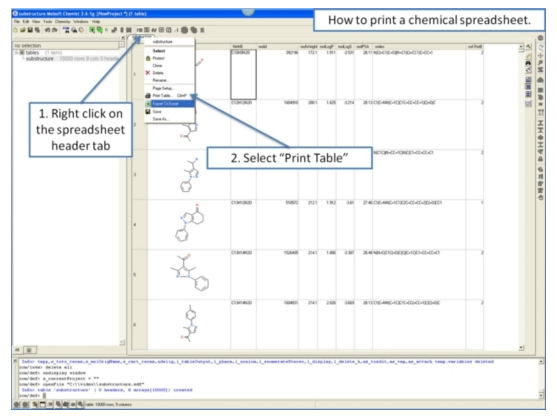
## 7.3.9 How to save a chemical spreadsheet in sdf format.



# 7.3.10 How to export your chemical spreadsheet into Excel.

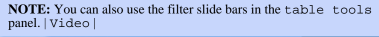


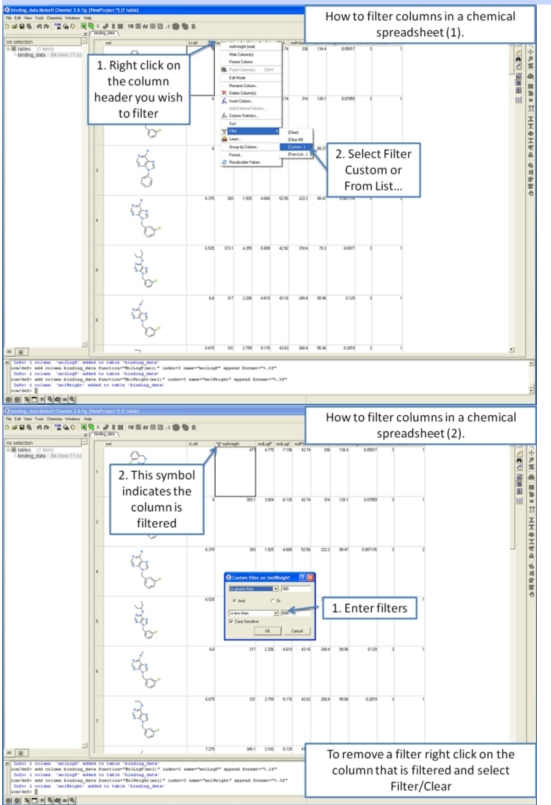
## 7.3.11 How to print a chemical spreadsheet.



7.3.10 How to export your chemical spreadsheet into Excel.

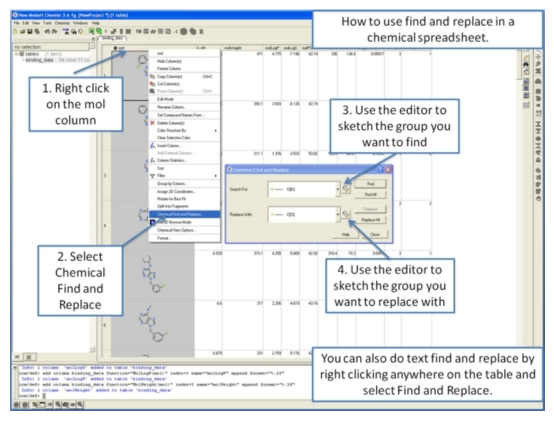
## 7.3.12 How to filter columns in a chemical spreadsheet.



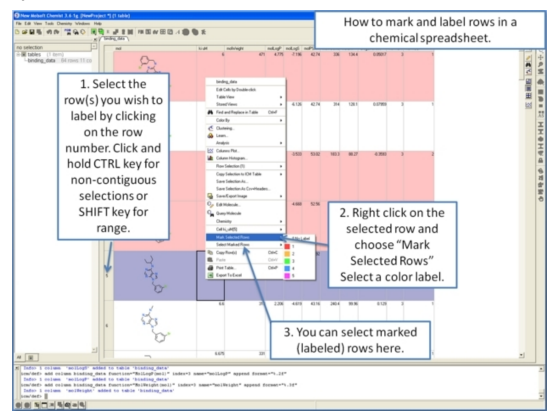


7.3.11 How to print a chemical spreadsheet.

# 7.3.13 How to use find and replace in a chemical spreadsheet.

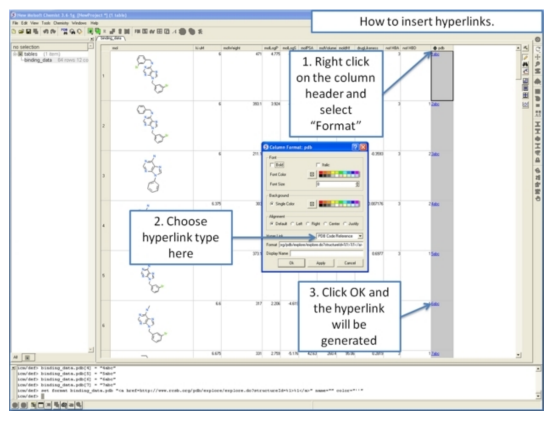


# 7.3.14 How to mark and label rows in a chemical spreadsheet.

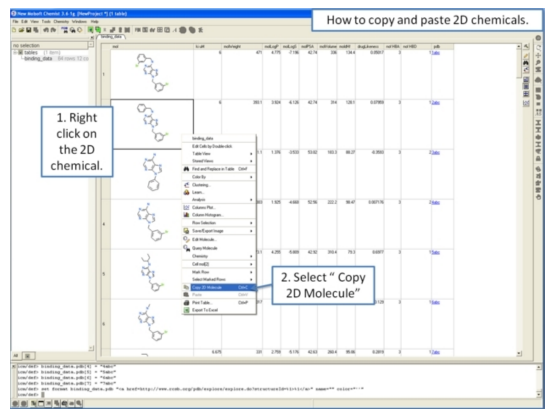


7.3.12 How to filter columns in a chemicalspreadsheet.

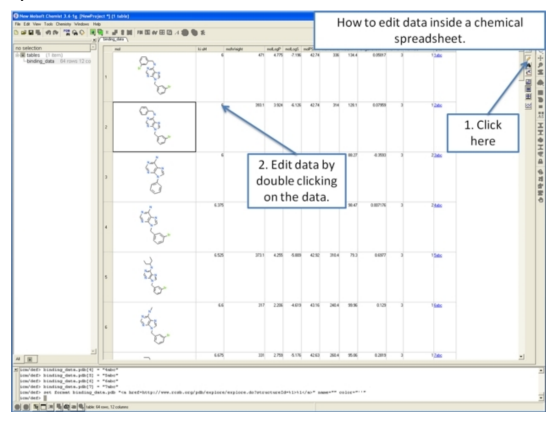
# 7.3.15 How to insert hyperlinks to the PDB, PubMed, and Uniprot.



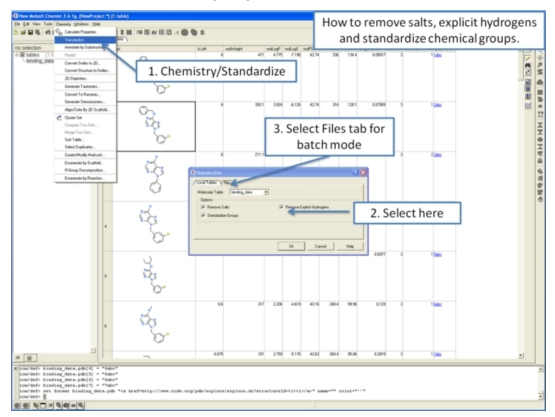
## 7.3.16 How to copy and paste 2D chemicals.



# 7.3.17 How to edit data inside a chemical spreadsheet.

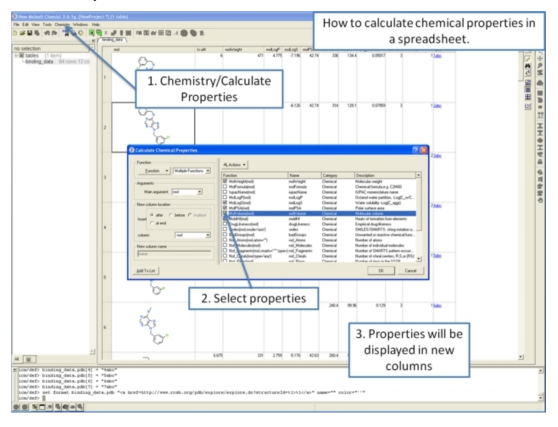


7.3.18 How to remove salts, explicit hydrogens and standardize chemical groups.



7.3.16 How to copy and paste 2D chemicals.

# 7.3.19 How to calculate chemical properties in a chemical spreadsheet.

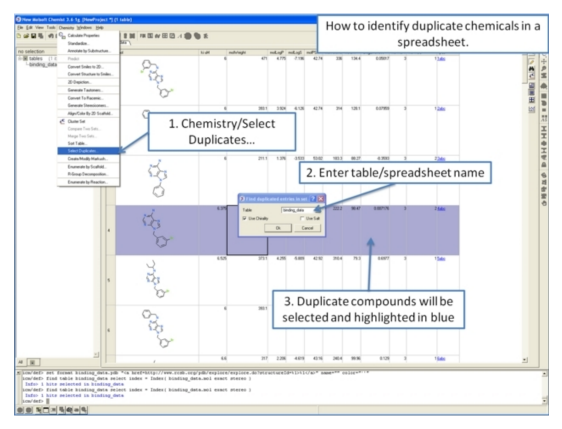


## 7.3.20 InChi

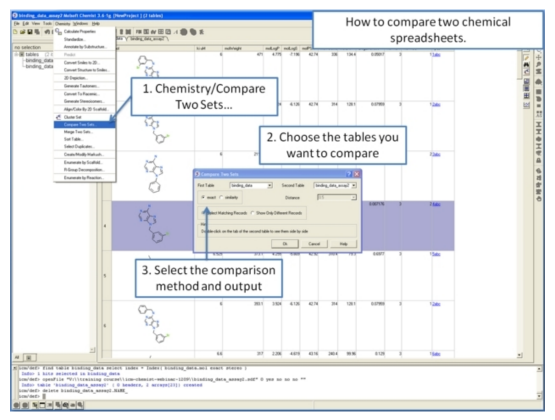
To convert a column containing an InChi key to 2D structure or vice-versa 2D structure to InChi key:

- Right click on a column and select Insert Column
- Choose Function -> Chemical -> Select 2DfromInChi or Inchi

# 7.3.21 How to identify duplicate chemicals in a chemical spreadsheet.



### 7.3.22 How to compare two chemical spreadsheets.



#### How to merge two chemical 285 A spreadsheets. io sele 医 田圃田 凶 rest Structure to S 1. Chemistry/Merge Two et To Ea Sets... 6.176 42.74 1.765.1 Marilla M 211.1 1.32 151 51.02 180.3 68.27 4.258 2 🗙 3. Select the way you want to merge them 2. Choose the tables you want to compare Cancel 6126 42.74 1261 43.1 -1 N N index = Index( binding data.mol find table hinding\_data select exact

## 7.3.23 How to merge two chemical spreadsheets.

## 7.3.24 Display and Convert Molecule

To display and convert a molecule from a molecular table in the 3D graphics display window:

- Select the molecule image or images in the molecular table.
- Right click and select the Chemistry/ Convert to 3D option.

	O N	🕞 🙀 Que	DB Record. <b>ry Molecule</b> mistry		Enumerate Stereoisomers (selected rows)
2		🖹 Cop	y Row(s)	Ctrl+C Ctrl+V Ctrl+P	Enumerate Tautomers (selected rows) Remove Salt (selected rows)
_	0 	209.024	2.38476	keyorganics:10R-0299	Convert to 3D and Optimize Load and Preserve Coordinates Conformation Generator (selected rows) Chemical Template Superposition (selected rows)

## 7.3.25 Copy Molecule

## To copy a molecule to paste into another application or into the ICM Molecular Editor:

- Right click on the molecule and a menu will be displayed.
- Select the option "Copy Molecule"

#### To copy a molecule or image to paste into another row within an ICM table or into the ICM Molecular editor:

- Right click on the molecule and a menu will be displayed.
- •
- Select the option "Copy Molecule" Right click in the cell into which you wish to paste the molecule.
- Select the option "Paste Molecule"

NOTE: To learn how to insert a row read the insert row section.

Molecules drawn in ICM can be cut and pasted into ISIS-Draw and molecules from ISIS-Draw can be cut and pasted into ICM.

To perform thes functions the correct settings need to be turned on in ISIS-Draw so that the compound drawing is saved in the clipboard.

IN ISIS-DRAW - Go to Options/Settings/General/Copy Mol Rxn file to the clipboard.

Compounds drawn in ISIS Draw can be cut and pasted into the ICM Molecular Editor and into ICM tables. Compounds can be copied in ICM by:

Right click on the compound in the chemical table and select Copy Molecule.

## 7.3.26 Edit Molecule

#### To edit a molecule:

- Right click on the molecule and a menu will be displayed. •
- Select the option Edit Molecule and the ICM Molecular Editor will be displayed.
- Edit the molecule.
- Click Exit in the ICM molecular editor.

## 7.3.27 Color Table Column

#### You can color your table based on values within a column by: You can

also color the compound according to specific values see Color Chemical Structure.

- Selecting the column.
- Right click on the column header and a menu will be displayed. •
- Select the option "Color By"

NOTE: You can remove the color from the table by right clicking on the column header selecting Clear Selection Color.

Links to other coloring options:

- Rows can be colored by marking them as described here.
- A 2D chemical sketch can be colored according to specific values
- (e.g. pharmacophore features) see Color Chemical Structure.

## 7.3.28 Chemical View Options

Different chemical view options in the ICM molecular table can be set.

- Right click on the "mol" column header.
- Select Chemical view options... and the following data entry box will be displayed

S Chemical View Options
Hetero-atom hydrogens
Terminal hydrogens
🧮 Atom stereo labels
Explicit hydrogens
T Aromatic rings
🔽 Show 'racemic' flag
🖵 Do not show 3D as 2D
🖵 Unique atom classes
T Atom numbers
🦵 Additional atom labels
Monochrome atom labels
🦵 No text atom labels
Atom labels font
Arial 💌 8 🜩
C Fixed size      Adjustable size
Ok Cancel

Options can be changed by checking the appropriate boxes or by entering the desired font and size.

## 7.3.29 Chemical Table Side-by-Side View

#### Chemical tables can be visually compared by placing them side-by-side. This can be done by:

- Double click on the table header for side-by-side view.Double click on the table header again to remove side-by-side view.

	mol	MolWeight	MolLogP	MolLogS
1		142.118	-0.589686	-1.70292
2	N N N	150.141	-0.070759	-2.11729

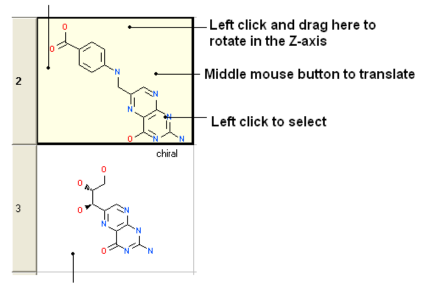
## Double click here for "side-by-side" view

## 7.3.30 Zoom, Translate and Z-rotate a Chemical in a table.

Sometimes you may want to get a better view of a chemical in a an ICM molecular table you can do this inside the chemical table by:

Double clicking on the chemical drawing and the background will ٠ turn yellow.

## Left click and drag here to zoom



Double click for yellow background to enable changes in chemical display

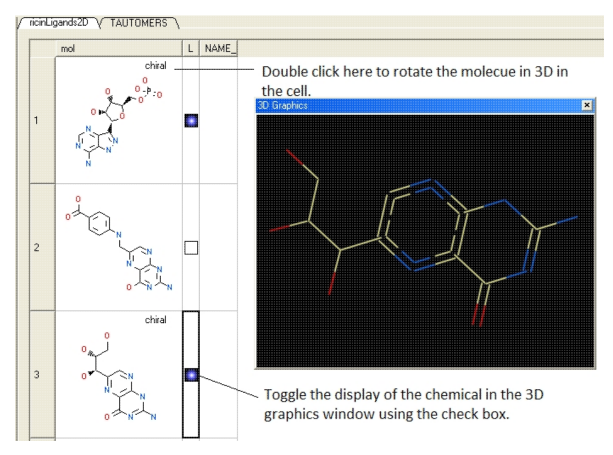
```
NOTE: You can also browse your structures in 3D. See section entitled
"Set Chemical Table 3D Browse Mode".
```

## 7.3.31 Set Chemical Table 3D Browse Mode

#### To browse the chemicals contained within a Chemical Spreadsheet in the graphical display:

- Right click on the "mol" column header.Select the option "Set 3D Browse Mode".

- Use the box in the column "L" to toggle the 3D display on and off.
  Multiple chemicals can be displayed by selecting the rows and then checking the box.



#### To remove 3D browse:

- Right click on the "mol" column header.Select UnSet 3D Browse Mode.

## 7.3.32 Chemical Find and Replace

Chemical Find&Replace tool allows you to find an arbitrary chemical fragment with one or more attachment point(s) and replace it with another fragment with the same number of attachment points.

#### To find a substructure and replace it with something else:

#### Right click here ricinLigands2D mol 1-column selection Hide Copy Column(s) Ctrl+C E) 1 🆄 Cut Column(s) Paste Column(s) Ctrl+V Rename... 🗙 Delete Column(s) Color Atoms by Contribution ۲ 援 Insert Column... 🐔 Column Statistics... 2 Sort Filter ۲ Assign 2D Coordinates... Rotate for Best Fit Clear selection Find and Replace. 3 Chemical View Options...

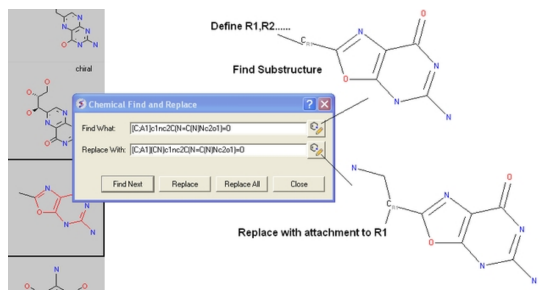
- Select the column in which the molecular structures are displayed. The column is usually called "mol".
- Right click on the "mol" column header and select Find and Replace. A data entry box as shown below will be displayed.

Chemical Find and Replace							
Find What: <empty>       Replace With:     <empty></empty></empty>							
Find Nex	t Replace	Replace All	Close				

- Click on the Molecular Editor button at the end of the Find What: data entry box.
- The ICM Molecular Editor will be displayed. Draw the substructure you wish to search for and replaced.
- Draw the pattern and mark attachment points with R1,R2.... R-groups can be added by right clicking at the attachment point and selecting the R-group from the drop down options.
- Close the ICM Molecular Editor and the string will be displayed.
- Repeat with the "Replace With:" data entry box. Make sure the same number of R1,R2... labels are drawn.
- Click the Find Next button and then Replace or Replace All. When a substructure to replace is identified it will be colored red.

**NOTE:** There are a number of keyboard shortcuts which can be used to draw chemicals. Also please note that an aromatic bond in the source molecule will not match a double bond in the replacement pattern.

Here is an example:

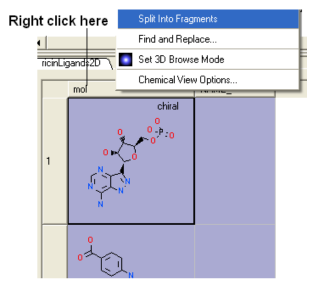


## 7.3.33 Split Chemical(s) into Fragments

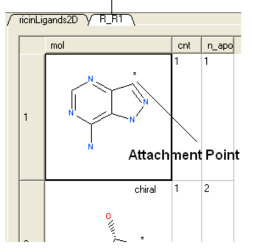
Chemicals displayed in an ICM Molecular Table can be split into fragments. This is useful for generating a series of R-groups to be added to a scaffold (See section describing reactions.

To generate fragments:

- Select the column or row(s) you wish to generate the fragment from.
- Right click on the "mol" column header and select "Split Into Fragments".
- A new table of chemical fragments will be displayed. Each fragment is assigned an attachment point which is flagged with an asterisk (\*).



## New table is generated containing fragments



## 7.3.34 Rotate Chemical for Best-Fit

To improve the display of a chemical within an ICM molecular table you can choose an option called "Rotate for best fit".

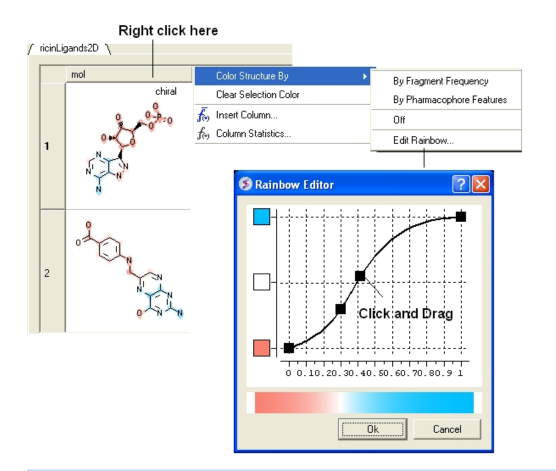
This option can be found by:

- Right click on the "mol" column header.Select "Rotate for Best Fit"

## 7.3.35 Color Chemical Structure

To color the structure of a chemical in an ICM molecular table by fragment of pharmacophore frequency:

- Right click on the "mol" column header.
- Select Color Structure By
- Select "By Fragment Frequence" or "By Pharmacophore Features"



**NOTE:** The coloring can be controled using the "Edit Rainbow" option and the coloring can be removed using the "Off" option

## 7.4 Molecular Editor

```
Available in the following product(s): \texttt{ICM-Chemist} \mid \texttt{ICM-Chemist-Pro} \mid \texttt{ICM-VLS}
```

#### |Video|

Draw new compounds by using the ICM Molecular Editor.

#### The molecular editor can be activated by:

• Clicking on the Open ICM Molecular Editor button shown below.

	00 7 9	0	🖳 R	83	8 1	2 <sup>11</sup> 111
B Search ▼	text or smiles		0	- 44		All Field

## OR

• Select Chemistry/Molecular Editor and the editor as shown below will be displayed.

## 7.4.1 Drawing a New Chemical Structure

To draw a new chemical structure the ICM Molecular Editor should be loaded.

To do this:

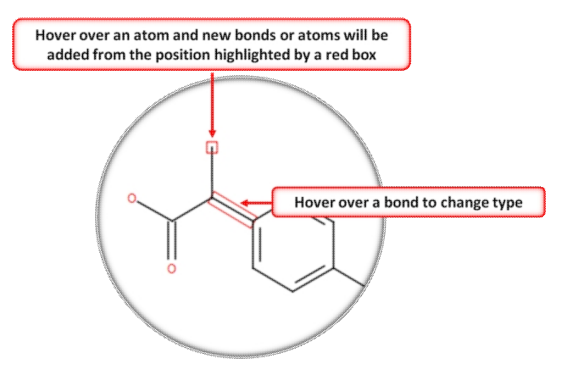
• Select Chemistry/Molecular Editor

Now you can start drawing your structure.

• First select an atom, bond or ring and then click in the white Molecular Editor Workspace to start skethching.

Ge ICM Molecule Editor [new file "] Eile Edit Yoew Jernplates Help		Chemical Sketchin	g
	K 💽 🖬 🗞		
Bond Types Atom Types Click anywhere in the canvas to start sketching	Name Name	Volue	

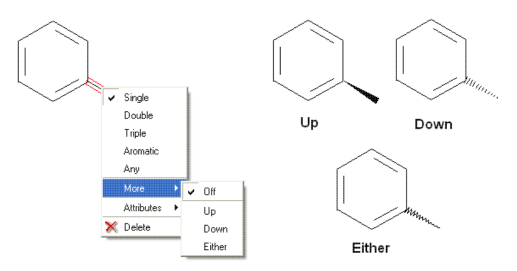
- You can extend your structure by selecting another atom, bond or ring.
- Select where on the structure you would like to add the new group by hovering the mouse over the desired position. The position you will add to will be highlighted in a red box.



## 7.4.2 Stereo Bond

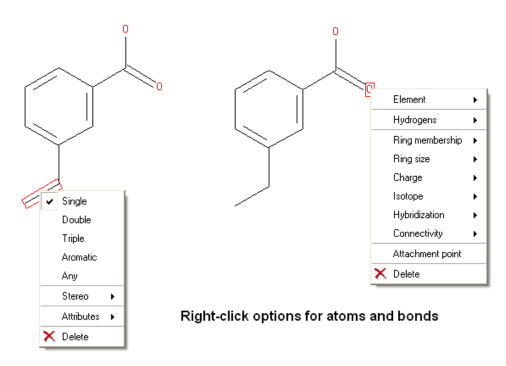
## To set the stereo bond type:

- Right click on the bond and a menu as shown below will be displayed.
- Select which bond direction you desire from up, down or either.



## 7.4.3 Right Click Options

There are a number of different options available when you **right click** on a **bond** or **atom**. These can be used for editing bond and atom types and there are additional options for Chemical Searching.



## 7.4.4 Chemical Dictionary

There is an inbuilt chemical dictionary which enables you to import a 2D sketch quickly. To use it:

- Click on the chemical dictionary button.Start typing the name of the chemical.Select the chemical from the drop down list.

ICM Molecule Editor (new file )		Chemical Dictionary
Eile £dit Yiew Jernplates Help □ @ ↓ 5 % ■ ↓ 5 % ■ ↓ 5 %	16 16 RIS A 9	C <sub>24</sub>
2. Type and select chemical name	a salicylid	0
	Salicylic add	1. Click here for dictionary
1	Salicylic acid beta-D-glucos	
000000	2-HydroxybenzaidehydeSalic	ylic
	2-Hydroxybenzyl alcoholsalic	ylic.
с	3-Hydroxybenzoic acid Salicy	k
N	4-Hydroxybenzoic acid Salicy	k
0	3-Aminosalicylic acid Salicylic.	. 69
F	Ethyl salcylate Salicylic	25.4
Р	Isobutyl salicylate Salicylic	
s	5-Sulfosalicylic acid Salicylic	ð/-
CI	Amyl salicylate Salicylic	
Br	Benzyl salicylate Salicylic	т
I	o-Tolyl salicylate Salicylic	
B	1-Salicylate glucuronide Salicy	dic
	2-Hydroxy-6-(8-tridecenyl)benz	roic acidsalicylic
14	1	
		and the second se

## 7.4.5 Dictionary of Chemical Groups

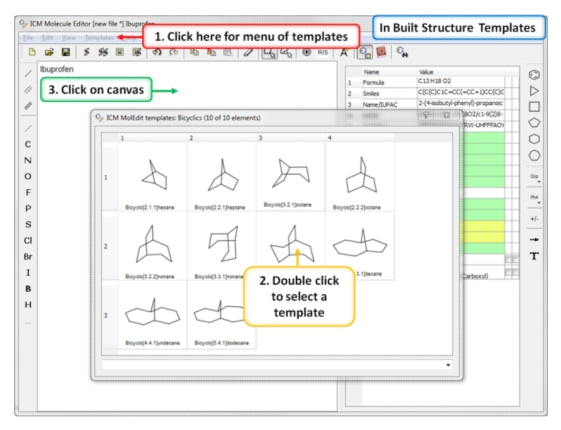
The ICM Chemical Editor has a dictionary of chemical groups. These groups are stored in a table stored in CHEM\_GROUPS.csv file. This file can be manually edited to remove or add new definitions. The group can be selected using the Grp button of the Editor.

M Molecule I	iditor [new file *] Salicylic	: acid				Chemical Group	s
9 K	M MolEdit templates: Di	ctionary (983 of 983 ele	ments)			8 23	
2 3	1 Ninic Dode Dimethyl suffacide	2 Selenium Sulfde	3 Ethanol H HO Hydroxyveas Formepizole	4 N S Cyssamine Cys	S O' Calolum Acetate As = 0 As = 0 Arsenic triacide O Foscamet	ick and hold	
4	Cycloserine	N Methimapole	N Betazole	Histamine	Mechiorethamine		

## 7.4.6 Template Menu

A set of chemical templates are available for:

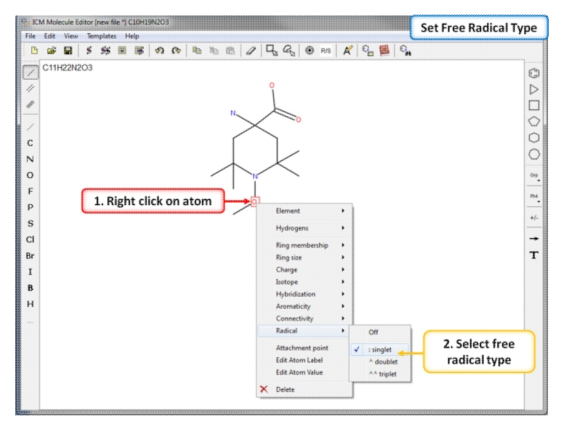
- amino acids
- bases
- bicyclics
- rings



## 7.4.7 Set Free Radical Type

To set a free radical type (singlet, doublet or triplet) to an atom:

- Sketch the ligand in the Molecular Editor.
- Right click on the atom and select the free radical type.
- Convert to 3D or add to a chemical table if needed for docking.



# 7.4.8 Adding and rotating a fragment in molecular editor by clicking-holding-and-dragging.

The Chemical Editor allows one to add bonds (click the bond button), or chemical groups (the Grp button), Often one needs to re-orient the added bond or fragment. In this case press on the atom to which you are going to append the bond or the fragment and **hold-and-drag** the mouse it until you see added fragment rotating around the attachment point. While you keep the mouse button pressed you can **rotate** by dragging in the preferred direction.

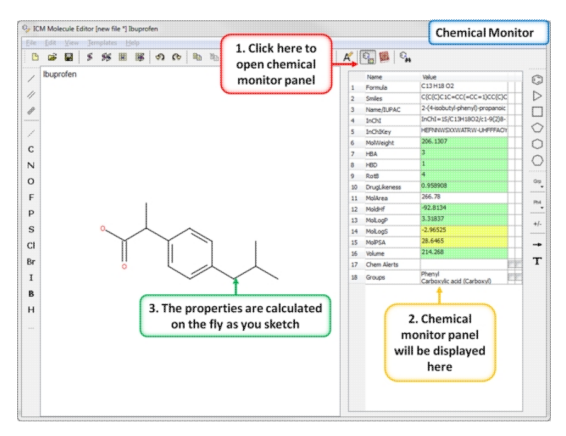
If you add templates (the Templates menu, or **paste** fragments, the mechanism is somewhat different because the group is added in its default orientation. In this case you can press **Ctrl** to rotate the fragment to to be able to attach the fragment in the desired orientation.

## 7.4.9 Property Monitor

When drawing a compound in ICM you can monitor important ADME-Tox and drug-likeness properties.

## To view the monitor

• ICM Molecular Editor/View/Chemical Monitor or click on the Chemical Monitor button and a window as shown below will be displayed.



The following properties are monitored.

- Chemical Formula
- SMILES string
- Name/IUPAC Name
- InChI
- InChI Key
- Molecular Weight (MolWeight)
- Number of Hydrogen Bond Acceptors (HBA)
- Number of Hydrogen Bond Donators (HBD)
- Number of Rotatable Bonds (RotB)
- DrugLikeness value Druglikeness is not based on other properties, this is separate model built using Bayesian Classifier (See: http://www.molsoft.com/mpropdesc.html). Normaly Bayesian classifier returns either 1 ot -1 (positive ot negative) therefore positive values means drug-like, negative - not.
- Molecular Area (MolArea)
- Prediction model build for 'delta Hf in gas' property using public NIST database (description can be found here: http://webbook.nist.gov). A low dHf value means that the compound is more 'stable'.
- LogP (MolLogP)
- LogS (MolLogS)
- Polar Surface Area (PSA)
- Volume
- Chem Alert will flag any reactive group, PAINS or Bad ADME-TOX groups.
- Groups option highlights chemical groups in the sketch.

The rows in the monitor window are colored from green (good) to red (poor). You can right click on the table to copy a value or add your own prediction model.

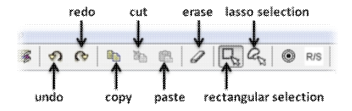
## 7.4.10 Editing structure using keyboard

You can select a fragment or simply position your mouse cursor over a bond or atom and use the following keystrokes for editing:

- Changing atom properties
- d set/unset heavy atom connectivity as drawn to avoid additional branches in chemical searches. Toggle.
- r set/unset the "in-ring" property of *atoms*. Toggle.
  y set/unset *atom* hybridization type (*sp1*,sp2,sp3). Toggle.
- A set *atomic* property to be **aliphatic**
- a set *atomic* property to be **aromatic**
- \* (asterisk) set *atomic* property to be **any atom**
- C Carbon
- H Hydrogen
- N Nitrogen
- 0 Oxygen
- F Fluorine
- P Phosphorus
- S Sulfur
- I Iodine
- **Changing chemical bonds** •
- 1,- (dash) single bond. •
- 2,= (equal) double bond. •
- 3,# (hash) triple bond. •
- 4,: (column) aromatic bond.
- 0,~ (tilde) any bond •
- Changing R-Groups
- Press 1 to mark an atom as R1
- Press Ctrl-1 to preserve its type and create [C;R1]
- Press Ctrl-0 to remove the mark.

## 7.4.11 Chemical Editing Options

Tools are provided to help editing your chemical sketch. These include buttons for selection, erasing, copy and paste.



## 7.4.12 Paste SMILES or InChi String

#### To paste a SMILES string into the Molecular Editor:

- Open the Molecular Editor.
- In the Molecular Editor window select Edit/Add Smiles/InChi. •
- Click on the canvas.

## 7.4.13 Save Chemical in Mol/RXN Format

#### To save the structure as a MOL or SDF file on your machine or server:

- Select File/Save As... in the Molecular Editor window
- Enter a filename and select where you wish to save the file.
- Save as type mol or rxn

## 7.4.14 Save to Chemical Spreadsheet

#### To save the chemical to a chemical spreadsheet:

• Click on one of the buttons shown below. You can append to a new table or an existing one.

## append molecule to a chemical spreadsheet



append molecule to a chemical spreadsheet and convert to 3D

## 7.4.15 Save as SMILES or InChi

#### To save the chemical in SMILES format

• Edit/Copy as Smiles

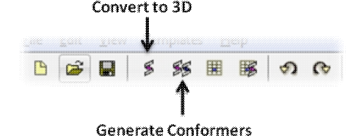
#### To save the chemical in InChi format

- Open the Chemical Monitor.
- Right click on the InChi row and choose copy.

## 7.4.16 Convert Chemical to 3D or Generate Conformers

#### To convert the chemical to 3D or to generate conformers:

- Click on the buttons shown below to convert to 3D or generate conformers in the Merck Molecular Force Field (MMFF)
- The 3D chemical will be listed in the ICM workspace. If conformers are generated a table showing the conformers will be created.



## 7.4.17 Isis Draw Copy and Paste

Molecules drawn in ICM can be cut and pasted into ISIS-Draw and vice-versa molecules from ISIS-Draw can be cut and pasted into ICM.

To perform these functions the correct settings need to be turned on in ISIS-Draw so that the compound drawing is saved in the clipboard.

 ${\bf IN}\ {\bf ISIS}\ {\bf DRAW}$  - Go to Options/Settings/General/Copy Mol Rxn file to the clipboard.

Compounds drawn in ISIS Draw can be cut and pasted into the ICM Molecular Editor and into ICM tables. Compounds can be copied in ICM by:

• Right click on the compound in the chemical table and select Copy Molecule.

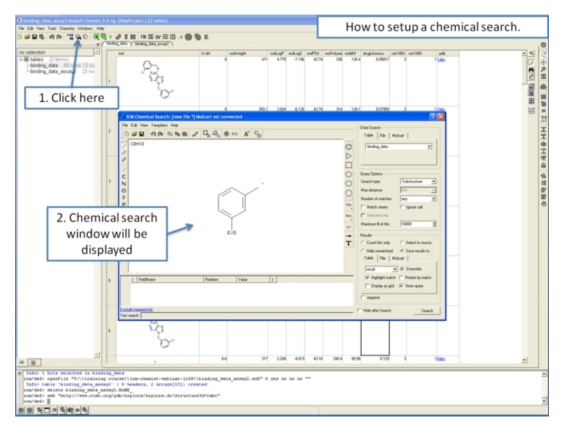
## 7.5 Chemical Search

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

Chemical similarity searching can be used to screen a database of compounds for structural similarity to a query chemical structure. The chemical similarity search window is shown below. To access this window use the Tools/Chemical Search menu or select the button shown below.

## Chemical search





## 7.5.1 Query Setup

To set up a query first you must have either drawn or loaded a chemical structure into ICM. Instructions for this are described in the Reading Chemical Structures and Molecular Editor sections of this manual. If for example the query molecule you want to search is in a chemical spreadsheet you can right click on chemical in the spreadsheet and

#### select Query Molecule.

At this point your query can be modified as described in the Molecular Editor sections of this manual. **However**, there are a number of ways to specifically modify your query and filter your search. The way to accomplish this is to right click on a bond or atom and a menu as displayed below will be displayed. The menus differ depending on whether you right click on a bond or atom.

If you wish to specify a filter at an atom.

• Right click on the atom and the menu shown below will be displayed.

🐾 ICM Chemical Search: [new file *] Molo	cart not connected	? 🛛
File Edit View Templates Help		Data Source
<u>]</u> ≧ ⊒	- C C R/S 🕺 🖸	Table File Molcart
C11H14		
1		
c		Query Options
N		Search type Substructure
0		Query Options Search type Max distance 0.4
F		Number of matches any
P	Right click on atom	☐ Match stereo ☐ Ignore salt
5		Selected only
	Element	Maximum # of hits 10000 🚖
Br	Hydrogens +	
B		Results
H	Ring membership ► Off Ring size ► A - Der	Count hits only C Select in source
	Charge	
Cap	RU	able File Molcart
Pit	V H1	result 🔽 🔽 Overwrite
	Aromaticity R3	✓ Highlight match
+/-	Connectivity	Display as grid 🔽 Store query
-	Attachment point	
*	× Delete	Append
( FieldName	Relation Value	) Hide after Search Search
•		•
Exclude fragment list		
Text search		

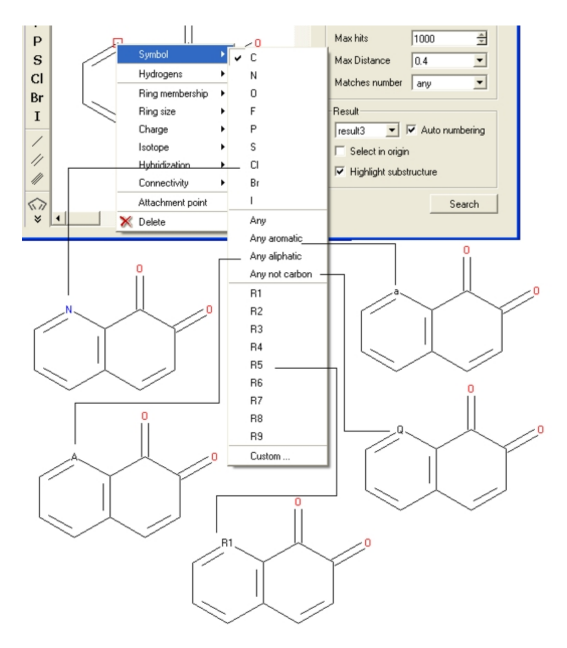
If you wish to specify a filter at a bond.

• Right click on the bond and the menu shown below will be displayed.

🐃 ICM Chemical Search: [new file *] Molcart not connected	2 🛛
File Edit View Templates Help	Data Source
[     FieldName     Relation     Value       Image: Constraint of the second	I Hide after Search Search

## To specify a particular atom type, aromatic, aliphatic or R-group at a particular atom site.

- Right click on the atom and select the "symbol" option as shown below.
- Select the desired atom type, aromatic, aliphatic or R-group and a symbol will be displayed as shown below.



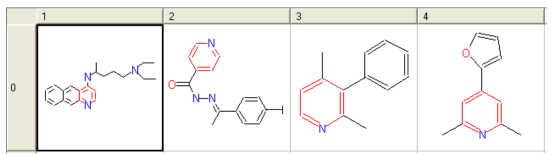
For example:

## Query:



Right click atom - Symbol - N

## Selection of chemical substructure search results



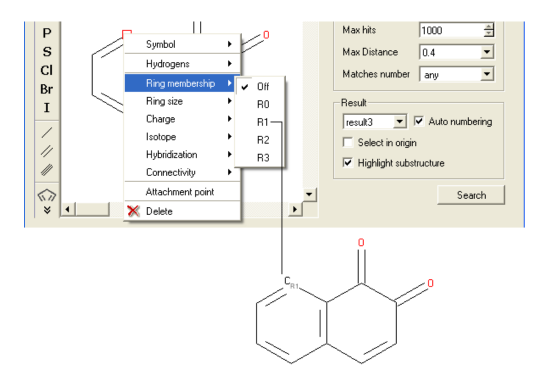
## To specify a particular number of hydrogen atoms at a particular site:

- Right click on the atom and select the "Hydrogens" option as shown below.
- Select how many hydrogens you wish to specify and a symbol will be displayed as shown below.

P S Cl Br I ∕// ∦ ©	Symbol Hydrogens Ring membership Ring size Charge Isotope Hybridization Connectivity Attachment point X Delete	<ul> <li>Off</li> <li>H0</li> <li>H1</li> <li>H2</li> <li>H3</li> <li>H4</li> <li>H5</li> </ul>	Max hits 1000 🚖 Max Distance 0.4 💌 Matches number any 💌 Result result3 V Auto numbering Select in origin V Highlight substructure Search

## To specify the number of rings a particular atom will be a member of:

- Right click on the atom and select the "Ring membership" option as shown below.
- Select whether the atom should be part of 1, 2 or 3 rings.



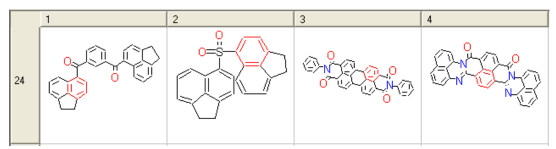
#### For example:

## Query:



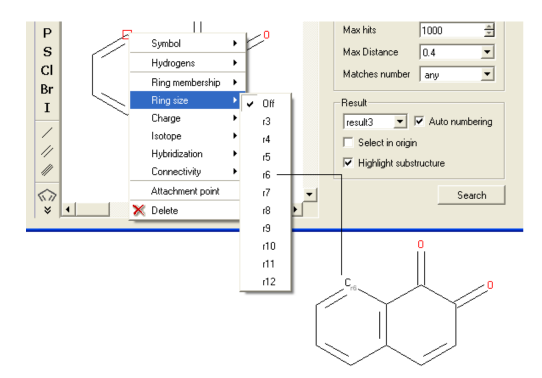
Right click atom - Ring Membership - R3

## Selection of chemical substructure search results



#### To specify the ring size connected to an atom:

- Right click on the atom and select the "Ring size" option as shown below.
- Select the size of the ring the atom should be connected to and a symbol will be displayed as shown below.



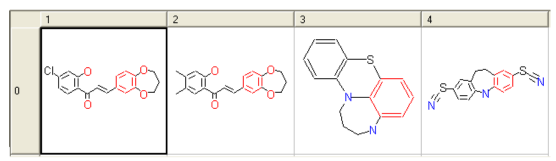
## For example:

## Query:



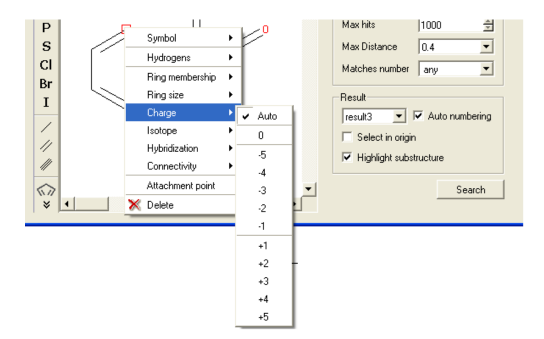
Right click atom - Ring size - R7

## Selection of chemical substructure search results



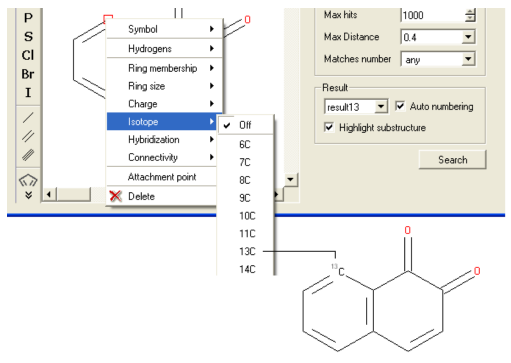
#### To specify the charge at a particular point:

- Right click on the atom and select the "Charge" option as shown below.
- Select the desired charge and a symbol will be displayed as shown below.



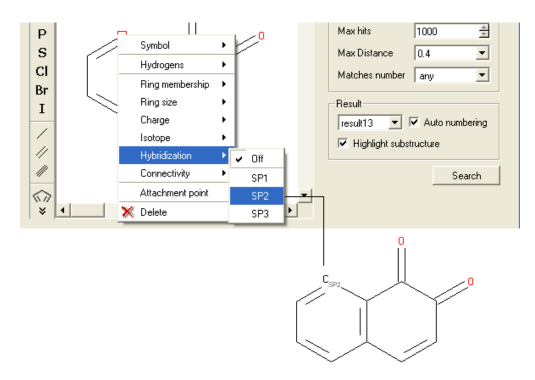
#### To specify an isotope at a particular atom

- Right click on the atom and select the "Isotope" option as shown below.
- Select the desired isotope from the list and a symbol as shown below will be displayed.



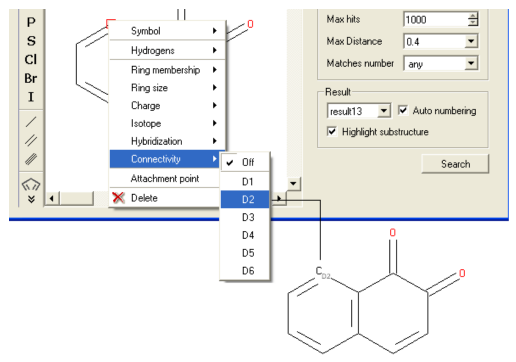
#### To specify the hybridization state of the atom:

- Right click on the atom and select the "Hybridization" option as shown below.
- Select the desired hybrization state and a symbol will be displayed as shown below.



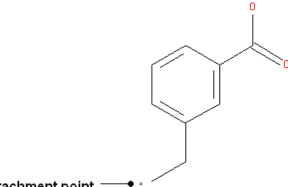
## To specify the number of atoms you wish an atom to be connected to:

- Right click on the atom and select the "Connectivity" option as shown below.
- Select the number of atoms you wish the atom to be connected to and a symbol will be displayed as shown below.



## To specify an attachment point - ie the position at which substituents will be added

- Right click on the atom and select the "Attachment point" option as shown below.
- An asterisk representing the attachment point will be displayed next to the atom.



## Attachment point

An attachment point means that the atom can be attached to zero or more bonds to heavy atoms.

## 7.5.2 Filter Search

## How to filter your query

To filter your query:

• Right click in the box shown below and select 'Add Condition'

( FieldName Relation Value	
( realise realise value	

**Right-click here and select Add Condition** 

To add conditions to your filter:

• Double click in the fields labeled "Name" and "Relation" and select the options from the drop down arrow or type in values.

	(	FieldName	Relation	Value	)
1		molid 📃 💌	==	î.	<u></u>
∎ Texts	earc	molid MOLSOFT_TABLE_ final_num int_num orig_num			<u> </u>

To remove a filter, right click on the filter and select 'Remove Filter'.

#### To exclude a fragment from your search

Click on the option Exclude fragment list:

Enter the SMART string of the fragment or use the Molecular Editor ٠ option to sketch the fragment.

• Click OK and the fragment will be listed at the bottom of the chemical search window as shown below.

🐐 ICM Chemical Search: [new file *] connected as andy@samba											
File	Edit View	Template	s Help						Data Source		
] B	📽 🖬 🛛	იი	16 <b>16 16</b>	a	<b>₽</b> , <i>Q</i> , ©	R/3	s 🕺 🔒		Table File Molcart		
=	C11H14				<b>I</b>						
11									,		
1	4 7	a adi									
Ż	4. 1	o eu	t or remov	veng							
c	See. 1	xclude	List				? 🗙	Ŷ	Query Options		
Ν								D	Search type Substructure		
0		ccccc1			C. Exclude	e Lis	st	?	distance 0.4 💌		
F								_	hber of matches any		
P					SMARTS:	-		-	😡 Match stereo 🦳 Ignore salt		
S Cl									Selected only		
Br							<u>ox c</u>	ancel	imum # of hits 10000 🚖		
I							/	-	- Results -		
В		Add			Ok	Á	Cancel	L	C Count hits only C Select in source		
н						$\neq$		1	C Hide unmatched     Save results to:		
						Table File Molcart					
Сф Ф					SMART						
PH T				ing o		result 🔽 🔽 Overwrite					
+/-				itor b etch		Highlight match FRotate by match					
-+			•			Display as grid 🔽 Store query					
×									F Append		
	( FieldNa	ame			Relation		Value	D		4	
	_								Hide after Search		
	1. Click	here				fr	agments are				
	/	nere									
•				-				F			
	de fragment list	c1cc	ccc1;;Cc1cccc	ic1							
l ext s	Text search										

## 7.5.3 Query Processing

To begin processing your query first you need to decide which database to search. The options are listed in the **Data Source** section of the ICM chemical search window.

## Data Source

You can either search a Table- Chemical Spreadsheet a File - Local Database or MolCart.

If you select **MolCart** you first need to setup the link to the correct database.

- Enter the Server Name in which the database is stored.
- Enter the database name.
- Enter your username and password for the server.
- You can save these details so you dont have to re-enter this information each time you use the chemical similarity search.

If you are searching a **Table**, click on the **Table** tab and then select the drop down button where the names of your currently loaded tables are stored.

If you are searching a **File** click on the **File** tab and then locate your local table database file .molt or if it is already loaded into ICM you can locate it with the drop down button.

## **Query Options**

#### Now select a search type:

- Click on the drop down arrow next to the "Search Type" option in the Query Options panel.
- Select the search type you want to use.

A **substructure** search is a search whereby only the defined molecule in the query will be searched against the database. Whereas, a **FP similarity** search which stands for fingerprint search enables any fingerprint within a structure to be searched for in the database. The similarity method calculates tanimoto distance using chemical fingerprints (1. -most distant, 0. - closest). The similarity = 1. - distance.

The **Max distance** option is available for use with the FP search and the **Matches number** option is for use with the substructure search. The option you do not require based on your search method will be blanked out. A "Max distance" value of 0 means that the search will only identify matches exactly the same as the fingerprint - the default is 0.4. The "Matches number" option allows you to stipulate how many times within a structure in the database your query can be found.

Additional options include:

- Match stereo option. With this option chirality will be taken into account in substructure and similarity searches.
- Ignore salts.
- Keep ring membership.
- Search tautomers. If this option is selected ICM will use different tautomeric forms of the query chemical during the search.

If you make a selection of your query ICM can use that selection to search. How to make selections in the Molecular Editor are described here. Enter the Maximum number(#) of hits you would like returned.

## Results

Before processing the query determine how you would like your results displayed in the **Results** section of the **Chemical Search** window.

**Count hits only** - this option will count the number of hits and display this number in a window once the searching has been completed.

**Select in source** - If you are searching a table you can select and highlight the query in the source table that you are searching.

**Hide unmatched** - Hide unmatched will hide the compounds that were not matched from view.

Save results to: - this option gives you the option to save the output results to a Table- Chemical Spreadsheet a File -Local Database or MolCart.

**Append** - this option will allow you to append to current table, file or Molcart database.

## Search

• Click on the **Search** button to execute the search. You can choose to hide the window after the search.

## 7.5.4 Search a Database by Text

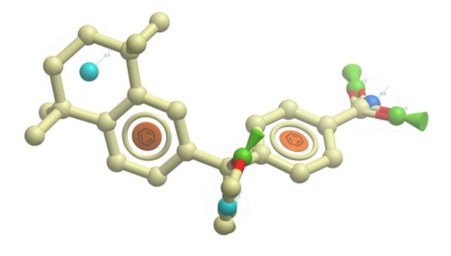
To search a database by text enter the text you wish to search in the **Text Search** data entry box at the bottom of the ICM Chemical Search window.

🐁 ICM Chemical Search: [new file *] Molcart not connected											
File Edit View Templates Help	Data Source										
File Edit View Templates Help	Data Source         Table       File       Molcart         Query Options       Image: Constructure       Image: Constructure         Query Options       Substructure       Image: Constructure         Query Options       Substructure       Image: Constructure         Max distance       0.4       Image: Constructure         Max distance       0.4       Image: Constructure         Maximum # of matches       any       Image: Constructure         Maximum # of hits       10000       Image: Constructure										
I B H H Gop Stereo > Attributes > */ V Elete V Stereo > Pit Y Stereo > Pit Stereo > Pit Y Stereo > Pit Stereo > Pit Pit Stereo > Pit Stereo >	Results         Count hits only       Select in source         Hide unmatched       Save results to:         Table       File       Molcart         result       Image: Overwrite         Image: Highlight match       Rotate by match         Display as grid       Store query         Append										
Exclude fragment list Text search											

Enter text here

## 7.5.5 Pharmacophore Drawing and Searching

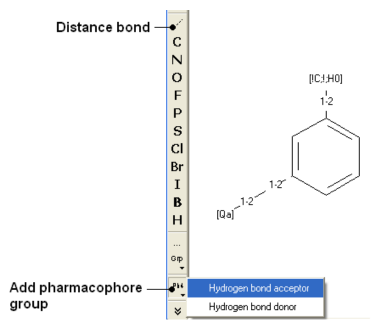
Pharmacophores can be drawn in 2D in the ICM Molecular Editor or in 3D in the Graphical Display. 2D pharmacophore sketches can be used to search chemical tables (spreadsheets) containing 2D or 3D coordinates. A 3D pharmacophore can be used to search chemical tables containing 3D coordinates only.



## 7.5.5.1 Pharmacophore Draw 2D

A 2D pharmacophore can be drawn using the ICM Molecular Editoror if you are going to use the drawing to search it is more efficient if you draw it in the Chemical Search window.

Use the distance bond button and the add pharmacophore group button to sketch the pharmacophore. The distance bond button represents the number of bonds between each pharmacophore point. You can edit the distance by right clicking on the bond and selecting edit. Other groups such as aromatic can be added using the standard molecular editor buttons.



**NOTE:** Do not mix the 2D and 3D pharmacophore environment. For example do not edit a 3D pharmacophore in the 2D editor.

## 7.5.5.2 Pharmacophore Draw 3D

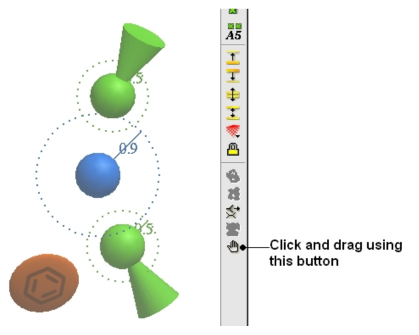
The easiest way to begin drawing a 3D pharmacophore is to draw a chemical in the ICM Molecular Editor which contains the key pharmacophore groups you want and then convert to 3D and extract the pharmacophore groups.

To draw a 3D pharmacophore this:

- Once the ligand is converted to 3D, right click on the ligand in the ICM Workspace.
- Select the Tools/Copy as Pharmacophore and choose the **pharm** centers option.
- The pharmacophore groups or centers will then be displayed in the graphical display and can be displayed and undisplayed in the ICM workspace.

#### To move a pharmacophore group:

- Use the **drag atom** button (picture of a hand. See below)
- Click on the pharmacophore group and then drag.



**NOTE:** Distances between groups can be monitored using the atom distance measurement tool. See Calculating the distance between two atoms.

#### To change a pharmacophore group:

- Right click on the pharmacophore group in the ICM Workspace or in the Graphical Display.
- Select Pharmacophore/Edit Point
- Choose the group from the drop-down list shown below.
- Enter the desired radius.

💙 Edit Pharmacophore Point	•	×
Туре	НВА 💌	
Radius	НВА	
	HBD	
	aromatic	
	hydrophobic	
	positive	
	negative	

#### To make a new pharmacophore group:

The easiest way to make a new pharmacophore group is to clone a pre-exisiting one. To do this:

- Right click on the pharmacophore group in the ICM Workspace or in the Graphical Display.
- Select Pharmacophore/Clone Point
- You can then move the new group as described above.

#### To change the direction of a pharmacophore group:

- Right click on the pharmacophore group in the ICM Workspace or in the Graphical Display.
- Select Pharmacophore/Assign Direction
- You can then move the new group using the drag atom button described above.

#### To remove the direction of a pharmacophore group:

- Right click on the pharmacophore group in the ICM Workspace or in the Graphical Display.
- Select Pharmacophore/Remove Direction

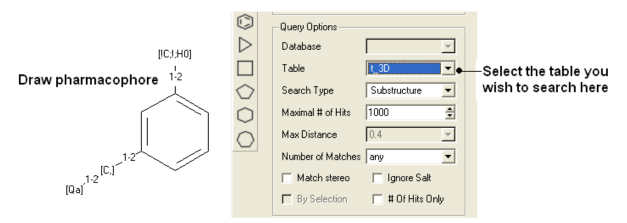
## a\_pharma.p9/9/Qh15 Filter Selection Edit ۲ Advanced ۲ Label • Center Connect to Molecule Disconnect Extract Ligand(s)... Delete Atom(s) Annotate Selection... Neighbors... Ň × Select ۲ Pharmacophore Edit Point... Clone Point Assign Direction

## Pharmacophore right click options

## 7.5.5.3 Pharmacophore Search

#### To perform a pharmacophore search using a 2D pharmacophore:

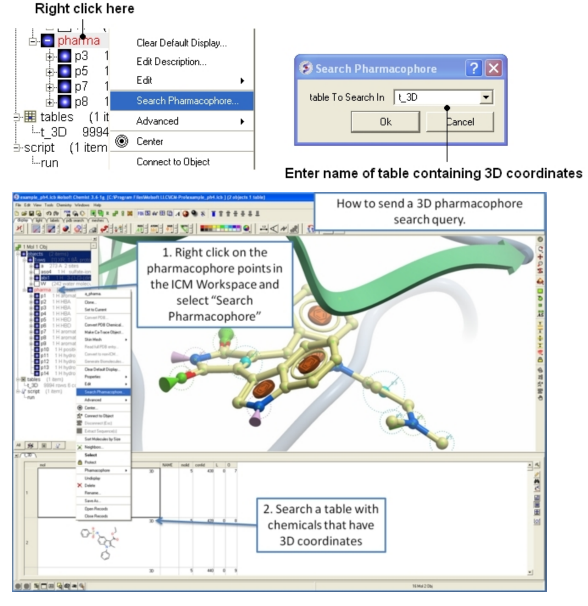
- Draw the 2D pharmacophore as described earlier in the Chemical Search window.
- Read in a molecular table to search or search a table in MOLCART.
- Select the chemical search options as shown below.
- Once the search has completed a new table with the results will be displayed.



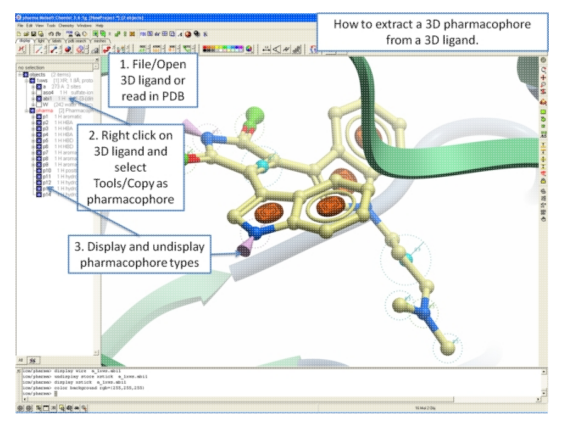
## To perform a pharmacophore search using a 3D pharmacophore:

- Right click on the pharmacophore in the ICM Workspace
- Select Search Pharmacophore

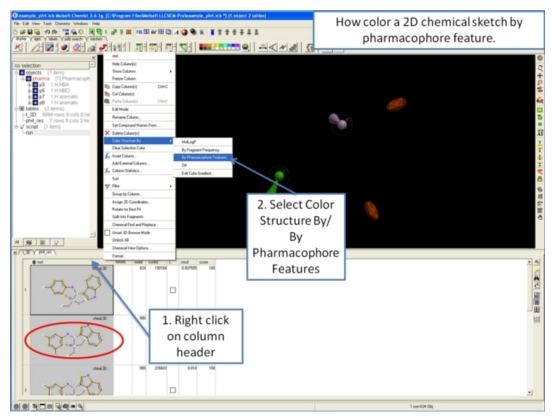
• Use the drop-down button to select the table you wish to search. The table must contain 3D coordinates.



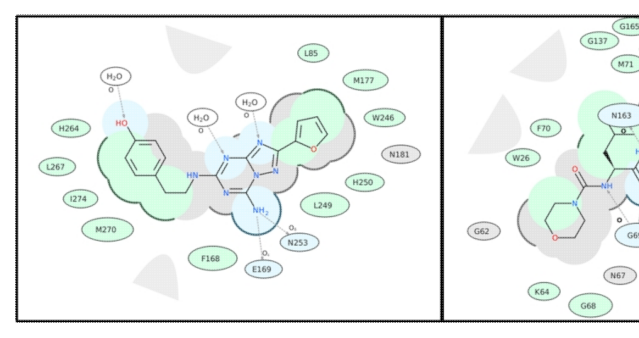
7.5.5.4 How to extract a 3D pharmacophore from a ligand.



## 7.5.5.5 How to color a 2D chemical sketch by pharmacophore feature.



## 7.6 2D Ligand Interaction Diagram



To create a 2D interaction diagram between a ligand and receptor:

- Read into ICM a PDB file containing the ligand and receptor. For correct hydrogen bond assignment you will need to convert to an ICM object.
- Right click on the ligand and choose Tools/Ligand 2D Diagram.
- A dialog box will be displayed. Select the receptor object from the drop down button and allow ICM to Auto Detect the ligand if needed.
- The Hydrophobic Distance Cutoff is <= to the minimum distance between ligand and the closest atom in the residue. The Hydrogen Bond Strength is based on the GRAPHICS.hbondMinStrength parameter so lowering the value will identify weaker hydrogen bonds.
- An option is available to match the 2D representation to a pre-calculated one saved in a chemical table loaded into ICM.

A guide to the coloring and representation of the 2D display:

- Green shading represents hydrophobic region.
- Blue shading represents hydrogen bond acceptor.
- White dashed arrows represents hydrogen bonds.
- Grey parabolas represents accessible surface for large areas.
- Broken thick line around ligand shape indicates accessible surface.
- Size of residue ellipse represents the strength of the contact.
- 2D distance between residue label and ligand represents proximity.
- Covalently bound residue represented with thick black line around ellipse. (C25 in the picture above)

To edit the 2D image (e.g. rotate a group):

• Right click on the image and choose Edit Molecule and edit using the Molecular Editor.

To save the 2D interaction diagram to an image file:

• Right click on the image and choose Save Export/Molecular Image

# 7.7 Convert Chemicals to 3D

Available in the following product(s): ICM-Chemist-Pro | ICM-VLS

#### To convert a chemical structure from 2D to 3D:

There are three ways in which to do this depending on whether you have a chemical in a chemical table or in the ICM workspace.

# From the Chemistry/Convert/2D to 3D option menu option - convert a chemical in a chemical table.

- Read in chemical spreadsheet table.
- Chemistry/Convert/2D to 3D
- Select the table from the drop down list.
- Select to keep hydrogens and/or fix amide bonds.
- Keep current table or overwrite.

#### If the compound is in the ICM Workspace:

- Select all of the structure to be converted by double clicking on it in the ICM workspace or by using other selection tools described in the Selection Toolbar Section of this manual.
- MolMechanics/ICM-Convert/Chemical and a data entry box as shown below will be displayed.

🦻 Convert small molecule object ? 🔀
The bond types and formal charges need to be correct
Object as_graph 💌
🔽 Display
🔽 buildHydrogens
🔽 fix amide bonds
🔽 overwrite original
Ok Cancel Help

If you have selected the compound as described above the "as\_graph" option in the Object data entry box will suffice. You can decide whether you wish to keep the chemical geometry or optimize it in a force-field. Other options include whether you wish to add hydrogens and fix amide bonds.

#### From an ICM chemical table:

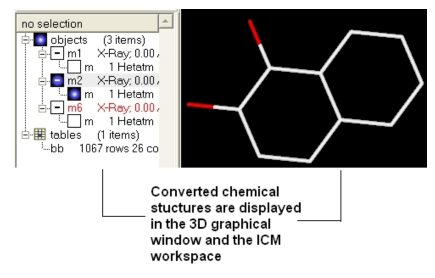
- Select which structures you wish to convert in the molecular table. For instructions on making selections within tables see the Making Table Selections part of this manual.
- Right click on one of the selections you have made and a menu as shown below will be displayed. Or use the Chemistry menu and select **Convert To 3D**.

×/	ricinLig	jands2D \		Save Molecule As Image		
		mol	C_	Edit Molecule		
		chiral 0	C,	Query Molecule		
				Chemistry	×.	Enumerate Stereoisomers (selected rows)
	1	••		Cell mol[2]	•	Enumerate Tautomers (selected rows)
		N N		Mark Selected Rows	•	Remove Salt (selected rows)
		N N		Select Marked Rows	<b>)</b>	Convert to 3D and Optimize
			₿ <u>₽</u>	Copy Row(s)	Ctrl+C	Load and Preserve Coordinates
		0		Paste	Ctrl+V	Conformation Generator (selected rows)
			8	Print Table	Ctrl+P	Chemical Template Superposition (selected rows)
		l ∽∕ N	Т			
	2					
				•		
		01101	N			
		chiral	2	53.219 -3 01583 -2.79511		
		0				
		0 m				

### Right click on selected rows

• Select the **Chemistry/Convert to 3D and Optimize** option and you will see the compounds being converted and minimized in the 3D graphical display window.

Once converted the compounds will be displayed in the 3D graphical display window and also in the ICM workspace.



Another way to convert all the ligands contained within a table (or a selection) into 3D coordinates :

- Chemistry/Convert to 3D..
- Use the drop down list to select the Molecular Table.
- Select whether you want to Keep Hydrogens, Fix Omegas and In Place. Select In Place if you want to overwrite current table.
- Click OK
- If you wish to run in **Batch** mode select the **Files** option.

**NOTE:** Use the 3D-Browse mode to view the chemicals in the graphical display.

### To convert 3D representation in a molecular table back to 2D:

- Chemistry/2D depiction
- Use the drop down list to select the Molecular Table.
- Select **In Place** if you want to overwrite the current table.
- Click OK.

### 7.7.1 Converting a Chemical from the PDB

The protein data bank has not been storing any information about covalent bond types and formal charges of the chemical compounds interacting with proteins! This oversight makes it impossible to automatically convert those molecules to anything sensible and requires your manual interactive assignment of bond types and formal charges for each compound in a pdb-entry. Therefore, if you apply the convert command to a pdb-entry with ligands, the ligands will just become some crippled incomplete molecules that can not be further conformationally optimized.

Therefore, follow these steps to convert a chemical properly from a pdb form to a correct icm object. There are two ways to do this either via the ICM Workspace (recommended) or via the Graphical Display.

# 7.7.2 Converting a Chemical from the PDB using the ICM Workspace

This is described in the Convert Protein and Chemicals chapter.

# 7.7.3 Converting a Chemical from the PDB using the Graphical Display

This is described in the Convert Protein and Chemicals chapter.

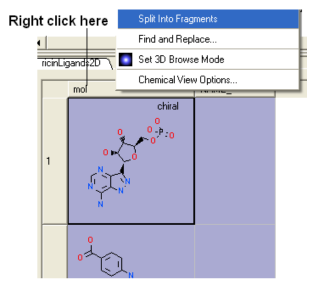
# 7.8 Generating Chemical Fragments

Available in the following product(s):  $\texttt{ICM-Chemist} \mid \texttt{ICM-Chemist-Pro} \mid \texttt{ICM-VLS}$ 

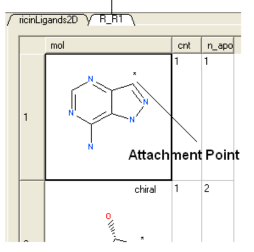
Chemicals displayed in an ICM Molecular Table can be split into fragments. This is useful for generating a series of R-groups to be added to a scaffold (See section describing reactions.

To generate fragments:

- Select the column or row(s) you wish to generate the fragment from.
- Right click on the "mol" column header and select "Split Into Fragments".
- A new table of chemical fragments will be displayed. Each fragment is assigned an attachment point which is flagged with an asterisk (\*).



### New table is generated containing fragments



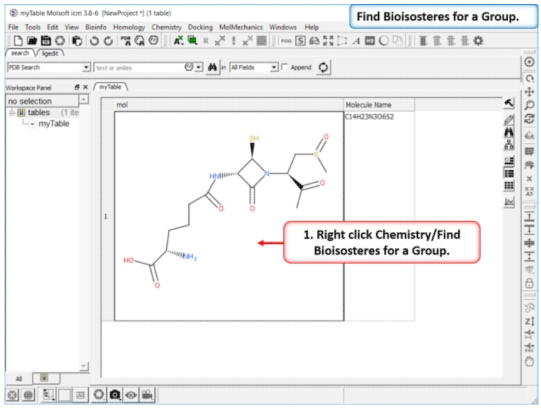
# 7.9 Find Bioisostere

|Bioisostere Webinar|

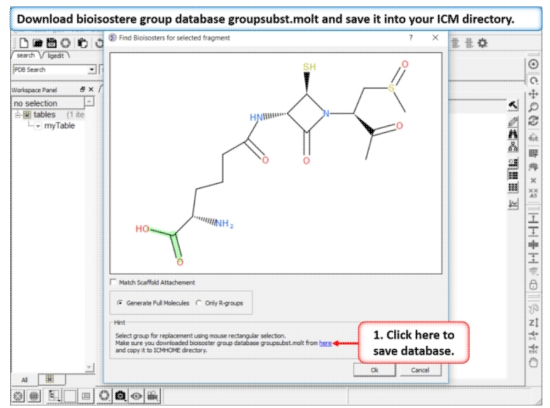
In drug design **bioisosteres** can be used to reduce toxicity, change bioavailability, alter metabolism and change activity of lead compound. Bioisosteres are chemical substituents with similar chemical or physical properties which produce broadly similar biological properties to another chemical compound.

### 7.9.1 2D Bioisostere

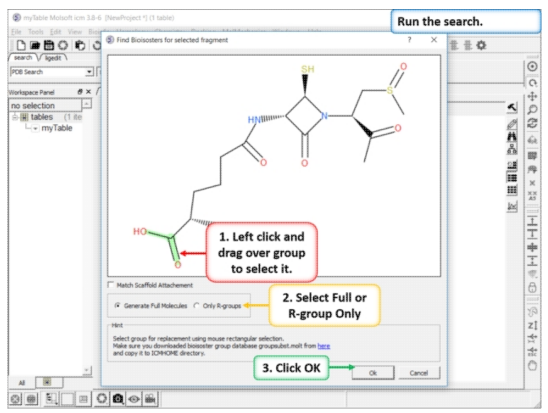
To find 2D Bioisosteres:



**Step 1:** Read in a chemical into a chemical table. Right click Chemistry/Find Bioisosteres for a Group.



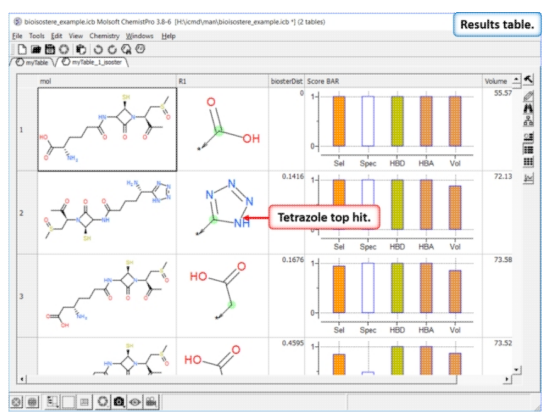
**Step 2: Download Database** If you do not see this link it means you already have the database in your ICM directory. If yo do not have it - download bioisostere group database groupsubst.molt and save it into your ICM directory. There is a link at the bottom of the dialog window



### which will direct you to the correct place to download.

Step 3: Setup the Search

- Select the tab for 2D Database at the top of the dialog window.
- Left click and drag over the group you wish to find a bioisostere for.
- Select if you want to "Match Scaffold Attachment". It matches the attachment atom from the R-group which you're trying to substitute with the the database. In this example the attachment is C^3 (SP3 Hybridized), so if you select this option it will only return hits where scaffold attachment column contains C^3.
- Select if you want to generate Full Scaffold or just show the R-group.
- Click OK to run the search.



**Step 4: Results Table** In this example we find tetrazole which is a classic isostere of carboxylic acid. The data in the columns represent:

- Mol is the query chemical with the bioisostere attached.
- **R1** is the bioisostere.
- **biosterDist** Bioisoster â distance' is calculated on the basis of five terms, two are knowledge-based and three are physicochemical. They are also reflected by the corresponding bars in the column Score BAR.
- **Score BAR**: For the bar display the terms are transformed into 0-1 range so that 1 corresponds to best match/identity.
- Sel selectivity, as estimated from frequency of replacement of query R- group by a retrieved substituent relative to total number of cases where this substituent replaces any -R group in the database
- **Spec** specificity, as estimated by the difference in activity between a database compound with a retrieved substituent and least active example of a different substituent (on the same scaffold and target) in the database
- **HBD** factor reflecting difference in the number of HB donor functionalities of the retrieved substituent and query -R group
- **HBA** factor reflecting difference in the number of HB acceptor functionalities of the retrieved substituent and query -R group
- Vol factor reflecting difference in volume of the retrieved substituent and query -R group
- **scfldAttach** indicates the atom(s) and it's hybridization in the original group attachment. It can be used to filter solutions in the query.

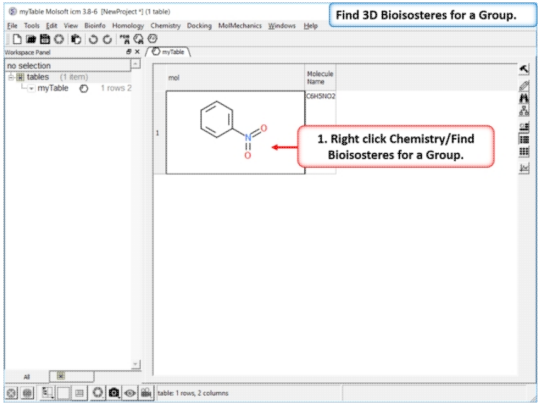
If you double click on a row in the table it will show specific examples of a particular substitution from ChEMBL.

### 7.9.2 3D Bioisostere

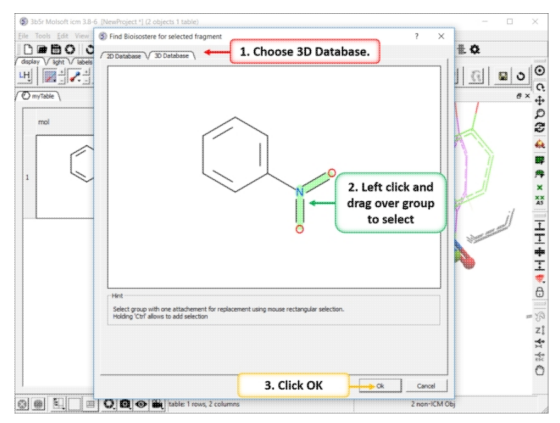
The receptor/ligand complex structures from the PDB contains relevant (bioactive) 3D conformation and the environment of the isosteric moieties. This can be used to find 3D bioisosteres. You can watch a

Bioisostere Webinar which explains how the 3D database was constructed.

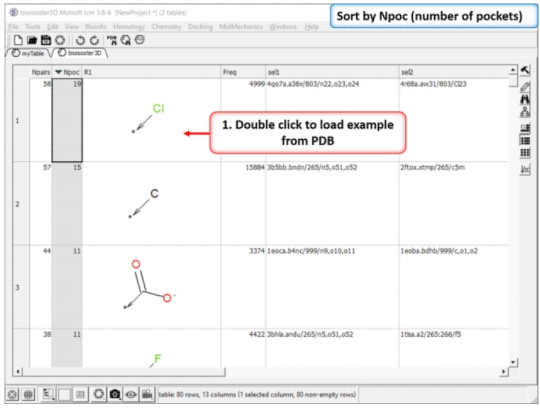
### To find 3D Bioisosteres:



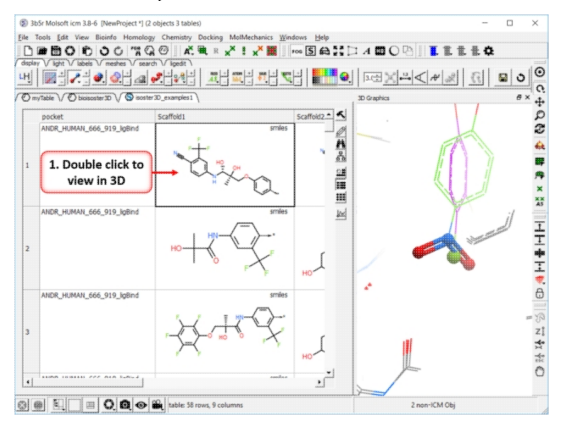
**Step 1:** Read in a chemical into a chemical table. Right click Chemistry/Find Bioisosteres for a Group.



# **Step 2:** Left click and drag over the group you wish to find a 3D bioisostere for and then click OK.



**Step 3:** The results table will be displayed the nPocs (number of pockets) column is a good indicator of a good bioisostere. As this is the number of times this replacement is seen in the PDB.



7.9.2 3D Bioisostere

Step 4: Click on any of the hits to view specific pair examples in 3D.

# 7.10 Molcart

Available in the following product(s): MolCart

Molcart is an enterprise wide chemical management system. Compound databases of any size can be stored in MolCart and analyzed and searched using ICM cheminformatic tools.

### 7.10.1 Molcart Installation

In order to run MolCart it is necesary to install the Free Open Source MySQL database on your machine. Please see your systems administrator or see http://dev.mysql.com/downloads/

### **Linux Installation**

Mysql: Check if mysql daemon is running :

/etc/init.d/mysql status

If you see that the MYSQL service is unused (not running), you need to start the mysql deamon. Become root and do the following:

/etc/init.d/mysql start

### Download and install Molcart files.

The MolCart package is a self-extracting executable file.

Installation Instructions:

- Download the MolCart file ( molcart-*version-platform*.sh ) from the Molsoft website.
- Type in a shell window: ./molcart-version-platform.sh -p=*THE\_PATH\_YOU\_WANT\_TO\_DOWNLOAD* (NOTE: You must be logged in as 'root' to install the 'molcart-version-platform.sh' to '/usr/molcart-version-platform'
- The following question will be displayed: Do you want to install the molcart-1.6-6 to "/usr/molcart-1.6-6" now? (y/n) [y]
- Answer YES and the unpacking process will begin
- You will now be prompted for a password.
- Select which default databases you wish to install.
- Make a note of the HOSTNAME, DATABASE NAME and USER NAME
- MolCart is now fully installed.

#### **Mac Installation**

System requirements: \* Mac OS 10.3 \* MySQL server for Mac OS 10.3

To install Molcart on the Mac just run this:

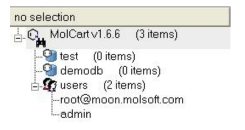
sudo /some/path/molcart-1.6-6-darwin.sh

### 7.10.2 Molcart Getting Started

### To start Molcart:

Tools/Connect Molcart

Once you have activated MolCart the loaded databases and users will be shown in the ICM Workspace as shown below.

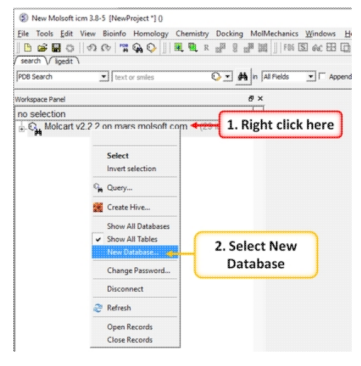


All the records and fields contained within each database can be viewed by expanding the tree structure in the ICM Workspace.

### 7.10.3 Creating Databases and Importing Data

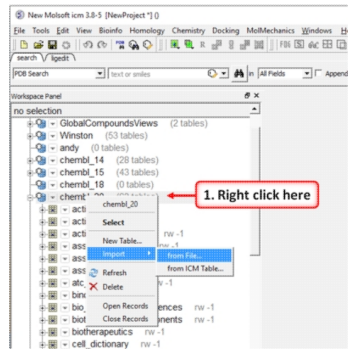
### To make a new database:

• Right click on the top of the MolCart tree and then choose New Database



To upload SDF, csv, SMILES or MOL files or an ICM table:

- Right click on the database where you wish to add the file.
- Select Import.



### 7.10.4 Molcart Search

### How to search the databases contained within MolCart

Click on the button shown below.



#### MolCart and Chemical Search Tools

To begin processing your query first you need to decide which database to search. The options are listed in the "Data Source" section of the ICM chemical search window.

Table	File	Molcart		
Databas	e test		ŀ	•
Table	cher	nbl		-

You can either search a local table (molecular table) or you can search MolCart.

If you select MolCart you first need to setup the link to the correct database - described earlier. Click on the button shown above (yellow pencil) and the Connect to Molcart window will be displayed as shown below.

S Connect	to Database	?	×		
Server Name	mars.molsoft.com				
Database	test				
User	molcart				
Password	*****				
Reconnec	at startup Port 53306				
	ОК	Can	cel		

- Enter the Server Name in which the database is stored.
- Enter the database name.
- Enter your username and password for the server.
- You can save these details so you dont have to re-enter this information each time you use the chemical similarity search.

See the Chemical Search section of this manual on the many different search procedures.

#### How to perform a text search

To perform a text search on one of the databases contained within Molcart you first need to index the text within the database and then search using the query option.

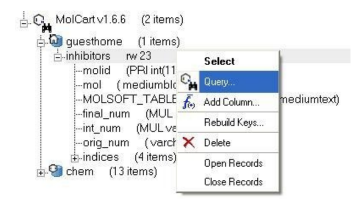
To make the database index, see text and picture below:

- Expand the tree of the database in the ICM Workspace.
- Select the column headers you wish to search which contain Full Text or Partial Text(the data type for each column is listed next to the column name). Multiple column headers can be selected by clicking and holding down the CTRL key. A range of column headers can be selected by holding down the shift key and clicking to select.
- Next, right click and select Create Index.
- Select 'Full Text' and you will notice an additional header in the ICM Workspace called 'indices'. The value in the items category represents the number of columns you have chosen to text search.

E. O MolCart ∨1.6.6 (2 i	tems)	
	ems)	
🛓 inhibitors 🛛 rw 23		
molid (PRI ir	nt(11))	
-mol (mediur		
MOLSOFT_T final_num (	Unselect	(MUL mediumtext)
int_num (M	Create Index	
<mark>orig_num (</mark> >	🗙 Delete	
indices (4 ine	ans)	-
🗄 🎱 chem 🛛 (13 items)		

#### To perform the text search:

- Right click on the database name in the ICM Workspace.
- Select 'Query' as shown below.



The ICM Chemical Search query window will be displayed as shown below. Type your query text in the space provided and hit the enter key or click on the search button.

ICM Chemical Search: connected as guest@www	
File Edit View Help	Data Source         C Local Tables       Image: MolCart         Query Options       Image: MolCart         Database       guesthome         Table       inhibitors         Table       Substructure         Maximal # of Hits       1000         Max Distance       0.4         Number of Matches       any         Image: Match stereo       Image: # Of Hits Only         Image: Search by Selected Fragment       Result
FieldName     Relation     Value       Text search	result3     ▼     Auto Numbering       Add to DB     guesthome     ▼       ▼     Highlight Substructure       □     Display as Grid

Type text search query here Filter your search here

#### How to add conditions to your query

• Right click in the box shown below and select 'Add Condition'. You can add as many conditions as you like.

( Fie	IdName	Relation	Value	)
( Fie	ldName	Relation	Value	

Right-click here and select Add Condition

• Double click in the fields labeled "Name" and "Relation" and select the options from the drop down arrow or type in values.

	l	FieldName	Relation	Value	)
1		molid 💌	==	î.	
<b>∢</b>   Texts	earc	molid MOLSOFT_TABLE_ final_num int_num orig_num			<b>}</b>

To remove a condition, right click on the filter and select 'Remove Filter'.

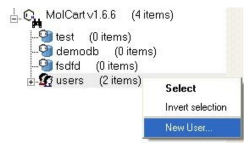
### 7.10.5 Molcart Administration

### **Edit Password**

• Right click on the MolCart header in the ICM Workspace and select Change Password.

### Add a New MolCart User - Root Only

• Right click on the User Section of MolCart in the ICM Workspace as shown below.



- Select the New User Option.
- Add new username and password in the data entry box.
- New user will be displayed in the ICM Workspace.

### Edit User Privileges - Root Only

- Right click on the user in the ICM Workspace.
- Select Edit Privileges and a data entry box as shown below will be displayed.



- Select the database name.
- Edit the privileges by checking or unchecking the appropriate boxes.

#### Add a New Database

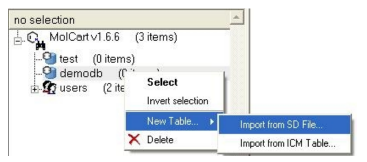
• Right click on the MolCart Header in the ICM Workspace as shown below.

no selection		<u> </u>
HolCart ∨1.6.6 9 test (0items) 9 demodb (0i	( 4 3)	Select Invert selection
	~	Show All Databases
iterriusers (2 iterr ⊡root@moon.m		New Database
admin		Change Password
		Open Records
		Close Records

- Select the New Database option.
- Enter a unique name for your new database.
- The new database name will appear in the ICM Workspace.
- Now you need to add data to your new database (See Instructions Below).

#### Add New Data to Database

• Right click on the database name in the ICM Workspace as shown below.



- Select New Table.
- Select either Import from SD file or Import from ICM table.
- Select the appropriate file and the records structure of your sdf or ICM table will be displayed as shown below.

🥩 MolCart Import		? 🛛
Table name melanin		
🔽 AE	String 💌	3769
🔽 APP	String 💌	0
🔽 CAS	String 💌	16
🔽 CI	String 💌	145
COMB_PREP	String 💌	1638
COMMENT	String 💌	0
IA I	String 💌	472
🔽 INN	String 💌	0
UI 🟹	String 💌	760
🔽 MA	String 💌	1082
₩ MF	String 💌	38
MOLNAME	String 💌	24
MOLSOFT_TABLE_HEADER	String 💌	192
MOL_WEIGHT	Real 💌	8

- The database name can be changed at this point and the fields contained within the database can be altered. Certain fields can be excluded by checking the boxes this will help in minimizing the size of a database. Caution must be taken if you want to change the field type or length.
- Click OK and your sdf file or ICM table will be added to the database. This can be seen by expanding the tree structure in the ICM Workspace.

### Delete a Database or User

- Right click on either the database or user in the ICM Workspace.
- Select Delete.

## 7.11 Calculate Properties

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

|Video|

#### To calculate chemical properties for compounds within a chemical table:

- Read in the chemical table.
- Select Chemistry/Calculate Properties and a window as shown below will be displayed.
- Arguments usually the column with the chemical structure is called mol. If this is not the case click on the drop down button and select the correct column.
- New column location select where you would like to insert the new columns.
- New column name This option is usually greyed out but the column name will be the same as the property name.
- Select the properties you wish to calculate using the 'tick' check boxes.
- Click OK and the properties will be added as new columns in the chemical table.

Function	Actions -			
,	Function	Name	Category	Description
Arguments	MolWeight(mol)	MolWeight (	Chemical	Molecular weight
	MolFormula(mol)	MolFormula 0	Chemical	Chemical formula,e.g. C2H6O
Main argument: mol 💌	MolLogP(mol)	MolLogP (	Chemical	Octanol water partition, -Log
	MolLogS(mol)	MolLogS 0	Chemical	Water solubility -Log(C_aggr)
New column location	MolPSA(mol)	MolPSA (	Chemical	Polar surface area
	MolVol(mol)	MoNol 0	Chemical	Molecular volume
Insert ( after C before C in-place	MoldHI(mol)	MoldHf 0	Chemical	Heats of formation from elem
	DrugLikeness(mol)	DrugLiken (	Chemical	Empirical drug-likeness
estant Ind	Smiles(mol,mode=normal)	Smiles 0	Chemical	SMILES/SMARTS: string no
column mol 💌	BadGroups(mol)	BadGroups (	Chemical	Unwanted or reactive chemi
	Nof_Atoms(mol.atom=""")	Nof_Atoms 0	Chemical	Number of atoms
New column name	Nof_Molecules(mol)	Nof_Molec 0	Chemical	Number of individual molecul
name	Nof_Chirals(mol.type=-1)	Nof_Chirals 0	Chemical	Number of chiral centers, R,
	Nof_Rings(mol)	Nof_Rings (	Chemical	Number of rings in the SSSR
	III T manual man			

Select properties here

### **Actions Button Helps with Selections**

If you click on the drop down button labeled Actions you will see 3 selection options:

- Select All properties.Uncheck selected options.
- Remove unchecked.

### 7.11.1 Number of Atoms

You can calculate the Number of Atoms using:

- Chemistry/Calculate PropertiesChoose the Function Chemical from the drop down list and then Nof\_Atoms.
- Add the SMARTS string for the atoms you are trying to count (e.g. sp3 : [\*;^3] halogen : X or [F,Cl,Br,I])

Nof SP3	example:
---------	----------

Function	į		K Action: •				
Eunction	Nof_Atoms	•	Function	Name	Category	Description	
			MolWeight(mol)	moliveight	Chemical	Molecular weight fr	
Arguments			C MolFormula(mol)	molFormula	Chemical	Chemical formula, e	
Nof Atoms( m	ol • ,atom= [(*;*	-31 • 1	Iupachame(mol)	iupacRame	Chemical	SUPAC nomenclatur	
			MolLogP(mol)	malLogP	Chemical	Octanol water partit	
New column location		C MolLogS(mol)	molLogS	Chemical	Water solubility -Log		
			C MalPSA(mail)	molPSA	Chemical	Polar surface area f	
6	after C before C	in-place	MolArea(mol)	molArea	Chemical	Total surface area fr	
nsert			MolVolume(mol)	molivolume	Chemical	Molecular volume fr	
	` at end		MoldHf(mol)	moldHf	Chemical	Heats of formation f	
	F		DrugLikeness(mol)	drugLikeness	Chemical	Empirical drug-liken	
olumn	lenal	<u>.</u>	C Smiles(mol,mode='unique')	smiles	Chemical	SMILES/SMARTS: E	
iew column nam	10		BadGroups(mol)	badGroups	Chemical	Unwanted or reactiv	
Reever Concernan Indefine		O Nof_Atoms(mol,atom='*')	nof_Atoms	Chemical	Number of atoms fr		
nof_SP3		C Nof_Molecules(mol)	nof_Molecules	Chemical	Number of individua		
			Nof_Fragments(mol,fragment*'*')	nof_Fragments	Chemical	Number of fragments	
Options		Nof_Chirals(mol,type='any')	nof_Chirals	Chemical	Number of chiral ce		
Don't store formula		Nof_Rings(mol)	nof_Rings	Chemical	Number of rings in t		
		May Rinn Citalmolt	may Dina Cina	Chamical	I amost infanantarit		

### 7.11.2 ToxScore

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

The **ToxScore** gives a prediction for how reactive or toxic a chemical is.

### How the ToxScore is Calculated

Around 1000 SMARTS strings associated with toxicity/reactivity were collected from various sources. They were assigned demerit score based on their perceived toxicity and frequency of appearance in approved drugs. The toxscore of any compound was calculated by summing up all the matching SMARTS strings. A toxscore  $\geq 1$  indicates likely toxicity based on substructure match.

### How to Calculate ToxScore

- Read into ICM an SDF file. A chemical spreadsheet will be displayed in the GUI.
- insert a column into the chemical spreadsheet. Right click on a column header > choose insert column > choose the chemical function > select Tox Score.
- A column containing a Tox Score and Tox Names will be displayed.

# 7.12 Standardize Table

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

**NOTE:** Before standardizing a chemical you may want to make a copy of the original so you do not lose any information. You can do this by right clicking on the name tab of the table and selecting **clone** or **save as**.

# To remove salts, explicit hydrogens and standardize groups in a chemical table:

- Chemistry/Standardize
- Select the table from the drop-down list. This will also work for selections made on the table.
- Select option Remove Salts, Keep Salt Info in Extra Column (NOTE: option availabe in 3.7-2b and higher), Remove Explicit Hydrogens or Standardize Groups
- Click ŎK.

	chimode		
🗐 Standardize			2
Local Tables V Fi	es		[
Molecular Table	result	*	
Options			
Remove Salts	5	📃 Keep Salt Info In Extra Column	
Remove Expl	icit Hydrogens	🔽 Standardize Groups	
		Ok Cancel H	lelp

### Batch mode

Standardize chemical groups will apply rules from ICMHOME/CHEMNORMRULES.tab For example some chemical groups may have different representations e.g. [N+] (=0)O versus N(=O)=O

### To run in batch mode:

- Chemistry/Standardize
- Select the **Files** tab
- Enter the path and name of the sdf file you wish to standardize or use the browse button.
- Enter the path and name of the output file or use the browse button.

#### To remove a salt from an individual row in a chemical table:

- Select the row or rows.
- Right click Chemistry/Remove Salt (Selected Row)

## 7.13 Annotate By Substructure

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

This feature allows you to annotate a chemical spreadsheet according to functional group. It also allows you to flag substructures which may have poor ADME properties.

S Annotate by Substructure	2 🛛
Existing Fragments Draw Fragments	
Molecular Table result Choose table	
● Functional Groups ○ Substructure Alerts ○ Custom Fragments	— Choose annotation option
Hint Default table with functional group will be used for annotation: C:/Program I	Files/Molsoft LLC/ICM-Pro_21/ FUNCGROUPS.sdf
	Ok Cancel

First read in a chemical spreadsheet or sdf file you wish to annotate. To do this:

• Information on how to read into ICM chemical spreadsheets and structure can be found here.

#### To annotate functional groups:

- Chemistry/Annotate by Substructure.
- Enter the name of the Molecular Table (Chemical Spreadsheet) or use the drop down button to locate it.
- Check the **Functional Groups** option.
- The functional groups will be listed in a new column in your chemical spreadsheet called **funcgroup**. The default table with functional group will be used for annotation called FUNCGROUPS.sdf in ICMHOME.

#### To annotate potentially poor ADME groups (Substructure Alert).

#### 7.12 Standardize Table

- Chemistry/Annotate by Substructure.
- Enter the name of the Molecular Table (Chemical Spreadsheet) or use the drop down button to locate it.
- Check the **Substructure Alerts** option.
- The alerts will be listed in a new column of your chemical spreadsheet called **alerts.** The default table with substructure alerts will be used for annotation called CHEMFILTER.sdf

#### To annotate by custom fragments

S Annotate by Substructure	×
Existing Fragments Draw Fragments	
Molecular Table result	
Functional Groups     Substructure Alerts     O     Custom Fragments	
Table With Fragments       result       Read in a table containing your custom fragments         Hint       Hint	
a column called ".LABEL" or ".LABELS" in either upper or lower case. Molecular formula will be used if no label column is found	
<u>Ok</u> <u>Cancel</u>	

- Read into ICM a table with annotated fragments. The fragments should be labeled in a column called ".LABELS" or ".LABEL" in either upper or lower case. If not "LABELS" column is found the molecular formula will be used instead.
- Chemistry/Annotate by Substructure.
- Check the Custom Fragments button

To draw a fragment and annotate.

Annotate by Substructure						?×
Existing Fragments Draw Fragment	ts					
Molecular Table result	~				Select checkbox	
	Click	here t	o sketch		to use fragment	
C(C(0)=0)c1ccccc1	-	label	1			
			Click her	e to label		
	v C/	label	2	×		
	v C/	label	3	¥ 🗌		
	~ ©,	label	4	~		
		Iabei	4			
First Fragment Match						
						cel

- Click on the **Draw Fragments** tab.
- Enter the SMILES string of the fragment or click on the ligand editor button in the dialog window to sketch the fragment.
- Enter a label name.
- Click the check box next to the label name.
- Click OK

# 7.14 Align/Color by 2D Scaffold

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

This option **aligns** a set of sketches in a chemical table in the same orientation according to a defined scaffold or **color** by a common substructure.

- Read in a chemical table.
- Chemistry/Align-Color by 2D Scaffold
- Choose a loaded molecular table from the drop-down arrow.
- You can callow ICM to auto detec the common scaffold or draw a new scaffold using the molecular editor or choose the scaffold from a table (Index = row number).

# 7.15 Set Formal Charges

```
Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS
```

To set formal charge:

- Read in a chemical spreadsheet File/Open SDF
- Chemistry/Set Formal Charges
- Enter the name of the chemical spreadsheet (table).
- Choose the option to either make Charge, Make Neutral, or Charge according to ICM pKa Model
- Select the groups you want to charge or neutralize.
- The charges will automatically be added to your table.

## 7.16 Enumerate Formal Charge States

To enumerate formal charge states.

- Read in a chemical spreadsheet File/Open SDF
- Choose Chemistry/Enumerate Formal Charge States
- Enter the name of the chemical spreadsheet (Molecular Table)
- Choose the pH at which you wish to evaluate the charge
- Choose a window value.
- Click OK and a new table "\_charge" will be displayed with the enumerated charge states.

### About the Method

For basic charges the logic is following:

- atoms with pKa > pH+window is always charged
- atoms with pKa inside the range [ pH-window : pH+window ] will be enumerated as charged and uncharged.

For example if you have two centers with pKa within the range then four states will be generated ( [0,0] [+1,0] [0,+1] [+1,+1] )

For acids it is similar.

## 7.17 Convert

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

### 7.17.1 Convert 2D to 3D

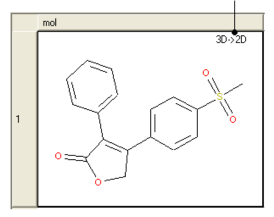
- Read a chemical table (sdf file) into ICM.
- Chemistry/Convert/2D to 3D
- Select the table from the drop down list.
- Select to keep hydrogens and/or fix amide bonds.
- Keep current table (In Place) or overwrite.

### 7.17.2 3D to 2D Depiction

If you have a chemical table displayed containing 3D coordinates or you wish to reassign the 2D coordinates in an sdf file you can use this option.

- Read a chemical table (sdf file) into ICM.
- Chemistry/Convert/2D Depiction
- Enter name of loaded chemical table.
- Choose In Place if you want to overwrite the table.
- Choose the **Files** tab to run in batch mode.

### Table contains 3D coordinates



### 7.17.3 Convert Smiles or InChi to 2D

#### To convert smiles or InChi strings to 2D sketches

- Read in a table containing the smiles or InChi strings in separate rows. For example the smiles strings maybe in an Excel file and you can load this into ICM by saving the Excel file as comma-separated (csv).
- Select Chemistry/Convert/Smiles or InChi to 2D.
- Select the table you want to convert using the drop down arrow and the name of the column containing the smiles string.
- Select whether you wish to keep the smiles column in the new table.
- Click OK and a table will be displayed containing the 2D structure.

ology       Chemistry       Docking       MolMechanics       Windows       Help         Image: Construction of the second se	
Standardize Annotate by Substructure Align/Color By 2D Scaffold Set Formal Charges Convert 2 2D To 3D -c(c(N+c(NH2)=0)N1)0)0 H H	1
Align/Color By 2D Scaffold         # × / Xdid           Set Formal Charges         # × / Xdid           Convext         *           2D To 3D         •C(C(N=C(N)H2)=0)N1)0)0           H         H	1
Set Formal Charges	1
Convert 2D To 3D •C(C(N+C(N)M2)=0)N1)0)0 H H	1
head A pulled Provide Model 3D to 2D Depiction	1
Predict Smiles to 20.	20H;
Generate 3D Conformers Generate Tautomers	
Generate Stereoisomers Generate Drug from Prodrug Generate Prodrug from Drug Enumerate Formal Charge States Ligand Energetics	)u
PCA Analysis     CC(C)(C)HCC(c1ccc(c(CO)c1)O)O       Compare Two Sets     CC(C)(C)HCC(c1ccc(c(CO)c1)O)O       Merge Two Sets     227       Select Duplicates     227	но
Combinatorial Chemistry	OH
Superposition     CC(C)/N=C(N)N=C(N)Nc1ccc(cc1)(0)       APF Tools	, P
MolScreen Model > 228	
Load Example Table	

### 7.17.4 Convert Structure to Smiles

### To convert an sdf file of 2D or 3D chemical coordinate in Smiles:

- Read a chemical table (sdf file) into ICM.
- Select Chemistry/Convert Structure to Smiles.
- Select the table you want to convert using the drop down arrow and the name of the column containing the 2D sketch.
- Select whether you wish to keep the 2D sketch column in the new table.
- Click OK and a table will be displayed containing the smiles string.

### 7.17.5 Convert to Racemic

# To remove stereo bonds and make all chemical centers R/S in a chemical table:

- Chemistry/Convert to Racemic
- Select the Molecular Table
- Select In Place if you wish to overwrite the table.

0	Convert To Racemic 🛛 🛛 🛛						
	Clears UP/DOWN bonds and makes all chiral centers R/S. You may use 'Generate Stereoisomers' after that.						
Molecular Table ricinLigands2D_3D 💌							
	Ok Cancel						

NOTE: To reassign stereo bonds use the Generate Stereoisomers option

# 7.18 Build Prediction Model

Available in the following product(s): ICM-Chemist-Pro | ICM-VLS

Structure-Activity Relationship (SAR) is a process by which the activity of a molecule is related to its molecular structure. If a significant ammount of structural and activity data is available a model can be made which can be used to predict the activity of a molecule or series of molecules.

In ICM SAR is undertaken using the Learn and Predict tools in a Molecular Table.

## 7.19 Predict

Available in the following product(s): ICM-Chemist-Pro | ICM-VLS

To make a prediction using a model you need either ICM-Pro + Chemistry or ICMChemistPro.

Read the table of data into ICM from which you wish to predict. Make sure the table contains the same columns used for the learn model.

- Tools/Table/Predict
- Select which table you wish to make the prediction on.
- Select which model you wish to use.
- Check that the required columns are in the table. If they are absent a red mark will appear against the column that is missing.
- Click Predict.

# 7.20 Generate 3D Conformers

Available in the following product(s): ICM-Chemist-Pro | ICM-VLS

To generate a series of conformers for a ligand(s):

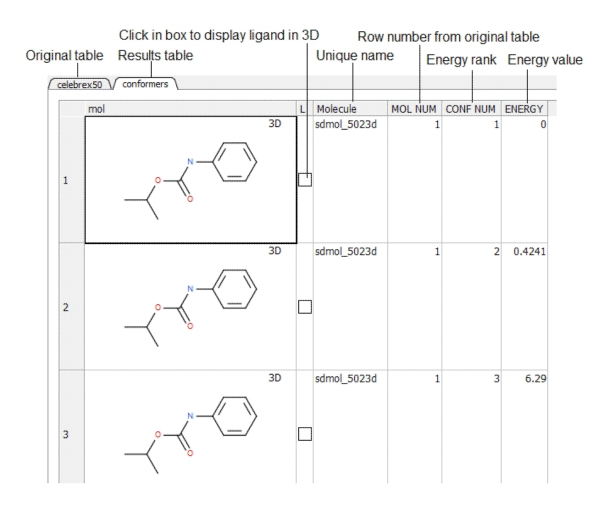
- Select the compounds (row(s)) you wish to generate conformers for in an ICM Molecular Table. Or to convert a whole table of compounds select **Chemisty/Generate 3D Conformers** menu.
- Right click on the selected row(s) and Chemistry/Conformation Generator (selected rows) and a data entry window as shown below will be displayed.

🦻 Generate 3D Conformers				? 🗙
/¡Local Tables; / Files \				
Molecular Table	•			
Max Number of Conformations	50	•		
vicinity 15.	thoroughness 1.	-		
🔽 Sample Rings	Sample CisTrans	;	🔲 Systematic Search	
🦵 Cartesian Refinement	┌─ verbose			
🔽 Keep Hydrogens	Pyramidal			
		Ok	Cancel H	elp

- Enter the maximum number of conformations you wish to generate.
- Enter a vicinity value. For more information on vicinity please see the command language manual http://www.molsoft.com/man/reals.html#vicinity.Density of conformers is controlled by vicinity= parameter. Currently it always operates in terms of torsion RMSD, default is 30 degrees angular RMSD. Note that we only generate local minima, therefore reducing vicinity below some level will typically not produce any new conformers.
- Enter a effort value (thoroughness in releases before 3.8-2). This relates to the length of sampling time. Effort multiplies the sampling length determined by default heuristic based on size and number of torsions.
- Check boxes for Sampling Rings, Systematic Search, Cartesian Refinement

(http://www.molsoft.com/man/reals.html#vicinity),Sample Cis and Trans, sample Pyramidal and Verbose (Display Warnings). The 'Cartesian' flag enables cartesian minimization/refinement of conformations (produced by torsion-only sampling). This should be checked whenever energetics is of importance. Torsion-only sampling is fast but only gives qualitative sample of accesible conformational space, relative energies of different conformers may not be accurate.

- Click OK and the sampling will be undertaken in the background see Windows/Background Jobs
- Once the sampling has finished a table as shown below will be displayed. Click in the toggle boxes in the column labeled "L". To display more than one 3D structure at a time use the row selection options.

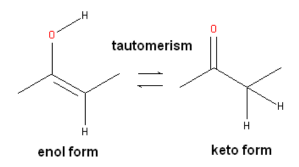


# 7.21 Generate Tautomers

Available in the following product(s): ICM-Chemist-Pro | ICM-VLS

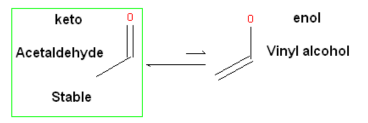
#### Theory

Tautomers are formed by an interconvertible reaction called tautomerization whereby there is a formal migration of a hydrogen atom along with a switch of a single bond and an adjacent double bond. A common example is the keto to enol tautomerism:



During tautomerization a chemical equilibrium of the tautomers will be reached based on several factors, including, pH, temperature and solvent. Tautomerizations are catalyzed by: bases (deprotonation, formation of a delocalized anion, and, protonation at a different position of the anion; and acids (protonation, formation of a delocalized cation, and deprotonation at a different position adjacent to the cation).

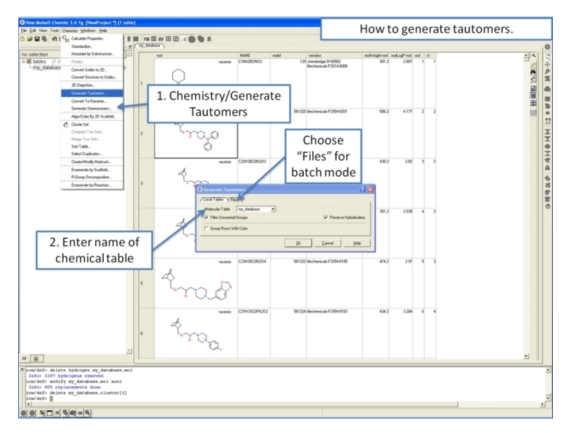
**ICM** will only generate energetically favorable tautomers. Generally tautomers that have a change in hybridization state are less stable and so ICM will not generate these thus reducing the number of scaffolds generated. For example the keto form shown below is more stable by ~14 kcal.mol than the enol therefore ICM will not generate the enol form.



The tautomer generation MoldHf (heat of formation) was trained on the data from http://webbook.nist.gov(e.g. http://webbook.nist.gov/cgi/cbook.cgi?ID=000050-00-0&Units=SI&cTG=on). We used PLS regression and our custom fingerprints to train the model.

### To generate tautomeric conformations of your compound:

- Select the compound(s) in the molecular table. Selected compounds are highlighted in blue.
- Right click in the table and select the option Chemistry/ Enumerate Tautomers. Or select the Chemistry menu/Generate Tautomers.
- If you select Chemistry/Generate Tautomers a dialog box will be displayed. There is also an option to run in batch mode (click the **Files** Tab).
- Choose the table containing the compounds using the drop-down list.
- **Filter Unwanted Groups** option will filter results from patterns in the TAUTOFILTER.tab file provided in the distribution. If results match any row from that table then the it will be excluded.
- **Preserve Hybridisation** Although generally a change in hybridisation state will generate less stable compounds in some cases this is not the case and so you can choose to change hybridisation for a single atom.
- **Group Rows With Color** option will color tautomers from the same compound with the same color to visually highlight groups.
- The compounds will be displayed in a separate molecular table called TAUTOMERS.



## 7.22 Generate Stereoisomers

Available in the following product(s): ICM-Chemist-Pro | ICM-VLS

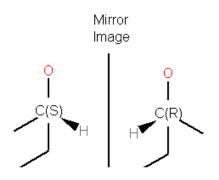
### Theory

Isomers are molecules which have the same chemical formula and sometimes the same kind of bonds but in which the atoms are arranged differently.

Structural isomers have different atom-to-atom connections e.g. propanol (C3H8O or C3H7OH) has two isomers Propan-1-ol and Propan-2-ol.

Diastereomers are not mirror images and have different internal dimensions (e.g. dihedral angles and distances between non-bonded atoms). They can be configurational diastereomers (which can be interconverted only by breaking bonds or by changing the configurations of stereocenters) or conformational diastereomers (which can be interconverted by rotation about bonds - including chair flips or by lone pair inversion .

Enantiomers have identical internal dimensions, and are nonsuperimposable mirror images. Enantiomers can be configurational and conformational.



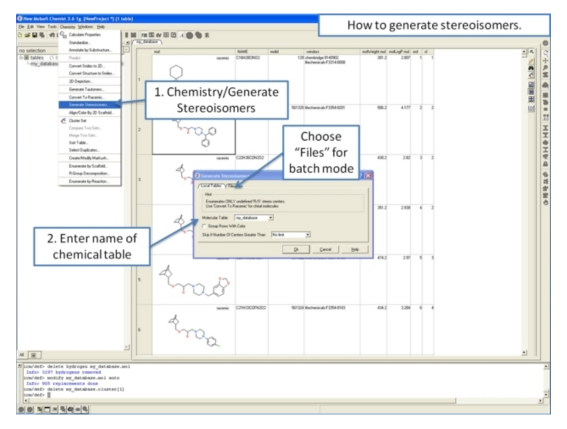
Enatiomers are distinguished based on the Latin terms for left (sinister) and right (rectus). In some cases where the handedness is unknown a chiral center can be labeled "RS" or unknown.

# To enumerate and display in a separate table the stereoisomers of selected compounds.

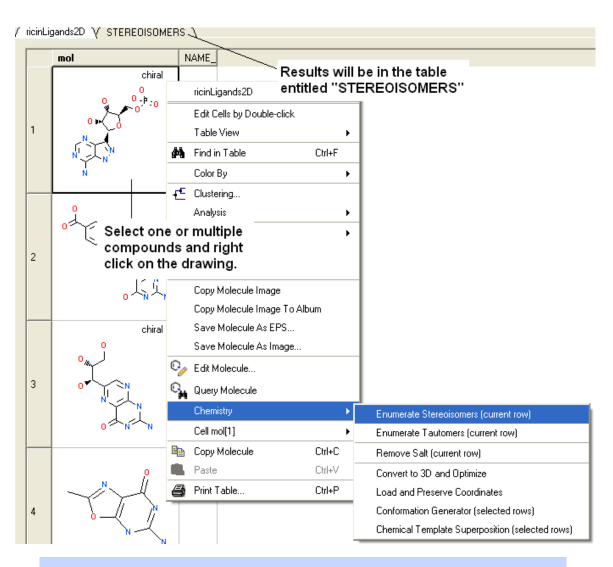
- Select the compound(s) in the molecular table. Selected compounds are highlighted in blue.
- Right click in the table and select the option **Chemistry**/ **Enumerate stereoisomers**. Or select the Chemistry menu and choose **Generate Stereoisomers**. If you generate stereoisomers via the Chemistry menu you will get a dialog box whereby you can run the process in batch mode. There is also an option to color stereoisomers from the same compound with the same color.
- Group Rows With Color option will color stereoisomers from the same compound with the same color to visually higlight them.
- Skip if Number of Centers Greater than option can be used to filter out compounds which have a defined number of stereo centers. If you do not wish to filter just use the default No limit option.
- The compounds will be displayed in a separate molecular table called STEREOISOMERS.

**NOTE:** Stereoisomer generation will keep pre-defined stereo centers and only enumerate racemic ones.

### From the Chemistry Menu:



From a chemical spreadsheet:



NOTE: Only centers with unknown chirality will be enumerated.

# 7.23 Prodrug

To generate a drug from a prodrug:

- Read in a chemical table (sdf file) into ICM.
- Choose the menu Chemistry/Generate Drug from Prodrug and a dialog box will be displayed.
- Enter the name of the Chemical Table.
- Select the reactions you wish to use to generate the drug. You can choose "Prodrug", "All", "Hydrolysis" "Reduction". You can choose the option "all" to use all the available reactions. You can see the reactions in each subset by clicking on the tab labeled "Load Reaction Panel". A chemical table containing the reactions will be displayed and the reaction is lebeled in a column called "subset".
  You can choose to output the results in the "in-addition" mode
- You can choose to output the results in the "in-addition" mode where the converted drug will be in the row below the prodrug or "replace original" whereby the prodrug will be replaced with the converted drug.

To generate a prodrug from a drug:

• Read in a chemical table (sdf file) into ICM.

- Choose the menu Chemistry/Generate Prodrug from drug and a dialog box will be displayed.
- Enter the name of the Chemical Table.
- Select the reactions you wish to use to generate the prodrug. You can choose "carboxylate", "hydroxyl", "amino", "diol", "sulfide". You can choose the option "all" to use all the available reactions. You can see the reactions in each subset by clicking on the tab labeled "Load Prodrug Panel". A chemical table containing the reactions will be displayed and the reaction is lebeled in a column called "subset".
- You can choose to output the results in the "in-addition" mode where the converted prodrug will be in the row below the drug or "replace original" whereby the drug will be replaced with the converted prodrug.

# 7.24 Ligand Energetics

Available in the following product(s): ICM-Chemist-Pro | ICM-VLS

### 7.24.1 Conformational Entropy

Entropy of the conformational ensemble is calculated according to the regular thermodynamic formula i.e.  $kT \log(Sum(Exp(-Ei/kT)))$ , Ei is the energy of the i-th conformation over the ground state, summation is over all conformations.

To calculate the conformational entropy of chemicals in an SDF file or a chemical spreadsheet:

- Chemistry/Ligand Energetics/Conformational Entropy
- Choose either the **Local Tables** (loaded chemical spreadsheet) or {Files} (SDF file) tab.
- Enter the maximum number of conformations you wish to generate.
- Enter a **vicinity** value. For more information on vicinity please see the command language manual
  - http://www.molsoft.com/man/reals.html#vicinity
- Enter a **effort** value. This relates to the length of sampling time.
- Check boxes for Sampling Rings or Systematic Search (instead of Monte Carlo).
- Click OK.

The calculation will then run and you will see a progress bar in the bottom right hand corner of the GUI or you can check the progress of the job go to **Windows/Background**. When the job has finished:

• A table will be displayed called **entropies** with a column called **Conf Entropy**. The energy is in kcal/mol.

### 7.24.2 Strain

To calculate the conformational strain of chemicals in an SDF file or a chemical spreadsheet:

- Chemistry/Ligand Energetics/Strain
- Choose either the Local Tables (loaded chemical spreadsheet) or {Files} (SDF file) tab.
- Enter a effort value. This relates to the length of sampling time.
- Check the **Systematic Search** option if you do not want to use Monte Carlo search.

The calculation will then run and you will see a progress bar in the bottom right hand corner of the GUI or you can check the progress of the job go to Windows/Background. When the job has finished:

• A table will be displayed called **strain** with a column called **Strain**. The energy is in kcal/mol.

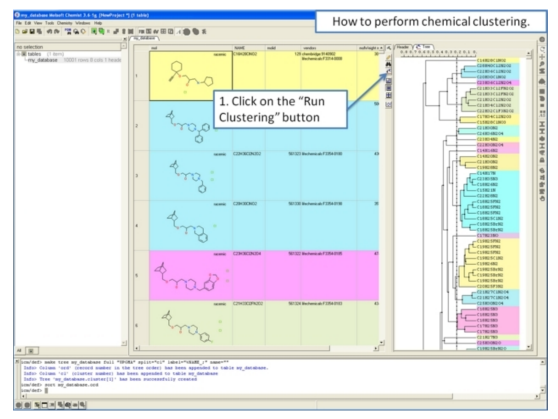
# 7.25 Cluster Set

Available in the following product(s): <code>ICM-Chemist | ICM-Chemist-Pro | ICM-VLS</code>

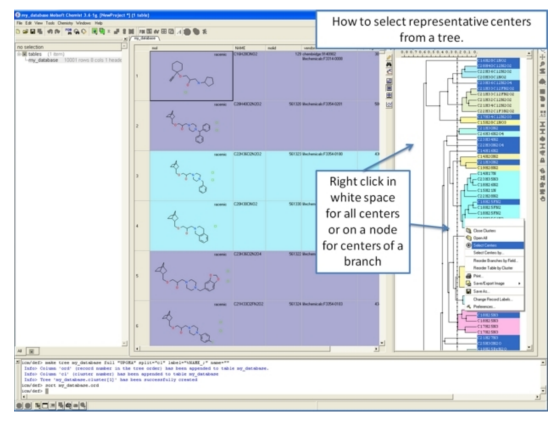
Clustering is described in more detail in the Tables Clustering section of this manual. To undertake chemical clustering choose:

• Chemistry/Cluster Set

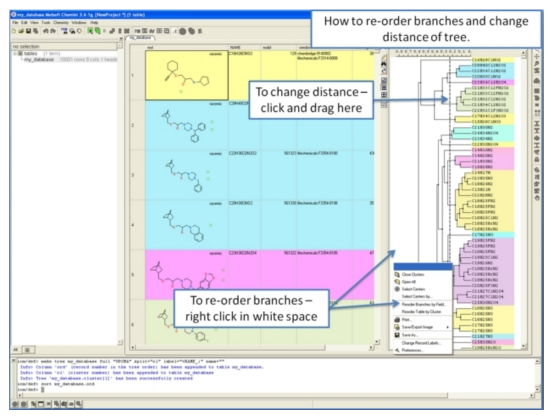
### 7.25.1 How to perform chemical clustering.



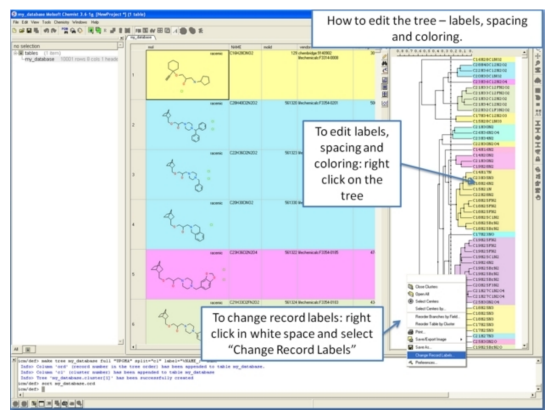
# 7.25.2 How to select representative centers from a tree.



# 7.25.3 How to reorder branches and change the distance of trees.



#### 7.25.4 How to edit the tree - labels, spacing and coloring.



## 7.25.5 Auto Close Tree Mode

To make working with large cluster trees a little easier you can activate the auto close mode which will close downstream clusters and make them more compact.

To do this:

- Right click in white space on the cluster and choose Preferences.
- Select "Auto Close Clusters Downstream".

#### 7.25.6 Calculate Maximum Common Substructure for Nodes

To calculate the Maximum Common Substructure (MCS) for nodes in a treee:

- Chemistry/Cluster Set
- Select the option Calculate Maximum Common Substructure for Nodes
- Hover over a node to view the MCS.
  Right click on a node and choose "Extract Maximum Common Substructure" to extract it to a table.

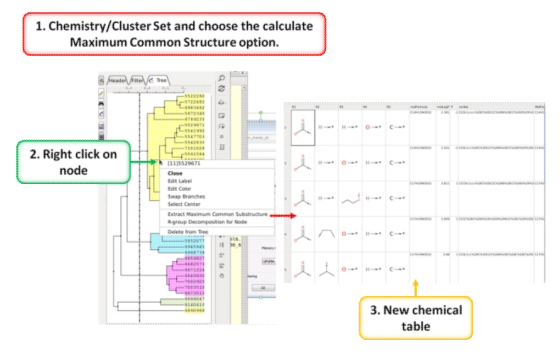
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East the existing trees	node to view	LARREINC
2 Add column with cluster numbers, name: d		R5
Calculate Maximum Common Substructure for Nodes	MCS	
Build for the selected © rows (at least 2 required)		Ri Ly Rydyns
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Color Column only (Ingerprints)     Column only (Ingerprints)     Method:     Tree Memory required: 0 MB     Parameters:     Linkage type:     Linkage		554016 5564016 7471392 5852077 6945643 6998738 6659827
Compressed herarchical dustering     Compressed herarchical dustering		5572455 6072214 5564016 7471392 5852077 5945643 6968738
Coly use existing distance     the "isof" column only (Ingerprints)     the mol column only (20 pharmacophore)  Method:     Tree     Memory required: 0 MB     Parameters:     Linkage type:     Linkage type:     Linkage distance matrix		5572455 6072214 5564016 7471392 5852077 6945645 6659827 6659827 6659827 6659827 6659827 6659827 6659827
Compressed hierarchical dustering     Compressed hierarchical dustering		5572455 6072214 5564016 7471392 5852077 5945643 6968738 6659827 66659827 66659827 66673224

#### 7.25.7 R Group Decomposition from Clustering Node

To perform R-group decomposition directly from a cluster node:

- Chemistry/Cluster Set
  Select the option Calculate Maximum Common Substructure for Nodes.
  Hover over a node to view the MCS.
  Right click on a node and choose "R-group decomposition for node".

The resulting table can be used for R-Group Enumeration or SAR.



## 7.26 PCA Analysis

```
Available in the following product(s): <code>ICM-Chemist|ICM-Chemist-Pro|ICM-VLS</code>
```

PCA analysis is described in the tables section here.

## 7.27 Visualize Chemical Space

There are two options to visualize chemical space:

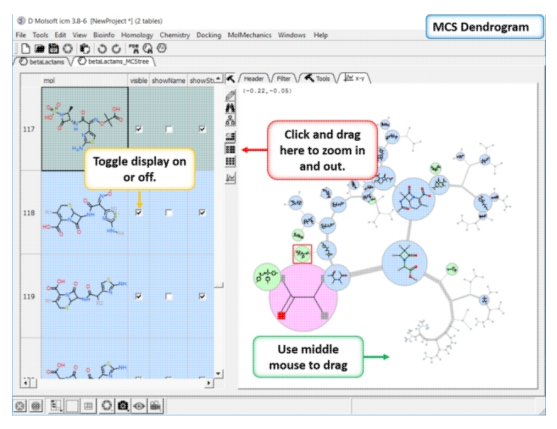
- Maximum Common Substructure (MCS) Dendrogram
- Self Organized Network

#### 7.27.1 Maximum Common Substructure (MCS) Dendrogram

The MCS method is K-greedy algorithm which simultaneously grows K different common substructures applying scoring and resorting on each step.

To display the interactive MCS dendrogram:

- Read in a chemical spreadsheet.
- Chemistry/Visualize Chemical Space/MCS Dendrogram
- Enter the name of the table you wish to analyze.
- Enter the name of the column you wish to use to label the points in the dendrogram.
- Choose **Optimize MCS** to minimize overlapping by branches in the tree.
- Click OK and the dendrogram will be displayed on the right hand side panel.



You can display and undisplay the points and labels by using the check boxes in the columns "visible", "showName" and "showStr". You can right click on the column and choose (un)check all. Points in the dendrogram are fully interactive and linked to the table. You can zoom and translate the plot around the space. The points are colored by > blue = common substructure, green = original compounds pink = root (most common)

## 7.27.2 Self Organized Map

A Self Organized Map (SOM) is a way to represent higher dimensional data in 2D (or 3D) such that similar data is grouped together. Nodes in 2D map are called neurons. Each neuron is assigned a weight vector of the same dimensionality as input space - you can read more here.

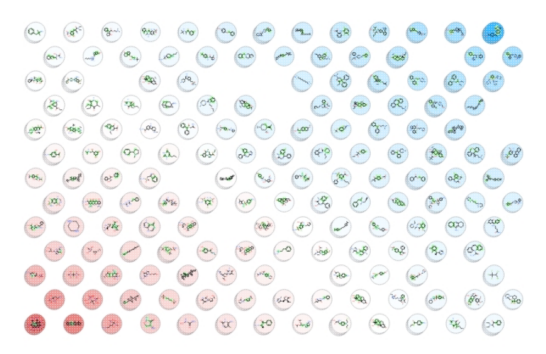
In ICM the initial placement can be either 1). random or 2). find four mutually remote points in the input dataset and put in the corners and uniformly distribute the rest. The training is undertaken in the following way: a vector is chosen at random from the training set, then ICM finds the node with the closest distance to the chosen vector. Next the radius of the Best Matching Unit (BMU) neighborhood is calculated and then the weights of the BMU neighborhood nodes are adjusted - this is repeated for N iterations. The final mapping stage assigns input vectors to their closest nodes.

To run the Self Orgnanized Map method in ICM:

- Read a sdf file into ICM.
- Chemistry/Visualize Chemical Space/ Self Organized Map
- Enter the name of the chemical table.
- You can choose to color the nodes by a particular column value and adjust the coloring by the Aggregate method (e.g. all members of the cluster will be colored by average, min or max).
- You can choose to select one or more columns of data to use as Descriptors.

- The map dimensions can be changed from 'auto' by adding an integer e.g. 4 will make a map size of 4x4
- If you do not check randomize the method will use the four mutually remote points in the input dataset and all other initial points will be generated so they spread uniformly between these four.
- Connectivity edges option is for visualization purposes only to show 'close' clusters.

A map containing the nodes be displayed as shown below. Each node contains a stacked cluster of input compounds. Sequential clicking on the node loops through the representatives.



## 7.28 Compare Two Sets

Available in the following product(s): <code>ICM-Chemist | ICM-Chemist-Pro | ICM-VLS</code>

#### To compare two chemical tables for similar compounds:

- Read the two tables into ICM.
- Chemistry/Compare Two Sets...
- Select the first table from the drop-down list and then select the second table.
- Choose whether you want to use **exact** or **similarity** comparison. If the **similarity** option is selected a **Distance** value needs to be entered. The distance value should be decreased below 0.5 (0.2 or 0.3 or even smaller if you want to find "strong" similarity differences. The distance relates to the Tanimoto distance as described here
- Choose whether you want to **Select Matching Records** or **Show Only Different Records**.
- Select OK
- Similar compounds will be highlighted blue (selected). A selection can be transferred to a new table by right-clicking on the table and select **Copy Selection to ICM Table**.

📀 Compare Two Sets		? ×						
First Table conformers	<ul> <li>Second Table</li> </ul>	conformers						
💿 exact 🔘 similarity	Distance	0.5 👻						
Select Matching Records Show	Select Matching Records Show Only Different Records							
Hint Double-click on the tab of the second tab	ble to see them side by sid	de						
	Ok ok	Cancel Help						

## 7.29 Merge Two Sets

Available in the following product(s): <code>ICM-Chemist | ICM-Chemist-Pro | ICM-VLS</code>

#### To merge two tables:

- Read the two tables into ICM.
- Chemistry/Merge Two Sets
- Select the first table from the drop down list (Table A) and the column you wish to use to merge the table by.
- Select merge method 1. **inner** only molecules present in BOTH A and B tables are kept; or 2. **left** ALL rows of A are kept; or 3. **right** ALL rows of B are kept.
- Select the second table from the drop down list (Table B) and the column you wish to use to merge the table by.
- Enter a name for the output table.
- Click OK and a new table will be displayed.

🧐 Merge Two	Sets		? 🔀
Table A	ricinLigands2D_tauto_1_tauto	by Column	mol 💌
inner	Cleft Cright		
Table B	ricinLigands2D_tauto	by Column	mol 💌
Result Name	T_join 💌		
left - ALL rou	molecules present in BOTH A and B tables are ke ws of A are kept ows of B are kept	pt	
	Ok	Cancel	Help

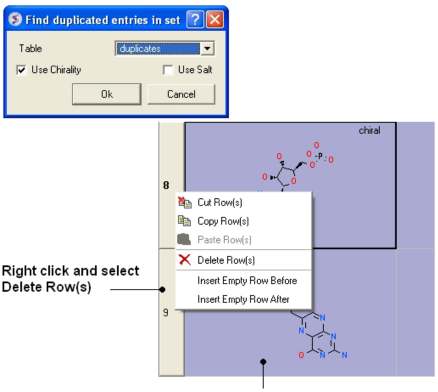
## 7.30 Select Duplicates

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

**NOTE:** Gui option is available in versions 3.5-10 and higher. The command line options for this function are described in the ICM Command Language manual at http://www.molsoft.com/man/icm-functions.html#Index-chemical

This option allows you to select and remove duplicate chemicals in a table.

- Read a chemical table into ICM.
- Chemistry/Select Duplicates
- Enter the table name you want to check for duplicates
- Enter whether you want chirality or the salts included in the analysis.
- Press OK
- Duplicate compounds will be **highlighted** blue in the table. You can delete them by right clicking on the row header ans selecing **Delete Rows(s)**



Duplicate rows are highlighted in blue

## 7.31 Multi Parameter Optimization

Multi Parameter Optimzation (MPO) is a method that can be used to derive a score for the relative importance of a number of different chemical properties. For example Lipinski Rules (LopP, MW, Nof(HBD), Nof(HBA) and CNS (LogP, PSA, MW, pKa, MW, Nof(HBD).

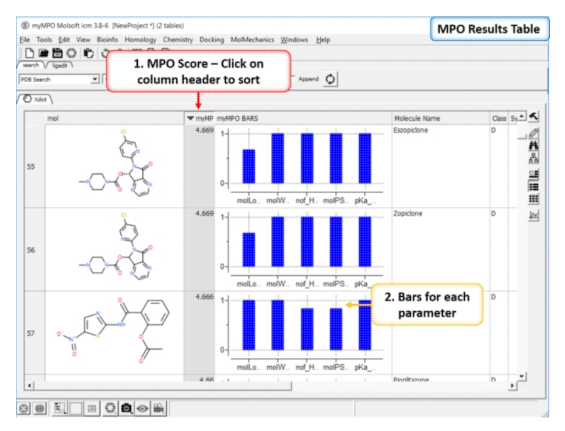
To run MPO:

- Read in a Chemical table.
- Chemistry/MPO/Create New MPO Score
- Select the inbuilt Lipinski or CNS parameters or make your own by choosing Empty.
- Choose whether you want to assign property colors and click OK.
- You will then see a panel on the right hand side of the spreadsheet under the table Tools tab.

- If you chose "Empty" you will need to add parameters either from MolSoft's inbuilt functions or from an existing column.
  If needed Adjust the High, Low, Slope, Value and Weight columns by clicking in the cell and editing.

in V igedit ) isearch	00	n va						
V CRMyr	• bad	1.	Chem	istry	/ MPC	>	Felds T Append	
Paramete 1 motoge 2 mol/vegh 3 nof_HBD 4 moPSA 5 pKa_mb	0 t ND 30 ND <b>3. Ad</b>	450 2 100 9	50 1.5 10 1 1		Weight		Header / Filer / Tools Edt Deta Table Xdet Add Parameter From Existing Column Function motiveight Add Parameter From Function Apply Input Table Xdet Calculate MPO Score Calculate Score	Column
							•1	

- Click on Calculate MPO Score and a results table will be displayed containing a Score and the parameter bars.
  Sort the Score column (myMPO).



## 7.32 Combinatorial Chemistry

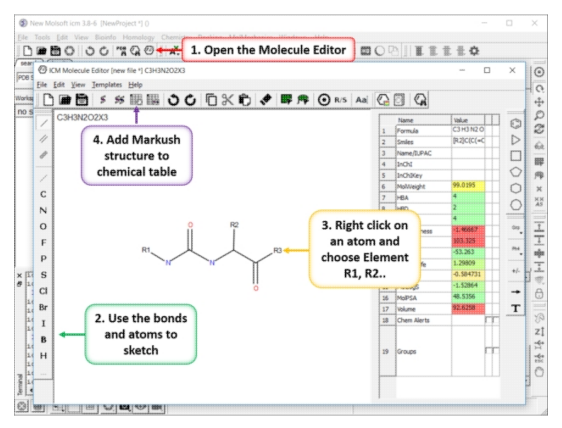
Available in the following product(s):  $\texttt{ICM-Chemist} \mid \texttt{ICM-Chemist-Pro} \mid \texttt{ICM-VLS}$ 

## 7.32.1 Sketch Markush Structure

Tutorial | Video

#### To create or modify a Markush Structure:

• Use the Molecular Editor to sketch the scaffold as shown below. Right click on an atom and choose -> Element-> R1,R2....



- Save the sketch to a chemical table by clicking on the "Append Molecule to Chemical Table" button in the Molecule Editor.
- Close the Molecular Editor window by clicking on the cross in the top right hand corner and the changes will be submitted to the table.

## 7.32.2 R-Groups

ICM has an inbuilt local database of R-groups generated from ChEMBL data. To open this file:

- Chemistry/Combinatorial Chemistry/ Load R Group Database
- A table of R-groups will be displayed attachment points are represented by asterisk. The 'freq' column represents the number of pairs found in ChEMBL for that scaffold.

You can sketch your own substituents:

- Open the Molecular Editor.
- Sketch the substituent and use > right click > attachment point to assign the attachment point.

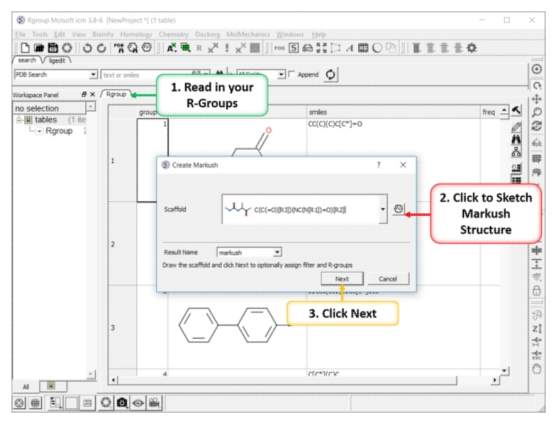
You can also split a chemical table into fragments as described here.

You can read in an SDF, mol, smiles file containing fragments. If you do not want the first atom of the substituents to be the attachment point you need to define the attachment point. Attachment points are automatically assigned when you extract fragments or you can define them manually by using the molecular editor (as described above).

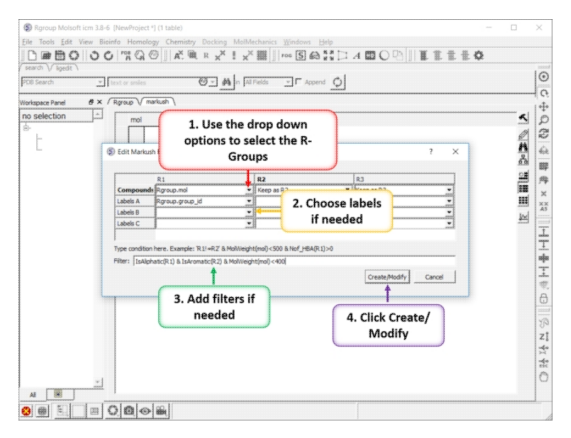
In many combinatorial chemistry options there is an option to use LigEdit Mofifiers. These are the substituent groups you see in the edit panel in the 3D Ligand editor.

## 7.32.3 Create Markush

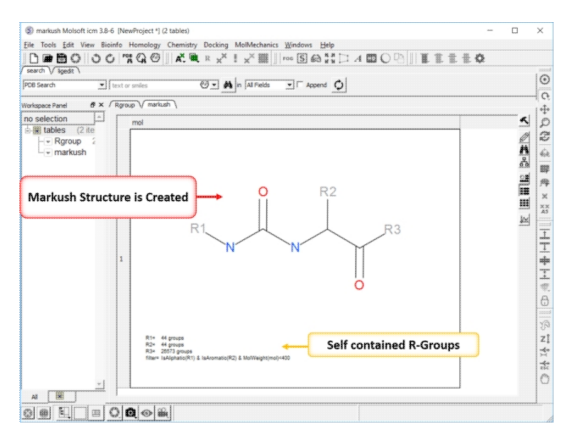
#### To create a Markush structure:



- Read in the R-groups for your Markush structure.
  Chemistry/Combinatorial Chemistry/Create Markush
  Click on the editor button and sketch the Markush structure.



- Use the drop down arrow to select a R-group table for each of your R-groups. You can choose to associate labels from the R-group table if
- needed.
- You can add filters if needed (e.g. IsAliphatic(R1) & • IsAromatic(R2) & MolWeight(mol)Click Create/Modify.



- The Markush structure will then be created with embedded R-groups and filters.
- You can extract the R-groups again to separate tables by: Right click on the Markush structure in the table and choosing the option "Extract Markush/Scaffold R-groups".

**NOTE:** Once a Markush is created you can right click on it in the chemical table you will see two convenient options 1. Edit/Markush Scaffold Properties and 2. Extract Markush Scaffold R-groups.

## 7.32.4 Enumerate by Scaffold

#### Tutorial | Video

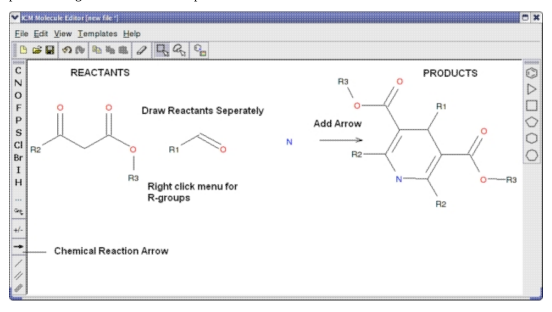
#### To enumerate a library based on R-groups:

- Sketch the Markush structure and save it in a chemical spreadsheet.
- Read in the R-Group substituents you wish to use to enumerate the library.
- Right click on the structure and select Chemistry/Enumerate R-groups or use the Chemistry/Combinatorial Chemistry/ Enumerate by Scaffold-Markush menu. If you use the menu option you will need to choose the table containing the scaffold from the drop down list of currently loaded tables. The index number refers to the row number in the scaffold table. Click OK.
- Select the R1, R2... table , labels and filters if necesary. As an example of a filter you could use R1 !=R2, OR MolWeightA new table will be produced called **T\_enum** with the Template structure highlighted in red.

### 7.32.5 Enumerate by Reaction

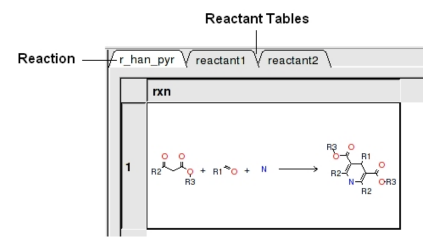
#### Additional Resources: Tutorial | Video | Command Line Docs

Reactions can be drawn using the ICM Molecular Editor. Reactants should be drawn side-by-side (no + sign is necesary) and separated from the product using the arrow. See example shown below:



This example is available in the ICM distribution as example\_reaction1.icb. The reaction is the Hantzsch Dihydropyridine (Pyridine) Synthesis. This reaction allows the preparation of dihydropyridine derivatives by condensation of an aldehyde with two equivalents of a beta-ketoester in the presence of ammonia. Subsequent oxidation (or dehydrogenation) gives pyridine-3,5-dicarboxylates, which may also be decarboxylated to yield the corresponding pyridines.

In this example we have two reactants therefore it is necessary to have two reactant substructure tables loaded into ICM. ICM will match the substructure drawn in the reaction with the chemicals in thereactant table.



reactant 1 table:

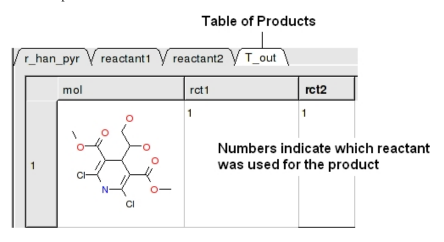
ſŗ	r_han_pyr V reactant1 V reactant2 \						
г			Substruc	ture search: Found 88 h	nits of 'O=C([C;D2]C(=O)O[R3])[		
		molid	MolW	mol	vendors		
	18	65236	170.094		asdi:500028335		
	19	87804	174.053		apolloscientific:12582 asdi:500014701		
	20	65360	182.058		apolloscientific:13718 keyorganics:11X-0925		
	21	87822	179.058		apolloscientific:2965 interchim:616		
	22	3341	194.058		asdi:500016383		

#### To apply a reaction:

- Chemistry/Enumerate by Reaction.
- In this example (example\_reaction1.icb) we already have the reaction drawn in a chemical table. Therefore select the **Choose Table With Reaction**. If you would like to draw a new reaction select **Draw New Reaction**.
- Enter the name of the table containing the reaction. If you have more than one reaction drawn you can select the row using the index option.
- Click OK and then you will be asked to enter the Reactants. Select the reactant tables from the drop down arrow for Reactant 1 and Reactant 2.

- You can transfer information to the reactant table by selecting columns in the **Labels** section.
- Unused reactants can be marked.
- Select what you want to do with multiple matches.

A table of Products will be then displayed in a table called T\_out. Columns in T\_out labeled "rct" display which reactants were used to build the product.



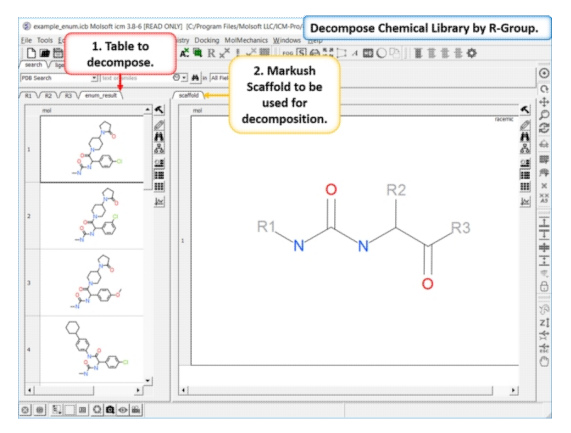
## 7.33 SAR Analysis

## 7.33.1 R-Group Decomposition

#### Tutorial | Video

To decompose a library into fragments based on a Markush scaffold (opposite of R-group (Markush) enumeration):

- Read the sdf file you wish to decompose into ICM and it will be displayed as a molecular table.
- Sketch the Markush structure you wish to use to decompose the table by and save it in a chemical table.



- Chemistry/SAR Analysis/R-Group Decomposition.
- Choose the table containing the Markush scaffold. Index refers to table row if you have more than one Markush structure in a table.
- Use the drop-down option to select the table you wish to decompose and the column containing the 2D chemical (usually called mol).
- If you have more than one R-group ICM can either generate a different table for each R-group or it can merge it into one single table whereby column will represent R1 and column two R2... This option is useful if you want to generate a SAR table with a column of activity data next to the R1 and R2 columns.
- If you check the box "Auto Add Missing R Groups" then unique R-groups will be extracted from the scaffold where hydrogens can be attached.

Sexample_enum.icb Molsoft icm 3.8-6 [READ ONLY Eile Tools Edit View Bioinfo Homology Chemi		hemistry/SAR	Analysis/R-G	roup Decom	position
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	1 R-Group Decompo	sition	R2	R3 ? ×	
<sup>3</sup> 2. Scaffold table	Scaffold	T	Index	1	é
			Auma mol Missing R-Groups P Add Ck Cancel 4. Click OK.	Original Compounds	
		1			<u> </u>

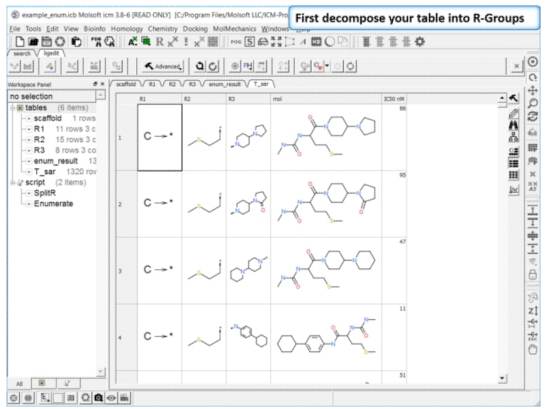
• The decomposed table will be reported in a new table with new columns for the R groups (R1,R2...). If you have an additional column with activity data you can sort the table by R-groups and see the effect of an R-Group on activity. This table can be used to build an additional SAR table as described here.

©e jile ∏n	Tools Edit	View Bioinfo	Homology Ch	NLY] [C:/Program Files/Molsoft LLC emistry Docking MolMechanics 	Windows	Help		mposition Results Table. 軍軍士士の
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		~			809	801	103	5

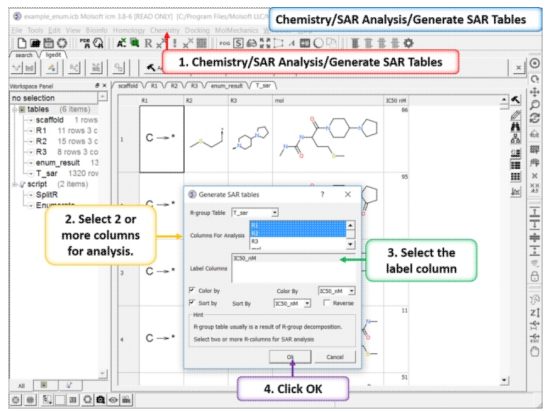
## 7.33.2 SAR Table

|Tutorial|

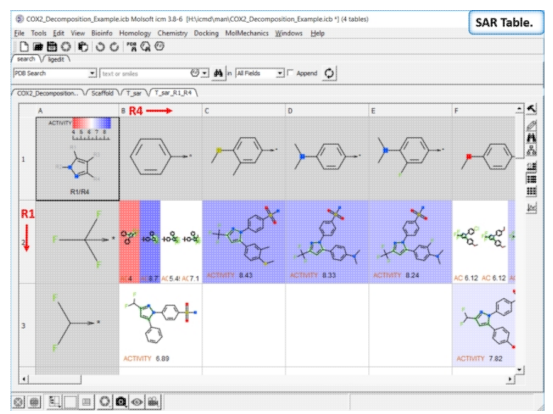
To generate a SAR table you first need to decompose the table into the R-Groups.



**Decompose your library.** Decompose the library to extract the R-Groups or read in a decomposed SDF file.



**Generate SAR table.** Chemistry/SAR Analysis/Generate SAR Tables. One or more columns can be chosen for analysis and multiple tables will be generated. Color the table by a selected column e.g. activity.

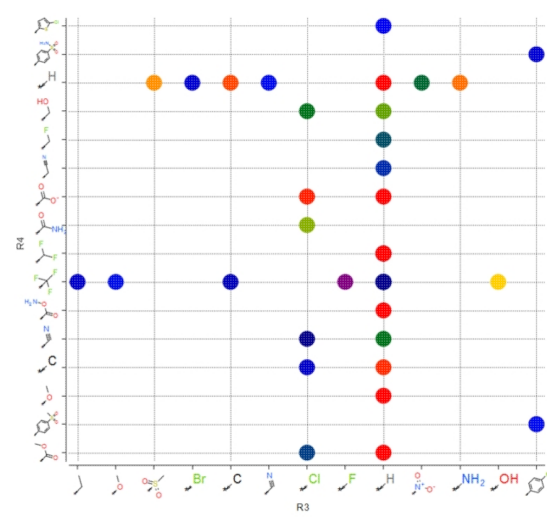


**SAR table.** The SAR table will be displayed with an activity scale and R-group scaffold in the first cell (A1) and the R groups along each

axis. You can use the table to investigate which R-group(s) contribute to activity.

## 7.33.3 Plot R-Groups

You can make a plot of R-groups from library decomposition. This is described in the plots section of the manual.



## 7.33.4 Generate SALI Table

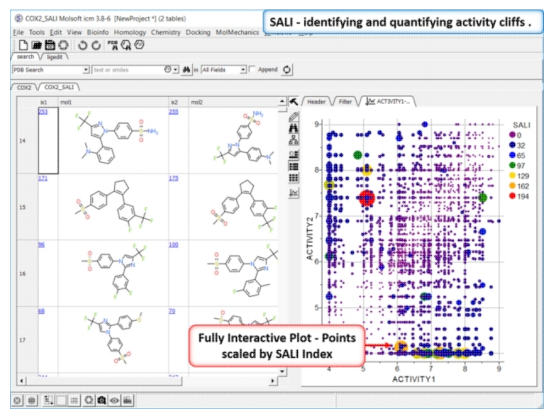
#### |Tutorial|

#### Structure--activity landscape index: identifying and quantifying activity cliffs.

This method produces a SALI score as described in the publications by John Van Drie and co-workers. It will identify "structure-activity cliffs": pairs of molecules which are most similar but have the largest change in activity.

- Read in a sdf file containing "mol" and "activity" column.Chemistry/SAR Analysis/Generate SALI table.
- •
- You can choose to use the Log of the Activity Column. The method will try to find pairs based on defined distance cutoff • (0.15 is a good threshold for larger datasets).

• A results table will be reported containing the pairs, tanimoto fingerprint similarity and SALI index.



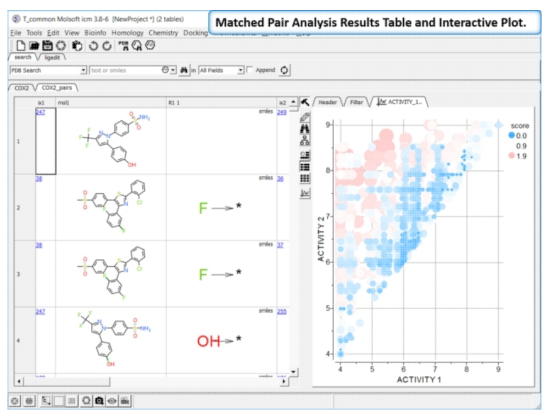
### 7.33.5 Matched Pair Analysis

#### |Tutorial|

Matched Pair Analysis (MPA)can be used to study changes in chemical properties based on small well-defined structural modifications to the chemical structure. The method tries to find the Maximum Common Substructure and clusters by similarity. The analysis is performed in two stages: clustering and then Maximum common substructure on each pair in each cluster. The output is similar to SALI output plus it gives the actual R-group pair variation. The pair table is sorted by score so top hits should show small variations and big score difference. To run MPA:

	T_common Molsoft icm 3.8-6 [NewProject *] (2 tables) Elle Tools Edit View Bioinfo Homology Chemistry Docking MolMechanics			Chemistry/SAR Analysis/Matched Pair Analysis		
			y Docking MolMechanics W			
		10				
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C00X2 \	CON2_pairs \					
	mpi	Molecule	ACTIVITY SET Min dist Avg di	and and a second s		
1	20G	1-3	7.22 1 999.9 995 Table CCO2 CO2 CO32 Cost Table CCO2 CO32 Cost Table CCO2	ysis ? × wityColumn ACTIVITY •		
2	10 f	1-10	Р-942 Р-54 Р Г-073	12 31 -14-2 F -0H F -F -0 F -8r F -1		
3	10 B	1-16	Custom Filter   Filter Chirality Hert Use Custom Filter field	for additional groups Ok. Cancel		
4	Sold	1-17	4 3 0.19 0.	-1		
0.0						
- m 1 - m 1				A		

- Read in a chemical spreadsheet containing a 'mol' column and activity data.
- Chemistry/SAR Analysis/Matched Pair Analysis.
- Enter the name of the Input Table.
- If the table has activity data or some other property use the option use Activity and select the column name from the drop down button.
- Enter a cluster split distance for determining similarity.
- The size of the R-groups for comparison can be set (default is 12 atoms).
- R-groups can be filtered from the analysis. or a custome filter using SMARTs.
- Chirality can be ignored in the comparison if needed.



The results table (tableName\_pairs) is fully interactive between the plot and the original table. The hits are sorted by the score column, so top hits should show biggest activity change with smaller groups. The plot is displayed activity 1 versus activity 2 colored and sized by score (red is better).

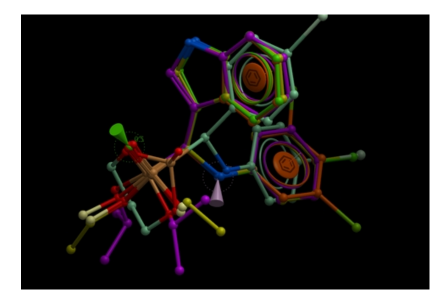
- **ix1** index number (row number) of first chemical which is hyperlinked to original table.
- **mol1** 2D structure of first chemical.
- **R1\_1** group difference between first (ix1) and second chemical (ix2).
- **ix2** index number (row number) of second chemical which is hyperlinked to original table.
- mol2 2D structure of second chemical.
- **R1\_2** group difference between second (ix2) and first chemical (ix1).
- **scfl** common scaffold
- **dvisible** ('d' delta 'visible' delta of your activity column: 'd' + original\_column\_name)
- **score** is calculated as normalized dActivity + (1 normalized weight of both groups).

## 7.34 Substructure Superposition

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

Chemical superposition can be undertaken in the following ways.

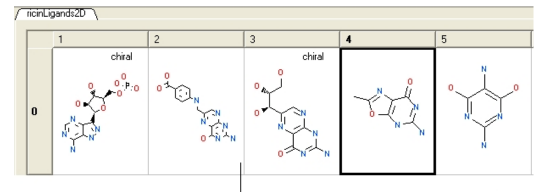
- Rigid Superposition of Compounds in a Table onto a Template in The Graphical Display
- Rigid Substructure Superposition of Chemicals in the Graphical Display
- Flexible Substructure Superposition
- See next section Atomic Property Field Superposition



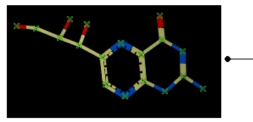
**NOTE:** The substructure superposition requires topologically exact match with the template. As long as there is a large consistent scaffold then the substructure superposition is the best approach. In cases where the structures differ topologically then the APF methods should be used to superimpose moieties which have similar properties.

# 7.34.1 Rigid Superposition of Compounds in a Table onto a Template in The Graphical Display

Here we describe how to superimpose chemicals from an ICM Molecular Table onto a 3D template displayed in the graphical display.



Superimpose from a molecular table onto a 3D template



Select chemical template in 3D display

position of Mompounds in a Table onto a Template in The Graphical Display

- Load the template chemical into the 3D display. It needs to be a non-ICM object. To convert an ICM object into a non-icm object: right click on the name of the object in the ICM workspace and select convert to non-ICM. You should see that the object is now labeled as a model in the ICM workspace.
- Select the chemical template. One way to do this is to double click on the chemical name in the ICM Workspace (selected=blue in ICM Workspace and green crosses in graphical display).
- Select the chemical(s) (row(s)) in an ICM Molecular Table.
- Right click on the table and select Chemistry/Chemical Template Superposition and a table a data entry window as shown below will be displayed.

🦻 3D Superimpose on a t ? 🔀							
template as_g	aph 💌						
thoroughness	. 💌						
🔽 Sample Rings							
Ok	Cancel						

- Enter the name of the template or use as\_graph if you selected the template as described above.
- The **effort** value (formerly 'thoroughness') represents the sampling length. The higher the value the longer the sampling takes.
- Select whether or not you wish the rings to be sampled.
- Click OK and the selected chemicals will be superimposed on the template in the chemical display.

## 7.34.2 Rigid Substructure Superposition

## Here we describe how to perform a rigid-superposition of chemical structures in the graphical display:

- Select the chemicals you wish to superimpose.One way to do this is to double click on the chemical names in the ICM Workspace whilst holding down the control button (selected=blue in ICM Workspace and green crosses in graphical display) or hold the right mouse button and drag over the chemicals in the graphical display.
- Chemistry/Rigid Substructure Superimpose
- A window will be displayed. Enter the name of the template structure using the ICM selection language. The ICM selection language can be found by right clicking on the molecule in the ICM Workspace first line of right click menu.
- Click OK

**NOTE:** Superimposed chemicals can be separated easily using the Arrange as grid option (Tools/Superimpose).

## 7.34.3 Flexible Substructure Superposition

## Here we describe how to perform a flexible-superposition of chemical structures in the graphical display:

- Select the chemicals you wish to superimpose. One way to do this is to double click on the chemical names in the ICM Workspace whilst holding down the control button (selected=blue in ICM Workspa ce and green crosses in graphical display) or hold the right mouse button and drag over the chemicals in the graphical display.
- Chemistry/Flexible Substructure Superimpose

- A window will be displayed. Enter the name of the template structure using the ICM selection language. The ICM selection language can be found by right clicking on the molecule in the ICM Workspace first line of right click menu.
- Click ÔK

**NOTE:** Superimposed chemicals can be separated easily using the Arrange as grid option (Tools/Superimpose).

## 7.35 Atomic Property Field Superposition

Available in the following product(s): ICM-Chemist-Pro | ICM-VLS

**NOTE:** The APF superposition method should be used when there is no common substructure between the chemicals that are being superimposed. If a common substructure is present then the substructure superposition methods described earlier should be used. The APF method will superimpose moieties that have similar 3D pharmacophoric properties.

The Atomic Property Fields (APF) superposition/alignment method was first reported by Maxim Totrov PhD (Principal Scientist - MolSoft) at the 2007 233rd American Chemical Society National Meeting, Chicago, IL USA and then published here.

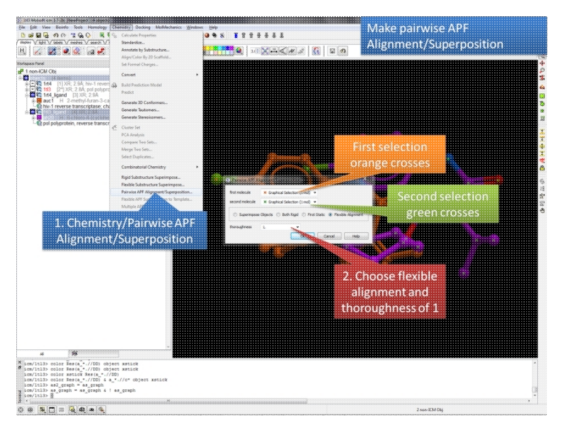
APF is a 3D pharmacophoric potential implemented on a grid. APF can be generated from one or multiple ligands and seven properties are assigned from empiric physico-chemical components (hydrogen bond donors, acceptors, Sp2 hybridization, lipophilicity, size, electropositive/negative and charge). Here we describe template APF

superposition whereby the APF is generated from a single or multiple template and is then globally optimized with the internal force-field energy of the ligand.

## 7.35.1 Pairwise APF Alignment/Superposition

#### This option allows you to superimipose two chemicals by APF.

- Read in two 3D chemical file or convert a chemical sketch to 3D.
- Select the first chemical by double clicking on it in the ICM workspace. It should be highlighted in blue in the ICM workspace and green crosses in the graphical display. Next, copy this selection to orange (as2\_graph)selection to save it your selection should now have orange crosses.
- Select the second chemical by double clicking on it in the ICM workspace. It should be highlighted in blue in the ICM workspace and green crosses in the graphical display.
- Chemistry/Pairwise APF Superposition/Alignment
- Double check that the first molecule and second molecule selections are correct.
- Choose whether you wish to superimpose both chemical by APF, have both chemicals rigid, first static, or flexible alignment.
- The results will output a new object of the superimposed ligands in the ICM workspace.



## 7.35.2 Flexible APF Superposition to Template

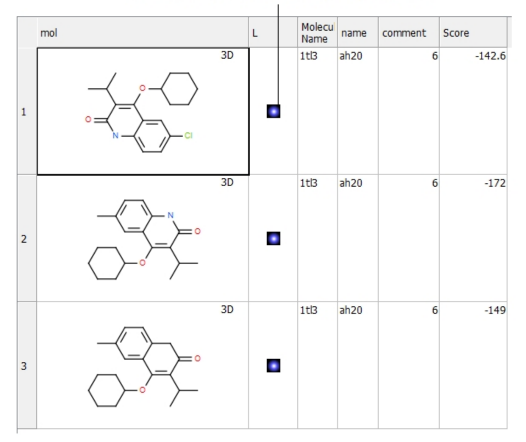
#### To perform Flexible APF Superposition to a template:

- Read a chemical table into ICM containing the compounds you wish to superimpose.
- Display in 3D the template structure you wish to superimpose on. See convert to 3D for instructions on how to generate a 3D template structure.
- Select Chemistry/Flexible APF Superposition to Template and a window as shown below will be displayed.

5 Flexible APF Superposition to Template							
Chemical Table	super_1rt4_1 💌						
template	× Graphical Selection (1 mol) ▼						
thoroughness	1. 💌						
💟 Sample Rings	Sample CisTrans						
Report APF Score Multiple Poses							
Sample Tautomers							
Score Envelope Fit	Score Envelope Fit						
Advanced	Advanced						
Weight Atoms by Oc	cupancy						
Lipophilic weight 1.	•						
Polar weight 1.	▼						
Ok	Cancel Help						

7.35.1 Pairwise APF Alignment/Superposition

- Use the drop-down arrow to select the chemical table containing the chemicals you wish to superimpose on the template.
- \* Select the 3D chemical **template** by double clicking on it in the ICM workspace. It should be highlighted in blue in the ICM workspace and green crosses in the graphical display.
- Enter an effort value. This represents how long the simulation will run for. A value of 1 has been validated as being a suitable length for this kind of superposition.
- Select whether you want flexible rings to be sampled by checking the appropriate box.
- Select whether you want cis and trans conformations of double bonds to be sampled by checking the appropriate box.
- Select whether you want the superposition to be scored in order to rank solutions by checking the appropriate box. About the APF SCORE the lower the score the better the fit with the template. No one score or threshold can be stated as being a "good" score it depends on the chemical types being superimposed. One way to tell whether an APF score is good or bad is to compare it to the score obtained by self superimposing the template.
- Select whether you would like to display multiple poses of the superpositions.
- Select whether you would like to sample tautomers.
- Select the Advanced button if you want the superposition to be weighted by occupancy of the atoms by checking the appropriate box. It is often desirable to preferentially superimpose parts of a ligand while ignoring other regions. This can be achieved by setting the occupancy to zero for regions you are not focusing on. See: command line manual http://www.molsoft.com/man/icm-commands.html#set-occupancy.You can also define the weights for lipophilic and polar matches.
- Click OK and the simulation will run in the background. Once the superposition is complete the molecules will be displayed in the graphical display.



#### Click here to view the chemical superimposed in 3D

# 7.35.3 Multiple APF Alignment of Compounds in a Table

#### To superimpose multiple chemicals in a chemical table by the APF method:

- Read a chemical table into ICM containing the compounds you wish to superimpose.
- Select Chemistry/Multiple APF Alignment and a window as shown below will be displayed.
- Use the drop down arrow to select the chemical table.
- Select the number of iterations for the simulation. This represents how long the simulation will run for. A value of 60 has been validated as being a suitable length for this kind of superposition.
- Click OK and the simulation will run in the background. Once the superposition is complete the molecules will be displayed in the graphical display.

🗐 Multiple APF /	? 🗙	
Chemical Table	result	•
Max. N of Iterations	60	•
	Ok	Cancel

The 3D coordinates can be extracted to a chemical spreadsheet as described here.

## 7.36 APF Tools

Available in the following product(s): ICM-Chemist-Pro | ICM-VLS

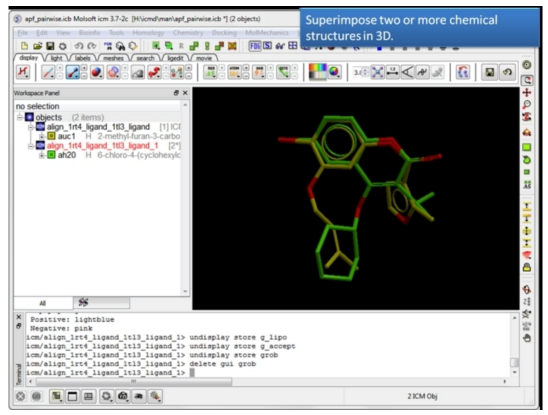
The Atomic Property Fields (APF) superposition/alignment method was first reported by Maxim Totrov PhD (Principal Scientist - MolSoft) at the 2007 233rd American Chemical Society National Meeting, Chicago, IL USA and then published here. This section shows you how to:

- Generate a consensus pharmacophore from the APF fields.
- Calculate a pairwise APF score.
- Perform ligand-based screening using APF fields.

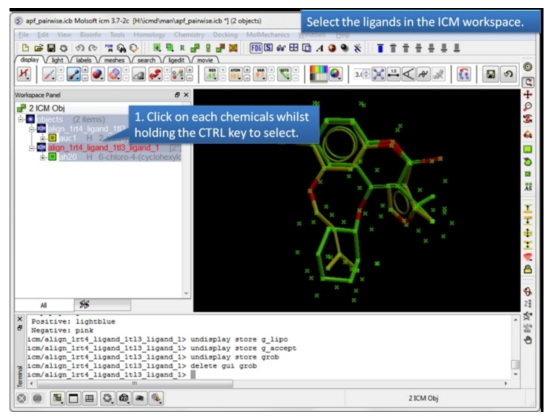
#### 7.36.1 Consensus Pharmacophore

To generate a Consensus Pharmacophore based on APF fields use the Chemistry/APF Tools/Consensus Ph4 menu option.

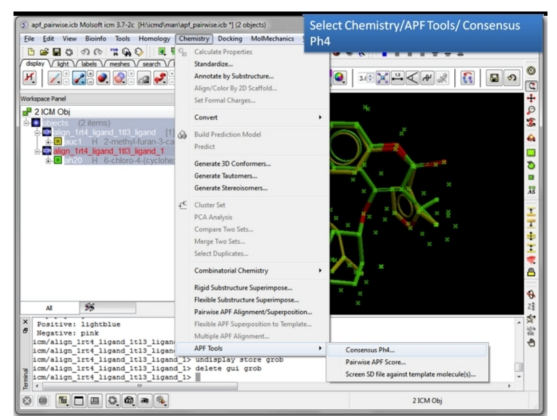
First convert your chemical into an ICM object.



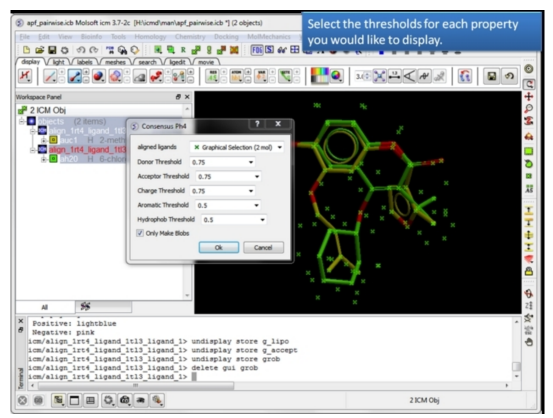
Load and display two or more superimposed chemicals. You can superimpose the chemcials using APF fields as described here or using substructure superposition.



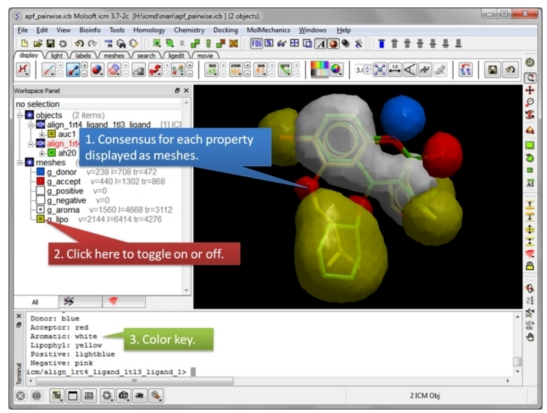
**Select the ligands.** You can select the ligands by clicking on them in the ICM workspace whilst holding down the CTRL key. Aleternative selection modes are described here.



**Choose the Consensus Ph4 menu option.** Chemistry/APF tools/Consensus Ph4.



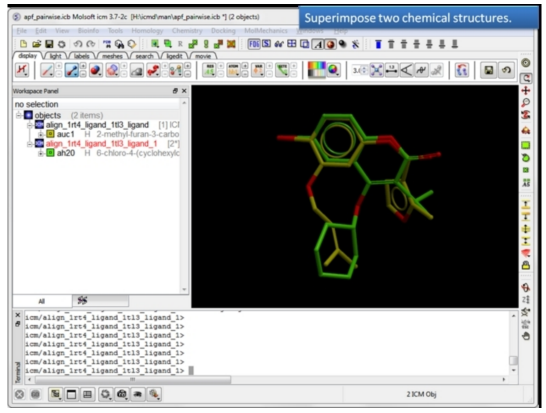
**Select the threshold for each property.** For example, if the setting is 0.75 then the pharmacophore will be displayed if the property is found in 75% or more of the ligands.



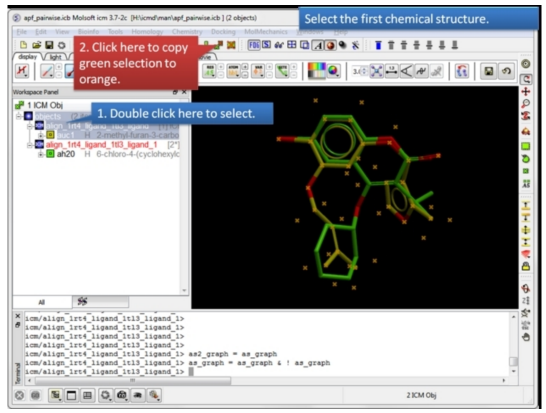
**Consensus is displayed as meshes.** Each mesh can be displayed or undisplay in the ICM workspace.

## 7.36.2 Pairwise APF Score

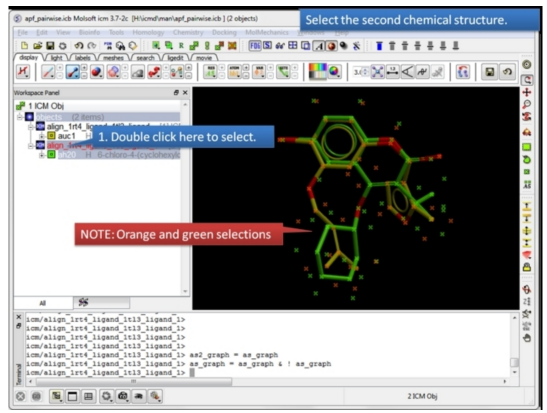
To determine how well the APF fields of two chemicals match you can determine the Pairwise APF Score using the Chemistry/APF Tools/Pairwise APF Score menu option.



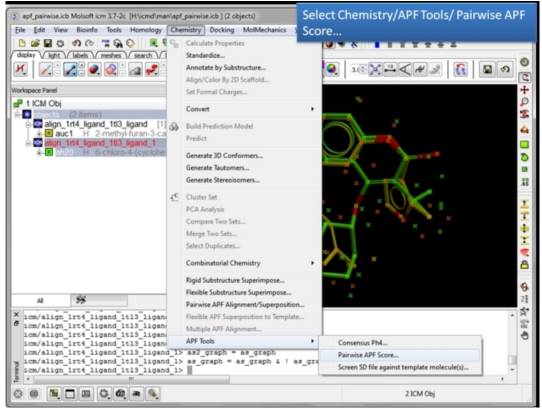
Load and display two superimposed chemicals. You can superimpose the chemcials using APF fields as described here.



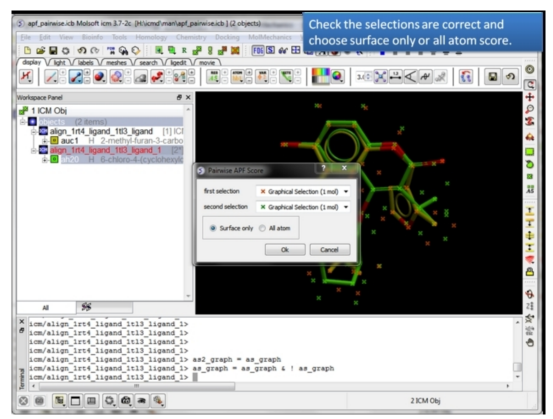
**Select the first chemical.** Use the green selection to orange button to select the first chemical.



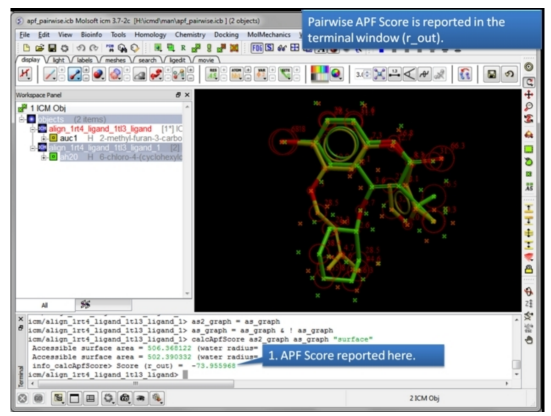
**Select the second chemical.** Double click the chemical in the ICM workspace to select it.



Choose the Pairwise APF Score menu option. Chemistry/APF tools/Pairwise APF Score



**Check the selections are correct.** Choose whether you wish to calculate the score based on all atoms or just the surface atoms.



**The APF Pairwise Score is displayed in the terminal window.** The score is also returned to r\_out. Do not see the Terminal window? Go to Windows menu and check the Terminal box.

# 7.36.3 APF Ligand Based Screen - Screen SD file against template molecule(s)

This option allows you to perform a ligand-based screen using the Atomic Property Fields of superimposed chemicals. SD files are screened and the ligands are scored according to their fit into the APF field. The documentation for this option is in the Screening Chapter.

### 7.36.4 APF 3D QSAR

|Tutorial|

This method uses 3D pharmacophores Atomic Property Fields.

### There are three steps to undertake APF 3D QSAR:

#### **Step 1: Chemical Superposition**

• The APF 3D QSAR method relies on a set of superimposed chemicals. You can superimpose chemicals using the tools here. The 3D chemicals in the ICM workspace then need to be extracted to a chemical spreadsheet as described here. Both the chemicals used for training and the chemicals that you wish to predict need to be superimposed.

### Step 2: Learning

• Use the chemical table containing the superimposed chemicals (Step 1). This table should contain a mol column and a column containing activity data.

- Chemistry/APF Tools/APF 3D QSAR and select the learn tab.
- Choose the name of the chemical table from the drop down list.
- Choose the column containing the activity data.
- The statistics related to the model can be seen in the model section of the ICM workspace left hand side.
- A table called **BASIS** will be created containing a column with the predicted value and cross-predicted value for each element of the training set.

#### **Step 3: Prediction**

- Read in a chemical table containing data you wish to predict. The column formats must match the model table and the chemicals must be superimposed on the training set (See Step 1).
- Chemistry/APF Tools/APF 3D QSAR/ and select the predict tab.
- Enter the name of the chemical table you wish to predict along with the activity column name.
- The table will be appended with a prediction column called **MolApfQsar**.

**NOTE:** The prediction model might be improved by removing some chemicals from the superposition in Step 1. It might be a subjective call as to which chemicals are good for the model and which ones should be removed. In some cases a crystal structure of a protein-ligand complex is available and the ligand pose would be a good template to superimpose onto.

## 8 Learn and Predict

Available in the following product(s): ICM-Chemist-Pro | ICM-VLS

#### In this chapter:

- Model building using the Learn method.
- Use a model to make a Prediction
- Fingerprint Methods
- 3D QSAR
- Learning Theory

There are four learning algorithms built into ICM

- Partial Least Squares (plsRegression)
- Principal Component Regression(pcRegression)
- Bayesian Classifier (bayesianClassifier)
- Random Forest (randomForest)

All 2D molecular property predictors are calculated using fragment-based contributions and 3D models use Atomic Property Fields.

For 2D fingerprints we developed an original method for splitting a molecule into a set of linear or non-linear fragments of different length and representation levels and then each chemical pattern found is converted into a descriptor.

In order to perform 'learn and predict' in ICM information must be stored in a table, molecular table or csv file. See the tables chapter for more information on ICM tables. Both chemical compounds and numeric data can be source for building prediction models.

### 8.1 Learn

|2D QSAR Tutorial|

#### To Learn from a data set and make a model:

First load in a table of data on which you wish to perform the learn and predict functions. See the tables chapter for more information on ICM tables.

- Read in your data e.g. chemical spreadsheet (e.g. File/Open SDF or .csv).
- Select Tools/Table/Build Prediction Model (Learn) and a window or use the Chemistry/Build Prediction Model option.
- Enter the name of table with which you want to perform the predictions. You may locate your table from the drop down arrow menu.
- Select the **Column** from which you wish to learn. Use the drop down arrow to select. **NOTE** If the table does not contain any numeric (integer or real) columns, there is nothing to predict, so the "Build Prediction Model Learn" button will be disabled.
- Enter a name for the learn model.
- Select which learning method you wish to use from the drop down menu (PLS, PCRegression, Bayesian Classifier, Random Forest).
- Select whether you would like ICM to estimate the number of latent variables or you can specify a range.
- Select **No Free Term Constraint:f(0)=0** if you would the PLS method to find a constant and add it to the model.

#### Descriptors

• Select which columns (descriptors) of your table you wish to use to 'learn'. You can select just the **mol** column or all numerical columns.

#### **Chemical Fingerprints**

- If you are using chemical desriptors select the fingerprint method want to use **Linear** or **Extended Connectivity Fingerprints (ECFP)**
- Select **Binary** if you want to use Binary fingerprints do not check if you want to use counted.
- **Minimal chain length** the minimal length of the chain of atoms, enumerated in each compound, usually it is 1 which means that you will consider an atomic composition numbers as descriptors. The typing of atoms can be customized with the map argument.
- Maximal chain length the maximal length of the chain atoms and bonds enumerated in each compound Usually it is 3 or 4. Larger values of this parameter will lead to a large number of possible combinations. To overcome that, either the typing needs to be simple (e.g. only sp2 and sp3 property, regardless of the atom number), or the data set needs to be really large.
- Length this is the total number of bins in a final fingerprint. Typically the size (either explicit or automatically estimated) is in a hundreds to throusands range. We recommend to check the **auto** option.

#### Customizing the fingerprint to fit the predicted property

	lev	Atom properties	Bond properties	1
1	1	element number,nHydrogens		Ø
2	2	element number,nHydrogens	bond order	0
3	3	element number,nHydrogens	bond order	0

• In this dialog you customize the atom properties for every chain length or ecfp fragment size. Click on the "pencil button" to add additional properties to a chain or fragment size (lev). Once the model is made you will see the weights of these properties in the model (right click on model in the ICM workspace and choose Weights).

#### Test

- Select the number of cross-validation groups you wish to use or selected rows can be used for cross validation. The number of iterations will impact the speed of the calculation. 5 is the default number of groups but 2 would be the least rigorous and selecting the 'Leave-1-out' would be the most rigorous calculation.
- **Bootstrapping** generates statistics (R2, RMSD) for a random prediction and you should aim for there to be a significant difference between the random R2 and the model R2.
- Click on the learn button and a model will be created and placed in the ICM Workspace panel (left hand side).

All models are then stored in the ICM workspace as shown below. A number of options are displayed in the right click menu.

🗄 🚱 models	(1 items	)
Apred	plsRegre	ession
		Select
		💥 Delete
		Rename
		Predict
		About
		Weights
		📕 Save As

### 8.1.1 About the Model

To see more details about your model (e.g. R2, RMSE etc....):

• Right click on your model in the ICM workspace and choose Model Data.

lame	Value		
	mod		
Туре	plsRegression	 	
-Nof latent vectors	13	 	
Dimensionality	768		
Fingerprint max chain le	ngth 3	 	
self: R2	0.997971		
self: RMSE	0.601485		
-self: MAE	0.391696		
-self: Spearman	0.997419		
test: R2	0.993538		
	1.07375		
test: MAE	0.505183		
	0.996567		
FUNC			
SCRIPT			

### 8.1.2 Model Weights

The **Weights** dialog can be helpful to see what is driving the correlation in your model or for troubleshooting. The key column is "w" which represents the weight of the fragment in the regression. You may see a particular fragment providing a high weight to the model so you may want to **add** similar fragments to your training set to see if it will improve the model. Removing fragments from the training set based on their weight is not recommended because each fragment is part of a multiple component and it is difficult to know precisely its importance.

- Right click on the model in the ICM Workspace and choose Weights
- A table will be displayed containing the following columns:

**BITS** = the number of the chain or fragment.

**name** = smiles string of fragment. You can copy and paste this into the molecular editor to view it in 2D.

 $\mathbf{w}$  = is actual weight of the fragment in the regression (used directly to get the result value).

mean = indicates the mean occurence of the fragment in the model. For example, a value of 6 means the fragment is used 6 times in the regression.

**rmsd** = a high value indicates high occurence of the fragment and influences the relative weight parameter.

**wRel** = relative 'importance' of the fragment based on the RMSD, mean and CorrY values.

### 8.1.3 Improve Model

The model can be improved by:

- Right click on the model in the ICM Workspace and choose Improve
- A dialog box similar to the one you used to make the model in Learn will be displayed.

### 8.1.4 Save and Share Model

You can save and share a model by:

- Right click on the model in the ICM workspace (left hand panel) and choose Save As... This will save the model in MolSoft's .icb format.
- The .icb file can be opened in another session using File/Open

### 8.2 Predict

To make a prediction using a created model.

Read the table of data into ICM from which you wish to predict. Make sure the table contains the same columns used for the learn model.

- Tools/Table/Predict or Chemistry/Predict
- Select which table you wish to make the prediction on.
- Select which model you wish to use.
- Check that the required columns are in the table. If they are absent a red mark will appear against the column that is missing.
- Click Predict.

### 8.3 Fingerprint Methods

**MolSoft's Fingerprints** MolSoft has developed a proprietary method for generating hashed chemical fingerprints used in chemical search, similarity calculations and QSAR. ICM fingerprints do not use any pre-defined set of fragments. Instead, ICM enumerates all possible linear and some non-linear fragments (circular and triangular) up to a certain size and hash them into a result vector. Equal fragments always get the same position in the vector and fingerprints can be either binary (presence or absence of a fragment) or counted (number of fragment occurrences is counted). The method includes 2D pharmacophore fingerprints whereby Euclidian 3D distance is replaced by number of bonds between atoms to make it applicable to the 2D structures directly.

Additional information on Chemical Fingerprint properties can be read here: http://www.molsoft.com/man/fingerprints.html

**Extended Connectivity FingerPrints (ECFP)** We also support Extended Connectivity FingerPrints (ECFP).

## 8.4 3D QSAR

The 3D QSAR method is described in the Cheminformatics Chapter Atomic Property Field Section.

### 8.5 A Little Theory on Learning

For a more detailed explanation of the theory behind Partial Least Squares (PLS) we suggest you read Geladi et al Analytica Chimica Acta (1986) 1-17.

**PLS (Partial Least Squares) Regression** PLS regression algorithm builds linear prediction model: in format y = (w, x) + b, where b is the **bias** - a real number, and w is the weights vector, which is scalarly multiplied by the data vector x. PLS uses the given learning y values very actively which allows it to produce fairly good models with respect to constraint of being linear. Although linear regression models have an advantage of weights for each descriptor which gives a useful information and allows feature selection in many cases.

The linear model simply is not able to predict higher order dependencies.

There are different ways to deal with it. By adding the second order columns into the descriptor set you can let PLS predict them. Actually if you have a lot of columns derived from basic data, the linear model built will be able to make a high-quality linear approximations of the actual functions. ICM has a powerful tool for automatical generation of such descriptors based on compound data -- molecule fingerprints generation algorithm. It generates hundreds of columns based on initial data. The withdraw is that analysing the weights given by PLS to generated descriptors is almost senseless. You will need a *mol* column in your table to use this feature.

ICM has built-in models for prediction of several significant molecule properties, like logP, logS, PSA based on fingerprints+PLS symbiosis, which have proven their quality.

#### PC (Principal Component) Regression

PCR also builds linear model in its simplest form, as PLS does, though it sets other weights to descriptors, and built models are usually worse in sense of predicting, because PCR uses *value* information of the *training data* only in secondary way. We recommend you to use PCR, when you want to build an ordinary regression (MLR - Multiple Linear Regression) model by using only some number of first principal components of X data matrix (ordered by decreasing eigenvalues) or even builing the full MLR model (by setting the number of PCs to value higher than the number of rows in matrix).

8.5 A Little Theory on Learning

## 9 MolScreen

#### Available in the following product(s): ICM-VLS |

MolScreen is a set of high quality 2D fingerprint and 3D pharmacophore models for a broad range of pharmacology and toxicology targets. The models can be used for lead discovery or counter screening. The models use MolSoft's 2D QSAR/Fingerprint and 3D Atomic Property Fields methods. The models can be screened directly using MolSoft's ICM-Pro + VLS software or via our contract research services. For more information please see: www.molsoft.com/molscreen.html

There are two screening panels in the Chemistry/MolScreen menu:

- 1. Brief Model Panel. This selection of models is provided free of charge from MolSoft.
- 2. Full Model Panel. Please contact MolSoft to use this this panel containing all available models.

In this chapter:

- MolScreen Model Types
- Brief Panel
- Full Model Panel
- Custom Model Panel
- Results Table
- Make Classification Model
- Make APF/SAR Model

### 9.1 Model Types

# 9.1.1 Kernel Chemical Classification/Regression (kcc) models

# **Hybrid Fingerprints**

# Dataset

IC50, EC50, Ki, Kd data was downloaded from ChEMBL18, they were combined and converted to pKd value. Compounds with pKd > 5. (i.e. 10 micro Molar) were classified as positives. 90% of ChEMBL18 compounds were assigned to training set. The remaining 10% compounds were assigned to external test set. To the training set, a subset of compounds with pKd>7. against any targets from ChEMBL18 was added as decoy. To the external test set, the approved drugs were added as decoy

# Training

A Kernel Chemical Classification/Regression (kcc) model was trained using the training set compounds. The performance of the model was evaluated using the external test set

## **Predicted Value**

The kcc model returns two scores:

The kcc score is the classification score: The positives have a median kcc score of 1. The decoys have a median kcc score of 0. The kcc score is renamed to MolClass in the pairwise table.

The kca score is the regression score of pKd value A kca score of 6. indicates microMolar activities. The kca score is renamed to MolpKd in the pairwise table.

In the pairwise table, a pPvalue is calculated for both the kcc and kca score. The maximum pPvalue was taken as the final pPvalue for that model. The pPvalue indicates -Log of probability that the compound belongs to the random decoy. A pPvalue of 1 indicates the compound is comparable to the top 90 percentile decoy. A pPvalue of 2 indicates the compound is comparable to the top 99 percentile decoy.

# 9.1.2 Docking to Ligand Field Z-Score (dfz) models

9.1.2 Docking to Ligand Field Z-Score (dfz)models

# APF

# Dataset

If there is available data from ChEMBL18: IC50, EC50, Ki, Kd data was downloaded from ChEMBL18, they were combined and converted to pKd value. Compounds with pKd > 5. (i.e. 10 micro Molar) were classified as positives. 80% of ChEMBL18 compounds were assigned to training set. The remaining 20% compounds were assigned to external test set. To the training set, a subset of compounds with pKd>7. against any targets from ChEMBL18 was added as decoy. To the external test set, the approved drugs were added as decoy

If ChEMBL data is not available, approved drugs were used as decoy to calculate Z-Score.

# Training

The dfz model was training in the following way:

1. For any target, all of its associated mammalian pocketome entries were used. 2. If ChEMBL data is not available, the pocketome entry with the highest number of co-crystallized ligand was used. 3. If ChEMBL training set is available, it was docked to each pocketome entries using APF method. 4. The best combination of template clusters were selected to maximize differentiation between actives and decoys. 5. Z-Score was calculated using approved drug's mean and standard deviation of APF score.

## **Predicted Value**

The dfz model returns one score:

The dfz score is the Z-Score: A score of 1 means the compound is 1 standard deviation above the mean score of approved drugs decoy. The dfz score is renamed to MolZScore in the pairwise table.

In the pairwise table, a pPvalue is calculated for both the Z-Score. The pPvalue indicates -Log of probability that the compound belongs to the random decoy. A pPvalue of 1 indicates the compound is comparable to the top 90 percentile decoy. A pPvalue of 2 indicates the compound is comparable to the top 99 percentile decoy.

### 9.1.3 Docking to Ligand Field Classification/Regression (dfa) models

374 9.1.3 Docking to Ligand FieldClassification/Regression (dfa) models

# Hybrid 4D/2D

# Dataset

IC50, EC50, Ki, Kd data was downloaded from ChEMBL18, they were combined and converted to pKd value. Compounds with pKd > 5. (i.e. 10 micro Molar) were classified as positives. 90% of ChEMBL18 compounds were assigned to training set. The remaining 10% compounds were assigned to external test set. To the training set, a subset of compounds with pKd>7. against any targets from ChEMBL18 was added as decoy. To the external test set, the approved drugs were added as decoy

# Training

The dfa model is trained in the following steps:

1. Either: a. Training set was docked to the 4D maps of all the pocketome entries associated with the target. b. If pocketome entry is not available. The training set compounds were aligned in 3D. 2. Combining the training set compounds with pocketome co-crystallized ligands (if available), cluster in APF. 3. Subset of ligands were selected from each cluster as APF template. 4. All training set compounds were docked using APF method to all clusters. 5. The best combination of clusters were selected to maximize recognition of actives from decoys. 6. For each selected cluster, a pKd regression model was trained using the 3D poses of the ligands above a certain APF score cutoff.

## **Predicted Value**

Any compound will be predicted using the dfa models in the following way: The compound will be docked using APF method to each of the template clusters. The compound will be assigned to the cluster that gives the highest normalized APF score. The 3D regression model of that cluster will then be used to predict the pKd value of that compound if the APF score is within the score cutoff.

The dfa model returns two scores:

The dfc score is the classification score: The positives have a median dfc score of 1. The decoys have a median dfc score of 0. The dfc score is renamed to MolClass in the pairwise table.

The dfa score is the regression score of pKd value A dfa score of 6. indicates microMolar activities. The dfa score is renamed to MolpKd in the pairwise table.

In the pairwise table, a pPvalue is calculated for both the dfc and dfa score. The maximum pPvalue was taken as the final pPvalue for that model. The pPvalue indicates -Log of probability that the compound belongs to the random decoy. A pPvalue of 1 indicates the compound is comparable to the top 90 percentile decoy. A pPvalue of 2 indicates the compound is comparable to the top 99 percentile decoy.

### 9.1.4 Docking to Protein Pocket Classification/Regression (dpc) models

3829.1.4 Docking to Protein PocketClassification/Regression (dpc) models

# Hybrid Docking

# Dataset

IC50, EC50, Ki, Kd data was downloaded from ChEMBL18, they were combined and converted to pKd value. Compounds with pKd > 5. (i.e. 10 micro Molar) were classified as positives. 90% of ChEMBL18 compounds were assigned to training set. The remaining 10% compounds were assigned to external test set. To the training set, a subset of compounds with pKd>7. against any targets from ChEMBL18 was added as decoy. To the external test set, the approved drugs were added as decoy

# Training

The dpc model is trained in the following steps:

1. The pocketome entry with the most co-crystallized ligands associated with the target is selected 2. The residues around the pocket were clustered, selected representative PDBs were retained 3. All training set compounds were docked to the 4D maps of the pocket in the presences of the co-crystallized ligands in the form of APF template 4. A score cutoff to maximize sensitivity and accuracy was selected 5. All compounds within that score cutoff were used to train a pKd prediction model using their 3D poses.

# **Predicted Value**

Any compound will be predicted using the dpc models in the following way: The compound will be docked to the 4D maps of the pocket in the presence of APF template The 3D regression model of the pocket will then be used to predict the pKd value of that compound if the docking score is within the score cutoff.

The dpc model returns two scores:

The dpc score is the classification score: The positives have a median dpc score of 1. The decoys have a median dpc score of 0. The dpc score is renamed to MolClass in the pairwise table.

The dpa score is the regression score of pKd value A dpa score of 6. indicates microMolar activities. The dpa score is renamed to MolpKd in the pairwise table.

In the pairwise table, a pPvalue is calculated for both the dpc and dpa score. The maximum pPvalue was taken as the final pPvalue for that model. The pPvalue indicates -Log of probability that the compound belongs to the random decoy. A pPvalue of 1 indicates the compound is comparable to the top 90 percentile decoy. A pPvalue of 2 indicates the compound is comparable to the top 99 percentile decoy.

# 9.2 Brief Panel

#### To use the Brief Model Panel:

- Read in a set of chemicals you wish to screen against the MolScreen panel (e.g File/Open sdf).
- Select Chemistry/MolScreen/Brief Model Panel
- Select the model type you would like to screen from the tab menu.
- Use the dropdown button to locate the chemical table you wish to screen
- Use the check boxes to select the targets you would like to screen or choose select all.
- There are some options which can be selected: **Color** will color the table by a rainbow scaled to the value range in the table. **LookupOnly** will search your chemicals to see if they are contained in the training and test set. **deleteEmptyCols** will clean up your results table by removing columns that do not have data. **PairwiseTable** will generate a model and target pairwise table.
- The results will be displayed in a separate table.

# 9.3 Full Model Panel

To run a full panel screen:

- Read in a set of chemicals you wish to screen against the MolScreen panel (e.g File/Open sdf).
- Select Chemistry/MolScreen/Full Model Panel
- Select the model type you would like to screen from the tab menu.
- Use the dropdown button in the table **Tools** panel (right hand side) to locate the chemical table you wish to screen.
- There are some options which can be selected: **Color** will color the table by a rainbow scaled to the value range in the table. **LookupOnly** will search your chemicals to see if they are contained in the training and test set. **deleteEmptyCols** will clean up your results table by removing columns that do not have data. **PairwiseTable** will generate a model and target pairwise table.
- Select the targets you wish to screen against by selecting the rows in the table. Click on the row number to select the row (the row will then be shaded blue). Use the CTRL key and click to

select a non-contiguous range or use Tab to select a contiguous range of models.

- Click Apply Selected Models
- The results will be displayed in a separate table.

To view the performance of a model select the model by selecting the row number and then choose **Check Model Performance** in the Tools panel of the table.

# 9.4 Custom Model Panel

MolScreen has some preprepared model panels available. To select a preprepared custom model panel:

- Chemistry/MolScreen/Select Custom Model Panel
- Choose the panel from the drop down Panel Name dialog box.
- Select the path where your MolScreen models are located (usually in .icm/models).

To create your own custom model panel:

- Chemistry/MolScreen/Make Custom Model Panel
- Give your panel a unique name.
- Enter a uniprot prefix (e.g. ESR1, ANDR)
- Enter a keyword (e.g. nuclear)
- Press OK and a table will be displayed containing the models.
- You can repeat the steps above to append to the custome model panel.
- Save your custom panel in your models directory e.g. (.icm/model)

To load your custom panel:

• Chemistry/MolScreen/Full Model Panel and check the option **Read Custom Panels** 

# 9.5 Results Table

The results table will contain the columns from the chemical table that you screened plus the following results columns: Depending on your selection you might have two tables **Model Pairwise** and **Target Pairwise** 

- mol your query chemical
- target the target model you screened against
- **pPValue** kcc/hybrid4D+2D models only: -Log(probability) relative to approved drug decoys capped at 5: 1: Top 10 percent 2: Top 1 percent 3: Top 0.1 percent
- **MolpKd** kcc/hybrid4D+2D models only: Prediction of pKd, pKi, pIC50, pEC50, etc >6. indicates submicromolar value, >9. indicates nanomolar, ND indicates disimilarity to known binders
- MolpKd Error kcc/hybrid4D+2D models only: Estimated total uncertainty/error for pKd prediction, including experimental error of ~1. pKd unit
- **MolClass** kcc/hybrid4D+2D models only: Classification Score: for kcc models: nProbability of being a positive, assume 0.5% hitrate for others: 1. Median value of actives 2.Median value of inactives
- TANIsim kcc models only: Tanimoto distance from model nearest compound
- **modelnearestCpd** kcc models only: nearest compound to the query in the model
- **MolZScore** apf models only: Z-Score 3. indicates 3 standard deviation above approved drugs decoy
- predictType model type
- **modelAUC** Area Under Curve (AUC) for model
- modelpKdAUC kcc models only: AUC for pKd model
- modelRMSE kcc models only: RMSE for model

- modelQ2 kcc models only: Q2 for model
- MolDockScore hybrid docking only ICM docking score, Score < -32. are in general significant, but may vary from structure to structure.

#### Fold Columns to Original Table

In the **Tools** panel of the table there is a convenient way of packaging your results into your original query table.

# 9.6 Make Chemical Classification Model

This option allows you to make your own kcc models either on your own data (local table) or from ChEMBL.

#### From your own data:

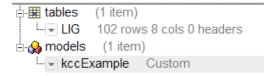
- Read in a chemical spreadsheet containing your chemical 2D structure (mol column) and an activity column containing data such as pKd, mM, uM or nM.
- Select Chemistry/MolScreen/ Make Chemical Classification Model
- Select the Local Table tab.
- Give your model a unique name.
- Find your chemical table from the drop down list.
- Find the column name containing **activity** data from the drop down list.
- Select the activity unit pKd, mM, uM or nM.
- Select whether you would want pKa charge prediction.
- Click OK.

#### From ChEMBL data:

- Give your model a unique name.
- Enter a Uniprot ID (e.g. 5ht1f)
- Select whether or not you want to include all mammalian data or just human.
- Check makemodel to make the model immeditately.

#### How to use the model to predict:

• When the model has been built you will see the model listed in the ICM workspace as shown below.



#### To run the model against a set of chemicals:

- Read in a chemical spreadsheet containing your chemical data.
- Right click on the model in the ICM workspace and choose Predict.
- Enter the table name and prediction model you would like to use.
- Once the prediction has finished you will see three additional columns: the kcc value, kca value and the tanimoto distance to the nearest compound in the model dataset.

# 9.7 Make APF Docking SAR Model

This option allows you to make your own dfa model either on your own data (local table) or from ChEMBL. If the chemicals are not superimposed then this option will automatically superimpose them. If you want to superimpose yourself before running then the chemical superposition

options are described here.

#### From your own data:

- Read in a chemical spreadsheet containing your chemical 2D structure (mol column) and an activity column containing data such as pKd, mM, uM or nM.
- Select Chemistry/MolScreen/ Make Chemical Classification Model
- Select the Local Table tab.
- Give your model a unique name.
- Find your chemical table from the drop down list.
- Find the column name containing **activity** data from the drop down list.
- Select the activity unit pKd, mM, uM or nM.
- Select whether you would want pKa charge prediction.
- Click OK.

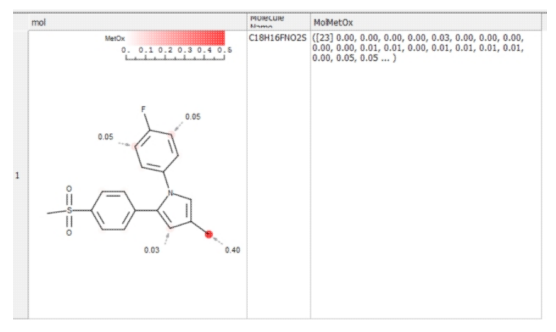
#### From ChEMBL data:

- Give your model a unique name.
- Enter a Uniprot ID (e.g. 5ht1f)
- Select whether or not you want to include all mammalian data or just human.
- Check makemodel to make the model immeditately.

# 9.8 Predict Metabolic Oxidation

To predict metabolic oxidation:

- Read in a chemical or chemical spreadsheet.
- Chemistry/MolScreen/Predict Metabolic Oxidation and select the table name.
- A results table as shown below will be displayed. Each atom is labeled by the predicted metabolic oxidation. The higher the number the more likely it is to be oxidized based on the model described below. The column MolMetOx is the atom by atom prediction.



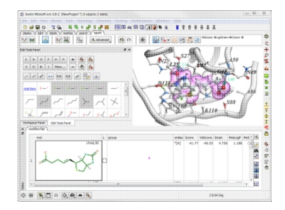
#### About the Method

Metabolic oxidation data from various sources was analyzed to determine possible positions where oxidation can occur. The ECFP fingerprints for

every atom in the training set were then calculuated. The frequencies of oxidation or non-oxidation for each atom based on calculated fingerprints was then calculated. We then trained a random forest regression model to predict the ratio of /(+) for every atom. The higher the number the more samples of oxidation over all samples at that position were found.

# 10 The 3D Fully Interactive Ligand Editor

The ligand editor is a powerful tool for the interactive design of new lead compounds in 3D. It allows you to make modifications to the ligand and see the affect of the modification on the ligand binding energy and interaction with the receptor.



Watch a webinar about the 3D ligand editor here.

There is a tutorial on how to use the 3D ligand editor here.

#### **Chapter Contents:**

- Setup Ligand and Receptor
- Ligand Editor Preferences
- Display Options
- Evaluate Ligand Score and Strain
- Edit Ligand
- Screen Replacement Group
- Dock or Minimize Ligand
- Dock Chemical Table
- Fragment Linking
- Impose Tethers and Distance Restraints
- Defined Flexible Receptor Groups
- Side Chain Refinement
- Dock to Multiple Receptor Conformations
- Covalent Docking
- Dock to Atomic Property Fields
- Save Docked Ligand as PDB
- Export Docking Project
- Close and Clean Up Ligand Editor Project

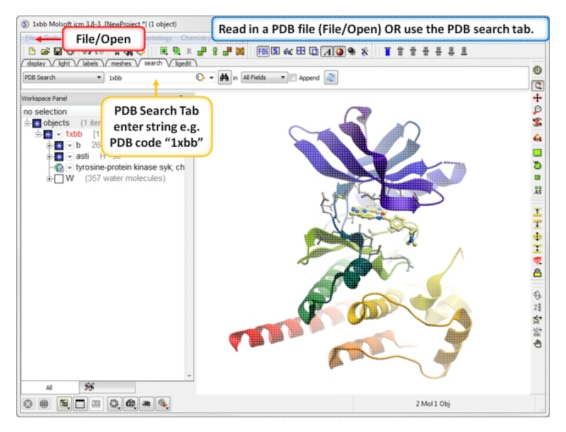
# **10.1 Setup Ligand and Receptor**

```
Available in the following product(s): ICM-Pro | ICM-VLS | ICM-Chemist-Pro
```

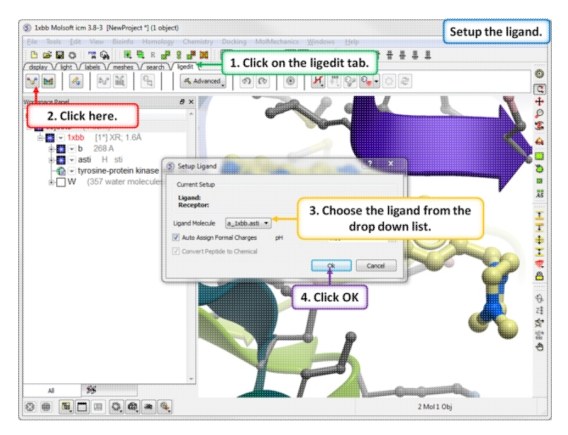
Video|

#### As an example we will use the kinase structure 1xbb.

• Click on the search tab and type 1xbb.



- Click on the **ligedit** tab
- Click on the **Setup Ligand** button.
- Enter the ICM selection language for the Ligand Molecule (a\_1xbb\_asti) or use the drop down button to locate it.
- Select whether or not you would like ICM to automatically assign formal charges and set the pH.

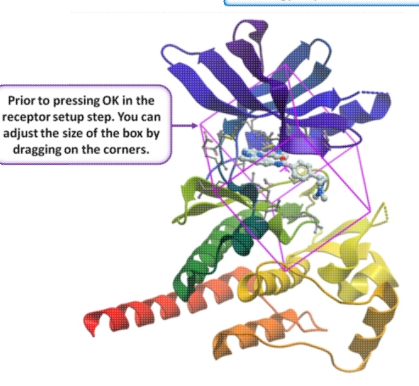


- Select the **Receptor Setup** button.
- Enter the ICM selection language for **Receptor Object** (a\_1xbb.) or use the drop down button to locate it.
- Select whether or not you wish to keep all waters in the receptor, keep tight waters, delete all waters, optimize existing hydrogens and optimize His, Pro, As, Gln, and Cys. The "mask" option will hide atoms not reported in the crystal strucure (e.g. occupancy 0). If this box is not checked ICM will build the missing atoms of the side-chain.
- Click on the option to select **Box Around Existing Ligand**. There are other options: **Identify Pocket Box** will run ICMpocketFinder and return a table of pockets. Click on the table to select the pocket you want and then press OK. You can also **Make Box Around Atom Selection**.
- Enter a box margin of 3. This option defines the size of the energy maps around the ligand. The value of 3. should encompass the whole site but if you have a binding pocket that is very elongated or unusual in any way it is recommended that you check that the purple box covers the site you are interested in.

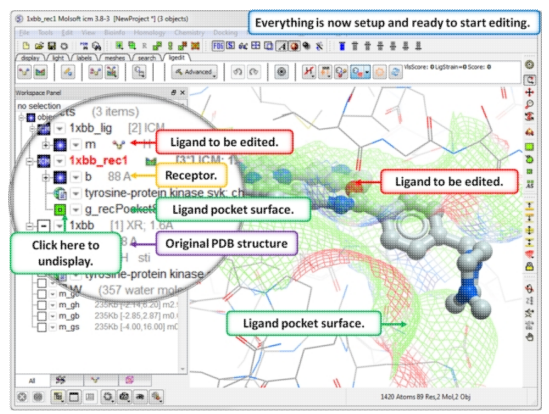
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	Current Setup Ligand: a_1xbb_jg. Receptor: Define Receptor Receptor Object a_1xbb. Keep Water In Receptor Define Pocket or Select Pocket Atoms		Mask Missing Sidechains
	A. If the ligand is Select Make Box Around Existing Ligand Make Box Around Atom Selection Box Margin 3. • More Options Create Ligand Pocket Surface	already in the this button.	e pocket
AI 55	1 · · · · · · · · · · · · · · · · · · ·	5. Click OK	

- At this stage you can change the size of the box by dragging on the corners of the purple box in the graphics display. Generally the default box is the appropriate size so no changes are needed.
- Click OK and the energy maps will be generated.

The energy maps are made inside the purple box.



• Once everything is setup you will see the ligand and receptor objects along with the original pdb file in the ICM workspace.



• If the ligand-receptor complex is a PDB structure and hydrogens have been added and optimized then it is important to run a minimization after setup.

# 10.2 Ligand-Editor-Preferences

Available in the following product(s): ICM-Pro | ICM-VLS | ICM-Chemist-Pro

This step is optional but you may want to tweak the default preferences a bit or add new modifiers and linkers. You can change the preferences by clicking on the **Advanced** button in the ligedit tab and then choose **"Setup Ligand Editor Preferences"**.

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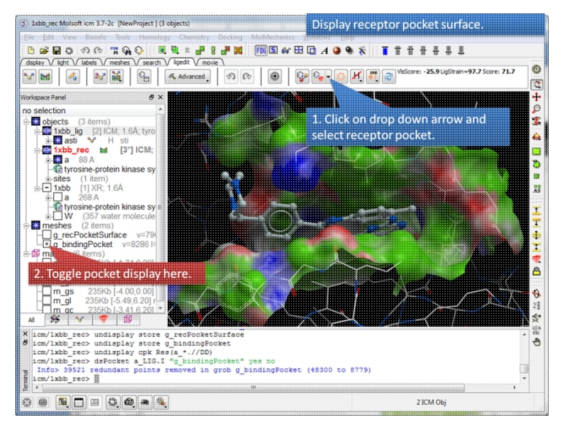
# **10.3 Display Options**

```
Available in the following product(s): \texttt{ICM-Pro} \mid \texttt{ICM-VLS} \mid \texttt{ICM-Chemist-Pro}
```

Video|

# 10.3.1 Display the receptor pocket surface

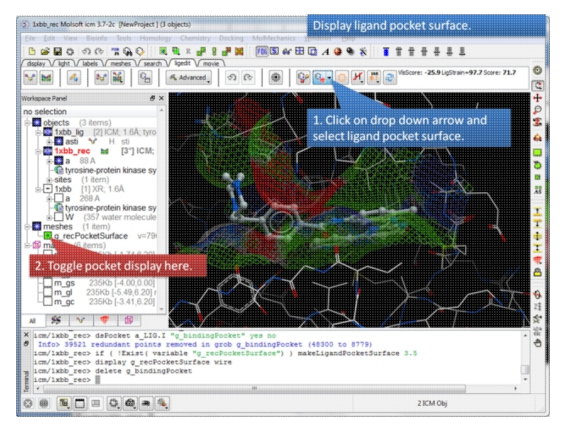
This option makes a surface based on the properties of the pocket.



About the Receptor Pocket Surface: White=aromatic lipophilic, Green=non-aromatic other (mostly aliphatic) lipophilic surface, Red=hydrogen bonding acceptor potential, Blue=hydrogen bond donor potential

## 10.3.2 Display the ligand pocket surface

The Ligand Surface option allows you to visualize cavities that are open for ligand modifications.

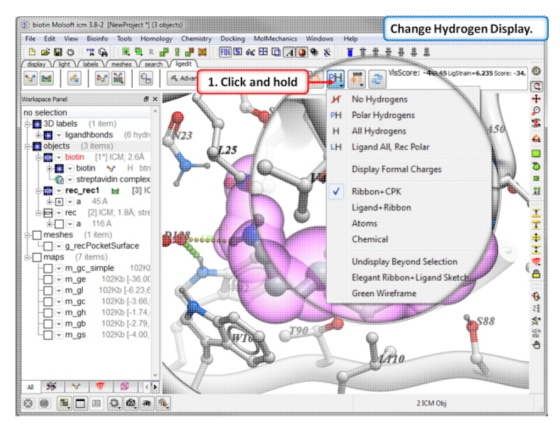


About the Ligand Pocket Surface: White=aromatic lipophilic, Green=non-aromatic other (mostly aliphatic) lipophilic surface, Red=hydrogen bonding acceptor potential, Blue=hydrogen bond donor potential

## 10.3.3 Display Hydrogen Atoms

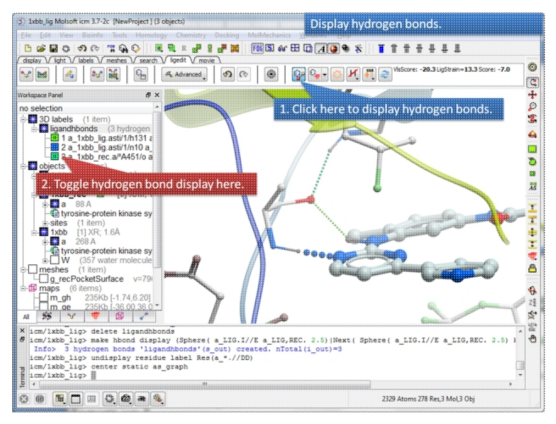
#### To display hydrogen atoms:

- Single click on the hydrogen atom display button will toggle the display between: no hydrogens, polar hydrogens, all hydrogens, all hydrogens in ligand and polar hydrgens in receptor.
- Click and hold on the button and you can choose the option rather than using the toggle.



# 10.3.4 Display Hydrogen Bonds

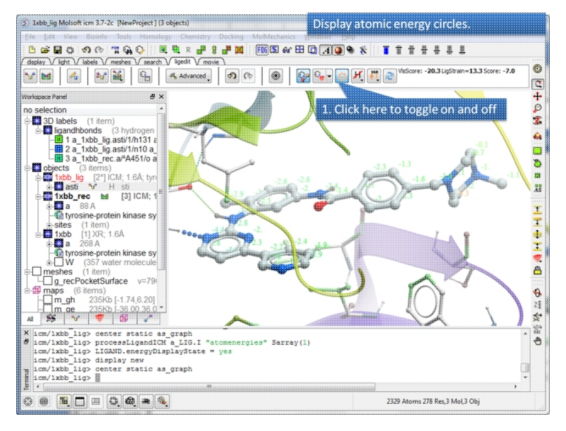
#### To display hydrogen bonds:



**About Hydrogen Bonds:** The coloring of the H-bonds are red (strong - thick spheres) to green (intermediate) to blue (weak - thin spheres). Once the hydrogen bonds have been displayed they can be displayed and undisplayed in the **3D labels** section of the ICM Workspace (left hand side of graphical window).

### 10.3.5 Display Atomic Energy Circles

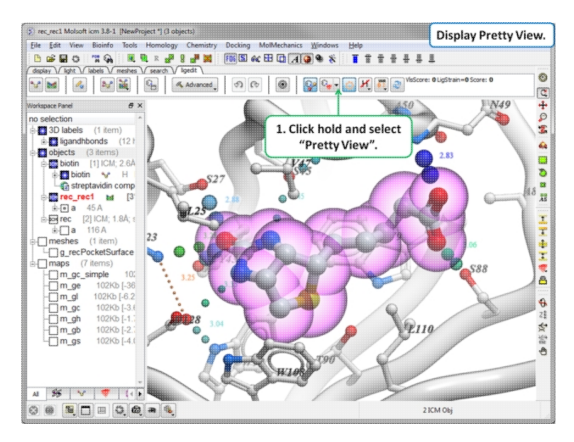
#### To display atomic energy circles:



About the Atomic Energy Circles: Good ligand-receptor interactions are highlighted by green spheres and low values. Poor energy interactions are displayed as orange-->red stars - red being a major clash and a very poor energy contribution. Each atom is given an energy value relating to its contribution to the total receptor-ligand interaction energy. Low values colored green are considered favorable. The green values on each atom are per-atom contribution to the total energy in the receptor grid ('show energy atom' command). The score/energy can not be partitioned into per-atomic contributions because several terms/contributions are not pairwise (in particular solvation-related, as well as the strain). However, grid-based approximate interaction energy is calculated per-atom and reflects, to an extent, favorable/unfavorable contributions from things like van der waals, hydrogen bonds, coulomb part of electrostatics and some approximation of lipophilicity.

### 10.3.6 Display Pretty View

To display the default "Pretty View":



- Setup the ligand and receptor.
- Click and hold on the button shown below and choose "Pretty View".

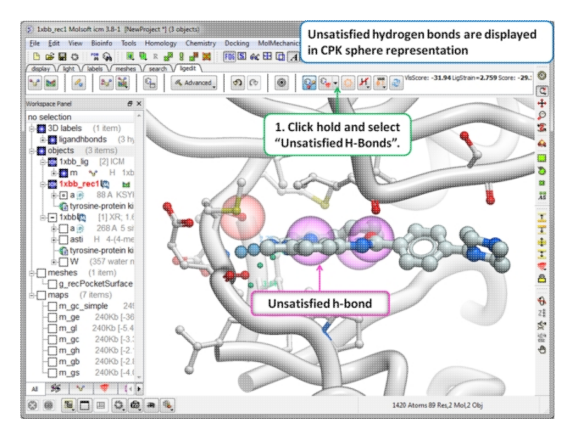
### 10.3.7 2D Interaction Diagram

To create a 2D interaction diagram - click on the 2D interaction diagram button shown below. More details regarding this option can be found here.

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2D	Intera	action	Diagram	Button

### 10.3.8 Unsatisfied Hydrogen Bonds

To display unsatisfied hydrogen bonds:

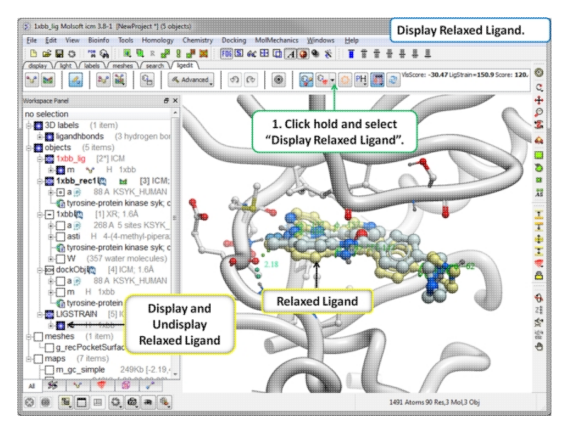


- Setup the ligand and receptor.
- Click and hold on the button shown below and choose "Unsatisfied H-bonds".

### 10.3.9 Display Relaxed Ligand

The relaxed ligand is is the closest local minimum after minimization in the Merck Molecular Force Field (MMFF) outside of the receptor.

#### To display the relaxed ligand:



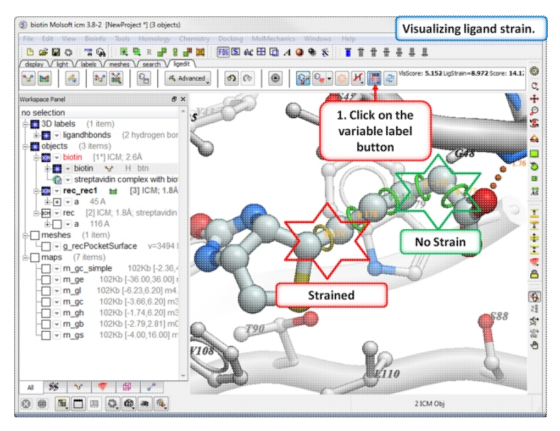
- Setup the ligand and receptor.
- Click and hold on the button shown below and choose "Display Relaxed Ligand".

## 10.3.10 Visualize Ligand Strain

If you are seeing a high ligand strain value when you calculate the score you can visualize the cause of this by viewing the variable angles.

#### To view the variable angles:

• Click on the variable angle buton in the ligand editor tool panel.

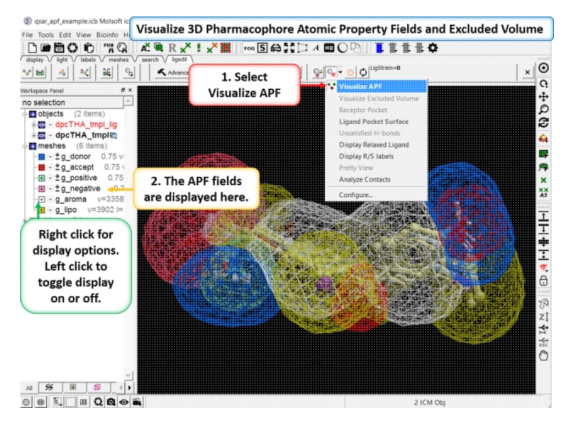


Rings of varying diameter and color are superimiposed on rotatable bonds. Green rings with large diameter are considered less constrained than rings with small green/yellow/red rings. Red rings are highly constrained. You can also compare your strained ligand with the relaxed ligand.

### 10.3.11 Visualize APF Field and Excluded Volume

If you are docking to Atomic Property Fields (APF - 3D Pharmacophore) as described here you can visualize the APF fields. Click on the button shown below and choose Visualize APF. The fields will be displayed and they can be toggled on and off in the ICM workspace - right click in the display box in the ICM workspace to change the surface display (e.g from mesh to transparent surface).

APF is a ligand based method but you can use a protein structure to represent an excluded volume - if you chose this option when performing APF docking then you can display the excluded volume.



## 10.3.12 Identify Ligand-Receptor Contacts

To display ligand-receptor contacts:

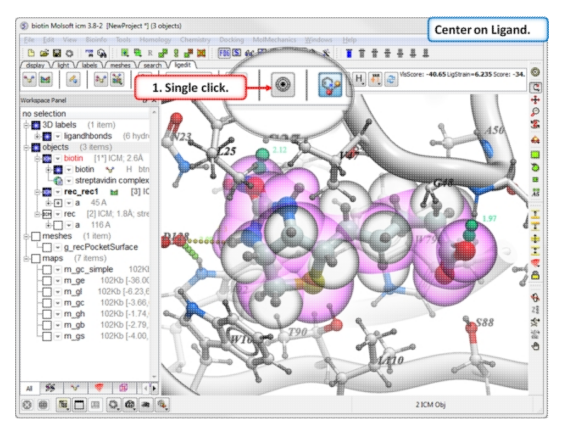
- Click on the button shown below.
- Select "Analyze Contacts".
- The xstick display in the region will be scaled according to the atom/residue contact area. For example residues making large contacts with a ligand will be displayed in thicker xstick representation (and colored yellow) than those making less significant contacts.
- A table as shown below will be displayed. The table lists the contact area, exposed area, percentage of contact area compared to exposed, the nearest atom of a residue and the distance.

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# 10.3.13 Center on Ligand

To center on a ligand in the graphical display:

• Click on the center button in the ligedit tab.



# **10.4 Evaluate Ligand Score and Strain**

Video|

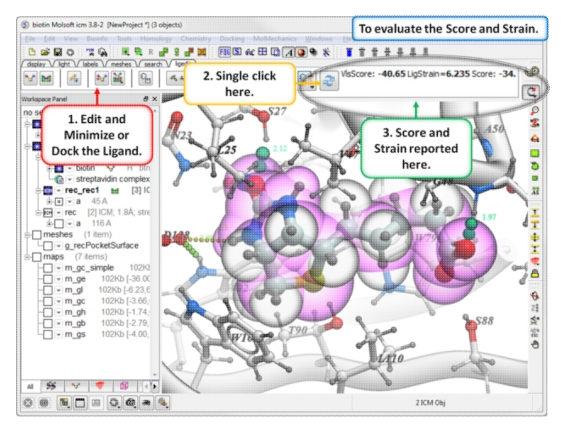
After you have edited a ligand and re-docked or minimized it you can evaluate the Score and Strain for the ligand. Ideally a good modification to your ligand will lower the score.

#### About the Score

- VIsScore = the ICM-VLS score used for ligand screening. The lower the Score the better the predicted interaction. VLS score is available with ICM-Pro + VLS and ICM-Chemist-Pro products. This score incorporates the strain of the ligand compared to the free state.
- **LigStrain** = The strain of the ligand. Lower values = less strain.
- **Score** = This score does not incorporate ligand strain. Therefore it is VlsScore LigStrain. The lower the Score the better the predicted interaction.

#### To evaluate the Score and Strain:

• Click on the "Recalculate Binding Score Button".



# 10.5 Edit Ligand

Available in the following product(s): ICM-Pro | ICM-VLS | ICM-Chemist-Pro

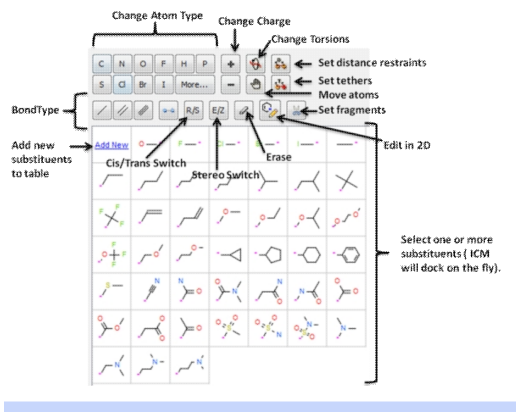
Video|

#### To edit a ligand:

- Setup the ligand and receptor.
- Display hydrogens using the hydrogen display button.
- Click on the **Edit Tools Button** and a panel of buttons will be displayed as shown below.
- To edit an atom or bond, first click on the desired atom, group or bond in the panel and the click on the atom or bond which you want to modify in the graphical display.
- Once you have made a modification you can evluate the score and judge how good the modification was.



• See graphic below to understand what each button does.

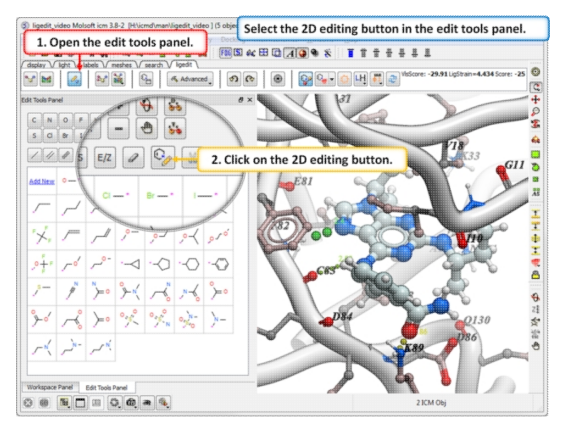


NOTE: After each modification a small mimimization is applied. To turn this off go to Advanced/Ligand Editor Preferences and remove the check mark from the option "Minimize After Modify Group".

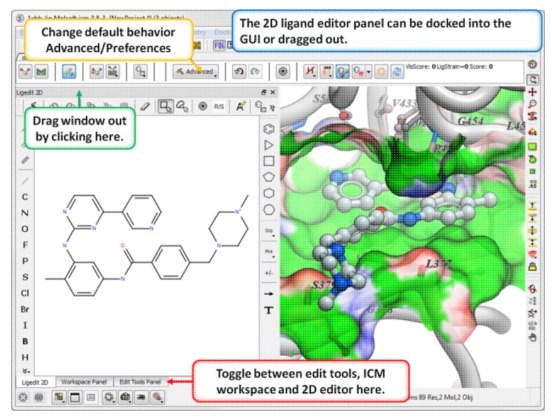
## 10.5.1 Edit Ligand in 2D.

#### To edit the ligand in 2D:

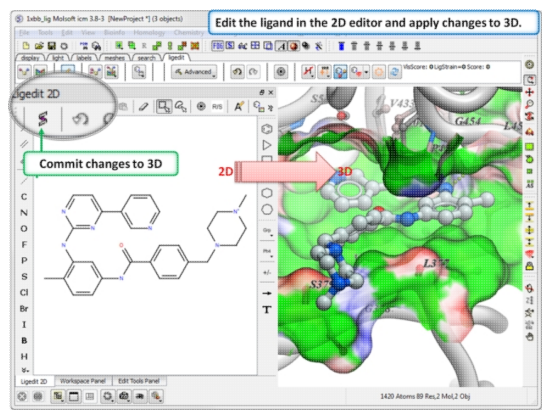
- •
- Setup the ligand and receptor. Click on the **Edit Tools Button** and a panel of buttons will be • displayed as shown below.
- Click on the 2D editor button and edit using the tools in the • molecule editor.



• The 2D ligand editor can be displayed in the panel on the left hand side docked into the GUI or you can change the location by dragging it. The default location can be controled by clicking on the Advanced button and choosing Preferences/ligedit2D Position.



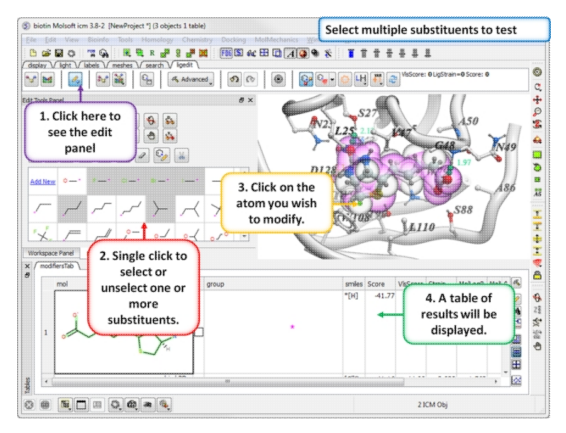
• Make changes to the ligand in the 2D editor and commit them to 3D as shown below.



### **10.5.2 Multiple Substituents**

You can select more than one group. ICM will sample the energy of each group and return a table of the results ranked by binding score. The table columns are described below.

- Click on one or more substituents (shaded grey when selected).
- Click on the atom you wish to modify.
- The results will be reported in the modifier table.



#### About the modifiers table

**mol** = 2D sketch of ligand with core substructure highlighted in green. Modifier group is not highlighted in green.

 $\mathbf{L} = \text{Click}$  in box to display ligand with modifier group.

**smiles** = smiles string of modifier group

**SubstScore** = Score for modifier group only

**Score** = Score of whole ligand including modifier group

**MolLogP** = Predicted LogP

**MolLogS** = Predicted LogPredicted LogSS

**MoldHf** = Preidction model build for 'delta Hf in gas' property. using public NIST database. Description can be found: http://webbook.nist.gov A low dHf value means that the compound is more 'stable'.

MolPSA = Polar Surface Area

**Volume** = Volume of ligand.

### **10.5.3 User Defined Modifiers**

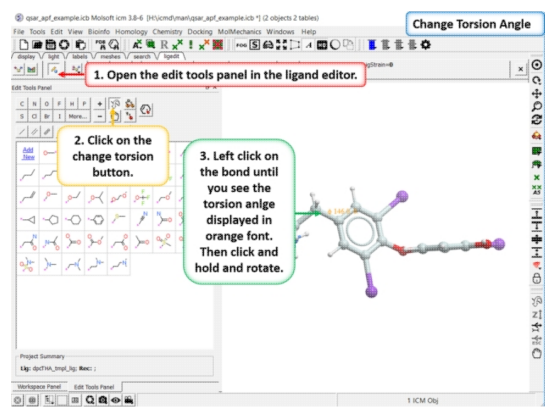
The default set of modifiers LIGAND.modifiers is just an sarray of smiles and it is hard-coded into ICM. You can add your own groups by clicking on the "Add New" button in the editor http://www.molsoft.com/gui/edit-ligand.html When sketching you need to add an attachment point - right click on the attachment point atom in the 2D editor and select attachment point.

You can also add manually through the command line:

LIGAND.userModifiers = LIGAND.userModifiers // "cl[c\*]nc(N)sl"
write system preference

### 10.5.4 Manually Change Torsion Angles

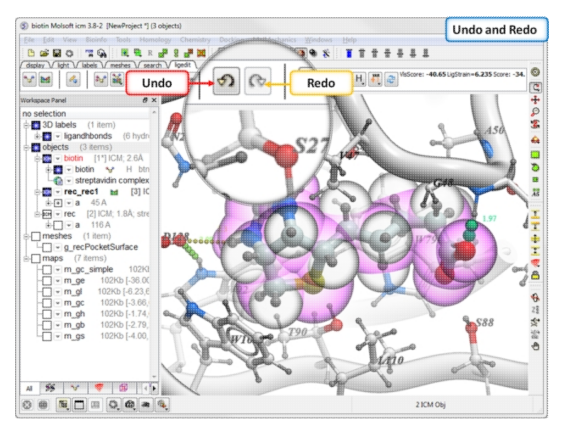
To manually change torsion angles - click on the torsion angle button in the Edit Tools Panel. Click on the bond you want to rotate until you see the torsion angle displayed in orange font. Then click and hold and rotate the torsion.



### 10.5.5 Undo and Redo

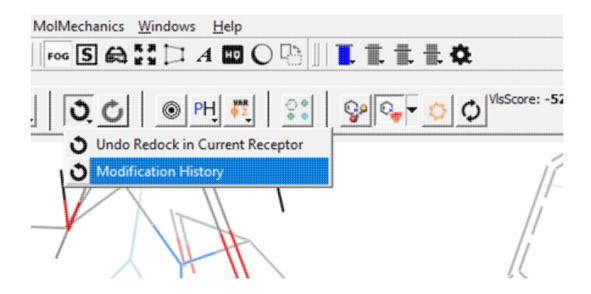
Every edit (even after docking) can be undone or redone.

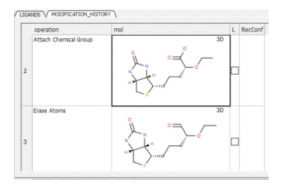
To undo or redo:



## 10.5.6 Modification History

You can view a table with a log of all modifications by clicking and holding on the "undo" button and choose "Modification history". This will produce a chemical table with a column (operation) describing the modification. Double click to load the modification into the receptor or use the "L" column to toggle the display on and off.



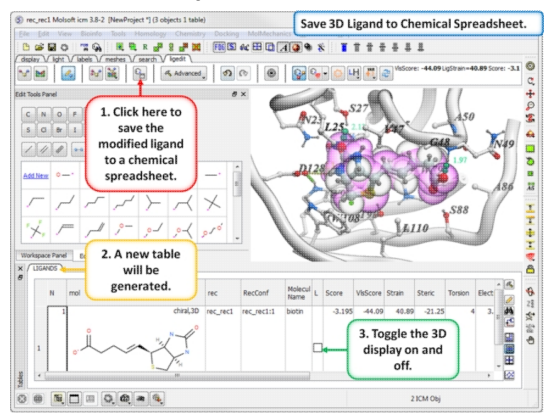


### 10.5.7 Save to Chemical Spreadsheet

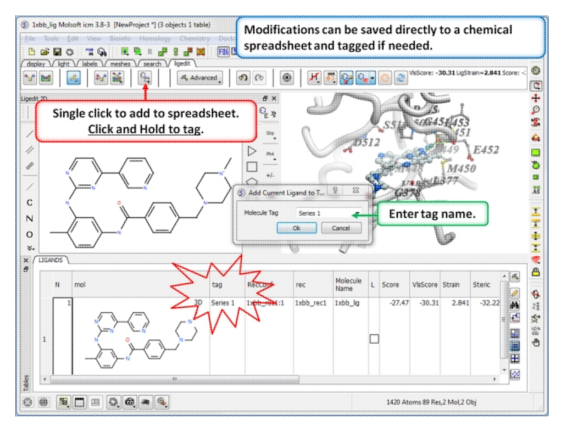
If you like a particular ligand and want to save the changes for use later, then you can save the 3D ligand to a chemical spreadsheet. You can then double click on the ligand in the spreadsheet to load it back into the editor or toggle the ligand on or off.

To save to a chemical spreadsheet:

- Edit the ligand.
- Click on the save to chemical spreadsheet button and a table called LIGANDS will be generated.



You can also tag a chemical with a unique name which is stored in the spreadsheet. To do this click and hold on the "Add Current Ligand to Table" button and enter a tag name. The ligand table can be grouped by this tag column (Right Click on the column and choose "Group View By").



10.5.8 After Editing the Ligand

#### After editing the ligand you can:

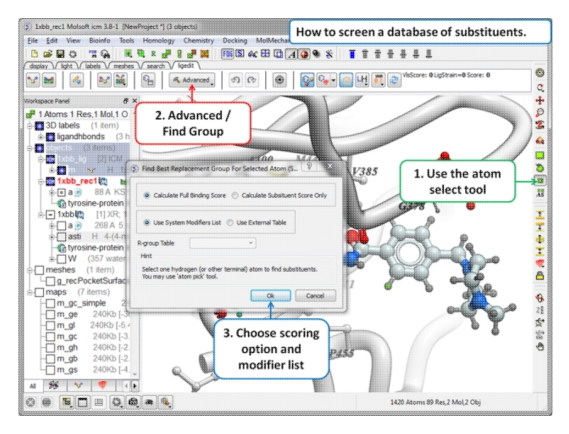
- Re-dock or minimize the ligand and evaluate the energy and score or view interactions such as hydrogen bonds and energy circles.
- Before docking if certain groups or distances need to be retained you can set distance restraints or tether atoms.
- You can try additional changes by screening a database of substituents.
- Add the ligand to a chemical spreadsheet.
- Save the docked ligand and receptor complex as a PDB file .

### **10.6 Virtual Screen: Find Best Replacement Group**

Available in the following product(s): ICM-Pro | ICM-VLS | ICM-Chemist-Pro

This option allows you to screen a database of substituents. You can use the ICM "in built" substituents or a database of your own.

#### To find best replacement group for selected atom:



- Setup the ligand and receptor.
- Select the atom you want to add a new replacement group to.
- Click on the Advanced/Find Group button.
- A dialog box as shown below will be displayed. Select whether you want the **substituent score only** to be evaluated (quick) or the **full binding** score.
- Select whether you want to screen the modifying groups built into ICM (see sarray of smiles called LIGAND.modifiers) or a table of your own modifier groups. If you choose your own table you will need to load the table (sdf file) into ICM and enter the name of the table into this dialog box or you can add modifiers to the sarray of smiles called LIGAND.userModifiers.
- ICM will add each fragment to the target atom and sample the energy and return a table ranked by score.

### **10.7 Multiple Position Group Scan**

This option allows you to scan groups and hetero atoms at multiple locations on the ligand. To do this:

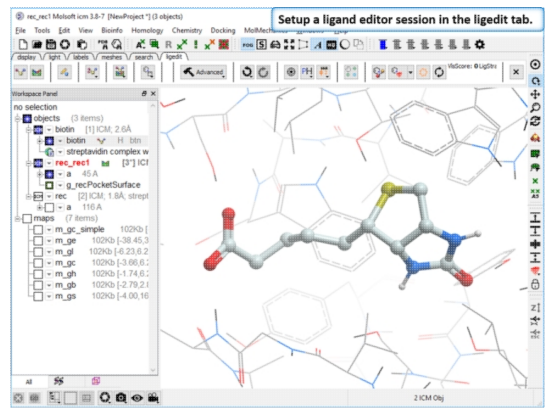
- Select the atoms on the ligand you wish to scan against.
- Select the groups from the R-group scan table (single click selects) and/or select the hetero atom scan.

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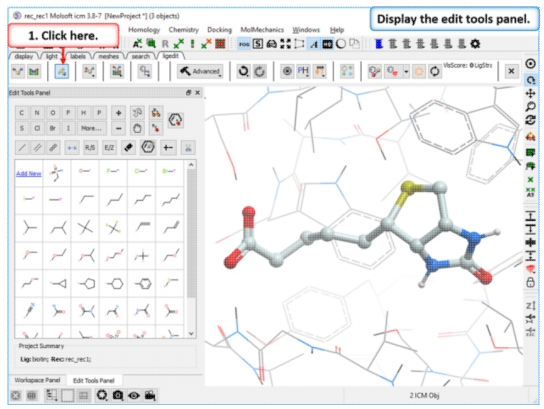
### 10.8 Bioisostere Scan

In drug design bioisosteres can be used to reduce toxicity, change bioavailability, alter metabolism and change activity of lead compound. Bioisosteres are chemical substituents with similar chemical or physical properties which produce broadly similar biological properties to another chemical compound.

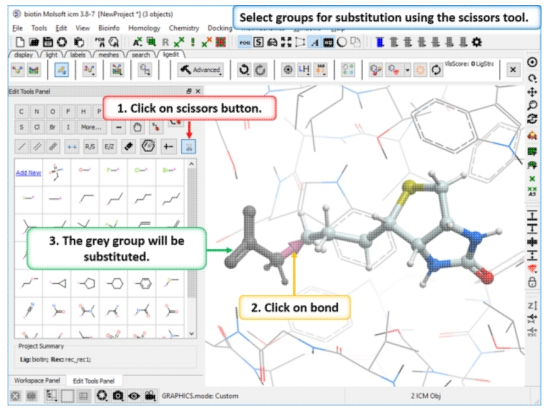
To scan for Bioisostere replacement groups:



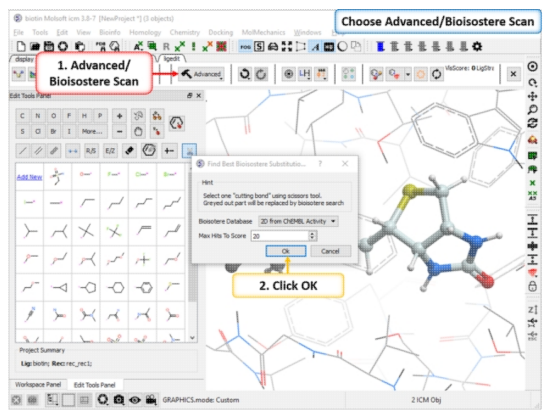
**Step 1:** Setup the ligand and receptor in the ligedit table.



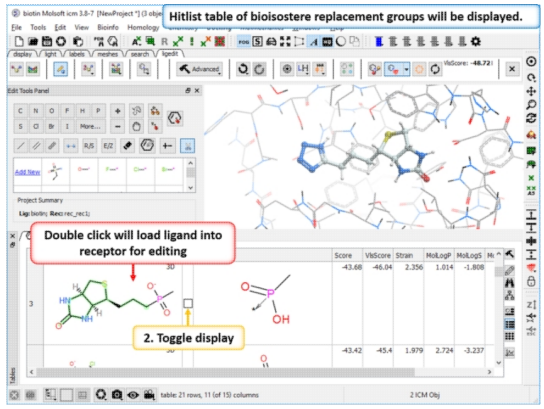
Step 2: Display the edit tools panel.



**Step 3:** Click on the scissors button and the click on the bond next to the group you want to scan against. The region that will be scanned will be greyed out.



Step 4: Select Advanced/Bioisostere Scan



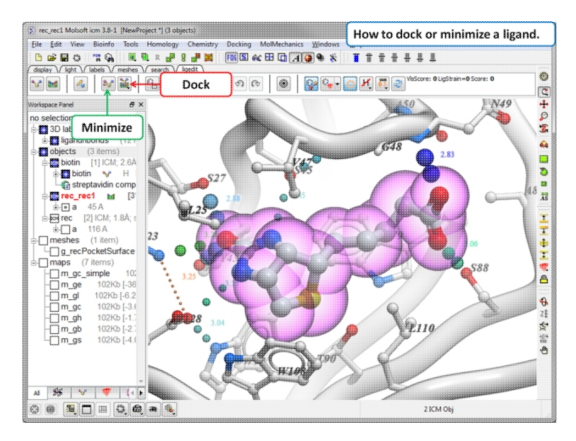
**Step 5:** A hitlist of bioisostere replacement groups will be displayed ranked by VLS score.

# 10.9 Dock or Minimize Ligand

```
Available in the following product(s): ICM-Pro | ICM-VLS | ICM-Chemist-Pro
```

Video|

How to Dock or Minimize a Ligand in the Grid maps:



- Setup the ligand and receptor.
- Click on the dock or minimize button (see image above).
- Check the Score and Strain in the top right panel.

If you dock a chemical spreadsheet will be displayed with the docking results. Use the check box in column 'L' to display the docked ligand or double click to load the ligand into the ligand editor.

A guide to the table columns:

- **Score:** This score does not incorporate ligand strain. Therefore it is VlsScore LigStrain. The lower the Score the better the predicted interaction.
- VLS Score: The ICM-VLS score used for ligand screening. The lower the Score the better the predicted interaction. VLS score is available with ICM-Pro + VLS and ICM-Chemist-Pro products. This score incorporates the strain of the ligand compared to the free state.
- Strain: The strain of the ligand. Lower values = less strain. (kcal/mol)
- **Steric:** Is the van der Waals interaction energy (sum of gc and gh van der waals). Current version of the score uses explicit van der Waals interaction energy calculation (no grids). (lower the better)
- Torsion: Number of torsions in ligand
- **Electro:** is the solvation electrostatics energy change upon binding (lower the better)
- Hbond: is Hydrogen Bond energy (lower the better)
- **Hydroph** is the hydrophobic energy in exposing a surface to water (lower the better)
- **Surface** is the desolvation of exposed h-bond donors and acceptors (lower the better)

#### How to undertake an all-atom minimization in cartesian coordinates:

- Note this option is available in ICM 3.8-4 and higher.
- Click and hold on the **Minimize** button.

• Choose Minimize in Receptor.

### 10.9.1 Change the Size of the Docking Region

#### How to change the size of the docking region:

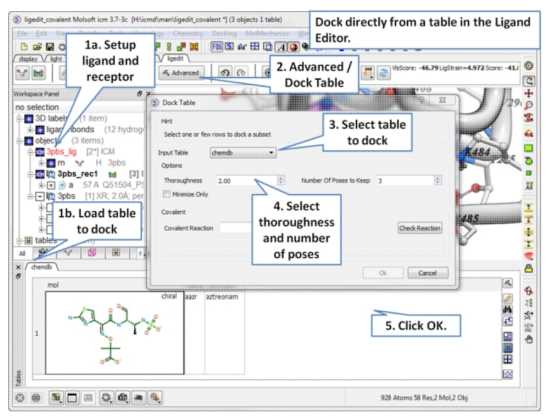
The purple box represents the region in which the energy maps are generated. If you want to change the size of this region you can do so by displaying the box (go to Advanced menu) and clicking and dragging on the corners of the pupple box. The maps will then be remade on the fly.

## 10.10 Dock from Table

Available in the following product(s): ICM-Pro | ICM-VLS | ICM-Chemist-Pro

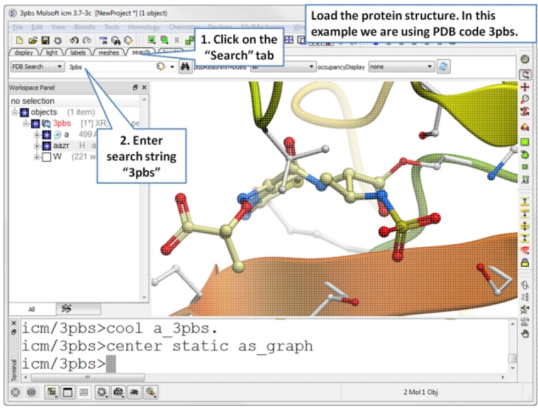
To dock from a table in the ligand editor.

- Setup the ligand and receptor in the ligand editor.
- Read in the sdf file or chemical spreadsheet which contains the chemicals you wish to dock.
- Select the **Advanced** button in the ligand editor and select Dock Table.
- In the dialog box select the thoroughness (length of simulation) and the number of conformations.

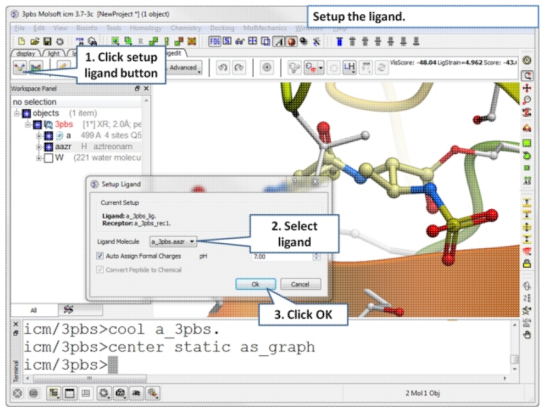


### 10.10.1 Covalent Docking from Table

To undertake covalent docking from a table:



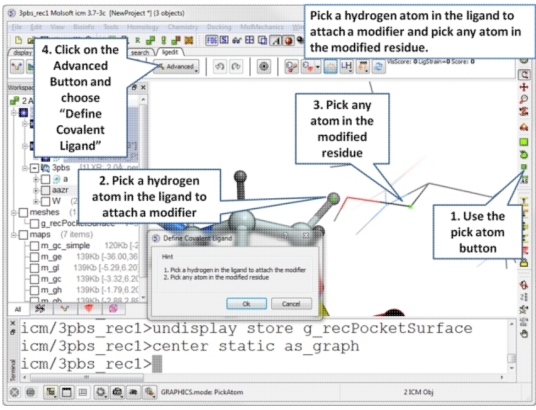
Read in the PDB file you wish to dock to containing the covalently linked ligand.



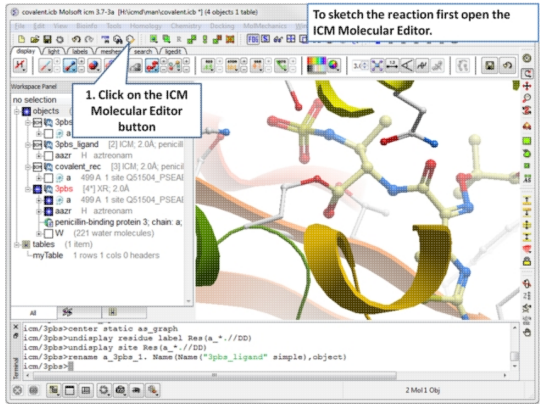
Setup the ligand in the ligand editor.

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<pre>* icm/: icm/: suicm/: %  %  %  %  %  %  %  %  %  %  %  %  %</pre>	More Options  Create Ligand Pocket Surface		4. Click OK	x ,

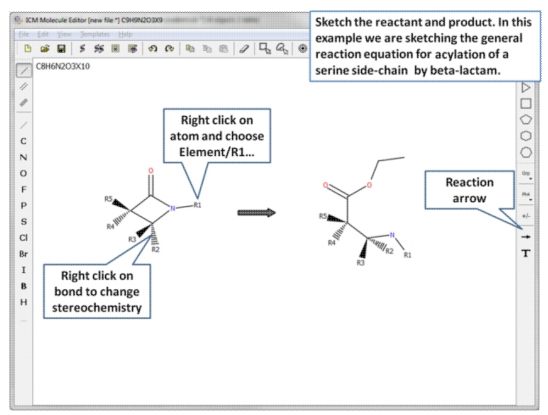
Setup the receptor in the ligand editor.



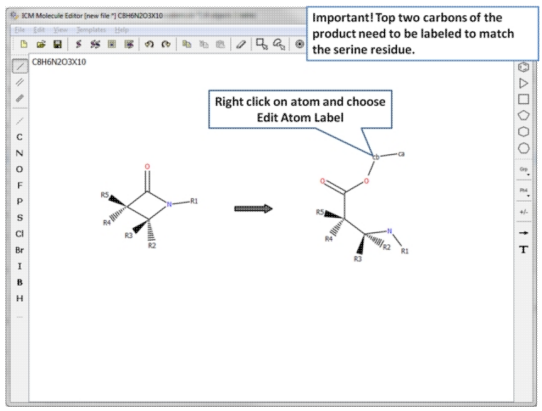
Pick a hydrogen atom in the ligand to attach a modifier and pick any atom in the modified residue. Next, click on the Advanced button and choose "Define Covalent Ligand".



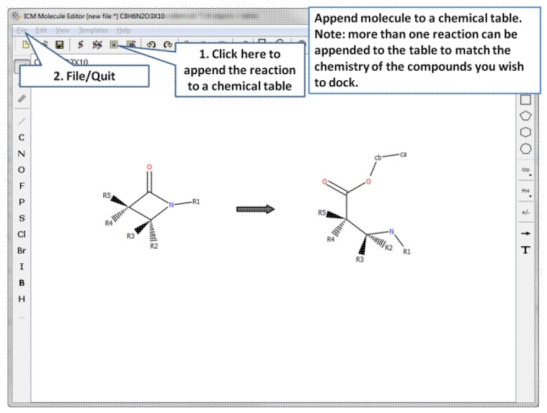
Open the molecular editor. Note you may first need to click on the display tab to see the Molecular Editor button.



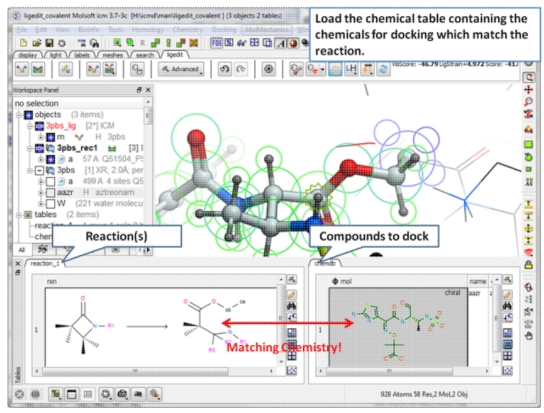
Sketh the reactant and product.



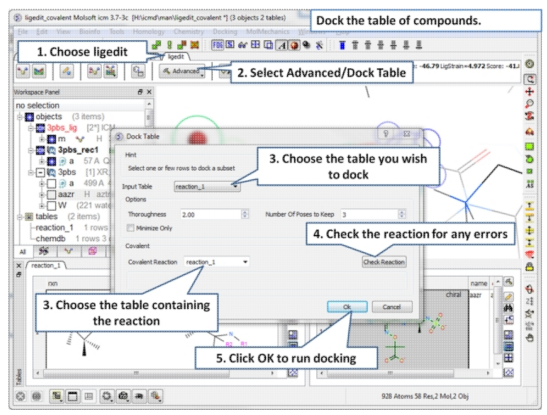
Two of the atoms need to be labeled to match the residue in the receptor.



Append the reaction to a chemical table. Note more than one reaction can be added to the table to match the ligands in your chemical table that you wish to dock.



Load the chemical table containing the ligands you wish to dock.

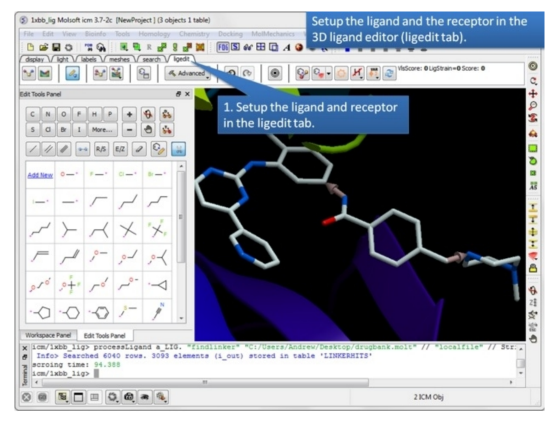


Choose Advance/Dock Table and select the covalent options shown.

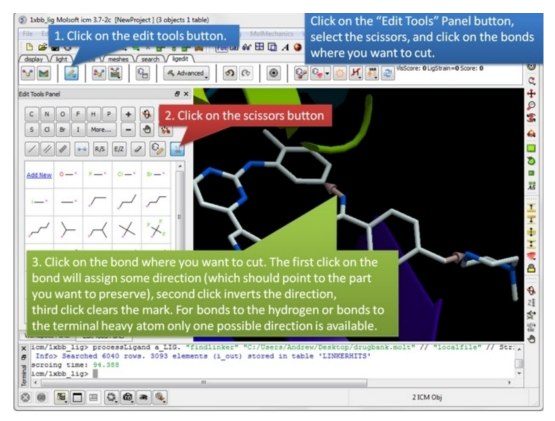
# 10.11 Fragment Linking

Available in the following product(s): ICM-Pro | ICM-VLS | ICM-Chemist-Pro

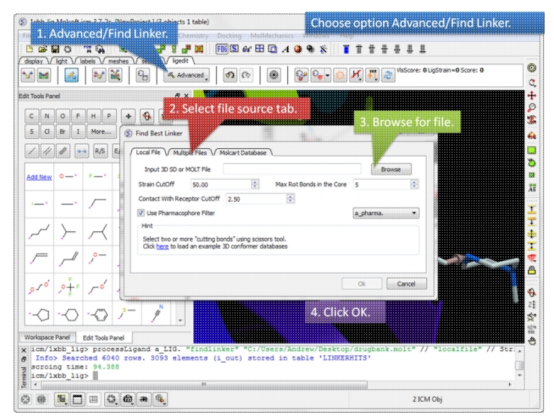
This feature finds the best scaffold replacement or linker fragment from a database. It lets you interactively select two or more points of connection or replacement and search for possible fragments in the 3D database (SDF or Molsoft local database MOLT). The cutting bonds used to define scaffold are matched against a 3D database with a certain RMSD cutoff (usually very small). The best superposition is taken for the original compound and the chemical is minimized. The result is returned as a standard Ligand Editor hit list where results can be easily browsed, sorted and compared. Linker prediction can be made in the presence or absence of the receptor. Filters such as energy strain, maximum number of rotatable bonds and contact distance to the receptor can be used.



**Step 1:** Setup the ligand and/or the receptor in the 3D ligand editor. If you have fragments already made you should select the fragments and then choose "Selection" in the drop down menu in the ligand setup step. If you do not wish for the receptor to be considered when sampling linkers then just setup the ligand only. Once the ligand and/or receptor are setup it is generally a good idea to minimize the ligand to remove any clashes and optimize the geometry.

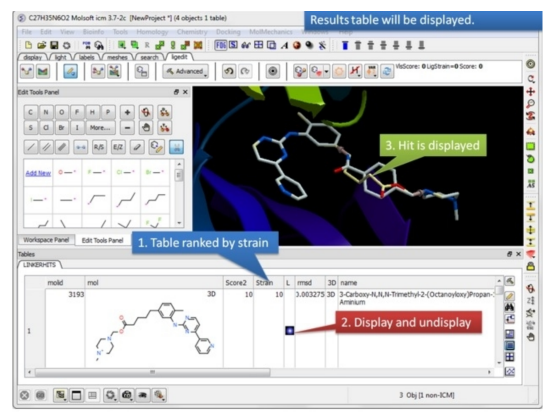


**Step 2:** Once the ligand and/or the receptor has been setup click on the edit tools button. A panel of editing tools will be displayed, click on the scissors button. Mark two or more bonds using the scissors tool. The first click on the bond will assign some direction (which should point the part you want to preserve), second click inverts the direction, third click clears the mark. For bonds to the hydrogen or bonds to the terminal heavy atom only one possible direction is available.



**Step 3:** Select the **Advanced** button and choose the option **Find Linker**. A dialog window will be displayed as shown above. You can choose to screen one ore more SDF files containing 3D coordinates or a molt file that is stored locally on your machine or on MolCart.

- Provide a path to the 3D SDF or Molt file. We have provided some examples of 3D conformer databases here ftp://ftp.molsoft.com/molcartdb/3D
- Strain Cutoff any ligand exceeding this Strain value will not be placed in the results table. If you are using a ligand directly from the PDB it is a good idea to minimize the ligand and check the initial strain or increase this cutoff.
- Max Rot Bonds in the Core any ligand exceeding this number of rotatable bond value will not be placed in the results table.
- **Contact with Receptor CutOff** any ligand exceeding this distance will not be placed in the results table. This value is significant if the receptor is included in setup.
- The **use pharmacophore filter** allows you to match linkers with a pharmacophore object. Learn how to make a 3D pharmacophore object here.



**Step 4:** Once the fragment fitting has completed a table of results will be displayed. The table is ranked by strain with the top hit in row 1. You can browse the hits using the toggle buttons in the "L" column.

- Strain is the energy strain of the ligand.
- **Score2** is the same value as strain (possible place holder for future release).
- **RMSD** is the RMSD to the original fragment before linker connection.

**Three Point Core Replacement** Three cores can be linked by selecting them with the scissors button as shown in Step 1 above. Number of hits will be significantly lower then in 2 point replacement because the extra anchor adds some restraint. We recommend to use larger database for the search (e.g. chembridge100K.molt) and make sure that ligand is minimized before searching.

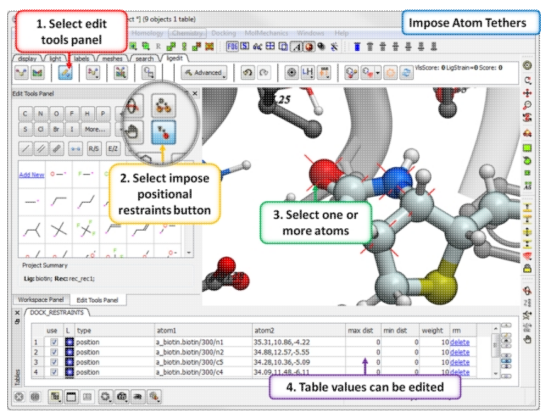


### **10.12 Impose Tethers or Distance Restraints**

### 10.12.1 Impose Ligand Atom Tether

#### To impose tethers to selected atoms of the ligand before redocking.

- Setup the ligand and receptor.
- Click on the edit tools panel button.
- Click on the impose positional restraints button.
- Select the atoms you wish to tether in the ligand.
- A table listing the tethers will be displayed and red crosses through the atom indicates a tether.
- Click on the re-dock button and the atoms selected will remain tethered in place.



**About the table:** The tethers can be edited and deleted using the table. Double click on a value in the table to edit it.

- Use Check "use" to use the tether in docking.
- L Toggle the tether display on or off.
- **Type** Type of tether "position" indicates a positiional restraint (tether).
- Atom1 ICM selction language for the tethered atom. Atom 1 can also be SMARTS expression which define the atom (for both distance and position) The matched atom should be marked with '\*'. For example oxygen in the double bonded to the 5-member ring in biotin can be defined as: NC(=[O\*])N
- Atom2 xyz position (3 comma separated numbers)
- max dist Sets the maximum distance for the tether.
- min dist Sets the minimum distance for the tether.
- weight the overall weighting factor for the tether penalty term.
- **rm** Delete the tether.
- err Deviation from the position after docking.

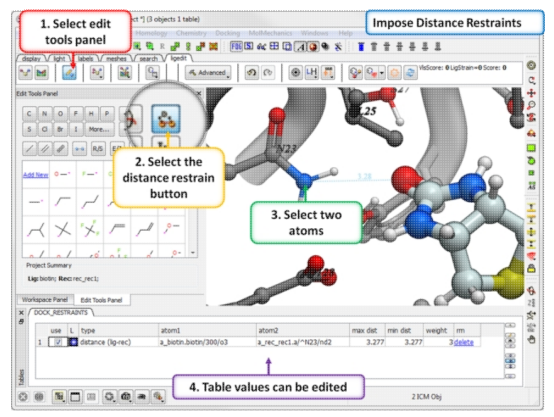
### 10.12.2 Impose Distance Restraint

There are three types of distance restraint you can make:

- 1. lig static rec : fixed side-chain (weighted tether)
- 2. lig flex rec : flexible side-chain (drestraint)
- 3. lig lig : drestrains

#### To impose distance restraints before redocking.

- Setup the ligand and receptor.
- Click on the edit tools panel button.
- Click on the distance restraint button.
- Select two atoms to form the distance restraint. The distance restraint can be between two atoms in a ligand or between the ligand and receptor
- A table detailing the distance restrints will be displayed and red crosses through the atom indicates a tether.
- Click on the re-dock button and the atoms selected will remain tethered in place.



**About the table:** The distance restraints can be edited and deleted using the table. Double click on a value in the table to edit it.

- Use Check "use" to use the distance restraint in docking.
- L Toggle the distance restraint display on or off.
- **Type** Type of restraint. There are three types of distance restraint
  - 1. lig static rec : fixed side-chain (weighted tether)
  - 2. lig flex rec : flexible side-chain (drestraint)
  - 3. lig lig : (drestraint)
- Atom1 ICM selction language for the first atom in the distance restraint.
- Atom2 ICM selction language for the second atom in the distance restraint.

- **max dist** Sets the maximum distance for the restraint.
- **min dist** Sets the minimum distance for the restraint.
- **weight** the overall weighting factor for the distance restraint penalty term.
- **rm** Delete the distance restraint.
- err Deviation from the position after docking.

**Note:** The area within min-max distance is zero (flat-bottomed). With a weight of 1. you will have fair coverage of conformations with minor violations, going up to 5 or 10 will pretty much force the system to always stay there, below 1. you will only have a sort of weak steering force.

## **10.13 Docking Templates**

You can use either substructure or Atomic Property Field Templates for docking.

- Setup the ligand and receptor.
- File/Open a 3D template structure and place in the pocket.
- Click on the Edit Tools panel button.
- Click on the **Impose Template Restraint** button.
- Select the template object from the drop down menu and choose whether you wish to impose substructure or 3D pharmacophore APF template.
- A table will be displayed called DOCK\_RESTRAINTS where you can turn the template on or off.

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# **10.14 Defined Flexible Receptor Groups**

You can allow certain defined residues in the receptor to be flexible during docking. To do this:

• Setup the ligand and receptor.

- Select the residues you wish to be flexible using the residue selection button.
- Select Advanced/Defined Flexible Receptor Groups
- Choose full side-chains or hydroxy and water.
- The side chains you have selected will then be displayed in thicker stick and will be flexible when you dock or edit a ligand.

**NOTE:** It is recommended to not select more than 3 residues to be flexible.

## 10.15 Side Chain Refinement

You can refine the side-chains around a docked ligand by:

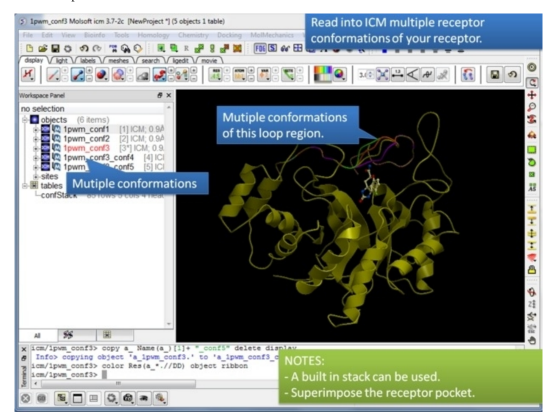
- Setup the ligand and receptor.
- Select the residues you wish to be flexible using the residue selection button.
- Advanced/Receptor Side Chain Refinement.

## **10.16 Multiple Receptor Docking**

Available in the following product(s): ICM-Pro | ICM-VLS | ICM-Chemist-Pro

|Video|

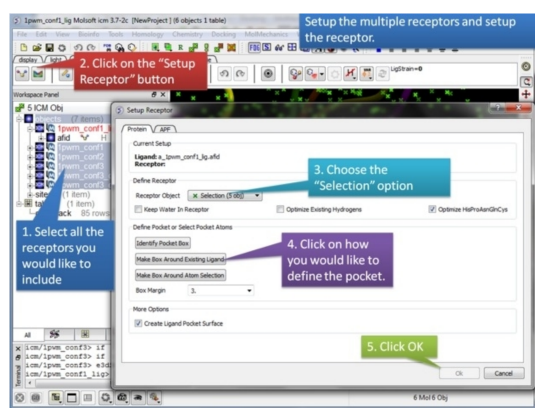
Multiple receptor conformation docking has been shown to be an efficient way of incorporating receptor flexibility into ligand docking (see Bottegoni et al J.Med.Chem 2010). You can dock to multiple receptor conformations in the 3D ligand editor by selecting the superimposed structures receptor setup stage. 4D maps will be generated encompasing all the receptors selected.



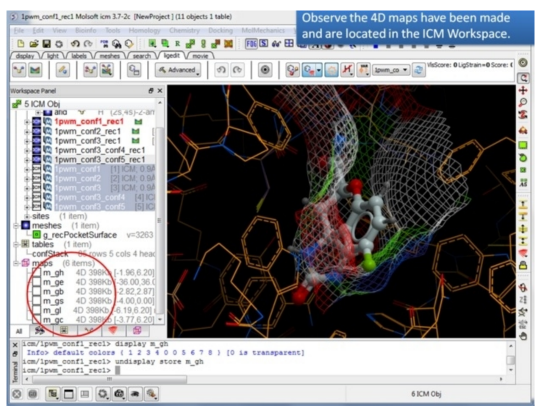
**Step 1:** Read into ICM the reeptor conformations you would like to dock to. If you have a conformation stack already loaded into ICM (e.g. after modeling a loop region) ICM can use that.

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<pre>one of the objects in the ligand binding pocket.  icm/lpwm_conf3&gt; copy a_ Name(a_)[1]+ "_conf5" delete display info&gt; copying object "a_lpwm_conf3.' to 'a_lpwm_conf3_conf5.' icm/lpwm_conf3&gt; color Res(a_*.//DD) object ribbon icm/lpwm_conf3&gt;</pre>	◆ 注 ☆ 塗 ●
3	6 Mol 5 Obj

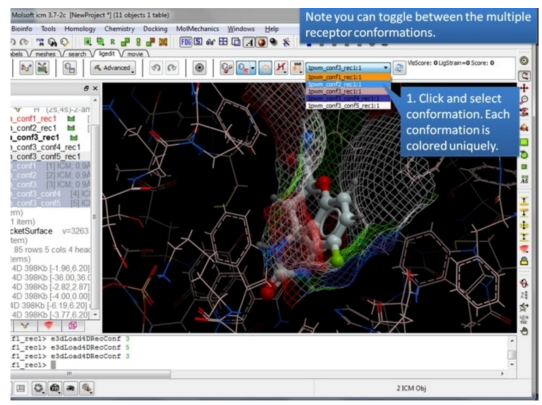
Step 2:Setrup the initial ligand you would like to dock.



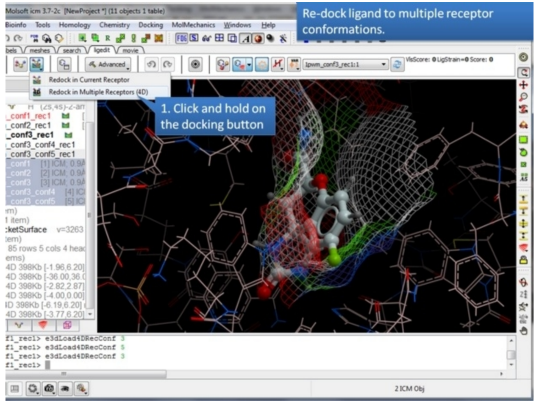
# **Step 3:** Select the receptors you would like to dock to. Make sure they are correctly superimposed.



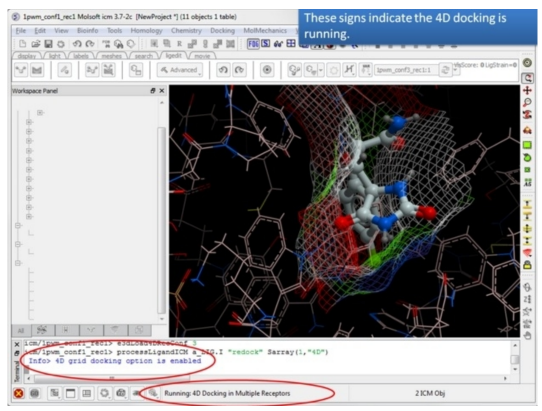
**Step 4:** You will see that 4D maps have been generated and stored in the ICM Workspace.



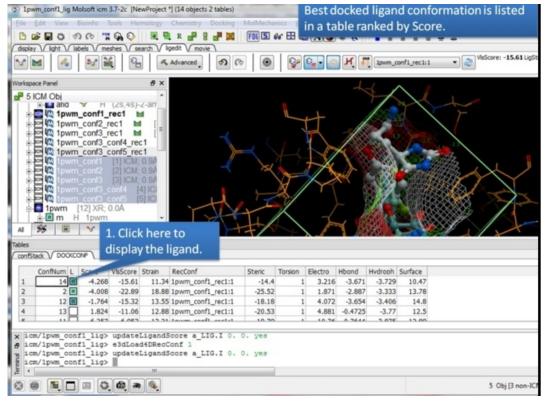
Step 5: You can easily toggle between each receptor to display them.



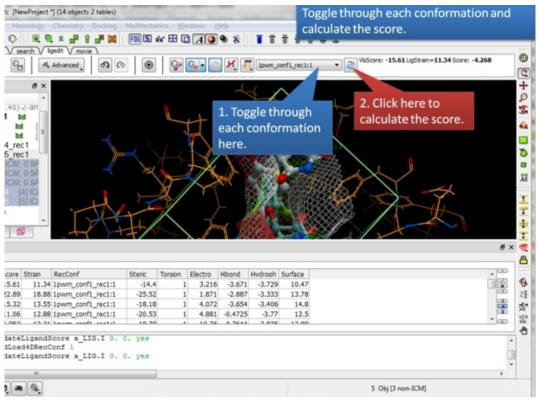
# **Step 6:** Click and hold on the docking button to select "Re-dock to multiple Receptors (4D).



Step 7: The docking will start running.



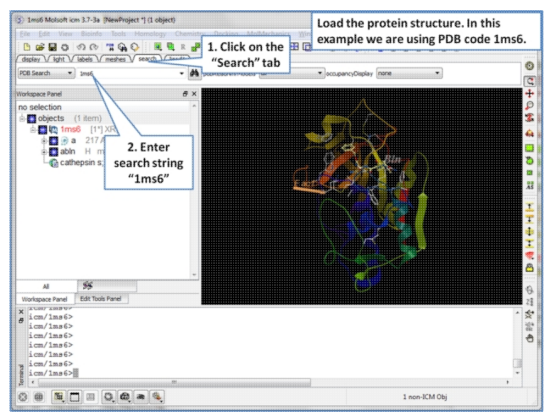
Step 8: Use the interactive table to view the results.



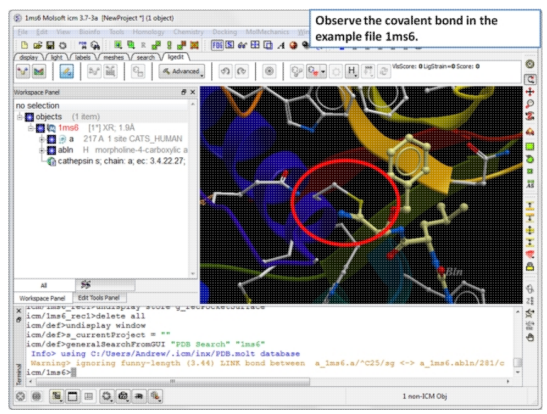
Step 9: Toggle through the results and view the score.

# 10.17 Covalent Docking in the Ligand Editor

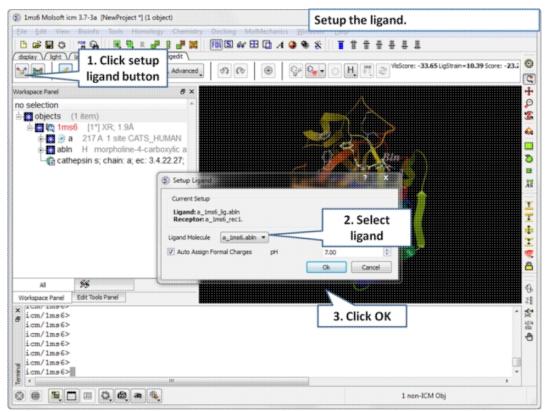
Available in the following product(s): ICM-Pro | ICM-VLS | ICM-Chemist-Pro



**Step 1:** Load the protein structure via the Search tab. In this example we will use PDB structure 1ms6.



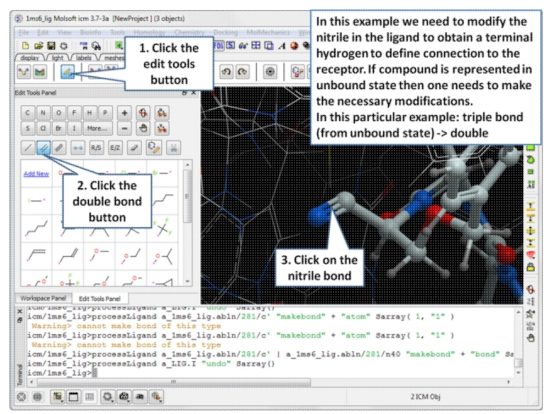
**Step 2:** The structure 1ms6 has a covalent bond between the ligand "abln" and the receptor residue C25.



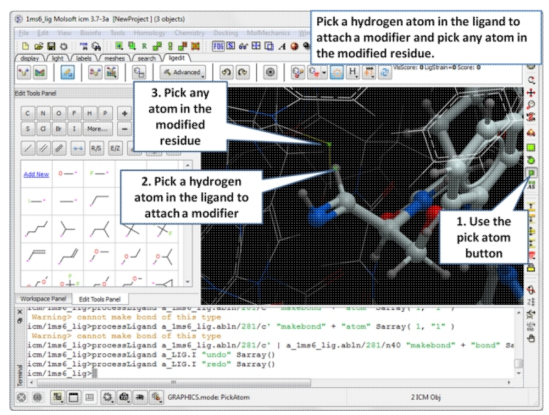
**Step 3:** Choose the ligand "abln" and setup the ligand in the regular way.

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Al 55	Receptor Object a_Ims6.	Coptimize Existing Hydroger 3. Define pocket	ns IV Optimize HisProAsnGinCys	
Al 55 Workpace Panel Ei Trumy ramsv- s icm/lms6>set icm/lms6>li( icm/lms6>if icm/lms6>if icm/lms6>if icm/lms6>li( icm/lms6]if	More Options Create Ligand Pocket Surface		4. Click OK	◆ 注 ☆ 常 令

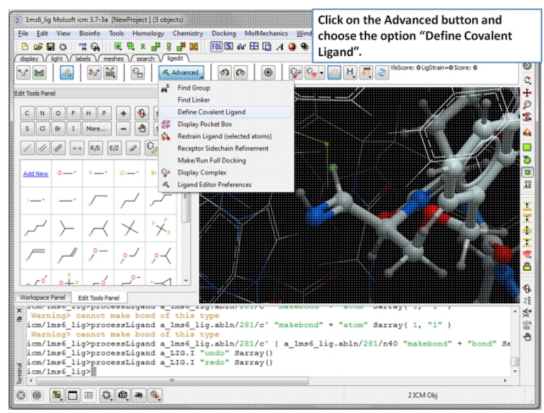
**Step 4:** Setup the receptor and make the maps. It is important to make sure the box encompasses the modified residue C25.



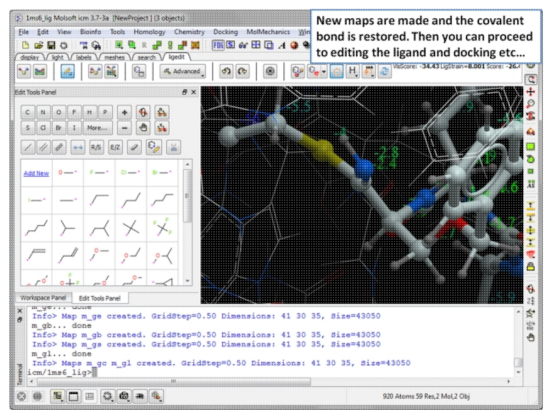
**Step 5:** Modify the nitrile in the ligand to obtain a terminal hydrogen to define the connection to the receptor.



**Step 6:** Pick the hydrogen atom in the ligand where the modifier will be attached and then pick any atom in the modified residue.



**Step 7:** Click on the Advanced button and choose the otpion "Define Covalent Ligand".



**Step 8:** New maps will be made and the covalent bond is restored. Then you can re-dock, minimize and edit the ligand in the usual way inside the ligand editor.

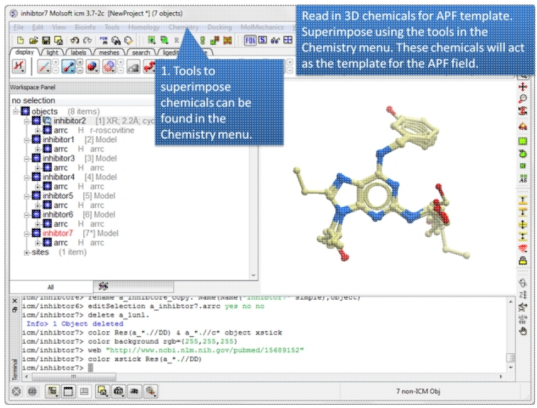
# **10.18 Dock to Atomic Property Fields**

Available in the following product(s): ICM-Pro | ICM-VLS | ICM-Chemist-Pro

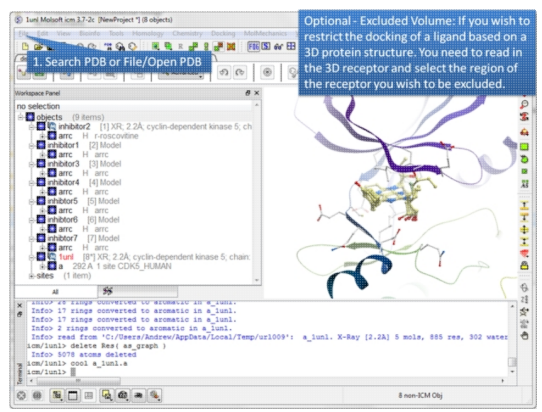
The Atomic Property Fields (APF) superposition/alignment method was first reported by Maxim Totrov PhD (Principal Scientist - MolSoft) at the 2007 233rd American Chemical Society National Meeting, Chicago, IL USA and then published here.

APF is a 3D pharmacophoric potential implemented on a grid. APF can be generated from one or multiple ligands and seven properties are assigned from empiric physico-chemical components (hydrogen bond donors, acceptors, Sp2 hybridization, lipophilicity, size,

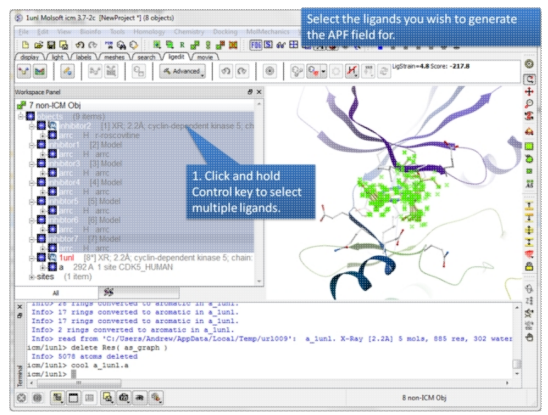
electropositive/negative and charge).Here we describe template APF superposition whereby the APF is generated from a single or multiple template and is then globally optimized with the internal force-field energy of the ligand.



**Step 1:** Read in the chemicals that will make up the Atomic Property Field you wish to dock to.



**Step 2:** Certain regions of the pocket can be excluded from docking by selecting an excluded volume.



Step 3: Select the ligands you wish to generate an APF field for.

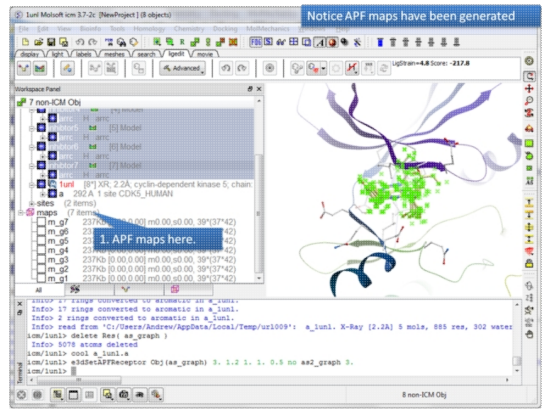
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Step 4: Setup the APF template. The APF parameters are:

• wAPF = relative weight of APF with respect to internal force-field energy. Adjusts tolerance to strain relative to

enforcement of chemical alignment.

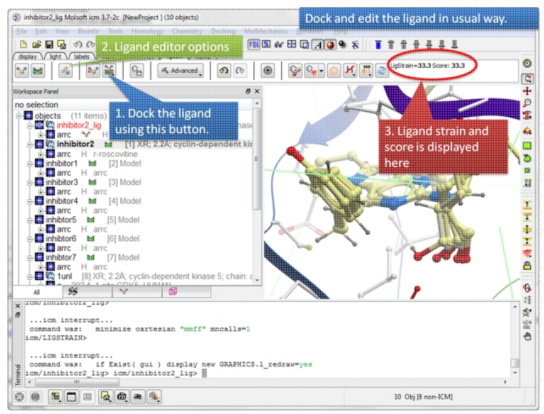
- wLipo = relative weight of lipophilic atoms.
- wPolar = relative weight of polar atoms.
- rAPF = radius of APF gaussians. Can be used to adjust 'softness'/tolerance of molecular alignment.
- grid step = use with caution, potentially can be reduced to allow very 'sharp' alignments.



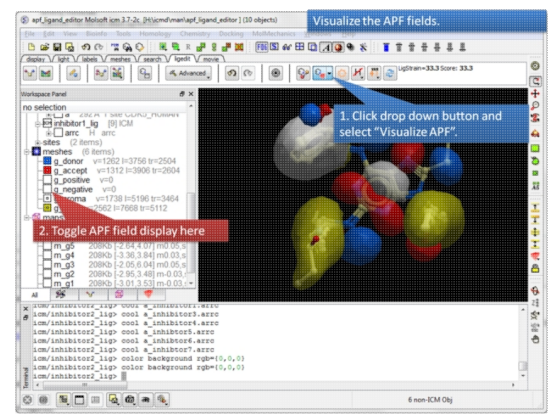
Step 5: The APF maps are listed in the ICM Workspace left hand side.

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Step 6: Select the ligand you wish to dock into the APF maps.



**Step 7:** Dock the ligand using the ligand docking button and note the Score.



Step 8: Visualize the APF maps.

# 10.19 How to Save Ligand Receptor Complex as PDB

```
Available in the following product(s): \texttt{ICM-Pro} \mid \texttt{ICM-VLS} \mid \texttt{ICM-Chemist-Pro}
```

To save a docked ligand as a PDB file in complex with the receptor:

Advanced/Export Complex

# 10.20 Export Docking Project.

Available in the following product(s): ICM-Pro | ICM-VLS | ICM-Chemist-Pro

This option makes it easy to switch from the 3D ligand editor environment (ligedit tab) to the docking menu. All the docking setup steps you have created in the 3D ligand editor are converted so you can use the docking and screening options in the docking menu.

- Setup the ligand and receptor in the ligedit tab. Perform edits to the ligand etc...
- Click on the Advanced button in the ligedit tab
- Click on Export to Docking Project
- You can choose to run a docking job immediately with the current ligand or just create the docking files. Check or uncheck the **Run Docking Job** button.
- The docking files will be placed in your working directory (Tools/Change Working Directory). The files will have the docking project name dock\_name\_of\_receptor and be stored in a directory

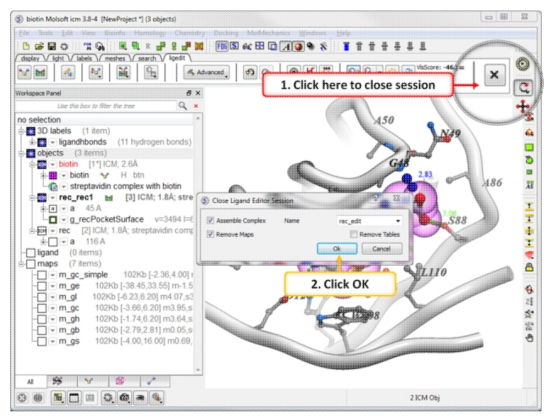
called dock\_name\_of\_receptor

• You can use these files to perform screening using the docking menu. For example you would set the docking project name as dock\_name\_of\_receptor.

## 10.21 Close and Clean Up Project

The Close Session option allows you to close and clean up your ligand editor environment. Click on the close button and you can:

- Assemble the complex
- Remove maps
- Remove tables



# **11 Working with Tables and Plots**

One of the easiest ways to store, sort and display data in ICM is by the use of a table. In most cases tables are automatically created, for example, if you search for a PDB file or when you load a compound database (SDF file). It is also possible for you to create your own table. Once a table is created, ICM provides easy to use tools to sort, add, edit and plot data.

Here we will concentrate on describing the actions you can perform on a table once it has been read into ICM. We will start by describing a simple table. Actions which can be performed on chemical tables are described in the section entitled Working with Chemical Spreadsheets.

- Standard ICM Tables
- Chemical Spreadsheets
- Insert Interactive Objects into Table Cell
- Plotting Table Data
- Principal Component Analysis
- Learn and Predict
- Cluster

#### A standard ICM table:

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2	101623	m1	-32.90	42	0	-5.78	-7.62	-38.36	📕 🔲 display Hbonds	
3	101662	m1	-34.12	36	1	-7.77	-6.08	-33.31	display docked struct	
4	101671	m1	-36.17	48	4	-5.90	-6.84	-42.93		
5	101722	m1	-34.70	36	0	-7.97	-6.45	-31.44	calculate distances	
6	101781	m1	-36.65	54	3	-6.55	-7.37	-46.09		
7	101784	m1	-35.07	38	2	-7.11	-6.26	-34.98		
8	101792	m1	-32.51	32	0	-6.11	-6.64	-35.07		
9	101813	m1	-47.90	39	0	-11.88	-6.29	-39.66		

## **11.1 Standard ICM Tables**

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

#### 11.1.1 Generate New Table

**NOTE:** To generate a new chemical table use the options in Chemistry/Create Empty Chemical Table.

#### To generate a new empty table:

- File/New and select the **Table** tab and a window as shown below will be displayed.
- Enter the number of rows and columns you wish to include in your table and whether you wish to add a column with chemical data.
- If you wish to make a chemical table (chemical spreadsheet) select the **Chemical Column** box.

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String Columns	2	*							
Integer Columns	0	<b>A</b>							
Real Columns	0	<b>A</b>							
🔲 Chemical Column									
					Ok		Cancel		Help

## 11.1.2 Reading a Table

A table can be read and saved as a \*.csv, \*.tsv or a .tab file. Saving or reading your table as a csv (comma separated value) file enables the table to be transfered or loaded from other applications such as Microsoft Excel. A chemical file such as an .sdf file can also be viewed as a chemical spreadsheet.

#### A table can be read into ICM by selecting:

• File/Open and then selecting the table you have saved (e.g. csv, sdf, tab).

**NOTE:** If you have loaded a table and it is not displayed it may be because the table window is hidden. To display the table, select the window menu and select table see the Window Menu Section.

## 11.1.3 Saving a table

To save the whole table:

• To save a table right click on the table header tab and select Save As..

#### To save a row selection:

- Select a row(s)
- Right click and choose Save Selection As or Save Selection As Csv + Headers

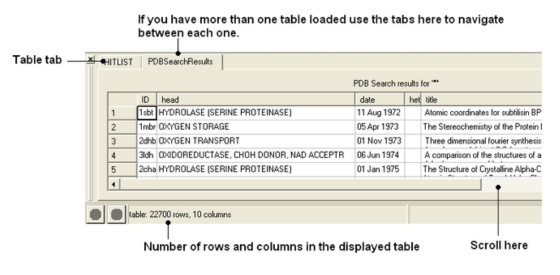
## 11.1.4 Basic Table Navigation

To view the contents of a table you can move the table up and down using the scroll bars on the side and bottom of the display.

**NOTE:** If you have loaded a table and it isnt displayed it may be because the table display isnt selected. To select the table display,

select the window menu and select table (See Window Menu Section).

If you have read more than one table in ICM you can select a table by clicking the tab on the top of the table (See Below).



**NOTE:** Double clicking on the tab allows two tables to be displayed at once. Double clicking again returns to the default table layout.

**NOTE:** Information regarding the number of rows and columns within a table is displayed at the bottom of the table.

## If you would like the table to be the main window in the graphical user interface:

• Select Windows/Table->Main

## 11.1.5 Table View (Grid Layout)

#### To change the table view (layout):

- Select the columns you wish to display in grid view. No selection will place all columns in grid view
- Right click on a table row and select Table View
- You can view the table in **Grid View** and toggle between grid and standard view. You can define your own grid using the **Custon Grid** option or display the table in **Form View**.

#### NOTE: You can save a table view.

## 11.1.6 Change Row Height

To change the row height:

- Click and drag on the row separator
- Hold the Ctl key to change the height of all rows.

## 11.1.7 Table View Save

Once you have a table view that you want to keep. You can save it by:

- Right click on a table row and select Store Views
- Select Save Current View
- Enter a name for the table view and you can return to that view by repeating the first two steps above.
- You can rename, delete or restore view by right clicking on the name of the table view.

## 11.1.8 Table Search

#### To search a table:

- Right click on a table row and select **Find and Replace**. You can also use CTRL F.
- Enter a search string.
- Press the **Find** button.

## 11.1.9 Table Color

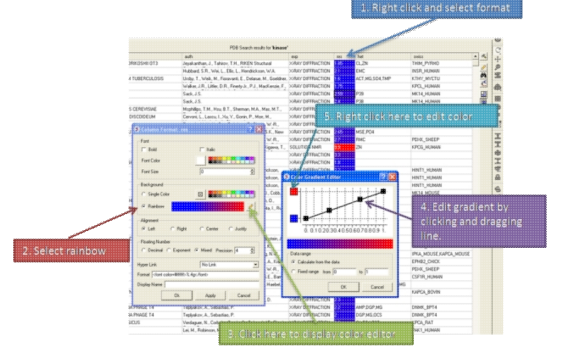
#### You can color your table based on values within a column by:

#### Single Color

- Right click on the column header and select Format, Color, Action.
- In the **Background** panel select the color you desire using the option **Single Color**.

#### **Color by Rainbow**

- Right click on the column header and select Format, Color, Action.
- In the **Background** panel select the option **rainbow.** To edit the range of values relating to each color click on the pencil (edit) button as shown below.



#### **Color Range - Advanced**

- Right click on the column header and select Format, Color, Action.
- In the Background panel select the option Range/Advanced

Column Format: molLogP									
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Comment									
Ok Apply Cancel									

• A dialog as shown below will be displayed where you can specify ranges and colors for each range. If two colors for range are equal then the whole range will be colored with single color, otherwise color will be interpolated within the range.

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## 11.1.10 Table Font

- Right click on the column header and select Format.Change the font using the options in the Font panel.

## 11.1.11 Table Alignment

- Right click on the column header and select Format.
- Change the font using the options in the **Alignment** panel.

#### Rows can be colored by marking them as described here

## 11.1.12 Mark a Row

A row in a table can be marked and grouped by a label which enables the row(s) to be selected easily at a later time.

#### To mark a row

- Right click on the row in the table you wish to mark. Or select multiple rows and then right click.
- Select Mark Row/ and then choose a number. In the GUI the number of rows that can be marked is limited to 5 but this can be

increased using the command line command.

- A row that is marked will be colored each number is assigned a
- color. The coloring can be changed in the gui tab in preferences.

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				Select M	arked Rows 🕨	· [ _	1		
	ID	head	6	Copy Cel	Ctrl+C		2		source
1	1hto	LIGASE		Paste	Dtrl+V		3	ine	MYCOBACTERIUM TUBERCULO
2	1hi1	RNA POLYMERASE		Print Tab	le Ctrl+P	1	4	acteriophage	BACTERIOPHAGE PHI-6
3	1htq	LIGASE	T	01701701	muncopy crystalograp	8		ked glutamine	MYCOBACTERIUM TUBERCULO
4	1htv	HORMONE/GROWTH FACTOR		01/01/01	crystal structure of des	day.		sulin	HOMD SAPIENS
5	1htw	STRUCTURAL GENOMICS, UNKNOWN FUNCT	ION	01/01/01	complex of hi0065 with	h ad	lp and magnesiur	n	HAEMOPHILUS INFLUENZAE
6	1kpr	HYDROLASE		01/01/02	first structural evidence	e of	a specific nhibiti	on of	DABOIA RUSSELLI PULCHELLA
7	1v8a	TRANSFERASE		01/01/04	structure of hydroxyeth	with	iazole kinase pro	tein from	PYROCOCCUS HORIKOSHII
8	1:0p	MEMBRANE PROTEIN	•	01/01/04	structure of the n-termine	inal	domain of the ad	enylyl	DICTYOSTELIUM DISCOIDEUM
9	2dcc	Hydrolase		01/01/06	x-ray crystal structure a	anal	ysis of bovine sp	een cathepsin	BOS TAURUS
		Right click here and	sele	ect Marl	Row		Coloring	elates to	numbers

#### To select marked rows

- Right click on the table and choose **Select Marked Rows** and choose a number which relates to the marked rows as described earlier.
- Selected rows will be highlighted blue once rows are selected a number of right click options are activated such as copy selection to new ICM table.

## 11.1.13 Table right click options

Right-click options vary according to where you click and what is selected. The options are intuitive, for example options that are performed on the whole table (eg Save and Delete) are performed by right-clicking on the Table tab. Other right-click options vary according to whether the row or column is selected or not.

## 11.1.14 Rename a Table

#### To rename a table:

- Right click on the table tab and select rename.
- Enter a new name and select OK.

#### Right click - / PDBSearchResults PDBSearchResults Select ID hea Clone 1hto LIG/ 1 Delete 2 1bi1 BNA Rename.. 3 1htg LIG/ Save As... 4 1htv HOF

## 11.1.15 Clone a Table

• Right click on the table tab and select clone.

### 11.1.16 Delete a Table

• Right click on the table tab and select delete.

## 11.1.17 Page Setup

Before printing a table you can change the orientation and scale.

To do this:

• Right click on the table header and select **Page Setup**.

## 11.1.18 Print a Table

#### A table can be printed by:

- Right click on the table tab and a menu will be displayed.
- Select the "Print" option. You may want to change the setup of the table (eg orientation and scale. You can do this using Page Setup option.

## 11.1.19 Export to Excel

To export a table to excel.

- Right click on the table header.
- Select the option to **Export to Excel**.

## 11.1.20 Save a Table

• Right click on the table tab and select Save As..

**NOTE:** You can save your table in comma separated format if you want to read it into another program such as Microsoft Excel.

## 11.1.21 Change Column and Row Width

#### To change the width of column and rows:

You can change the width of a row or column by clicking on the separating line and dragging. You can make each row the same width by holding down the **Shift** key and dragging one of the row edges.

## 11.1.22 Making Table Selections

#### To select one column of a table:

• Click on the column header

#### \_ Click here to select a column

HITLIS	r   PC	)BSearchResults							
			PDB Search results for ***						
	ID	head		date	het	title			
1	1sbt	HYDROLASE (SERINE PROTE	EINASE)	11 Aug 1972		Atomic coordinates for subtilisin BPN			
2	1mbr	OXYGEN STORAGE		05 Apr 1973		The Stereochemistry of the Protein M			
3	2dhb	OXYGEN TRANSPORT		01 Nov 1973		Three dimensional fourier synthesis of			
4	3ldh	OXIDOREDUCTASE, CHOH D	ONOR, NAD ACCEPTR	06 Jun 1974		A comparison of the structures of ap			
5	2cha	HYDROLASE (SERINE PROTE	EINASE)	01 Jan 1975		The Structure of Crystalline Alpha-Ch			
•	_								
	able: 22	2700 rows, 10 columns							

Г

#### To select one row of a table:

• Click on the row header

	ID	head	date	het	title
1	1sbt	HYDROLASE (SERINE PROTEINASE)	11 Aug 1972		Atomic coordinates for subtilisin BF
2	1mbr	OXYGEN STORAGE	05 Apr 1973		The Stereochemistry of the Protein
3	2dhb	OXYGEN TRANSPORT	01 Nov 1973		Three dimensional fourier synthesis
4	3ldh	OXIDOREDUCTASE, CHOH DONOR, NAD ACCEPTR	06 Jun 1974		A comparison of the structures of a
5	2cha	HYDROLASE (SERINE PROTEINASE)	01 Jan 1975		The Structure of Crystalline Alpha-C
•					

To select a row click here

#### To select more than one row or column:

- Click on one row or column whilst pressing the Ctrl keySelect multiple number of rows or columns whilst still pressing the Ctrl key

	ID	head	date	het	title
1	1sbt	HYDROLASE (SERINE PROTEINASE)	11 Aug 1972		Atomic coordinates for subtilisin B
2	1mbr	OXYGEN STORAGE	05 Apr 1973		The Stereochemistry of the Protein
3	2dhb	OXYGEN TRANSPORT	01 Nov 1973		Three dimensional fourier synthes
4	3ldh	OXIDOREDUCTASE, CHOH DONOR, NAD ACCEPTR	06 Jun 1974		A comparison of the structures of
5	2cha	HYDROLASE (SERINE PROTEINASE)	01 Jan 1975		The Structure of Crystalline Alpha-
•	-				

#### Select multiple rows and columns by clicking and selecting whilst pressing the Ctrl key.

#### NOTE: The Ctrl key acts as a toggle enabling select and unselect.

#### To select a range of columns or rows:

- Click on the first row or column in the range whilst pressing the Shift key.
- Click on the last row or column in the range whilst pressing the Shift key.

# To select a range of columns or rows - click on the first member of the range and the last whilst pressing the shift key.

		ID	head	date	het	title
7	ſ	3lyz	HYDROLASE (O-GLYCOSYL)	01 Feb 1975		Real-space refinement of the
8 1		1lyz	HYDROLASE (O-GLYCOSYL)	01 Feb 1975		Real-space refinement of the
9		6lyz	HYDROLASE (0-GLYCOSYL)	01 Feb 1975		Real-space refinement of the
10		5lyz	HYDROLASE (0-GLYCOSYL)	01 Feb 1975		Real-space refinement of the
11		2lyz	HYDROLASE (0-GLYCOSYL)	01 Feb 1975		Real-space refinement of the
12		1chg	HYDROLASE ZYMOGEN (SERINE PROTEINASE)	01 Mar 1975		Chymotrypsinogen: 2.5-angst
13		2cna	LECTIN (AGGLUTININ)	01 Apr 1975		The covalent and three-dime
14		1hip	ELECTRON TRANSFER (IRON-SULFUR PROTEIN)	01 Apr 1975		Two-Angstrom crystal structu
15		1gpd	0XID0-REDUCTSE(ALDEHYDE/DONR,NAD/ACCPT)	01 Jul 1975		Studies of asymmetry in the th
•						

## — Click here hold the shift key

#### Click here hold the shift key

#### To invert a selection:

- Right click on the original selection and a menu will be displayed.
- Select the Row Selection/Invert selection option.

**NOTE:** Invert selection can only be used on rows.

#### To select the whole table:

- Right click in the table and a menu will be displayed.
- Select the Row Selection/Select All option.

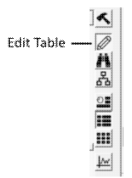
#### To remove a selection:

• Click anywhere within the table.

A selection can also be made from a plot select('table-plot{See Select plot section}).

## 11.1.23 Editing a Table

**To edit a table:** Click on the Edit Cells by Double Click button on the right hand side of the table. You can then edit each cell by double clicking on it.



OR

#### To edit the text or values within a cell:

• Right click on the table and select Edit Cells by Double-click .

## 11.1.24 Rename Column

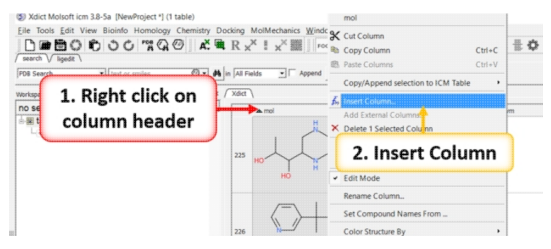
#### To edit the name of a column:

- Right click on the column header and a menu will be displayed.
- Select the option "Rename Column..." and enter the appropriate new text.

## 11.1.25 Inserting Columns

#### To insert a column:

- Identify the position within the table where you wish the column to be inserted.
- Right click on the column header and a menu will be displayed.
- Select "Insert Column"



A dialog box will then be displayed as shown below.

Column located     In the table?     New column name     Siring     Siri	

Use Add To List to add more than one column

Click OK to add column(s)

- Select the function you wish to add to the new column. Functions can be applied to many columns e.g. add etc..
- A set of arguments related to the function selected will then be displayed.
- Enter the appropriate arguments related to the function selected.
- Select where you want the new column to be located in the table.
- Enter the new column name
- If you wish to add multiple columns then use the **Add to List** option.

Many different functions are available:

- New Add a new column containing a real number, integer, string, or random number.
- **Transformations** A number of transformations can be selected and applied to a table column as shown below.
- **Mathematical** A number of mathematical functions
- **Text** Apply a number of different functions to the text in a column.
- Chemical Calculate a number of different chemical properties.
- Convert Units Radian to Degrees and Degree to Radian

#### Once the function and the correct arguments have been entered:

- Select whether you wish the new column to be added before, after or in place of this column.
- Enter the name of the new column.

**NOTE:** If you want to add more than one column choose **Add to List** and the action will be added to a list on the right hand side of the dialog box.

## 11.1.26 Increment ID Number

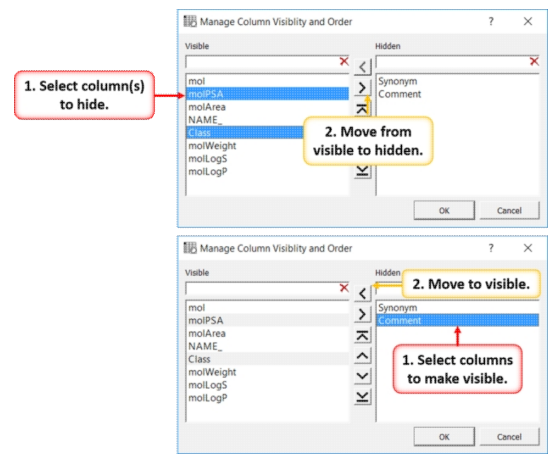
To add and Increment name ID in cell as a column (e.g. mol1, mol2, mol3 $\hat{a}f$ ):

- Right click on a column and choose Insert Column.
- Select Function New -> ID.
- Enter name and starting number and press OK.

## 11.1.27 Column Visibility

To hide a column:

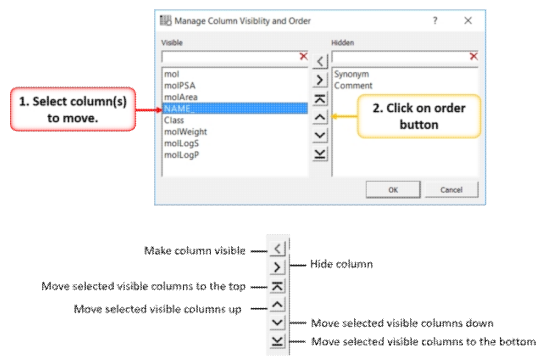
- Right click on any column header and choose "Manage Column Visibility and Order"
- A dialog box as shown below will be displayed.
- Click on the arrows to move a column from hidden to visible and vice-versa. Multiple columns can be selected by holding the Ctrl key and clicking.



## 11.1.28 Column Order

To re-order the columns in a table:

- Right click on any column header and choose "Manage Column Visibility and Order"
- A dialog box as shown below will be displayed.
- Click on the arrow buttons to re-order. Multiple columns can be selected by holding the Ctrl key and clicking.



## 11.1.29 Freeze Column

NOTE One application of Freeze column is to rename rows.

To freeze a column:

• Right click on the column you wish to freeze and select **Freeze** Column.

To unfreeze a column:

• Right click on the column you wish to unfreeze and select **Unfreeze Column**.

## 11.1.30 Freeze Row

To freeze a row:

• Right click on the row number of the row you wish to freeze and select **Freeze Row**.

To unfreeze a row:

• Right click on the row number of the row you wish to unfreeze and select **Unfreeze Row**.

## 11.1.31 Column Statistics

To calculate various statistics describing columns and inter-column relationship:

- Right click on the column header and a menu will be displayed.
- Select "Column Statistics"

The output is printed into the ICM Terminal window and the Column Statistics Window.

## 11.1.32 Inserting Rows

#### To insert a row:

- Identify the position within the table where you wish the row to be inserted and select the row.
- Right click on the row name (eg the number of the row) and a menu will be displayed.
- Select Insert Row Before or Insert Row After.

A blank row will be inserted. You can add data to this row by following the instructions in the edit table section.

				00100				
	(P	DBSear	chRes	ults				
						PDE	3 Search results for ***	
	Γ		ID	head		date	title	Γ
Right click _		1		LIGASE		01/01/01	crystallographic structure of a relaxed glutamine synthetase from mycobacterium tuberculosis	٨
here	-	<u> </u>	Cut F	low(s)				
		2	Сору	Row(s)		01/01/01	rna dependent rna polymerase from dsina bacteriophage phi6 plus bound ntp	E
			Paste	e Row(s)			philo pius bound nip	
		, ×	Delet	e Row(s)		01/01/01	multicopy crystallographic structure of a relaxed glutamine synthetase from mycobacterium tuberculosis	٢
		5	Inser	Empty Row Before				
			Inser	Empty Row After	FACTOR	01/01/01	crystal structure of destripeptide (b28-b30) insulin	F
	4	4						
	-		41		NOMEC	01./01./01		Υ.

#### Select row

## 11.1.33 Insert Image

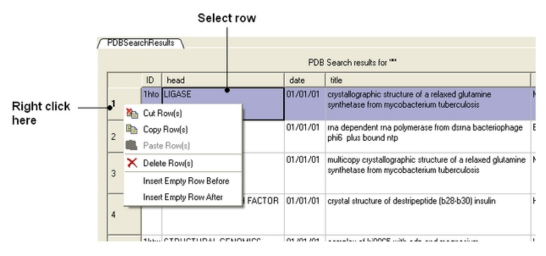
To insert an image into a table:

- First create an image column in the table. Use the Insert Colun option and choose Function > New > Image.
- Right click on a cell in the new image column you have created and choose **Set Image** and choose the image you wish to import.

## 11.1.34 Copy Cut and Paste Row

#### Copy, Cut and Paste Row:

- Select the row(s) See table selection section.
- Right click on the row header
- Select Copy Row(s).
- To paste a row select the row header under which you wish to paste the row. Right click and select **Paste Row**(s)



## 11.1.35 Copy Cell

#### To copy a table cell:

- Right click on cell.
- Select Copy Cell you can then paste it into a new table.

## 11.1.36 Copy Selection to an ICM Table

#### To copy a selection to a new table:

- Select the row(s) See table selection section.
- Right click on the row header
- Select **Copy Selection to ICM Table** and then choose Auto (ICM will name the table or New and you can enter a new table name.

## 11.1.37 Deleting Columns and Rows

#### To delete a column or row:

- Select the column(s) or row(s) you wish to delete. See the select table section for information on how to make table selections.
- Right click on the row number to delete a row or right click on the column header to delete a column and select the delete option from the menu.

## 11.1.38 Hide and Show Columns

If you have a large table you may wish to only show and display certain columns and hide others. By default any loaded table will have all the columns displayed.

#### To select which columns you wish to hide:

- Select the column(s) you wish to hide. See the select table section for information on how to make table selections.
- Right click and select the hide option from the menu.

IX	NAME	Score	Natom	Nflex	Hbond	Hphob	VwInt	Eintl	Hide
03476	m1	-35.47	33	0	-8.31	-4.97	-33.81	1.49	1.0
103485	m1	-44.18	36	1	-10.40	-7.35	-34.44	5.61	Column histogram
103522	m1	-36.40	46	1	-6.74	-7.44	-38.30	1.01	Rename
103526	m1	-37.21	31	1	-11.60	-5.32	-34.68	9.26	🗙 Delete column(s)
103547	m1	-33.21	36	0	-7.53	-5.89	-33.12	1.13	Color By
103566	m1	-35.13	49	4	-5.13	-7.93	-44.10	4.48	Insert column after
					1				Insert column before
									Sort
le: 12923 row:	s, 13 (of 14) co	lumns							Filter 🕨

#### Select column(s), right click and then select the hide option.

#### To show hidden columns:

- Right click on the column header and a menu will be displayed.
- Select the Show Columns options.
- Select which column you wish to show from the drop down list.

IX	NAME	Score	Natom	Nflex	Hbond	Hphob	Vv	• •	Hide	0.151		
03476	m1	-35.47	33	0	-8.31	-4.97	-33		Rename			
03485	m1	-44.18	36	1	-10.40	-7.35	-34	v	Delete colu			
03522	m1	-36.40	46	1	-6.74	-7.44	-38	~	Color By	annioj		
03526	m1	-37.21	31	1	-11.60	-5.32	-34					
03547	m1	-33.21	36	0	-7.53	-5.89	-33		Insert colu			
103566	m1	-35.13	49	4	-5.13	-7.93	-44		Insert column before			
103592	m1	-36.01	47	2	-4.39	-7.99	-49		Sort			
103614	m1	-36.25	41	5	-8.71	-5.47	-29		Show colu	mns		All
103615	m1	-34.95	49	1	-7.84	-7.68	-38		Filter		•	Dsolv
103621	m1	-34.46	31	0	-6.87	-5.01	-39.	21	2.08	9.51		mfScore
103626	m1	-32.70	34	0	-7.00	-4.72	-35.	59	2.15	6.61		FILE
03648	m1	-36.11	31	3	-7.40	-5.63	-35.	08	4.93	2.93		POS
103707	m1	-32.11	50	4	-5.91	-7.89	-49.	16	2.12	16.18	_	FUS

table: 12923 rows, 10 (of 14) columns

**Note:** In versions 3.8-6 and higher there is an option called "Manage Column Visibility and Order" as described here.

## 11.1.39 Column Format and Custom Actions

To change the **column format**, **font** color or size, the **alignment** of the column data, the background color, add hyperlink, or script:

• Right click on the column header and select Format

		J()	Column Statistics	
			Sort	
			Filter	•
_		🚕	Learn	
			Group by Column	
		-	Format	
	res 🄶	100	C.	OTTIOO
	2.4	Ał	MP,CIT,MN	GLNA1_MYC1

#### Right click on the column header

- A window as shown below will be displayed.
- Make the desired changes and click Apply

🗮 Column Format: A	? 🛛
Font	
Bold	Italic
Font Color	
Font Size	0
Background	
<ul> <li>Single Color</li> </ul>	
O Rainbow	
Alignment	
💿 Default 🔘 Left	🔿 Right 🔿 Center 🔿 Justify
Hyper Link	No Link 💌
Format	
Display Name	
Ok	Apply Cancel

#### To add a hyperlink to PubMed, PDB or Uniprot:

- Click on the drop down hyper link button.
- If the data in the column is a PDB, Uniprot/SwissProt or PubMed • code then choose the built in format from the menu.
- Click Apply and the data in the column will become blue hyperlinks.

#### To add a user-defined hyperlink:

- Click on the drop down hyperlink button.Choose "Simple link" See below for details on how to access cell data.
- Click Apply and the data in the column will become blue • hyperlinks.

#### To add an internal ICM link:

- Click on the drop down hyperlink button.Choose "Internal ICM link" See below for details on how to access cell data.

- Enter ICM scripting language in the panel that opens (see image below).
- Click Apply and the data in the column will become blue hyperlinks.

The value of the clicked cell is accessed as %1. To refer to the other cells and/or table itself the following shortcuts can be used:

- %@ # table name
- %# # clicked row number
- %^ # clicked column number

#### **Example:**

To add a an internal ICM link that reads a pdb dile and then displays it you could use the following ICM commands in the link:

			🗒 Column Format: ID 🛛 💽 🔀
			Font Italic
			Font Color
			Font Size
			Background
			Single Color
			Alignment           Oefault         Left         Right         Center         Justify
			Hyper Link Internal ICM link
• 11			Format <a href="#_">%1</a>
All			read pdb %@.%^[%#] cool
t V PDE	3		
	ID	heac	
1	<u>1hi1</u>	RNA	
2	1hto	LIGA	
3	<u>1hta</u>	LIGA	
4	<u>1htv</u>	HOR	
5	<u>1htw</u>	STRU	
6	1kpm	HYDF	
7	1s0p	MEME	Display Name
8	<u>2dca</u>	Hydro	Ok Apply Cancel
<			

## 11.1.40 Column Table Annotation

- Right click on the column you wish to annotate and choose **Format**, **Color Action**.
- Enter the comment in the bottom panel labeled Comments.
- The comments will be displayed when the user hovers the mouse over the column header. Other table comments can be added using the HTML panel.

## 11.1.41 Table Sorting

#### To sort a table by a column value:

- Right click on the column header.Select the Sort option.

PD	3SearchResults hitlist					
		F	PDB S	earch res	ults for 🐃	
ID	head	date 🗸	he	title		1
1uot	REGULATOR OF COMPLEMENT PATHWAY	23 Sep 200	3	Hide		1&4
1r1c	ELECTRON TRANSPORT	23 Sep 200	3 🚹	Column h	istogram	NOSA
1o5j	UNKNOWN FUNCTION	9 Sep 200	3	Rename		F PERIPLASMIC DIVAL
1o5h	STRUCTURAL GENOMICS, UNKNOWN FUNCTION	7 Sep 200	3 🗙	Delete co	olumn(s)	F PUTATIVE SERINE (
1o5i	OXIDOREDUCTASE	7 Sep 200	3	Insert co	lumn after	F 3-0X0ACYL-(ACYL
1qzr	ISOMERASE	7 Sep 200	3	Insert co	lumn before	F THE ATPASE REGIO
1qyq	LUMINESCENT PROTEIN	1 Sep 200	3	Sort		F THE CYCLIZED \$650
1qyo	LUMINESCENT PROTEIN	11 Sep 2003	3	Filter		ION INTERMEDIATE
1000	IMMUNE SYSTEM	10.5ep 2001	2	Filter	,	1 M82G2 COMPLEXED
le: 22	700 rows, 10 columns					1 non-:
R	ight click in the column header 🛛 —				row represe cending ord	ents ascending der.

## 11.1.42 Table Filtering

|Video|

There are two ways to filter table content one is via the table panel or by using the right click column header options.

#### To filter a table using the extra panel sliders:

S Xdict Molsoft icm 3.7-3b [NewProject *] (1 table)		How to apply filters to a table.
File Edit View Bioinfo Tools Homology Cher	sistry Docking MolMechani	
B <b>BB</b> O 00 % <b>G</b> O <b>R</b> R R	1. Click here 🧌	
ROB Search     Vigedit       POB Search     Vigedit       text or smiles     Vigedit	to view extra panel (right)	🔹 occupancyDisplay Inone 💌 🥥 🦉
Xáct		2. Click on the
nol	nof RotB nof HBA nof H	Filter tab
	2 3 5	> nd G
6. Symbol	2 3 🖸	nof Rotts
indicates		30 C X 13
65 column has	3. Click here	4. Drag to 33
been filtered	to enter	make changes
	filters	<u>                                      </u>
chiral	2 2	150 500 <b>X</b>
		> mol.ogP
		> mok.og5
		5. Click to
		apply or 🔗
chral		remove filters
C Trist del	4 4	
		۳. ۳.

Open the extra panel and choose the filter tab. Click and drag on the sliders to conveniently choose the filtering values or enter the values directly. Click on the green 'tick' to apply filters. To filter a table using the column header:

- Select the column you wish to filter. See the select table section for information on how to make table selections.
- Right click on the column header. •

• Select the **Filter** option.

				PDB Search results f	or '**
	ID	head	d-t-		L
1	1sbt	HYDROLASE (SERINE PROTEINASE)	1	Hide	ites for subtilisin BF
2	1mbr	OXYGEN STORAGE	05	Column histogram	istry of the Protein
3	2dhb	OXYGEN TRANSPORT	01	Rename	hal fourier synthesis
4	3ldh	OXIDOREDUCTASE, CHOH DONOR, NAD ACCEPTR	06 🔀	Delete column(s)	the structures of a
5	2cha	HYDROLASE (SERINE PROTEINASE)	01	Insert column after	Crystalline Alpha-C
6	4lyz	HYDROLASE (O-GLYCOSYL)	0.	Insert column before	nement of the struc
7	3lyz	HYDROLASE (O-GLYCOSYL)	0.	Sort	nement of the struc
8	1 lyz	HYDROLASE (O-GLYCOSYL)	0.	Filter	
۹ 	6luz		0.00		(Clear) (Clear All)
					(Custom)

- Select the "Custom" option and a data entry box as shown below will be displayed.
- Enter the appropriate operations and filter values for your ٠ search.
- Click OK.

💈 Custom filter on	'date'	? 🗙
equals	•	
⊙ And ⊂ Or		
equals	-	
🔽 Case Sensitive		
	Ok	Cancel

#### NOTE: When a column has been filtered a symbol as shown below will appear in the header of the column.

			PDB Search results for "*"				
	ID	head	∀date	het	title		
21260	1njo	RIBOSOME	02 Jan 2003	PPL	THE CRYSTAL STRUCTUR		
21261	1njt	HYDROLASE	02 Jan 2003	ACE	COMPLEX STRUCTURE OF		
21262	1njs	TRANSFERASE	02 Jan 2003		HUMAN GAR TFASE IN COL		
21263	1njq	METAL BINDING PROTEIN	02 Jan 2003	ACE	NMR STRUCTURE OF THE		
21264	1njp	RIBOSOME	02 Jan 2003	PPL	THE CRYSTAL STRUCTUR		
21265	1nju	HYDROLASE	02 Jan 2003	DNI	COMPLEX STRUCTURE OF		
21266	1oa7	HYDROLASE	02 Jan 2003		STRUCTURE OF MELANOC		
21267	1oa6	HYDROLASE INHIBITOR	02 Jan 2003		THE SOLUTION STRUCTUR		
01000	10.5	HYDROLASE INHIRITOR	02 Lan 2003		THE SOLUTION STRUCTUR		

#### This symbol means that the table has been filtered according to data within this column.

#### To append the filtered information into a new table:

- Select the whole table either by right clicking or pressing Ctrl A.
- Right click on the table and select "Append to other table".
- Enter a new name for the table you are appending with your filter • results.

#### OR

#### Selected rows can be appended to a new table by:

- Right clicking on the selected rows and a menu will be displayed. Selecting the "copy selection to ICM table" option. •
- ٠

#### A table can be filtered by a cell value:

- By clicking once in a cell. •
- Right click and a menu will be displayed. ٠
- Select the option "Filter by cell value".

#### A filter can be cleared by:

• Right clicking on the column selection and selecting Filter/Clear or Filter/Clear All

## 11.1.43 Append Table

To append rows to a new table:

- Select the rows you wish to copy to a new table.
- Right click on the table and choose **Copy/Append Selected Rows to** table.

Or you can use the Tools menu option here.

## 11.1.44 Mark and Select Rows

A row in a table can be marked and grouped by a label which enables the row(s) to be selected easily at a later time.

#### To mark a row

- Right click on the row in the table you wish to mark.
- Select Mark Row/ and then choose a number. In the GUI the number of rows that can be marked is limited to 5 but this can be increased using the command line command.
- A row that is marked will be colored each number is assigned a color. The coloring can be changed in the gui tab in preferences.

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				Select M	arked Rows	•	-	1	-		
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3	1htq	LIGASE	1	01701701	muncopy cry	statogrape				ed glutamine	MYCOBACTERIUM TUBERCULO
4	1htv	HORMONE/GROWTH FACTOR		01/01/01	crystal struct	ure of dest	- property -			sulin	HOMD SAPIENS
5	1htw	STRUCTURAL GENOMICS, UNKNOWN FUNCT	ION	01/01/01	complex of hi	0065 with	adp a	nd magn	esium		HAEMOPHILUS INFLUENZAE
6	1kpr	HYDROLASE		01/01/02	first structural	levidence	ofas	pecific n	hibilio	n of	DABOIA RUSSELLI PULCHELLA
7	1v8a	TRANSFERASE		01/01/04	structure of h	ydroxyeth	thiazo	ole kinasi	e prot	ein from	PYROCOCCUS HORIKOSHII
8	1:0p	MEMBRANE PROTEIN	•	01/01/04	structure of th	he n-termin	nal don	nain of th	e ade	nylyl	DICTYOSTELIUM DISCOIDEUM
9	2dcc	Hydrolase		01/01/06	x-ray crystal s	structure a	nalysis	of boyn	e sple	en cathepsin	BOS TAURUS
						. •					

Right click here and select Mark Row

Coloring relates to numbers

#### To select marked rows

- Right click on the table and choose **Select Marked Rows** and choose a number which relates to the marked rows as described earlier.
- Selected rows will be highlighted blue once rows are selected a number of right click options are activated such as copy selection to new ICM table.

## 11.1.45 Group View by Column

To group a column by value:

• Right click on the column header and choose "Group View by Column". Chemicals in the same group are grouped in the same row. You can expand the view by clicking on the "+" sign".

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Xdict				1. Right click and choose	+
	mol	Molecule Name	🗄 dass ┥	"Group view by column".	
+ 48 (1 of 2)		Histamine			
+ 50 (1 of 3)		Lamotrigine		32	i i e a
expanded	group can be I here to see all s of the group.	Pyrazinamide		33	ゆ 23 次 21 の
0	0.0.				

To remove grouping:

• Right click on the column header and choose "Undo the Group View"

## 11.1.46 Mouse and Cursor Actions on a Table

The actions resulting from a mouse click or cursor on a table can be changed by:

- Right click on a table and select Table View/Show Extra Panel
- A panel as shown below will be displayed.

PDB Search results for <b>'kinase</b>	•'			
	source 🔺	X Header		
thiazole kinase protein from	PYROCOCCUS H	Name	Value	
rosine kinase domain of the	HOMO SAPIENS	tableTitle	PDB Search results for ' <b>kinase'<!--</td--></b>	
bacterium tuberculosis thymidylate	MYCOBACTERIU	doubleClick	nice "%1" no no no no	
ndp kinase	DICTYOSTELIUM		delete a_"MiniObj.	
eta	HOMO SAPIENS	CUISOF	cursorFindPDB %# "%@" yes	
eta	HUMU SAFIENS	separateT ab		
sse, mutation r65q 🛛 🕈	SACCHAROMYCI			
ine monophosphate kinase (thil)	AQUIFEX AEOLIC			

## Right click here and select Table View to display and undisplay extra panel

Double click here to edit actions

• Double click in the **Value** column and the column can be edited. Add ICM commands for the action you want. A value in a column can be referred to using "%" e.g. column two would be referred to as "% 2". In the example shown above the function nice is acting on the contents of column one for the double click action.

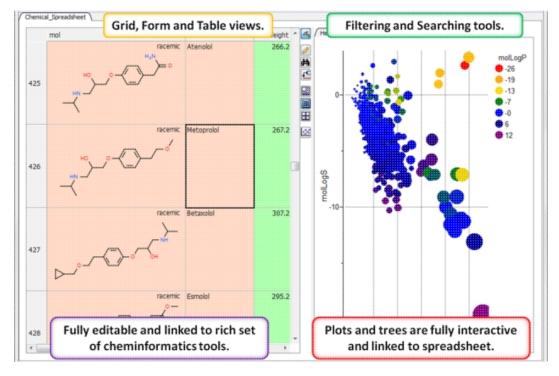
NOTE: The action associated with cursor and double click is placed in a variable name TableName.cursor and TableName.doubleClick

## 11.2 Molecular Tables

```
Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS
```

An ICM molecular table is created when an SDF or Mol file is read into ICM. To read and open a mol or sdf file go to File/Open (See Open an ICM file section) All of the table functions described in the previous section Standard ICM Table can be applied to molecular tables. Molecular tables are described in more detail in the Cheminformatics chapter.

An example of an ICM molecular table:



# 11.3 Insert Interactive Objects into Table Cell

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

NOTE: This option is available in Version 3.7-2b and above.

To store objects in a table please use the following:

Example:

```
read pdb "lcrn"
display a_
add column t Parray( object a_ preview )
```

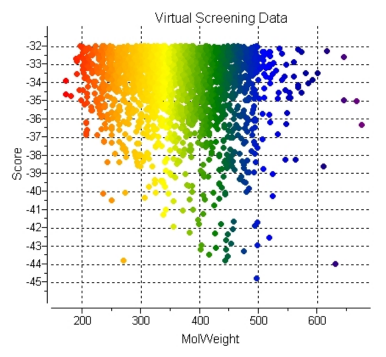
**Viewing** Objects from the table can be dragged and dropped into the ICM Workspace to display them again. **How to make the objects in the table fully interactive** 

• Right click on the column header and select "Live Rotatable View"

## **11.4 Plotting Table Data**

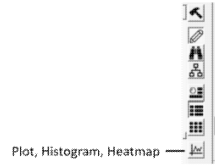
```
Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS
```

The data within a table can be plotted graphically. A histogram can be made for the data within one column or a plot can be constructed for the data within two columns.



To make a plot, histogram or heatmap:

- Read in a chemical spreadsheet or other data in spreadsheet form (e.g. csv).
- Click on the plot button at the side of the table (shown below).

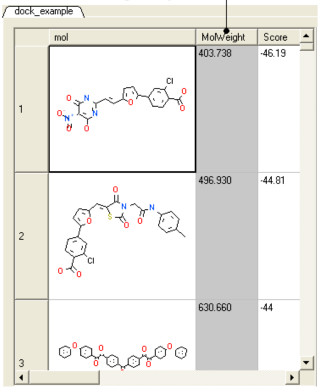


## 11.4.1 Column Histogram

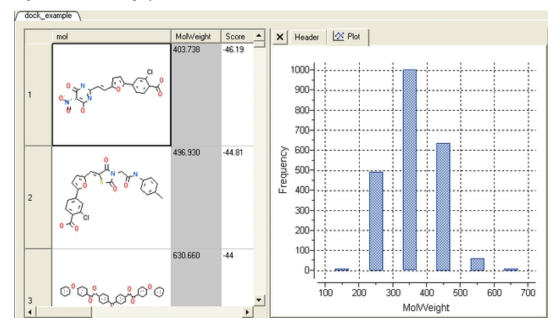
#### To plot a histogram of the data within one column:

- Select the column by clicking on the column header.
- Right click on the column header.
- Select the Column histogram option.

# Click here to select the column and then right click and select column histogram option



A plot will then be displayed next to the table.

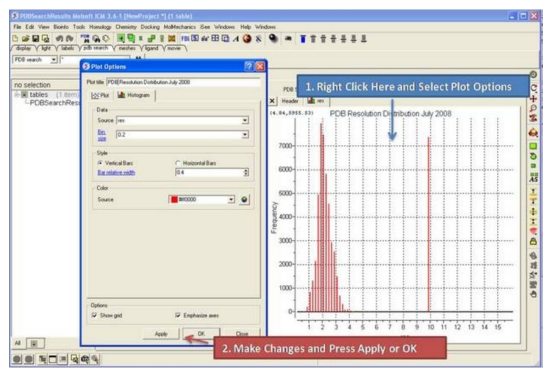


## 11.4.2 Histogram Options

Once you have created a histogram you can change the following parameters by right clicking on the plot and selecting options

Options:

- Change plot title.
- Change the data source using the drop down button and select another column in the table.
- Change the histogram bin size.
- Change the bars positioning from vertical to horizontal.
- Change the bar relative width compared to bin size. Bigger values give thicker bars.
- Color the bars



## 11.4.3 Histogram Bins

There are two ways to change the bin size. 1. Using the options dialog box or 2. interactively by left clicking and dragging at the top of the plot as shown below - this will allow you to find the best density estimation picture.

	ath	Click and Drag Here to Chang	er 🔒 ins
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4	Ye.J., Chang, W., Liang, D.	X-RAY DIFFRACTION 11 FO	
5	Teplyakov, A., Gilland, G.L., Structure 2 Function Project	X-RAY DIFFRACTION 1:	
6	Chandra, V., Jasti, J., Kaur, P., Betzel, C., Srivivasan, A.,	X-RAY DIFFRACTION 11 50	<sup>00</sup> ]
12 2	Ksiazek, D., Brandstetter, H., Issael, L., Bourenkov, G.P.,	XRAY DIFFRACTION 1. E 40	00
8	Jepakanthan, J., Tahirov, T.H., RIKEN Structural	XRAY DIFFRACTION 11 탈 30	ool
9.5	Watavabe, D	X-RAY DIFFRACTION 21 LL	
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15	Watanabe, D.	X-RAY DIFFRACTION 11 10	00
12	Watanabe, D.	X-RAY DIFFRACTION 21	
13	Krishna, R., Rajan Psabu, J., Manjunath, G.P., Datta, S.,	X-RAY DIFFRACTION 3.1	
14	Martick, M., Scott, W.G.	X-RAY DIFFRACTION 2	0 1 2 3
3	Color D.D. The D. Delaure U. MR.P.D.	V DAV PUEEDAP TURK	res

## 11.4.4 Plots

To construct a scatter plot from data within two columns:

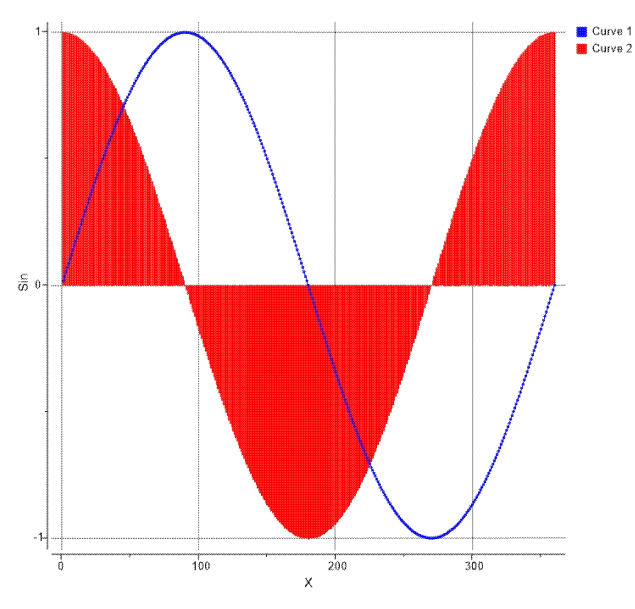
• Click on the plot options button (right hand side of table).

- B Plot Options 世世众 Plot title 🕍 Plot 🛛 🕅 Histogram 🔹 🚟 Heatmap Cata 2. Select data x molivieight • C Logarithmic - 4 here mol.ogP • C Logarithmic Swap X and Y <none > \* å Balloon (Example: mol. ID) -Filter (Example: colliame > 10) . III 1. Open plot 152 0.813 options dialog 14 • Shape • Plotting style dots \* window. Point labels T Show labels for selection only Color <ford> • • Source 172 0.2324 nsity 🔳 T Inte Grade 0 Confidence Inh (none) • Y conf. \* X conf. <none> Least squares fitting ine 169 2.099 @ Lines C Xer e Options 🔽 Show grid Emphasize aves T Antialias -OK Close
- Select the data from the drop down options in the Data section of the dialog box. There are additional options to change the color, mark shape or size, and grid and axis display.

## 11.4.5 Multiple Curves

Multiple curves can be added to a plot:

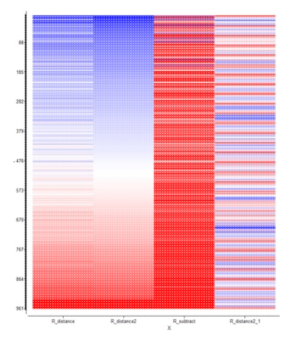
- Make a plot of the first curve.
- Right click on the plot and choose **Options.**
- Select the "+" button next to Curve data entry box.
- Enter new X and Y data for the second Curve.
- Shading as shown in the image below (red shading) can be achieved by selecting Plotting Style > Vertical bars.



## 11.4.6 Heatmap

To generate a heatmap:

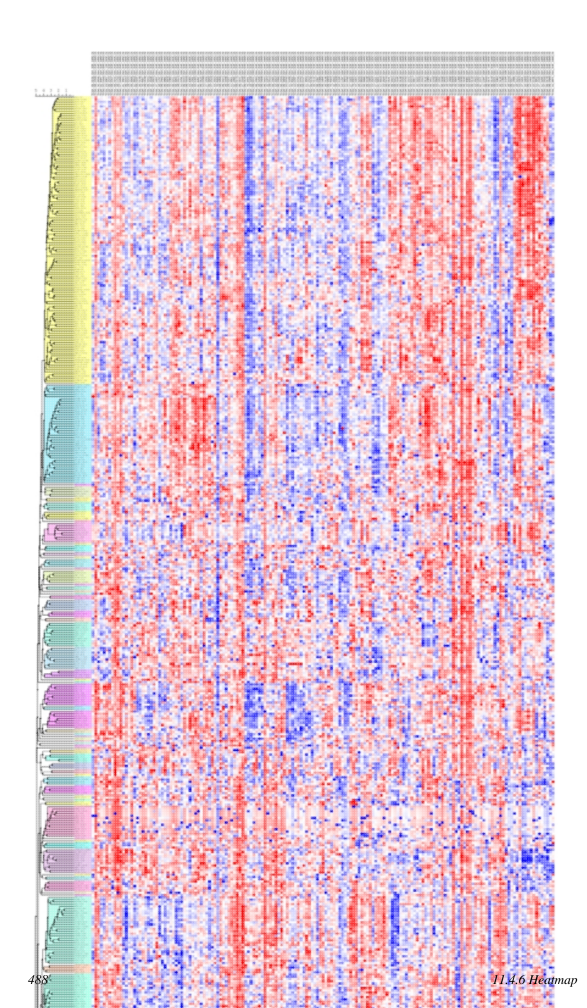
- Read in a table containing the data.Click on the Plot/Histogram button on the side of the table.
- Click on the heatmap tab at the top.
  Add the data sources you want in the heatmap based on the column headers in the table.
- Check how you want to label the rows.Edit the color gradient if necesary.
- Click OK.



#### 11.4.6.1 Heatmap Example

This example shows how you can place a fully interactive heatmap next to a tree.

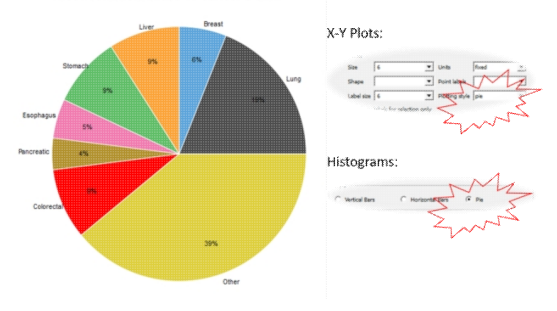
- Read in table d.tsv from here: ftp://ftp.molsoft.com/pub/workshop/ (1. cut and paste the address into a web browser. 2. Right click on d.tsv and choose "Save Link As". 3. Open the .icb in ICM File/Open).
- Select numerical columns by clicking on the column header and holding CTRL key.
- Right click on the column header and choose "Reorder by Clustering".
- Click on the cluster button on the right hand side of the table and use the default options.
- Right click in the cluster and "Change Record Labels"
- Check color check box and click 'Add All Numerical'
- Remove cluster related columns from the end ('cl' 'ord', etc ...)
- Click OK a tree and heatmap as shown below will be displayed.
- CTRL +/- allows you to zoon in and out.



## 11.4.7 Pie Chart

To generate a Pie Chart:

- Read in a table containing the data.
- Click on the Plot/Histogram button on the side of the table or right click choose **Options** on a plot.
- For Histogram choose Plotting Style > Pie radio bin. Number of bins/sectors can be controlled the through the 'Bin Size' parameter. For Plots choose Points > Plotting Style > Pie 'X' should be discrete (integer or string column with categories), 'Y' any numerical column.



Most Common Cause of Cancer Death

## 11.4.8 Add a title to a plot

#### To add a title to a plot:

• Right click on the plot and select Edit Title or choose Options

## 11.4.9 Axis Options

Each axis has a set of options which can be accessed by right clicking on the axis and selecting **Options.** 

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4	Entre DA This A Debinen W MPPD	V DAV PREEDAPTICAL		44					res				- II

#### To change the title of the X or Y Axis:

• Right click on the axis and select options

#### To change the data range:

- Right click on the axis and select **options**
- Change the **From** and **to** values in the **Range** box

#### To change the Grid steps (ticks) on the X or Y axis:

• Select either a fixed step e.g. 10 and you can define the number of subdivisions (ticks) in each step. Choosing 1 will display zero ticks between divisions.

#### To change the axis to logarithmic

- Right click on the axis and select **options**
- Select the Logarithmic scale check box

## 11.4.10 Change Axis Data

#### To swap the X and Y axis:

- Right click on the plot and select **Options**.
- Select the Swap X and Y button.
- Click OK.

#### To change the data source for either the X or Y axis:

- Right click on the plot and select **Options**.
- Select the drop down arrow as shown below and select a different column from the table.
- Click OK.

## 11.4.11 Logarithmic Plots

#### To change the scale of the axis to logaritmic:

- Right click on the plot and select **Options**.
- Select the **Logarithmic** check box.

## 11.4.12 Change Mark Shape or Size

#### To change the plot mark, shape, style or label:

- Right click on the plot and select **Options**.
- Select the desired size and shape using the drop-down buttons in the **Marks** section of the window.

#### To add point labels:

- Right click on the plot and select **Options**.
- Select the drop down arrow in the **Point labels** dialog box
- If you only want to label selected points check the **Show labels** for selection only option. Making plot selections is described here.

## 11.4.13 Change Mark Color

#### To change the color of the plot marks:

• Right click on the plot and select **Options**.

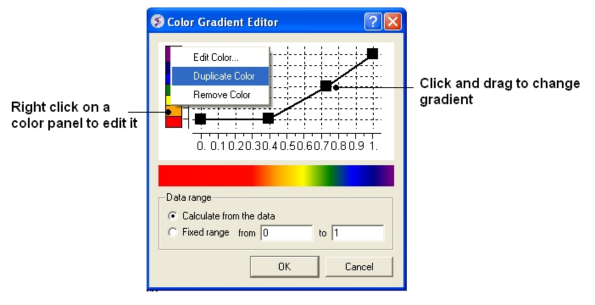
- In the Color section of the window select the Source (column name plotted as X or Y) you wish to color. Select the color palette and choose the desired color or you can
- choose a Gradient of colors.

	Color palette button
Color	
Source	MolWeight 👤 🔹
Gradient	red/orange/yellow/green/blue/darkblue/purple 💽 🏒
L	

Color gradient editor

## To edit the color gradient

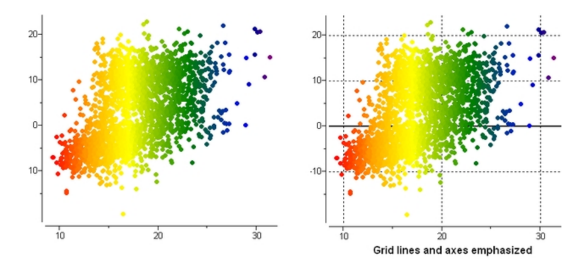
- Click on the Color gradient editor button and a window as shown below will be displayed.
- Click and drag on a mark in the gradient plot to change the color gradient.
- Right click on a color in the Y-axis to Edit, Duplicate or Remove Color.
- The color gradient can be applied to all points in the data or for a fixed range.



## 11.4.14 Grid and Axis Display

To remove the grid display and/or highlight the axes:

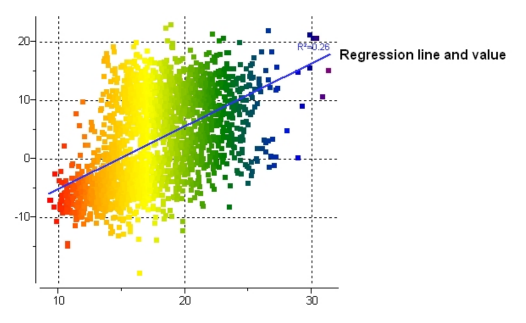
- Right click on the plot and select **Options**.
- Check the **Show grid** or **Emphasize axes** options.



11.4.15 Least Squares Fitting

# To fit the data to a straight line using least square fitting and report the equation:

- Right click on the plot and select **Options**.
- Select the check box for **Least squares fitting line**.
- Choose Linear, Logarithmic or X=Y.



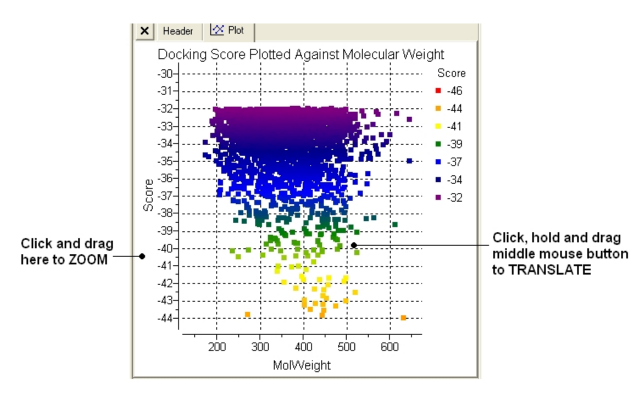
## 11.4.16 Zoom, Translate and Center

### To zoom into a plot:

• Click outside the plot on the left-hand-side and drag the mouse or use the middle mouse wheel to zoom in and out.

### To translate a plot

• Click, hold and drag using the middle mouse button on the plot.



### To center onto a plot

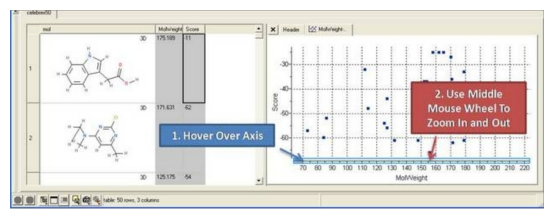
• Right click on the plot and select **Center all** or **Center Selection**. Making selections in a plot is described in the next section.

### To center into an axis

• Right click on the axis and select center.

#### To zoom into an axis

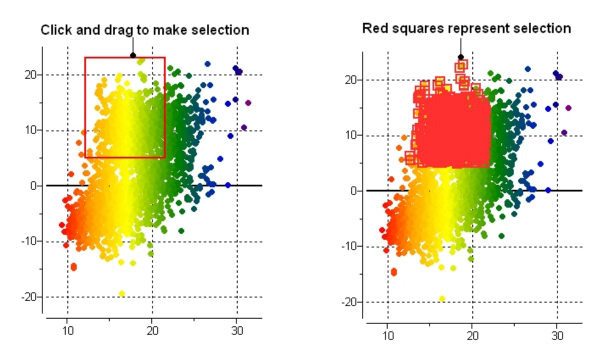
- Hover the mouse over the axis until you see a blue rectangle surrounding the axis.
- User the middle mouse wheel to zoom in and out as shown below.



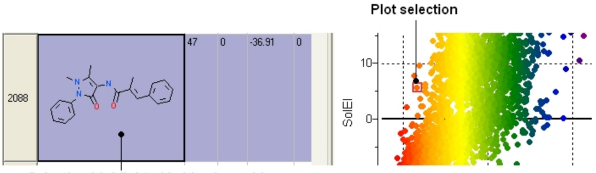
## 11.4.17 Plot Selection

## To make a selection in a plot:

• Click and drag in the plot to make a selection. Individual points can be selected with a single click.

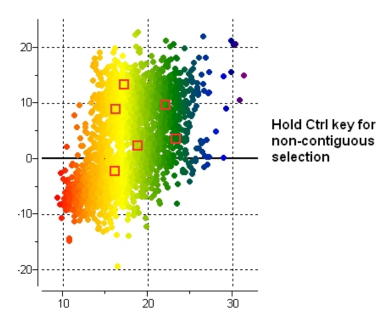


All selections are directly linked to the table from which the plot was made. Selections in the table are highlighted in blue.



Selection highlighted in blue in a table

Non-contiguous selections in the plot can be made by holding the CTRL key.



## 11.4.18 Point Label

To add a label to one or more points:

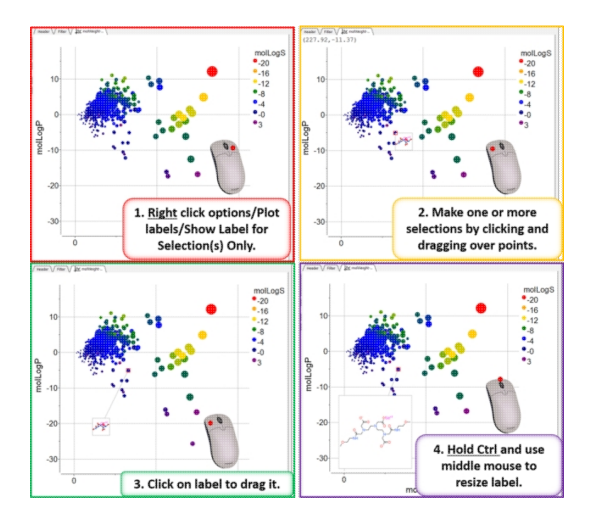
- Right click on the plot and choose options.
  Look for the section Points and the dialog box \*{Points Labels}
- For this example we will use the drop down button and choose "mol".Check the option show label for selection only.

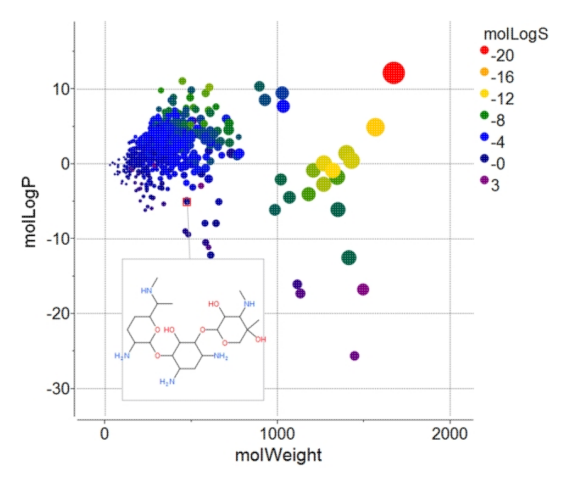
To drag the label:

• Left click on the label and drag.

To make the label display bigger

• Hold the Ctrl key and use the middle mouse button.





## 11.4.19 Print Plot

## To print a plot:

- Right click on the plot and a menu will be displayed.
- Select the print option.

## 11.4.20 Saving a Plot Image

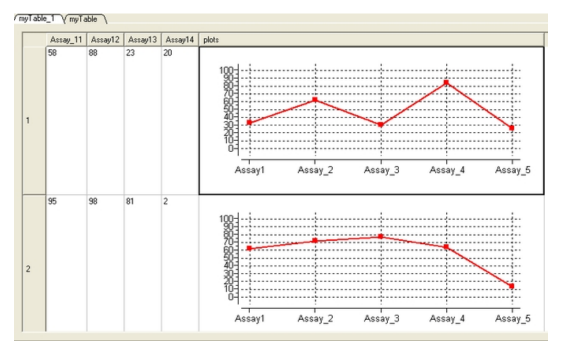
#### To save a plot image or copy to clipboard:

- Rifht click on the plot and select Options and then check the antialias button. Click Apply and OK.
  Right click on the plot and a menu will be displayed.
- Select the Save/Export Image option. •

## 11.4.21 Table Inline Plots

#### Plots can be inserted into a table row by:

- Select the columns you wish to plot.
- Right click on the column header and select **Inline Plots**
- The plot will then be displayed in each row of the table. •



## 11.4.22 Annotate Plot

You can add text, lines, rectangles, and ellipse to a plot. To annotate a plot:

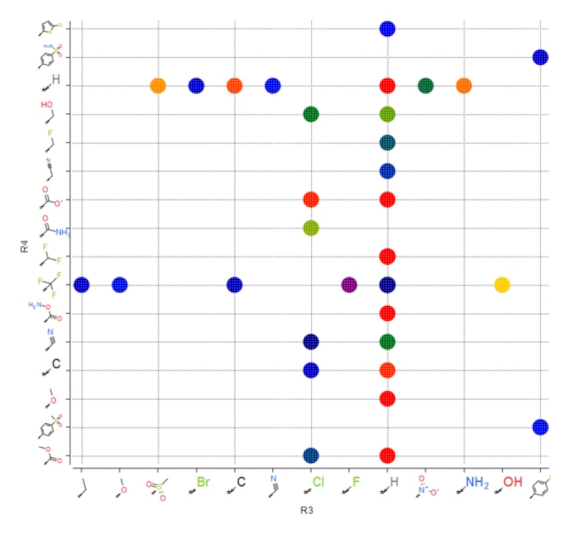
- Right click on a plot and select Annotate.
- Select the draw element and then click on the plot.
- Change the draw element properties if neeeded (e.g. font size).

Annotations			? ×
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Aa[ Text		Outline Color	8 💌
Draw Elements		Draw Properties	Text Properties

## 11.4.23 Plot R-Groups

To plot R-groups e.g. from the results table of library decomposition.

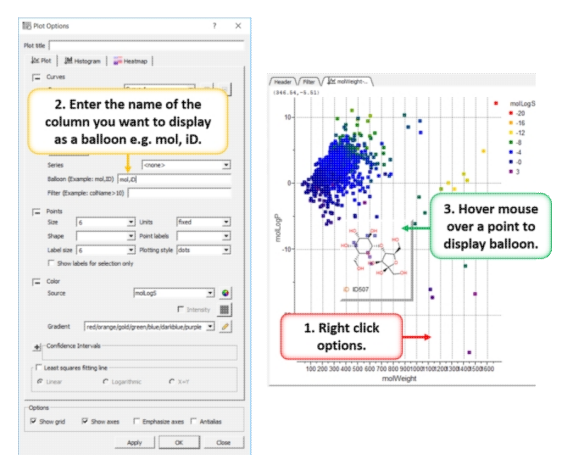
- Click on the plot button on the side of the chemical spreadsheet.
- Select the R-group column for the X and Y axis.



11.4.24 Tooltips Balloons

When you hover over a point in a plot it is sometimes useful to be able to visualize some data regarding that plot. You can do this by setting a ballooon.

- Right click on the plot and select Options.Enter the name of the columns containing the data you wish to display in the balloon (e.g. mol, ID).
- Hover over a point in the plot and the balloon will be displayed.



## **11.5 Principal Component Analysis**

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

Principal Component Analysis (PCA) is the younger brother of ICM's more powerful data analysis tools, like property prediction and clustering, though it still may give a good description of the data with a few columns or even chemical compounds. PCA is a mathematical procedure that transforms a number of correlated variables into a number of smaller uncorrelated variables known as Principal Components The first component accounts for as much of the variability as possible with the rest of the components accounting for the remainder. PCA may be very helpful when you believe the data actually contains only a few meaningful components. Principal components are linear combinations of the provided data columns.

To perform a PCA analysis a table (either chemical or standard ICM table) needs to loaded into ICM. For information regarding ICM Tables and ICM Chemical Tables please follow these links.

## To begin the PCA procedure

- Right click on a ICM Tables and ICM Chemical Table and select the PCA option. It is important to right click inside the data table and not on a column or row header in order to see the correct menu on which pca is listed.
- Select which columns you wish to incorporate into the PCA analysis.

🦻 PCA Parameters		?×
Table:	melanin	-
Column Prefix:	PC	
PC Number Limit:	3	*
Explain Data Variance (%):	99	
Descriptors: C the "mol" column and C the selected columns ( the mol column only		
Plot principal componets:		
O Do not plot		
<ul> <li>Build scatter plot for the</li> </ul>	he first 2 components	
<ul> <li>Build colored scatter (</li> </ul>	plot for the first 3 compone	ents
ОК	Cancel He	elp

- Enter the table name on which you wish to perform the PCA analysis. If only one table is loaded this option will be greyed out.
- Enter the number of Principal Components (PC number limit) you wish to generate. Generally 3 principal components may be effectively visualized and it will be enough often to fulfil the data variance percentage requirement (see next option). The value displayed in the terminal window under the heading "cumulative explained data variance" will show what percentage of data relates to each PC.
- Enter a value in the "Explain Data Variance (%)" data entry box (99% is the default value) if you prefer this indirect way of limiting number of PC. The algorithm will stop when either PC number or explained variance limit is reached, so if you want only one of this criteria to work, make sure that the other limit is weak (by assigning accordingly the number of PC limit a high value, e.g. 50, or setting data variance to 100%).
- Select which descriptors you would like to include in the PCA analysis.
- Select which plot you would like to display. If you choose to display a plot use the color key on the side of the plot and the information contained within the ICM terminal window to relate which axes and points relates to which PC. PC3 is usually the color in the plot with the values displayed in the plot key.
- Click OK and if selected a plot will be displayed on the right-hand-side of the table. Points within a plot are linked to the table and can manipulated as other plots contained within a table.

## 11.6 Learn and Predict

Available in the following product(s): ICM-Chemist-Pro | ICM-VLS

Please click here to go to the Learn and Predict chapter.

## 11.7 Data Clustering

ICM allows you to create hierarchical clusters for chemical and other objects. Cluster trees can be used for:

- Navigation through large data sets.
- Selecting group representatives (taxons).
- Filtering tables to exclude redundancy.
- Finding similar elements, and more.
- Creating hierarchical views of data sets in many different styles, with subsequent image export/printing ability.

## 11.7.1 Cluster

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

To perform clustering based on chemical similarity or any other data you must have an SDF file or table loaded into ICM.

• Right click on the table and select the Clustering option OR select the Chemistry menu and choose the option entitled **Cluster Set**.

### OR

• Tools/Table/Clustering.

A data entry box as shown below will be displayed.

Clustering Paramete	rs		?	×
Table:	Xdict			•
Name:	[			
Take Labels from Column:	NAME_			•
Keep the existing tree	S			
Add column with duste	r numbers, name: d	******		
<b>Build</b> for the selected (	) rows (at least 2 require	ed)		
Descriptors:				
← the "mol" column and	d all numerical columns	C	the mol column only (2D pharmacophore) dimt_UPGMA	<b>.</b>
C the selected column	s only	C	the seq column only	
C the "mol" column onl	y (fingerprints)	C	use existing distance matrix:	
Method:				
Tree     Parameters:			Memory required: 1 MB	
Linkage type:			UPGMA	
Keep distance ma	trix			
C Compressed hierard	hical dustering			
L			OK Cancel Help	

- Select the table you want to cluster from the drop down menu.
- Give the tree a name.

- Select the column you would like to use to label the tree (this can be changed or appended to once the tree has been made right click on tree and choose "Change Record Labels".
- You can choose to keep the existing tree.
- The cluster number names will be added to your table you can change the label of the columns (default = cl).

Now select the descriptors you wish to use for clustering.

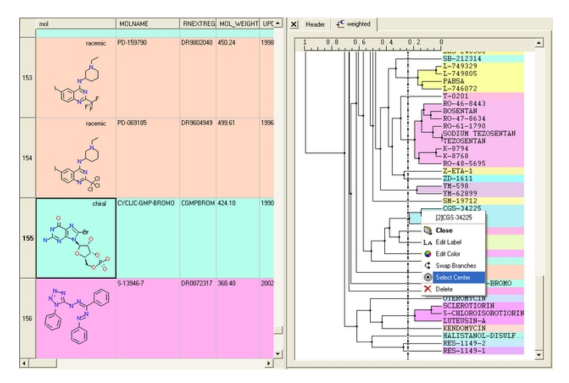
- The "mol" column and all numerical columns.
- The "mol" column only (2D pharmacophore)
- The selected columns only.
- The sequence column only.
- The "mol column only (fingerprints).
- Use existing distance matrix. UPGMA is a linkage method which is used to group tree nodes together. (they are described in the link below). dimt\_UPGMA is used when 'use exiting matrix' mode is selected. It can be any exiting matrix. This usually comes from previous clustering if 'Keep distance matrix' option was selected

Select the method you wish to use for clustering.

- UPGMA (unweighted pair group method using averages): Distance calculated is the average of all elements (recommended). https://en.wikipedia.org/wiki/UPGMA
- Single linkage: Nearest neighbour linkage https://en.wikipedia.org/wiki/Single-linkage\_clustering
- Complete linkage: Furthest neighbour linkage https://en.wikipedia.org/wiki/Complete-linkage\_clustering
- WPGMA: (weighted pair group method using averages): Rough
- approximation of weighted (slightly faster)Farthest
- https://en.wikipedia.org/wiki/Complete-linkage\_clustering
   Ward: https://en.wikipedia.org/wiki/Ward%27s method
- **UPGMC:** Centroid linkage clustering
- **UPGMC-corr:** Correlation clustering provides a method for clustering a set of objects into the optimum number of clusters without specifying that number in advance.

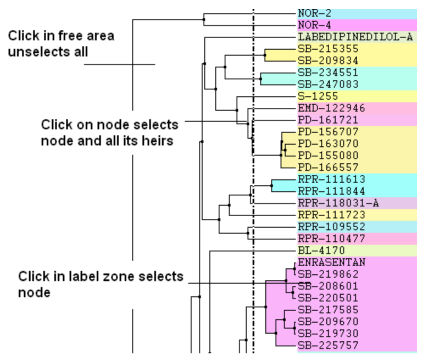
**NOTE** If you wish to cluster a selection of a table; first select the compounds you wish to cluster (Ctrl A will select all or see the section of the manual entitled making table selections.

Check the option "Keep Distance Matrix" if you wish the Distance Matrix to be saved.



## 11.7.2 Tree Selection

A number of different selections can be made once the tree has been created.



- Click in free area unselects all.
- Click on node selects node and all its heirs.
- Ctrl+click adds to (if not selected yet) or removes from (if already in selection) current selection.
- Shift plus click allows a range to be selected.
- Click in label zone selects node.

- Keyboard "Up"/"Down" cursor keys move selected node up/down in the tree. With Shift held the selection expands in the according direction.
- Keyboard "Escape" unselects all.

## Select Center Representatives From a Node.

To select the "center" representative compound or value from a cluster.

• Right click in the free area of the tree and select the option "Select Centers".

<b>NOTE:</b> Selections in the t	ree will be highlighted in blue in th	ne tree
	LSB-247083	
	S_1255	
	EMD-122946	
	PD-161721	
and in the table.		

#### Copy selection to new table.

All selections can be copied to a new table by:

- Right clicking on the selected rows in table and a menu will be displayed.
- Selecting the "copy selection to ICM table" option.

## 11.7.3 Save and Print Tree

#### To save a tree

Option 1:

• Save the whole session as an ICM project. See Saving an ICM project

#### Option 2:

• Save the table as an .icb file. Right click on table header and select "save as".

#### To save a tree as a picture

- Right click in the "free area".
- Select "Save Image ... "

The image of the tree can also be saved to the clipboard.

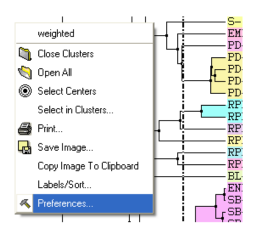
- Right click in the "free area".
- Select "Copy Image to Clipboard"

#### To print the tree

- Right click in the "free area".
- Select "Print"

## 11.7.4 Tree View

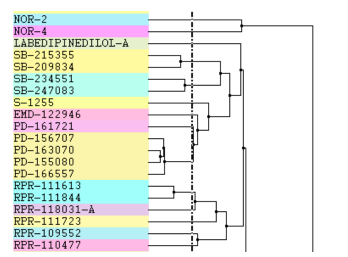
A number of changes to the tree view can be made by right clicking in the tree "free area" and selecting preferences as shown below.



A preferences window as shown below will be displayed.

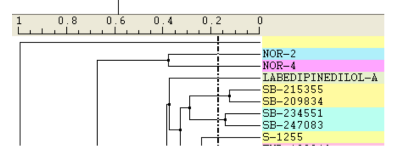
Preferences	? 🛛
🔲 Reverse Growth	
🔲 Reverse Order	
Use color for	Color lines 🗾
Distance range: from 0 to 1	Fill Color lines
Minimal line spacing	Color lines to parent
Line width	Only labels
Left margin 20 🍨	Right margin 🛛 20 🚔
OK Cancel	Label Font

## **Reverse Growth:**



## **Distance Range:**

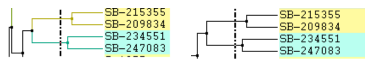
## Distance range changes scale here



## **Color Preferences:**

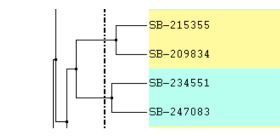
SB-215355 SB-209834	
SB-234551	- SB-234551
SB-247083	- SB-247083

Left: Fill Right: Color Lines

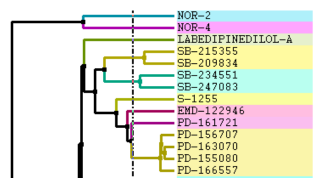


Left: Color lines to parent Right: Only labels

## **Increase or Decrease Line Spacing:**



## Increase or Decrease Line Width:



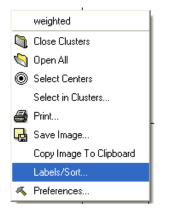
**Change Font and Margins:** 

🦻 Preferences	? 🛛	
🔲 Reverse Growth		
🗖 Reverse Order		
Use color for	Color lines 📃	
Distance range: from 0 to 1	Fill Color lines	
Minimal line spacing	Color lines to parent	
Line width	Only labels	
Left margin 🛛 👤 🚔	Right margin 20 🚔	
OK Cancel		— Change Label Font
Chanc	ge Margins	-

## **Change Label**

To change the tree labeling (GLOBAL):

\* Right click in the "free area" and select the Label/Sort option.



A window as shown below will be displayed:

≶ Labels/Sorting	? 🔀
Edit Object Label Format: Format:  %MF/%MOLNAME/%MOL_WEIGHT; Choose column: MOL_WEIGHT _ App Update Labels	end to Format
Rank tree by column:	Reorder
Sort Table by Tree	
	Exit

- Choose the column name you wish to label the elements of your tree.
- Click the Append to Format button. You may wish to delete some of the information in the Format data entry box if you do not wish that column header to be displayed. Note more than one column header can be used as a label.
- Click the update labels button to view the changes.

To change the tree labeling (NODE):

• Right click on the node you wish to change the label of and select "Edit Label" and a data entry box as shown below will be displayed.

🦻 Node Label		? 🛛
Enter Label		
Enter New Label Here		
	ОК	Cancel

• Enter the new label.

## **Change Node Coloring**

To change the color of a node:

- Right click on the node you wish to change the color of and select "Edit Color". Select the desired color and click OK. •
- •

# **12 Working with Local Databases**

ICM tables are capable of storing tens of thousands records. However, some problems operate with data sets so large that they do not fit in computer's memory. To work with such large amounts of data ICM uses the concept of Molsoft database (MOLT) files. Unlike many other table file formats, such as SDF, CSV and others, database files are optimized for fast search and other operations, like unique entry addition and diverse subset selection.

Database files do not provide all the functionality available for tables but they allow the user to organize large amounts of data, search data using various advanced criteria and share created data collections with other users. For large amounts of chemical data database files provide specialized chemical functionality.

ICM provides the following database file operations:

- create database files from SDF, CSV/TSV, SMILES files and from ICM tables;
- impose unique constraints on certain columns upon table creation to avoid redundancy;
- store multiple tables in a single file; rename, delete tables in a database file;
- search fast using advanced conditions, including advanced chemical search;
- select diverse subsets from chemical database files;
- browse database tables using flexible filtering and sorting conditions;
- directly edit/delete/insert entries in the database;
- export in popular formats, such as SDF (for chemistry) and CSV;
- export and import tables to and from Molcart.

## **Chapter Contents:**

- How to make a local database
- Browse Database
- Edit Database
- Query Local Database

## 12.1 How to make a local database.

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

To convert a file into MOLT format.

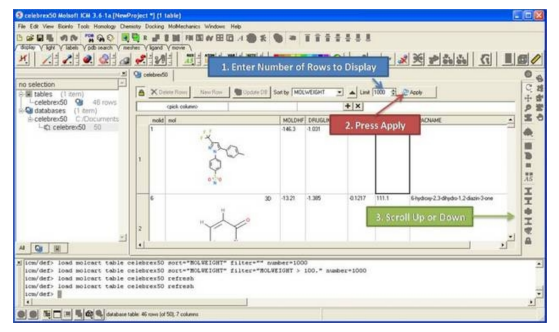
- File/Convert large SDF, CSV to Database (molt).
- A **Database Import** dialog box will be displayed. Select the fields you wish to import.
- Once the file has been converted to MOLT and imported then it will be displayed in the ICM Workspace.
- Double click on the file name in the ICM workspace and the database browse mode will activated.

## 12.2 Browse Database

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

To browse a database first decide how many rows of your database you wish to view

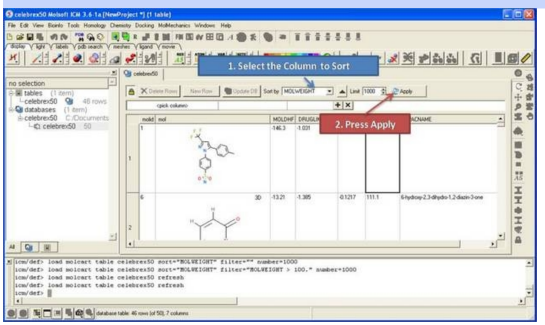
- Double click on the file name in the ICM workspace and the database browse mode will activated.
- Select the number of rows you wish to be displayed in the Limit data entry box.
- Click on the Apply button.
- Scroll up and down the displayed rows using the scroll bar on the right hand side of the table.



#### To sort a database by a value in a column

- Click on the drop down arrow next to the **Sort by** data entry box and select a column.
- Click on the **Apply** button.

## **NOTE:** The database will be sorted globally not just the displayed rows.

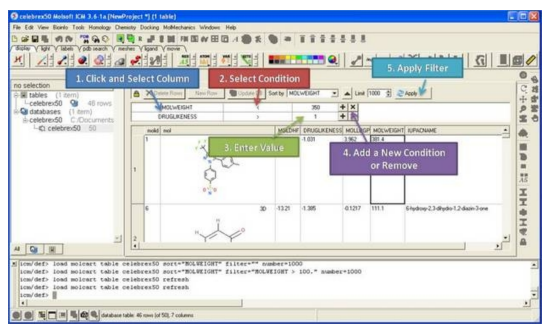


#### To filter a database:

• Double click on the panel labeled and the filtering tools will be activated.

H / / / @ @	uble click h	ere to activ			-	] <u> </u>	<u>**   *   X</u>	<b></b>	
databases (1 tem)	cpick o	okevo		_		+ ×			1
⇒celebrex50 C/Documents	molid mol			MOLDHE	DRUGUKENE	SS MOLLOO	MOLWEIGHT	IUPACNAME	
	1	:0 . Y	0-		-1.031	3962	281.4		
	2	"	**************************************	43.21	-1.385	0.1217		64ydoy-23dfydo-12daan-3one	

- Click and select a column to filter (as shown below).
- Enter a condition.
- Enter a value.
- Add or remove another condition
- Click the **Apply** button



To view a diverse set of rows from your database:

• Right click on the database in the ICM Workspace and select **Select Diverse Set**.

## 12.3 Edit Database

```
Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS
```

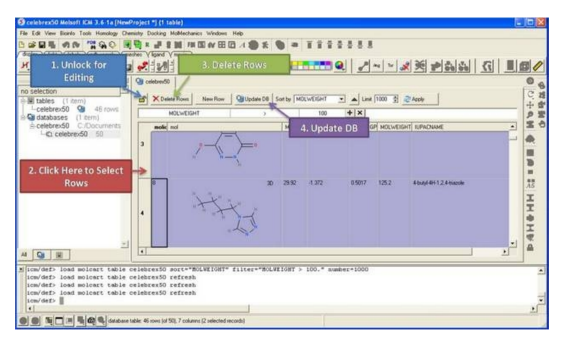
To make any changes to a database you must first unlock it:

- Click on the **Allow editing rows** button which has a picture of a lock on it.
- Once this button has been selected the **Delete Rows**, **New Row**, and **Update DB** buttons will become activate.

File Edit Verre Biorio Tool Homologi Der Sei Elle Vite Vite Vite Vite Vite Vite Vite Vit		n this butto	18 .	the d	atabası	• • • • •		× 2444 ()		0 0.
l-celebrex50 ♀ 46 rows ⊮Q databases (1 tern)	MOLWER	and the second sec	4		350 + ×				_	+ 9 5
i÷celebrex50 C./Documents i–©: celebrex50 50	motest and motest		20	-146.3 -1.1	0RUGLIKENESS -1.001	0.1237	P MOLWEIGHT 1381.4	T JUPACNAME	-	A BBS IN HHAHE
		NOLWEIGHT" f:				aber=1000			J <sup>I</sup>	HWa .

## To delete rows

- Select the row(s) by clicking on the row numbers. A range of rows can be selected by holding the shift key and clicking a non-contiguous set of rows can be selected by holding down the control key.
- Click the **Delete Rows** button.



#### To insert a new row

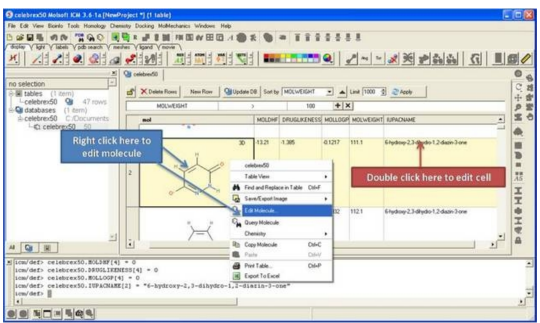
- Select a row.
- Click on the **New Row** button. A new row will be inserted underneath the selected row.
- Click Update DB

#### To edit strings and numbers in a row

- Double click on the cell you wish to edit and then enter a new value.
- Click Update DB

## To edit a chemical (2D sketch):

- Right click on the cell and select Edit Molecule
- Make changes using the Molecular Editor
- Click Update DB

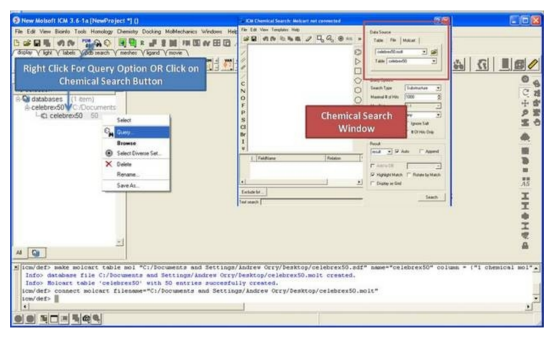


## 12.4 Query Local Database

Available in the following product(s): ICM-Chemist | ICM-Chemist-Pro | ICM-VLS

#### To query a local database:

- Right click on the database in the ICM Workspace and select Query.
- Query using the chemical search tools.



# 13 ICM in KNIME

|Video - Installation|Video - MolCalcProp Node|

KNIME (Konstanz Information Miner) is a user-friendly and comprehensive open-source data integration, processing, analysis, and exploration platform. Currently we have KNIME nodes for the following ICM functions:

## Available Nodes

### **Docking and Screening**

- **MolIcmDockPrep** Prepare receptor and ligand for docking. Covalent docking support for MolIcmDockPrep node (ICM 3.8-1 or higher is required)
- **MolIcmDock** Ligand Docking and Screening (\_dockScan and \_dockProjPrep). Recently we have added "Output Type" for MolIcmDock node. It allows to choose between SDF and interactive ICM HitList which can be viewed using the MolView node.
- **MolScreen** MolScreen is a set of high quality 2D fingerprint and 3D pharmacophore models for a broad range of pharmacology and toxicology targets.

## Cheminformatics

- MolConverTo3D Chemical 2d to 3d conversion.
- MolConfGen Conformation generator (\_confGen).
- MolAPFSuperimpose Flexible APF superposition (\_chemSuper).
- **MolCalcProp** Allows to calculate various chemical properties and desciptors (e.g. Calculate MolArea, MolWeight, InChi...). For a complete list of available functions please click here.
- **MolStandardize** Chemical Table/Spreadsheet standardization remove salt, remove hydrogen, standardize groups, apply 2D depiction, assign formal charges.
- MolUnique removes duplicate molecules.
- MolCenters Performs chemical clustering and selects a specified number of representatives.
- **MolCluster** Cluster either by fingerprints or by numerical column selection.

## Visualization

• MolView - Connect this node to the output of other Molsoft nodes. (e.g. MolScreen, MolIcmDock...).

## To use ICM in KNIME

- 1. Download KNIME (Analytics Platform) from www.knime.org
- 2. In KNIME go to File/Install KNIME extensions and search for and select the KNIME Base Chemisty Types and Nodes. When you do this you will see a Chemistry directory in the Node Repository.
- Download MolSoft's KNIME Java (see instructions below). Note, that if you have an old version of Molsoft Nodes installed manually you need to remove the jar file and restart KNIME with "-clean" option.

## MolSoft Knime

• To enable automatic update of molsoft nodes: go to File/Preferences. Open Install/Update - Available Software Sites, Click on "Add..." button and enter information below:

🔺 Preferences		
type filter text	Available Software Sites	
<ul> <li>General</li> <li>Help</li> <li>Install/Update</li> <li>Automatic Updates</li> <li>Available Software Sites</li> <li>KNIME</li> <li>Molsoft</li> </ul>	type filter text         Name       Location         Image: Construct of the state	Add Edit
⊞ Run/Debug ▲	minduicy-cc	Remove
_	ame: molsoft Local	Reload
Ŀ	ocation: http://molsoft.com/knime	Enable
		Import
	OK Cancel	Export,
	OK	Cancel

• Then, goto "File/Install update extensions", uncheck "Group items by category" If you don't have "Base Chemistry" package installed, then install it first (type 'chem' to locate package)

🔺 Install				_ 🗆 🗙
Available Software Check the items that you wish to install.				
Check the items that you wish to install.				
chem				æ
Name		Version		
🔲 🕼 ChemAxon/Infocom Marvin Extensions Featu	ıre	2.6.3.v01	35	
🗹 称 KNIME Base Chemistry Types & Nodes		2.9.1.004	1076	
🛛 🗖 🖗 KNIME CDK Chemistry Feature (deprecated)		2.9.1.004	1076	
KNIME Chemistry Add-Ons		2.9.1.004	1076	
🗌 🔲 🏧 Source for KNIME Base Chemistry Types & N		2.9.1.004	1076	
🗌 🔲 👰 Source for KNIME CDK Chemistry Feature (de	eprecated)	2.9.1.004		
Source for KNIME Chemistry Add-Ons		2.9.1.004	1076	
Select All Deselect All 1 item selected				
Details				
☑ Show only the latest versions of available software	🔽 Hide ite	ems that are alre	ady installed	
Group items by category	What is	already installed?		
🔲 Show only software applicable to target environment				
	< Back	Next >	Finish	Cancel

• Next install 'Molsoft' nodes. Please uncheck the option "Group items by category" other wise it will not list the nodes.

🔺 Install		
<b>Available Software</b> Check the items that you wish to install.		
molsoft		R
Name	Version	
Molsoft Knime Nodes	1.0.0.19	
Select All Deselect All 1 item selected		
Details		=
$\overline{\buildrel {\it V}}$ Show only the latest versions of available software	$\overline{ullet}$ Hide items that are already installed	
Group items by category	What is <u>already installed</u> ?	
Show only software applicable to target environment		
	< Back Next > Finish	Cancel

• After this point everything is setup. You can either go to File/Update KNIME to get the updates manually or setup automatic update.

## How to run MolSoft in Knime:

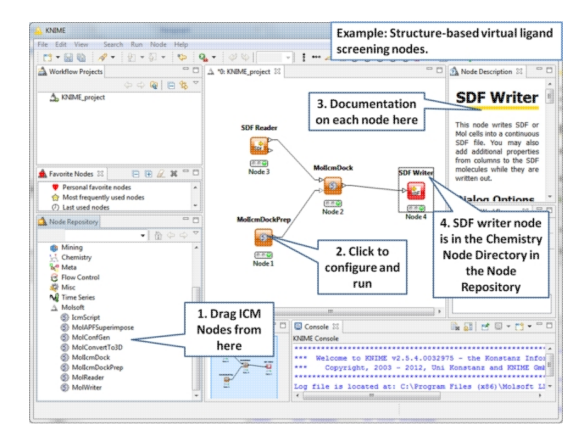
**Setting Path to ICM Executable** Usually KNIME can figure out where the Molsoft ICM executable is located. In some cases when there are multiple versions of ICM it might return an error as shown below:

Execute failed: Cannot run program "/icm": CreateProcess error=2, The system cannot find the f

To fix this:

- File/Preferences
- Select MolSoft and then browse for the location of ICM e.g.C:\Program Files (x86)\Molsoft LLC\ICM-Pro

Here is an example for Structure-based virtual ligand screening in KNIME:



# **14 Tutorials**

Graphics Tutorials Malapula Paragantetian, Coloring, Lobaling		
Molecule Representation, Coloring, Labeling and Annotation	HTML	
Creating Fully Interactive Slides for PowerPoint and the Web	HTML	Reference Guide  Webinar
Sequence and Alignment Tutorials		
Read Sequences and Align	HTML	Video
Link Sequence to Structure	HTML	
Alignment Annotation	HTML	
Protein Structure Tutorials		
PDB Search	HTML	
Convert PDB to ICM Object	HTML	
Superimpose Proteins	HTML	
Structure Analysis	HTML	
Calculate contacts	HTML	
Display hydrogen bonds	HTML	
Identify ligand binding pockets	HTML	
PDB Preparation: Symmetry	HTML	
PDB Preparation: Occupancy & B-Factors	HTML	
PDB Preparation: Alternative Orientation	HTML	
PDB Preparation: Biomolecule	HTML	
Protein Modeling Tutorials	1111111	
Building an Homology Model	HTML	
GPCR Modeling Example	HTML	
Loop Modeling	HTML	
Predicting the Effect of a Mutation on		
Binding	HTML	
Predicting the Effect of a Mutation on	HTML	
Stability		
Cheminformatics Tutorials		
Chemical Sketching using the Molecular Editor	HTML	Reference Guide
Chemical Search	HTML,	Refrence Guide
Chemical Clustering	HTML,	
Working with Large Chemical Space	HTML,	
3D Pharmacophore Search	HTML	Video
2D Pharmacophore Search	HTML	Video
How to Create a Markush Structure	HTML	Video
How to Enumerate a Markush Library	HTML	Video
How to Decompose a library based on a Markush Library	HTML	Video
How to Enumerate a library by reaction	HTML	Video
Detecting Activity Cliffs	HTML	
Atomic Property Fields (APF)		
APF Superposition, Score, Consensus, Screening and Clustering.	HTML	
Ligand Editor Tutorials		
Working with the ICM Interactive 3D Ligand	T T(T) N A T	
Editor Ligand Docking Tutorials	HTML	Reference Guide.
Dock Biotin to the Streptavidin Receptor	HTML	Video
Re-Dock an Inhibitor to Ricin Crystal		
Structure	HTML	Video

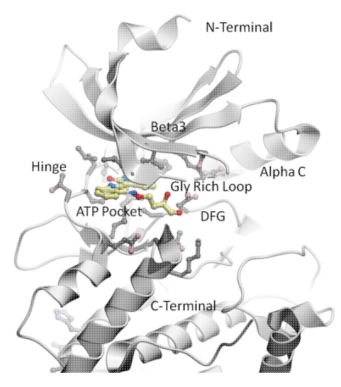
Protein-Protein Docking Tutorial
Protein-Protein Docking using FFT Method HTML
Virtual Screening Tutorials
Virtual Ligand Screening to Ricin Receptor HTML
Virtual Ligand Screening to Cyclooxygenase HTML Video-1   Video-2
Docking a Markush Library HTML
Induced Fit Docking Tutorials
Multiple Receptor Conformation Ensemble HTML
Explicit Group Docking HTML
QSAR Tutorials
2D FingerPrint QSAR Model HTML
2D 3D Pharmacophore QSAR Model HTML

## **14.1 Graphics Tutorials**

## Introduction

The following examples are focused on the structure of the kinase catalytic domain. The catalytic domain is nestled between the N- and C-terminal and has high sequence conservation between kinase families. The adenine moiety of ATP interacts with the hinge region which links the N- and C-terminals of the catalytic domain. The ATP sugar binding region is on the "floor" of the pocket where ribose makes a hydrogen bond with a polar residue. A conserved lysine residue on the beta3 strand is part of the "roof" of the pocket where the triphosphate group of ATP binds. A flexible glycine-rich loop moves in and out of the pocket depending on the ligand bound state of the PK and is regulated to some extent by the movement in and out of the pocket by the AlphaC helix. A buried region at the "back" of the pocket is protected by a "gatekeeper residue" forming a variable hydrophobic cavity. The hydrophobic cavity along with the DFG motif region are of interest for drug design because it opens up regions of the pocket which are not conserved and do not bind ATP.

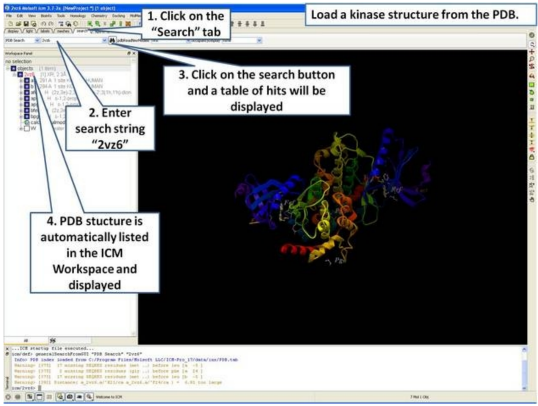
To learn the basics of the graphical user interface we will annotate the key structural regions of a protein kinase.



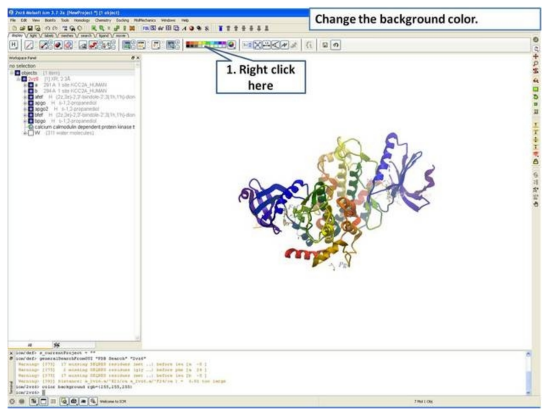
## 14.1.1 Change Molecule Representation and Color

This tutorial shows you how to:

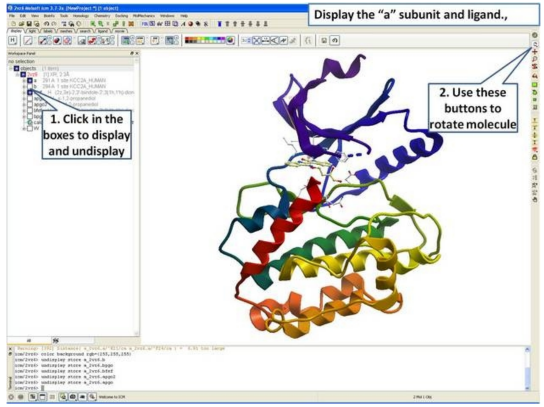
- How to load a PDB structure into ICM.
- How to change the background color.
  How to display and undisplay a molecule.
  How to color ribbon representation.



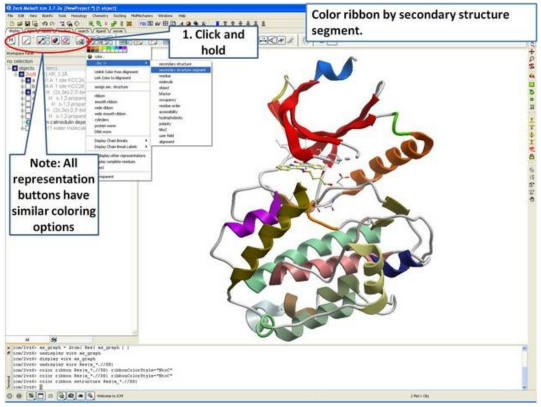
**Step 1:** Click on the search tab to load and display the PDB structure 2vz6



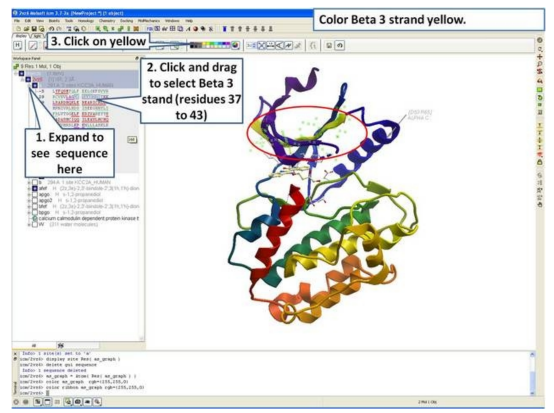
**Step 2:** Change the background color by right clicking on the color palette.



Step 3: Display the "a" subunit of the kinase.



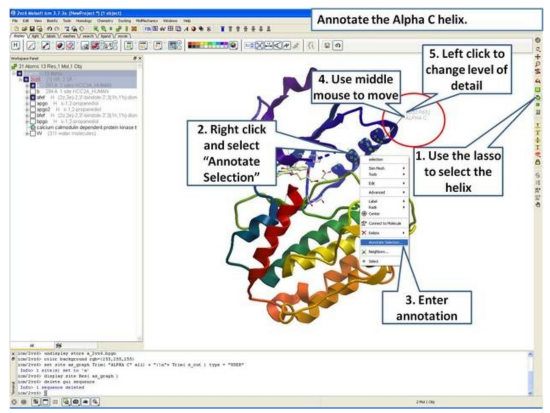
Step 4: Use the options in the "Display Tab" to color the ribbon.



Step 5: Select the beta 3 strand (residues 37:43) and color yellow.

#### 14.1.2 Annotation

This tutorial shows you how to add user defined annotation to a particular region of a protein structure.

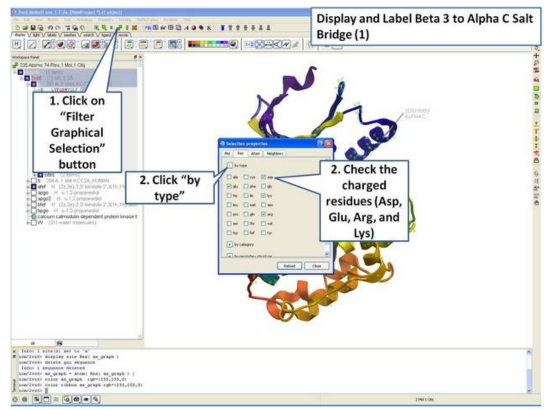


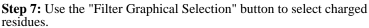
**Step 6:** Select the region you wish to annotate and right click on the selection for options.

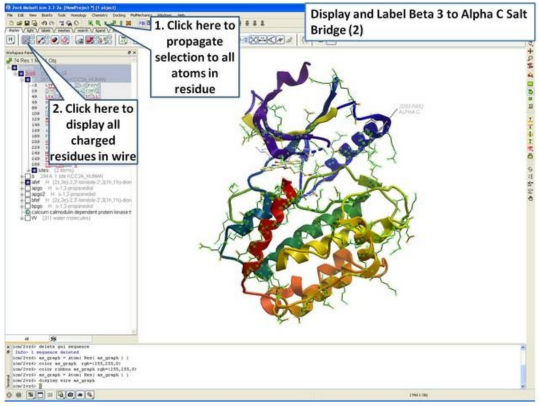
### 14.1.3 Labels

This tutorial is a continuation from the previous tutorials in this section and shows you how to:

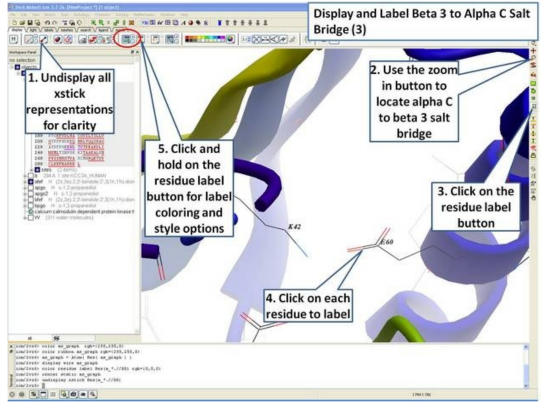
- How to select individual residue types.
- How to propagate a selection from one atom to all.
- How to use the residue label button.
- How to make a spherical selection.
- How to label a residue selection.



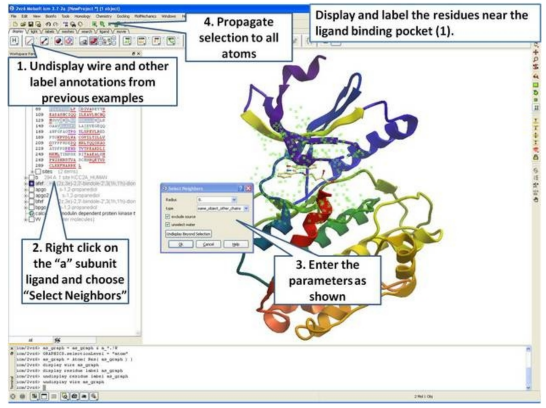




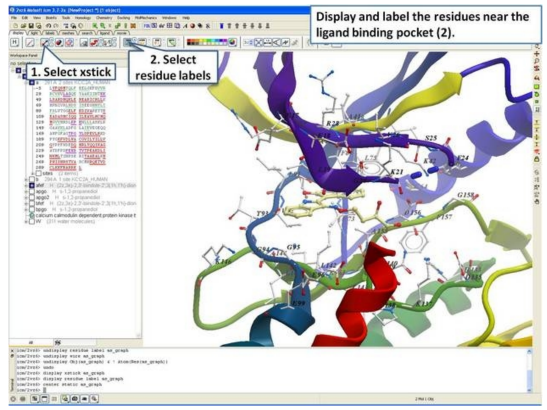
**Step 8:** Propagate selection to all taoms to display all atoms of the side chain.



**Step 9:** Locate the B3 to AlphaC salt bridge (K42- E60) and label using residue label button. **Note:** Once you have selected the two residues you can invert selection (click on invert selection button) and then undisplay all the other charged residues.



Step 10: Select neighboring residues to the ligand.

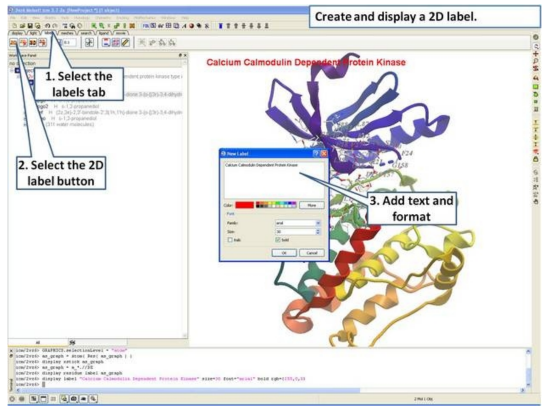


Step 11: Display the residues and label them.

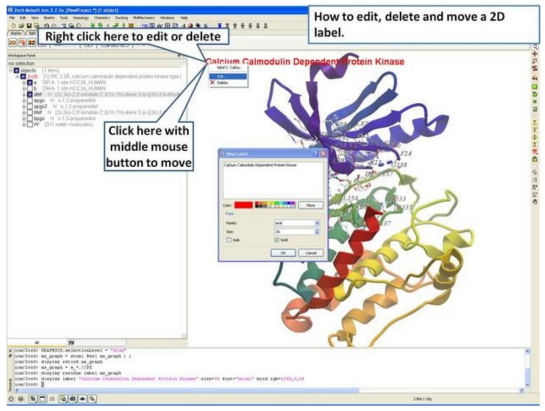
### 14.1.4 2D and 3D Labels

This tutorial is a continuation from the previous tutorials in this section and shows you how to:

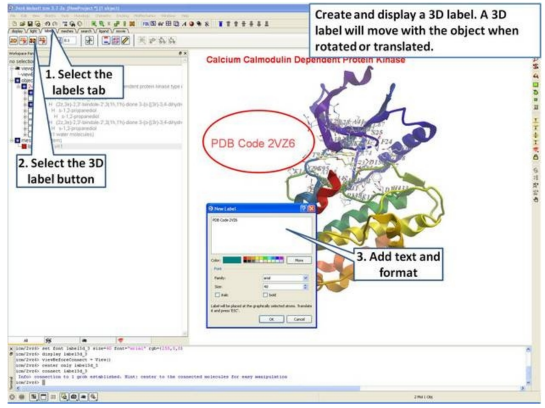
- Create and display a 2D label.
- Delete and move a 2D label.
- Create and display a 3D label.
- Delete and move a 3D label.



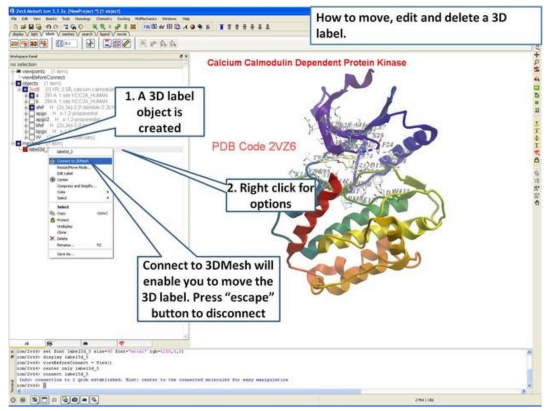
Step 12: Create a 2D label using the options in the "labels" tab.



Step 13: Right click on the 2D label to edit or delete it.



Step 14: Create a 3D label using the options in the "labels" tab.



Step 15: Right click on the 3D label to edit or delete it.

### 14.2 Creating Fully Interactive Slides for PowerPoint and the Web Tutorial

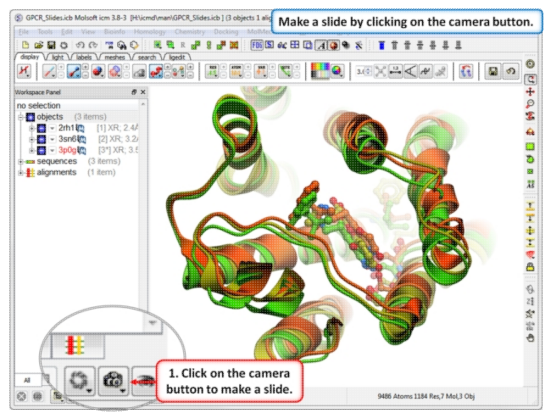
## Introduction

A slide enables you you to conveniently store a large number of different 3D visualization properties (e.g. color, viewpoint, representation etc...) along with text and window layout. ActiveICM enables you to view and display ICM graphical slides and animations interactively inside Windows Microsoft PowerPoint and web browsers such as Internet Exporter and Mozilla Firefox.

In this example we are going to look at the effect of an agonist binding to the Beta-2-Adrenergic GPCR (B2AR) and prepare a molecular document. A prepared ICM file containing three B2AR GPCRs superimposed can be downloaded here ftp://ftp.molsoft.com/pub/workshop/ (1. cut and paste the address into a web browser. 2. Right click on GPCR\_slides.icb and choose "Save Link As". 3. Open the .icb in ICM File/Open). The file contains pdb files 2RH1 (R inactive state), 3POG (R\* active state, with G-alpha mimic), and 3SN6 (R\*G G-protein signalling state). The agonist ligand causes Helix 5 to shift inwards and Helix 6 to swing outwards. This agonist binding poses were predicted using ICM a year before the crystal structures were solved, you can read about this here and a good review of the structure and function of GPCRs here.

# Tutorial

In this tutorial we will build a series of slides and then annotate them using HTML to create hyperlinks to the slides. The resulting molecular document can be exported in movie format or in fully interactive 3D files.



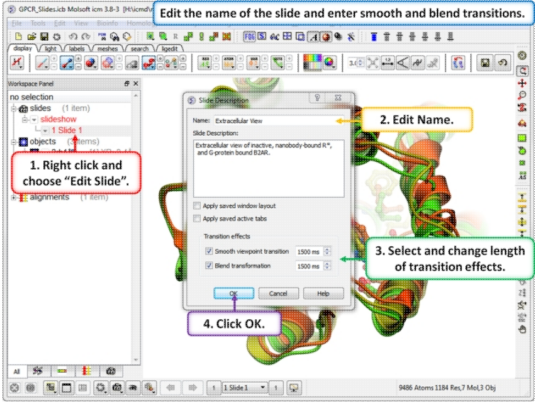
**Step 1:** Download the example file here

ftp://ftp.molsoft.com/pub/workshop/ (1. cut and paste the address into

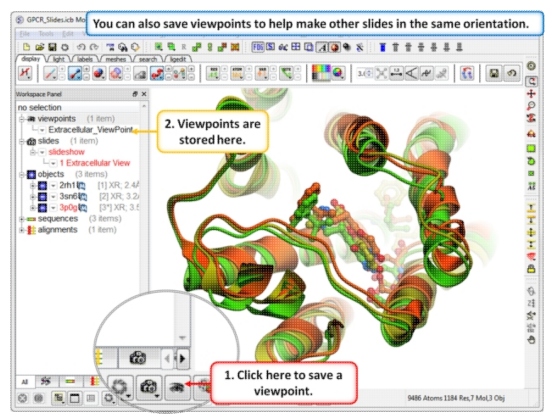
a web browser. 2. Right click on GPCR\_slides.icb and choose "Save Link As". 3. Open the .icb in ICM File/Open). Make a slide by clicking on the camera button at the bottom of the GUI.



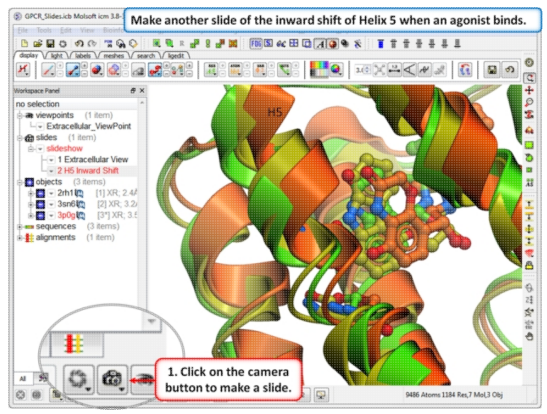
**Step 2:** The slide is stored in the ICM Workspace panel (left hand side of the GUI). You can make changes to the 3D display and return to the display in the slide by clikcing on the slide in the ICM Workspace.



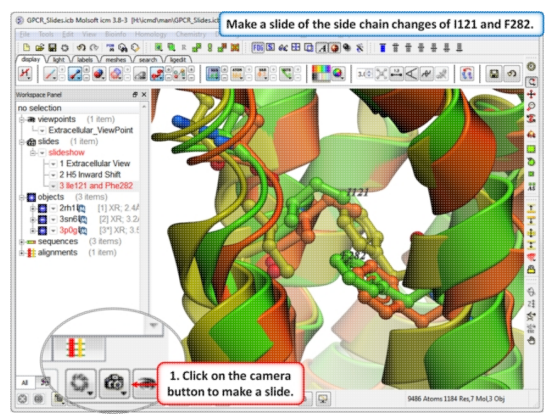
**Step 3:** Properies of the slide can be edited by right clicking on the slide in the ICM workspace.



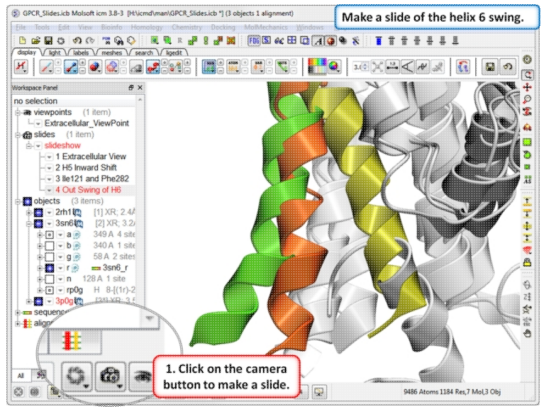
**Step 4:** Saving a viewpoint is useful if you are making many slides from the same viewing position.



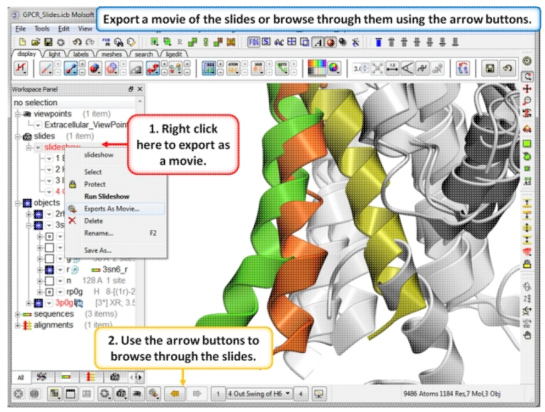
**Step 5:** In this slide we show that Helix 5 makes an inward shift when an agonist binds.



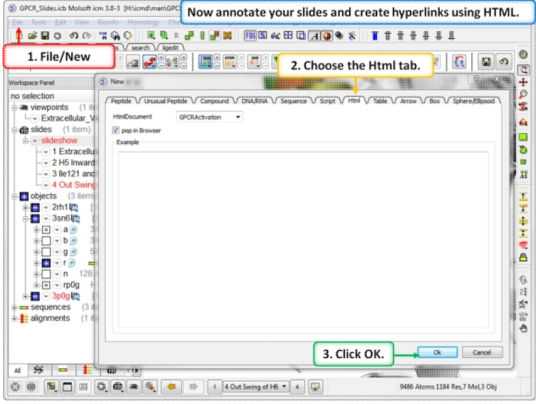
**Step 6:** In this slide we show that upon binding an agonist Ile121 switches its rotameric state and Phe 282 moves.



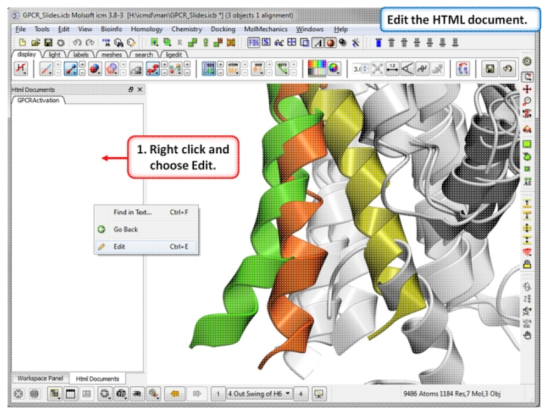
**Step 7:** In this slide we show that agonist binding causes Helix 6 to swing out.



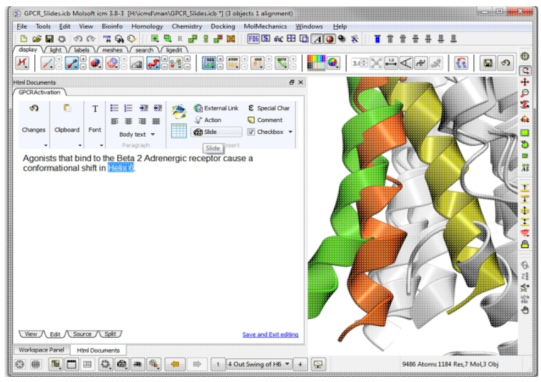
**Step 8:** You can export the slides as a movie or you can browse through them using the arrow buttons.



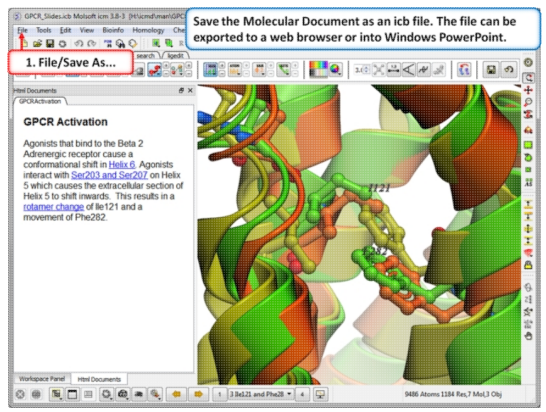
**Step 9:** To build a molecular document you can annotate the slides using HTML.



**Step 10:** Edit the document by right clicking on the HTML panel and choosing **Edit.** 



**Step 11:** Hyperlinks can be made to the slide. Other functions such as adding check boxes, ICM scripts and external links are also available.

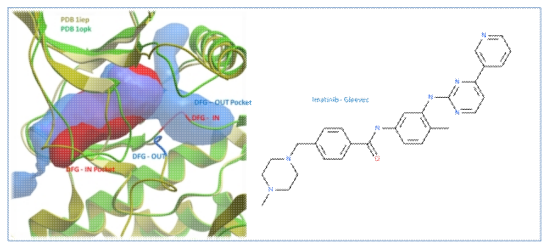


**Step 12:** Save the molecular document in icb format and view in ICM-Pro or ICM-Browser. You can also export to a web browser or view the fully interactive slides in Windows Powerpoint using a plugin called ActiveICM.

# 14.3 Protein Structure Tutorials

# Introduction

The positioning of the DFG motif in the activation loop of a protein kinase has an effect on the size and properties of the ATP binding pocket. Most kinase inhibitors target the kinase with the DFG inwards to the ATP binding site. Type-II inhibitors target the site where the DFG motif is in the out position which opens up the pocket and provides additional hydrophobic binding sites. Targeting DFG-out conformations can improve inhibitor specificity and slower-off rates.

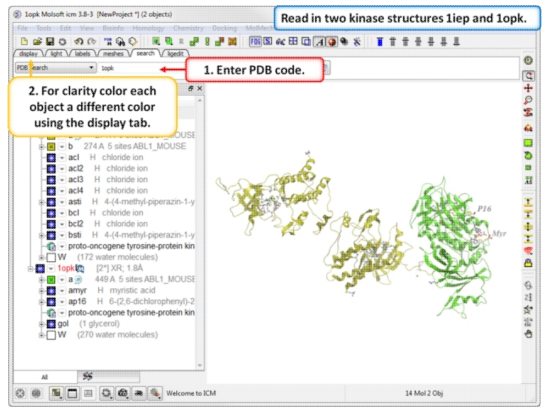


Comparison of the crystal structures of Abl kinase with DFG-IN and DFG-OUT (Left). Imatinib (Gleevec - right) binds to the DFG-OUT conformation.

# Tutorial

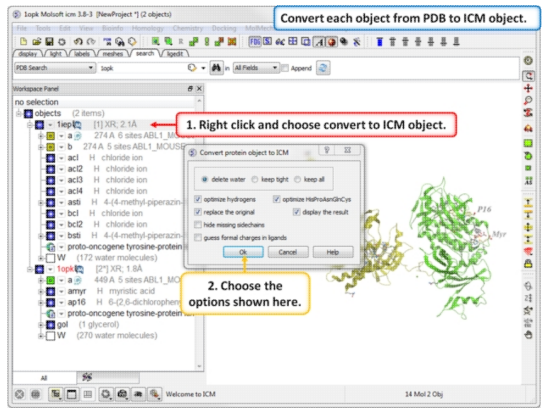
In this example we will look at two kinase structures and compare their binding pockets. One of the structures is bound to the Type-II inhibitor Imatinib (Gleevec) a drug used to treat Chronic Myelogenous Leukemia. We will use ICM tools to look at the side-chains that interact with the inhibitor and compare the pocket with a DFG-in structure using ICMPocketFinder. Please click on the links provided to learn more about the methods used.

### 14.3.1 PDB Search



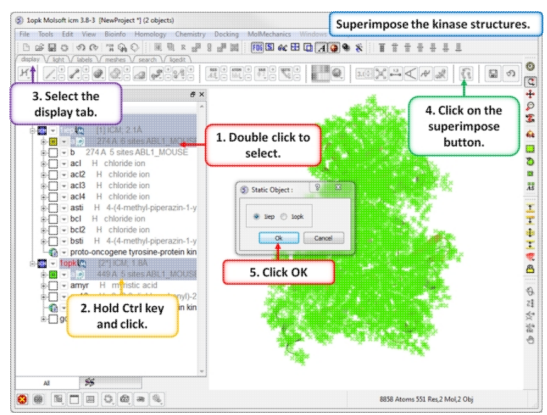
**Step 1:** Click on the Search Tab and choose "PDB Search" and oad the PDBs structures liep and lopk. You can color each object a unique color by clicking on the Display Tab and clicking and holding on the ribbon button and selecting Color by>Object.

#### 14.3.2 Convert PDB to ICM Object



Step 2: Convert each PDB file into an ICM object.

### 14.3.3 Protein Superposition

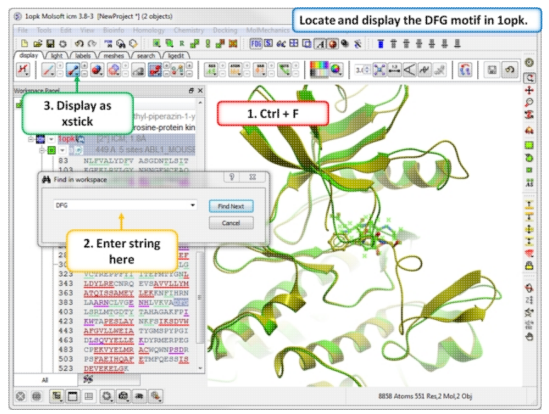


Step 3: Superimpose the two kinase structures.

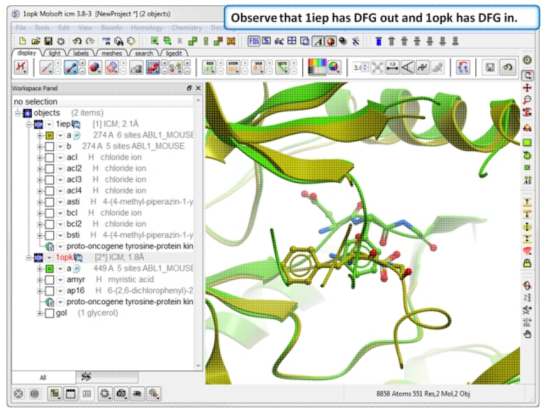
### 14.3.4 Protein Structure Analysis

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265 SLTVAVETLE EDTMEVEEPL	
285 KEAAVMKEIK HPNLVQLLGV	
305 CTREPPFYII TEFMTYGNLL	
325 DYLRECNROE VSAVVLLYMA	1 LET
345 TOISSAMEYL EKENFIHEDL	F382282
365 AARNCLYGEN HLVEVA	and the second
385 SRIMTGDTYT AHAGAKFPIK 405 WTAPESLAYN KFSIKSD	
405 HTAPESLAYN KFSIKSpann 425 FGVILWEIAT YGMSPY 44 Find in workspace	
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465 PEKVYELMRA CWOWNE	
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**Step 4:** Search the "a" subunit of 1 iep for the "DFG" motif. You can press Ctrl F and expand the sequence in the ICM workspace. Display the DFG region in xstick.

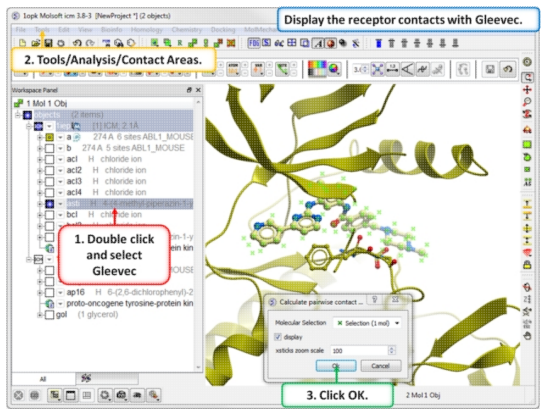


**Step 5:** Search the "a" subunit of lopk for the "DFG" motif. You can press Ctrl F and expand the sequence in the ICM workspace. Display the DFG region in xstick.

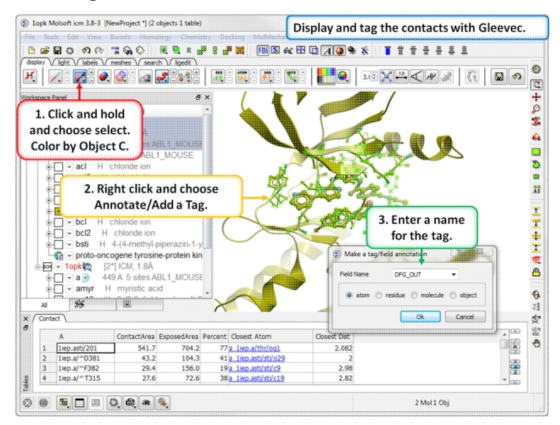


Step 6: Observe the DFG location in both structures.

### 14.3.5 Contact Areas

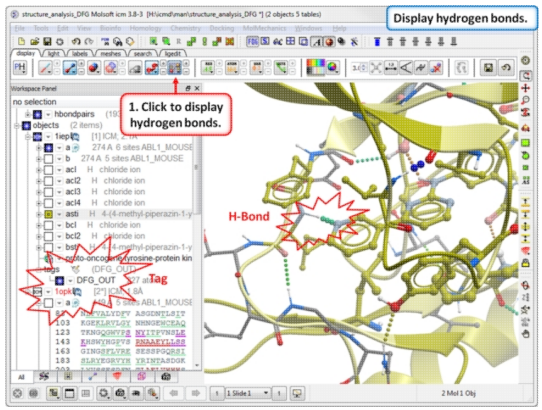


**Step 7:** Use Tools/Analysis/Contact Areas to display the contacts with the Gleevec ligand.



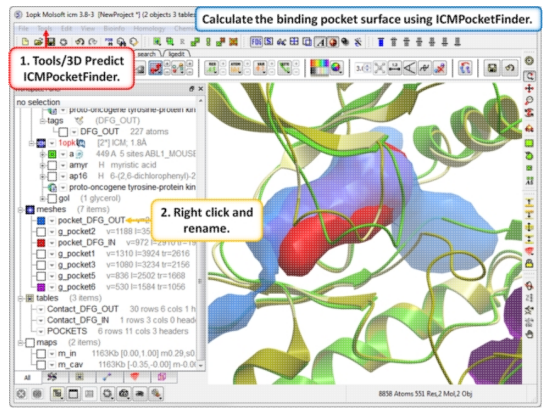
**Step 8:** Tag the residues surrounding the ligand. Repeat Steps 7 and 8 for the 10pk structure and ligand.

### 14.3.6 Hydrogen Bond



**Step 9:** Compare the hydrogen bond interaction of the Type I and Type II inhibitors with the hinge region (liep - residues 317:322) of the kinase.

### 14.3.7 Pocket Identification



**Step 11:** Use ICMPocketFinder to compare the pockets of DFG-IN and DFG-OUT. Run ICMPocketFinder for each structure - rename the pocket so as not to overwrite it.

#### 14.3.8 Protein Preparation and Crystallographic Analysis Tutorial

### 14.3.9 PDB Preparation - Symmetry

**Background** When inspecting a ligand binding pocket it is important to check that the true pocket is formed by chains which are not explicitely present in a PDB entry. Therefore it is necesary to use **Tools/X Ray/Crystallographic Neighbor** to find all molecules/subunits or chains involved in the interaction with the ligand. Molecular objects and 3D density maps may contain information about crystallographic symmetry. It consists of the following parameters:

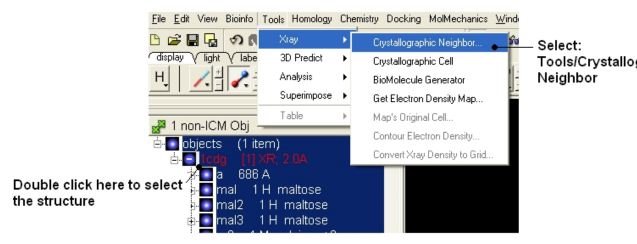
- 1. Crystallographic group eg. P2121 that determine N (depends on a group) transformations for the atoms in the asymetric unit.
- 2. Crystallographic cell parameters A, B, C, Alpha, Beta and Gamma

To generate the coordinates within one cell one needs to apply N transformations and then to generate neigboring cells the content of one cell needs to be translated in space according to the cell position.

**Example** As an example let us look at Cycloldextrin glycosyltransferase (PDB Code: 1CDG). The problem with docking to this receptor is that the true pocket is formed by chains which are not explicitly present in the PDB entry. Site mb1 includes serine 382. This cannot be predicted just by looking at the structure. Therefore we need to identify symmetry related molecules to this protein.

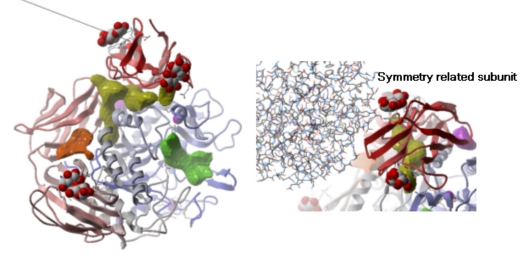
- Use the PDB search tab to load the crystal structure 1cdg.
- Inspect the ligand binding pocket of **maltose** (mal)

• To identify if there are any other chains involved in the interaction with the ligand select the whole structure in the ICM Workspace.



- Tools/Crystallographic Neighbor
- Select a 7A radius
- Check "create symmetry related molecules" and "display symmetry neighbors".
- Inspect the neighbors surrounding **maltose(mal)**. Each symmetry related subunit can be colored by object by clicking and holding the representation button in the display tab and selecting color-by.

site mb1 includes residue ser 382 for symmetry-related molecule. site mb3 includes the following residues for symmet



#### 14.3.10 PDB Preparation - Occupancy and B-Factors

**Background** When preparing a PDB for analysis (eg docking or modeling) it is important to check the reported occupancies and b-factors. The occupancy is a fraction of atimic density at a given center. If there are two eqally occupied conformers both will have an occupancy of 0.5 - the normal value is 1 range 0-1. The \*{B-Factor} is the mean-square displacement of atom from its position in the model - the normal range is 5-50.

One way of visualizing the occupancy and b-factor is by coloring the structure by these values. You can do this by clicking and holding on a representation button in the **display** panel and selecting Color-by.

As an example let us look at the crystal structure 1ATP

- Type in the PDB search tab 1atp and the structure will be displayed in the graphical display.
- Use the ICM workspace to undisplay everything except for the "e" subunit. You can do this by clicking in the blue boxes in the ICM Workspace.
- Display the "e" subunit in wire representation using the wire button in the **display** tab.
- Click and hold on the wire button and select Color-by B-Factor. Regions of high B-factor are colored red.

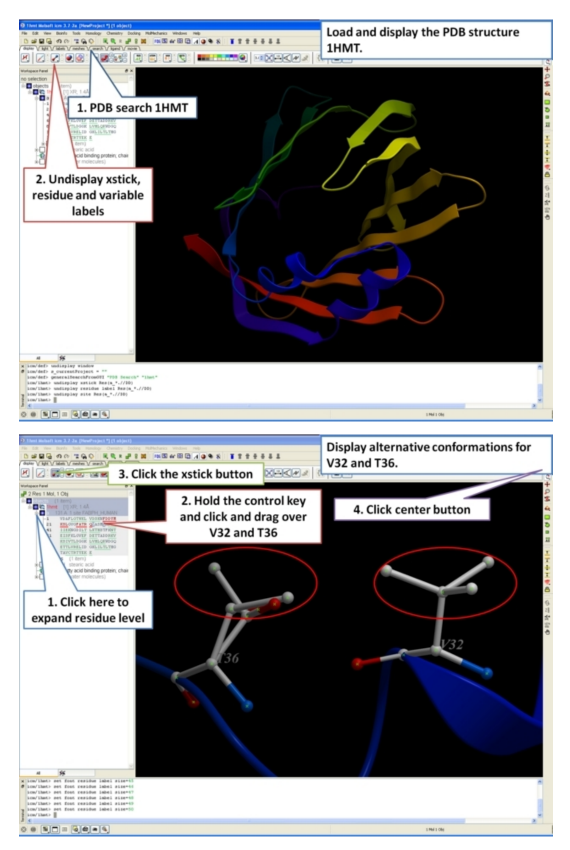
#### Click and hold display labels V odb search V meshes √ movie н no sele 🥯 color.. ė-**o** o .by -> atom type ÷... wire residue sc chemsitry residue tree molecule thin molecule\_C normal object bfactor thick occupancy undisplay other representations reset to default accessibility hydrophobicity select polarity sec. structure NtoC user atom field user res. field alignment

#### Residues with high B-Factor are colored red

# 14.3.11 PDB Preparation - Residue Alternative Orientation

For some very high resolution structures two alternative conformations for a residue are provided. Therefore for docking you need to decide to use one conformation of the residue or generate seveal separate docking models. This could be performed using multiple receptor conformation docking.

Here is an example of alternative residue orientations found in a crystal structure of a Fatty Acid Binding protein in complex with stearic acid.



### 14.3.12 Biomolecule Generator

#### Objective

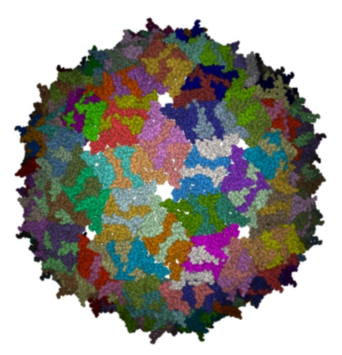
Here we will investigate the biological environment of a virus protein . PDB code 1DWN.

#### Background

It is very useful to know how a protein from the PDB may look in a biological environment. The PDB entries solved by X-ray crystallography and deposited in the PDB contain the information about the crystal structure rather than the biologically relevant structure. For example, for a viral capsid only one instance of capsid protein complex will be deposited and only one or two molecules of haemoglobin that is a tetramer in solution maybe deposited. In some other cases the asymmetric unit may contain more than one copy of a biologically monomeric protein. ICM reads the biological unit information and has a tool to generate a biological unit. Not every PDB entry has the biological unit information.

#### Instructions

- Read and load the PDB file 1DWN
- Tools/Xray/Biomolecule Generator
- Tick the makeAllBiomolecules box (Warning this may take a few minutes to generate)
- The generated molecules will be listed in the ICM Workspace. Each one can be selected and displayed. The biomolecule is shown below.



**NOTE:** Please note that right clicking on a PDB file in the ICM Workspace will tell you whether there is any **Biomolecule** information available for the structure. If this information is not present then the option will be greyed out.

#### Manual References (Web Links)

Biomolecule Generator

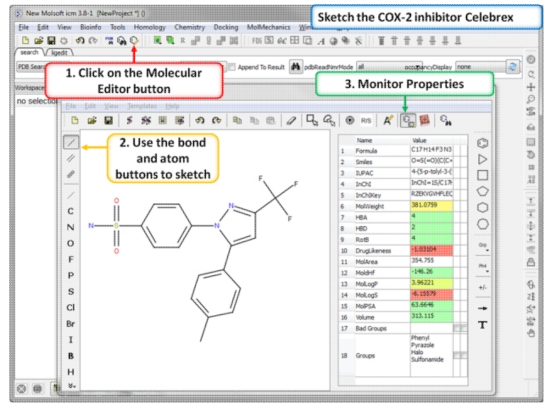
# **14.4 Cheminformatics Tutorials**

### 14.4.1 Chemical Draw and Spreadsheet Tutorial

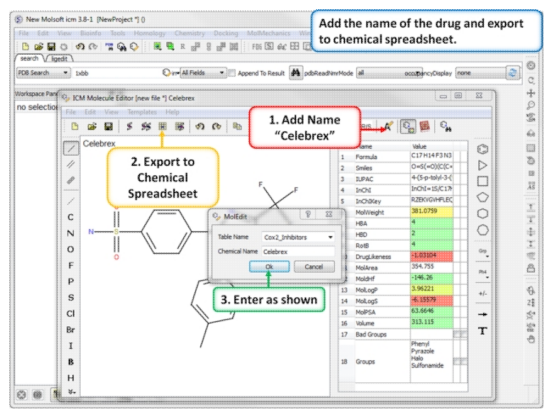
# Background

Cycloxygenase 2 (COX-2) inhibitors can target inflammation and pain and are non-steroidal anti-inflammatory (NSAID) drugs. Some of the COX-2 inhibitor drugs have an interesting history because some of them cause a significant increase in heart attacks and stroke in patients. Probably the best known COX-2 inhibitor is Merck's drug Vioxx which was voluntarily removed from the market in 2004. In this example we are going to build a small database of COX-2 inhibitor drugs, learn how to import COX-2 inhibitor data from ChEMBL and how to manipulate chemical spreadsheets in ICM.

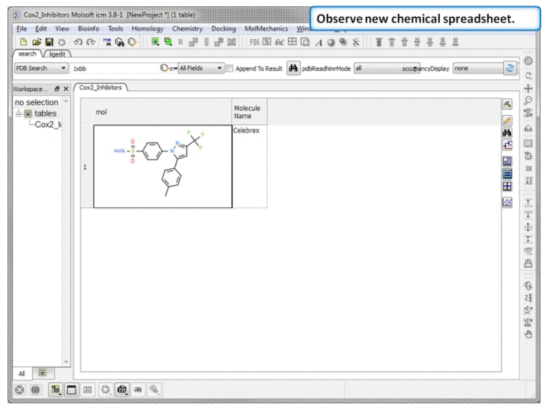
# Tutorial



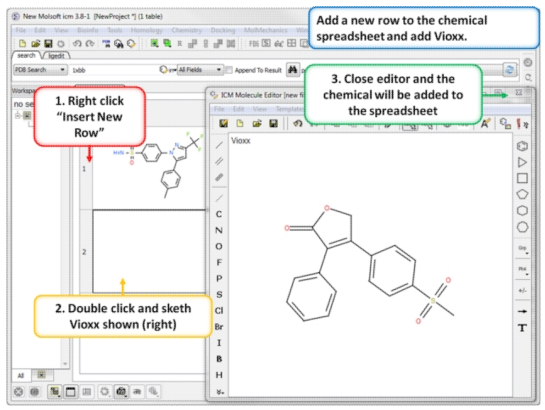
**Step 1: Sketch Celebrex.** The first COX-2 drug we are going to draw is Celebrex (Celecoxib) from Pfizer. Click on the Molecular Editor button and use the bond and atoms button to skethc Celebra (shown). Note you can monitor key chemical properties of the molecule as you sketch it.



**Step 2: Export to Chemical Spreadsheet.** Rename the chemical to its brand name (Celebrex) and export to a chemical spreadsheet.



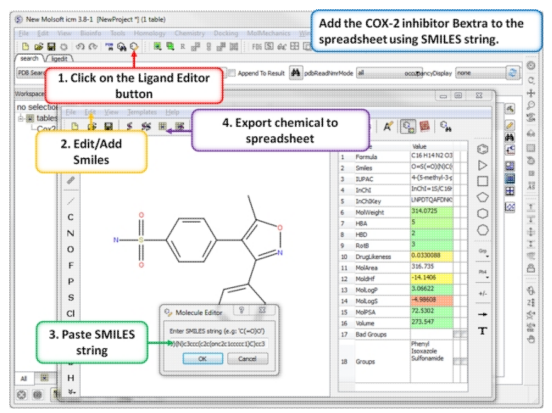
**Step 3: Observe Chemical Spreadsheet.** You should now have one chemical in your Cox2\_Inhibitors spreadsheet.



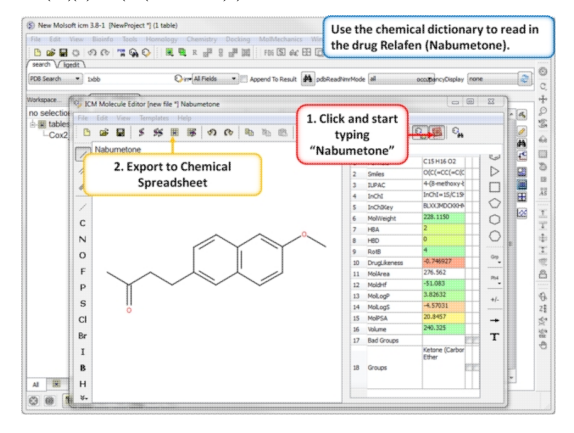
**Step 4: Add a new row.** Add a new row to the spreadsheet and then sketch the Vioxx (Rofecoxib) drug (shown).

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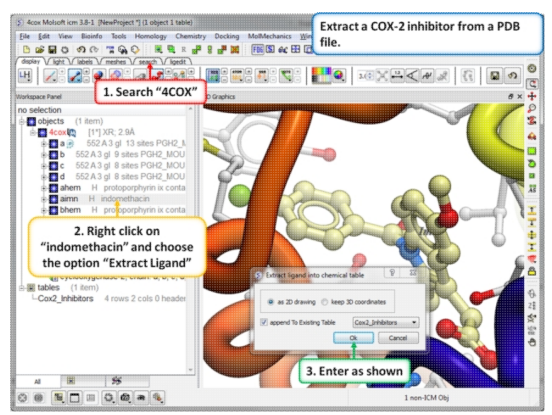
**Step 5:Observe Chemical Spreadsheet.** Now you should have 2 chemicals in your spreadsheet.



**Step 6: Using SMILES.** A convenient way to import chemical data is by using SMILES. Bextra (Valdecoxib) is a COX-2 inhibitor that was removed from the market by Pfizer in 2005 due to concerns of an increased risk of heart attack and stroke. Bextra's SMILES string is: O=S(=O)(N)c3ccc(c2c(onc2c1ccccc1)C)cc3



**Step 7: Using the Chemical Dictionary.** Another convenient way to import chemical data is to use the in built chemical dictionary. Relafen is Nabumetone is an NSAID that is rapidly metabolized in the liver to a major active metabolite, 6-methoxy-2-naphthyl acetic acid.



**Step 8: Extracting Chemicals from the PDB.** You can extract a chemical from a PDB file into a chemical spreadsheet by right clicking on the ligand in the ICM workspace and choosing the option "Extract Ligand".

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**Step 9: Calulate Properties and Insert Columns.** Use the Chemistry/Calculate Properties to calculate properties (e.g. logP, logS, MW) for the drugs in the spreadsheet.

In this example we will import more COX-2 inhibitor information from ChEMBL. ChEMBL is a database of bioactive drug-like small molecules curated from the primary scientific literature.

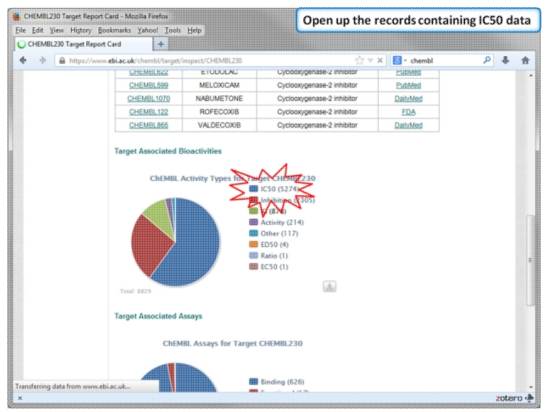
Note: In version 3.8-4 and higher you can search ChEMBL directly from the GUI interface.

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Step 10: Search for COX-2 in ChEMBL.

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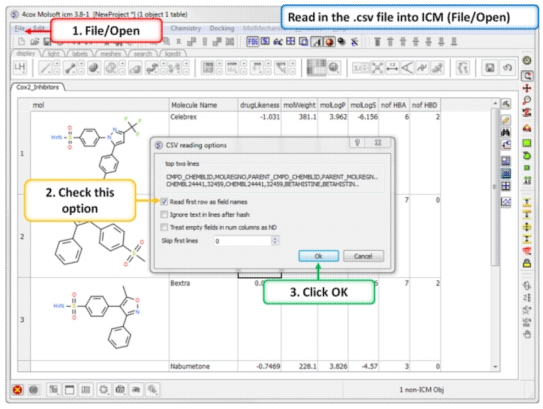
Step 11: Select Human COX-2 entry.



Step 12: Select chemicals annotated with Ki values.

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Step 13: Download in XLS format.



**Step 14: Convert to csv format.** Open the .xls file in Excel and save in comma separated value format (.csv). Read in the .csv file into ICM using File/Open. Rename the table Cox2\_Inhibitors - right click on table header tab and choose rename.

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**Step 15: SMILES to 2D.** Convert the SMILES column into 2D. This may take a little bit of time because the table is quite large. We can use this

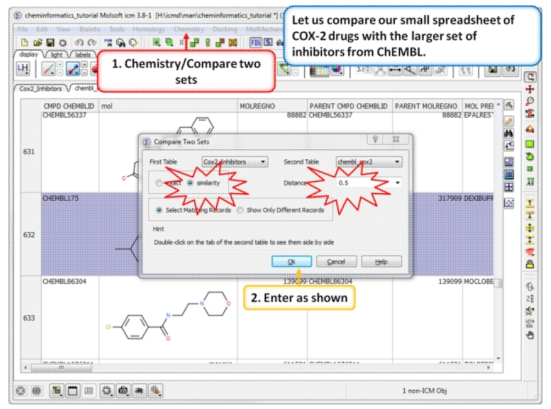
table in the following tutorial.

## 14.4.2 Chemical Searching

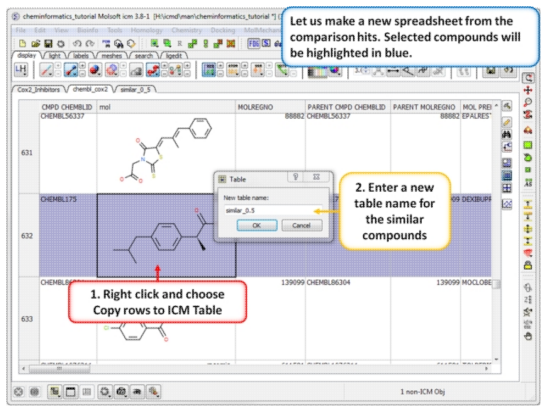
# Background

This tutorial follows on from the previous tutorial where we made a database of COX-2 inhibitors (Cox2\_Inhibitors) and read in a chemical spreadsheet containing IC50 data from ChEMBL (chembl\_cox2). In this tutorial we will perform chemical spreadsheet comparison and explore some of the chemical search options inside ICM.

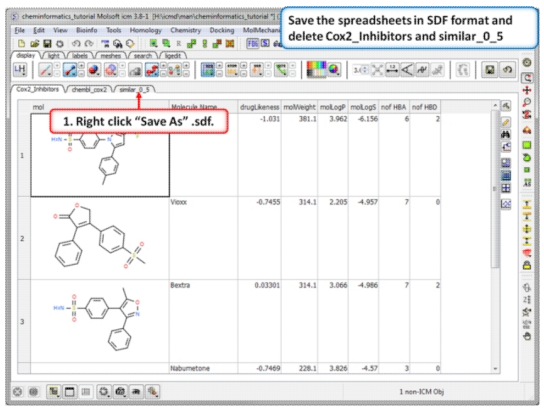
# Tutorial



**Step 1: Similarity search - compare two spreadsheets.** Here we can compare our two chemical spreadsheets using Chemistry/Compare two sets using a similarity search.



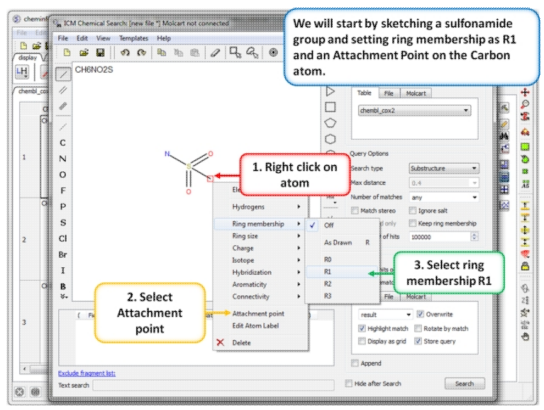
**Step 2: Output the similar chemicals to a new spreadsheet.** Similar compounds are selected (shaded blue) we can extract them to a new table using the right click menu.



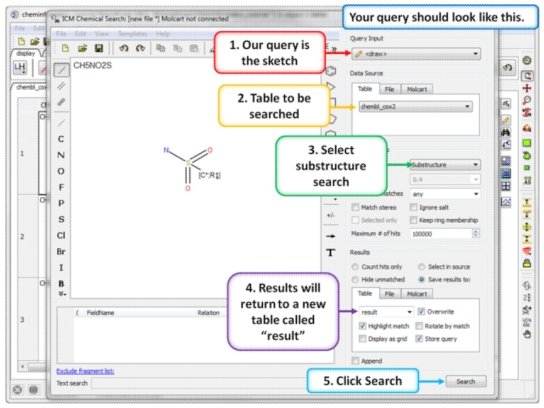
**Step 3:Save as SDF.** You can save any chemical spreadsheet by right clicking on the table header tab.

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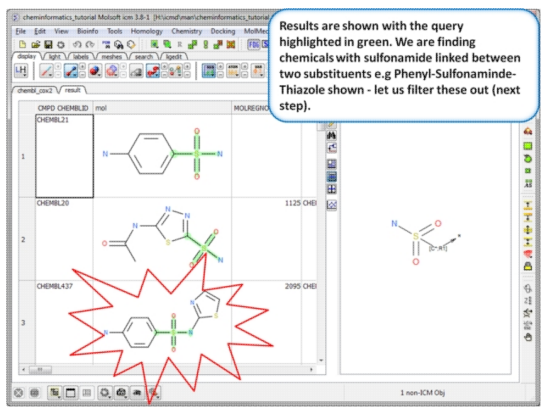
**Step 4:Chemical Searching.** Now let us explore the chemical search options.



**Step 5: Sketch search query.** We will start by sketching a sulfonamide group and setting ring membership as R1 and an Attachment Point on the Carbon atom.



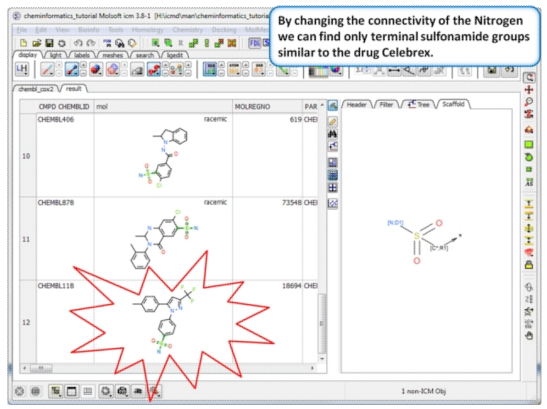
**Step 6: Chemical search window.** Choose the options as shown in the screenshot and press search.



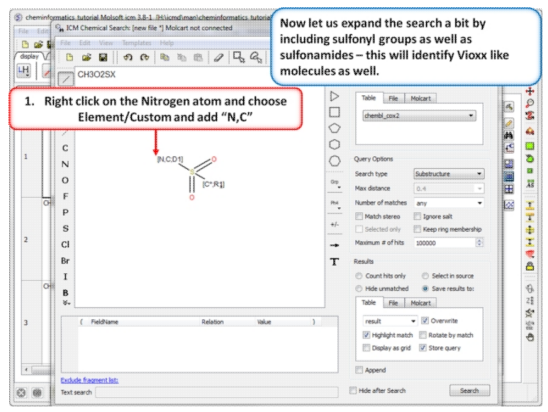
**Step 7: Search query results.** The results will be displayed in a table as shown.

chemint chemint Chemical Search: [new file "] Molcart not connected File Edit     View Template: Help     Cassiary     Cassiary     CH5NO2S	By changing the connectivity of the Nitrogen we can find only terminal sulfonamide groups similar to the drug Celebrex.
CH5NO2S CH5NO2S Chemb.co 1. Right click on the Nitrogen atom and choose Connectivity "As Drawn" C C Br I C Br I C Br I C C C C C C C C C C C C C	Data Source     Data Source     Table File Molcart     dembl_cox2     Cost File Molcart     Query Options     Search type Substructure     Max distance     Act      Act     Ac
3 ( FieldName Relation Value	) result    ⑦ Overwrite 5
Exclude theorem tist: Text search	Hide after Search Search

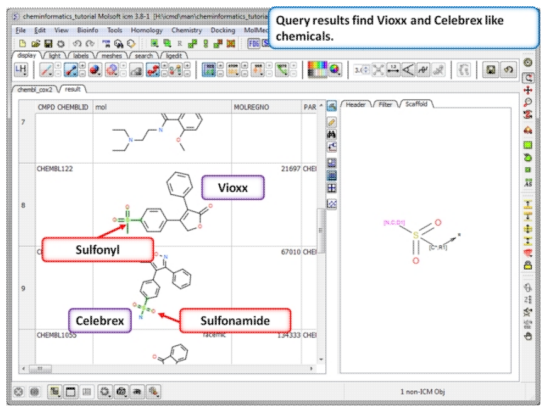
**Step 8: Query filter - connectivity.** By changing the connectivity of the Nitrogen we can find only terminal sulfonamide groups similar to the drug Celebrex.



**Step 9:Query filter results** By changing the connectivity of the Nitrogen we can find only terminal sulfonamide groups similar to the drug Celebrex.



**Step 10:Query filter - Custom Element.** Now let us expand the search a bit by including sulfonyl groups as well as sulfonamides - this will identify Vioxx like molecules as well.



**Step 11: Query filter results.** By adding to the filters we can identify Vioxx-like chemicals containing sulfonyls and Celebrex-like chemicals containing sulfonamide groups.

### 14.4.3 Chemical Clustering

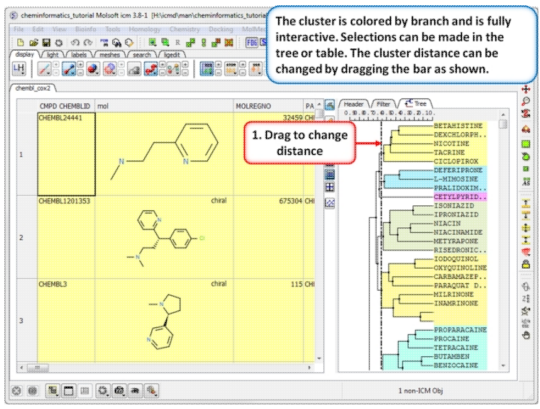
# Background

This tutorial follows on from the previous tutorial where we made a database of COX-2 inhibitors (Cox2\_Inhibitors) and read in a chemical spreadsheet containing IC50 data from ChEMBL (chembl\_cox2). In this tutorial we will perform chemical clustering on the ChEMBL chemical spreadsheet.

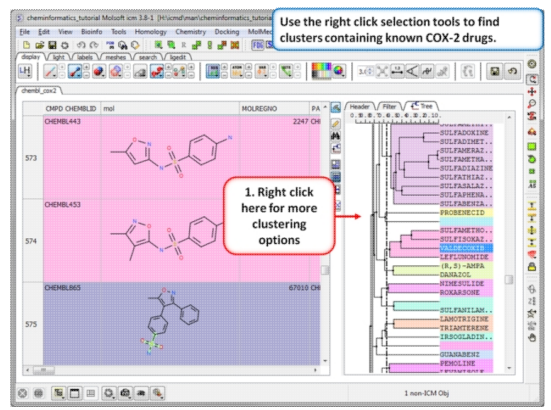
# Tutorial

Ele Edit View Biointo ២៤មែលឲ្យលុ	Tanàn kéompinan Chamistra Darkina kénikén	Cluster the ChEMBL_Cox2 table by chemical substructure.
H Contraction Chemble Cox2	Clustering Parameters	
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-	Add column with duster numbers, name: d     Build for the selected 72 rows     Descriptors:     the "mol" column and all numerical columns	
CHEMBL1196	the selected columns only     the "mal" column only (linear fingerprints)	Use existing distance matrix:
CHEMBL1073	Parameters:	PGMA 139286 GLIPIZIDE
3	-{}.	Cancel Help
	option C. @	1 non-ICM Obj

**Step 1: Cluster** Cluster the ChEMBL\_Cox2 table by chemical substructure using the options shown in the screenshot.



**Step 2: Change the distance of the tree.** The cluster is colored by branch and is fully interactive. Selections can be made in the tree or table. The cluster distance can be changed by dragging the bar as shown.

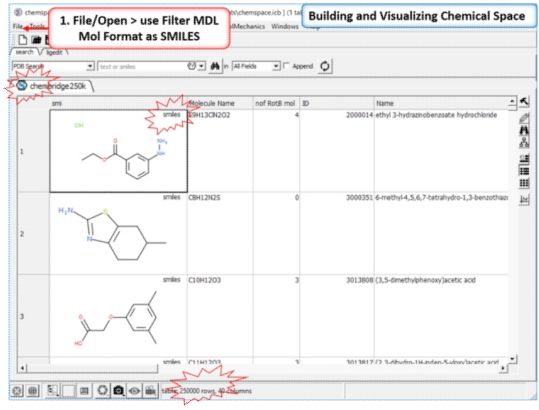


**Step 3: Cluster options.** Use the right click selection tools to find clusters containing known COX-2 drugs. For example, right click and

choose select centers, explore other options in the right click menu.

### 14.4.4 Working with Large Chemical Space

#### 14.4.4.1 Build and Visualize Chemical Space



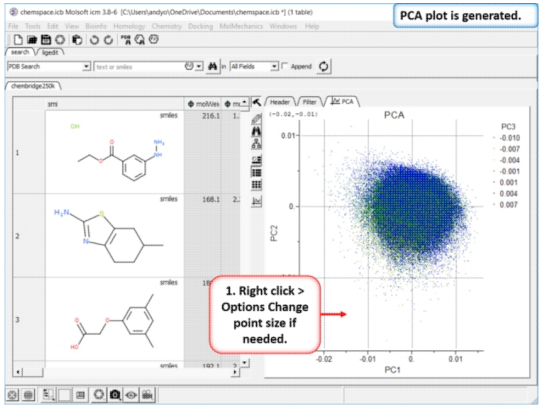
**Step 1: Read in a large SDF file.** In this example we are reading in an SDF file containing >250K chemicals. This is too large for the memory of most desktop computers and so we read it in using the File/Open filter MDL Mol **Format as SMILES**.

rch V ligedit Search Chembridge25	1. Chemistry/Calculate Prop	perties		
smi	Molecule Nar	ne nof Rotß mol ID	Name	*
o	Calculate Chemical Properties	K Actione •	? ×	< chloride
H <sub>2</sub> N	Function • Multiple Functions • Arguments 2. Select properties.	Function           Punction           MolFormula(sm)           Inparliame(sm)           MolcopP(sm)           MolcopS(sm)           P MolcopS(sm)	Name moliVeight moliFormula kuparName mol.ogP mol.ogS molPSA	benzothiazi
	Insert (* atter ( before ( mipsile C at end column (smi) •	MolArea(sm) MolAume(sm) MolAume(sm) MolAf(sm) Drug(Serress(sm) Smiles(sm,mode="unique") BadGroups(sm) Nof Atoms(sm,atom="")	molikrea molitikrme molitikrf drugi, keness smiles badüroups nof, Atoms	
	New column name	Nof_Nolecules(smi)     Nof_Fragments(smi, fragment***)     Nof_Fragments(smi, fragment***)     Nof_Chrad(smi, type="any")     Max_Ring_Size(smi)     Min_Ring_Size(smi)     Max_Fused_Rings(smi)     4	nof_Molecules nof_Fragments nof_Orivals max_Ring_Size max_Flugg_Size max_Flugg_Rings	808

Step 2: Add some chemical properties to the chemical spreadsheet.

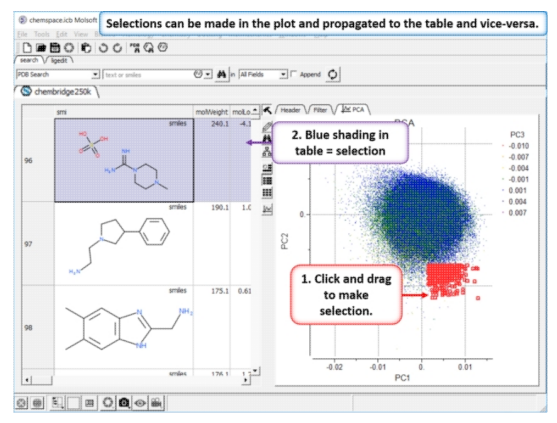
	emistry/PCA	dyo\OneDrive\Documents\chemspace.icb *] (1 table) hemistry Docking MolMechanics Windows Help	Genera	te Princip	oal Compone	nts
search V lope		1. Select Columns	:   <b>9</b> 9	<u>।</u>		×
5	mi	Aliom	🗢 molVolu Mo	lecule Name	nof RotB mol ID .	• <
249998		smiles         425.1         4.267         4.132         72.35         430.3          O         EE         PCA Parameters         Table:         Chembridge250k         Column Prefix:         PC           PC Number Limit:         [3         3         Column Prefix:         PC         Column Prefix:         Colu	405.7 C19 ? ×	#120CINO65	9	
249999	3. Use selected columns only.	Smile: Explain Data Variance (%): 99 Descriptors: C the "mol" column and all numerical columns C" the s C the selected columns only C the "mol" column only (fingerprints) C the mol column only (2D pharmacophore)		410CI2FNO	4	<u>354</u>
250000	4. Build scatter plot	smile:     Plot principal components:       C     Do not plot       Build scatter plot for the first 2 components       Build colored scatter plot for the first 3 components       DK     Cancel	Mejp	418CNO2	6	-
4		Landon and			Þ	•
0015						

**Step 3: PCA** Select 6 of the new properties you added to the table. Generate Principal Components using the options in Chemistry/PCA Analysis.



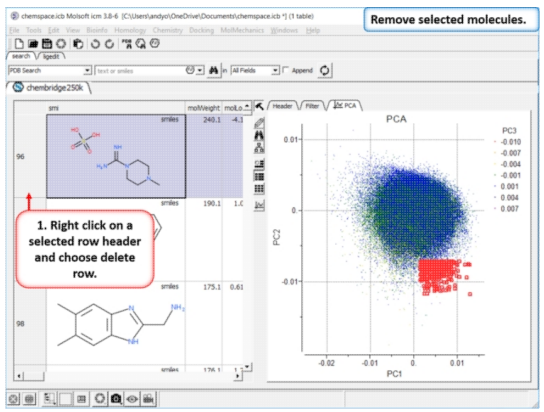
**Step 4:PCA Analysis Plot is Generated.** Right click on the plot and choose Options to change parameters such as point size.

#### 14.4.4.2 Select Molecules in Chemical Space



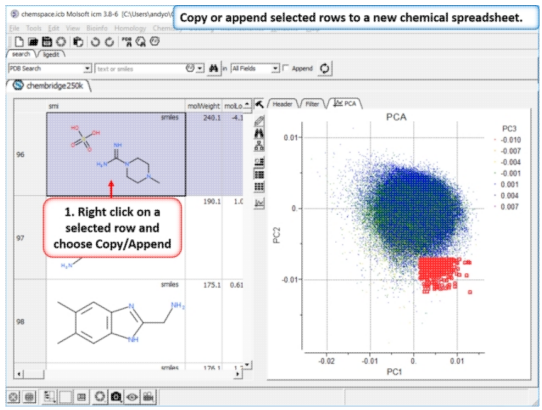
**Step 5: Selections.** Selections can be made in the plot and propagated to the table and vice-versa.

#### 14.4.4.3 Delete Row



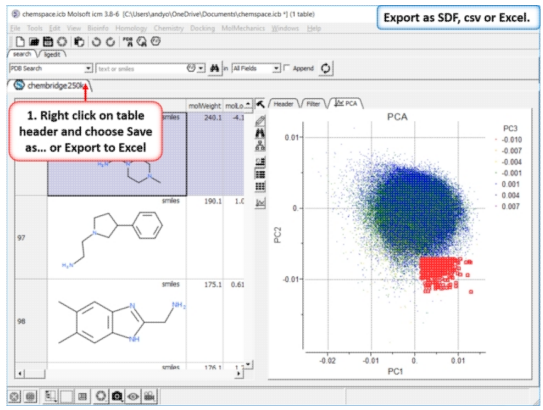
Step 6: Delete Rows

#### 14.4.4.4 Append to New Workspace (Spreadsheet)

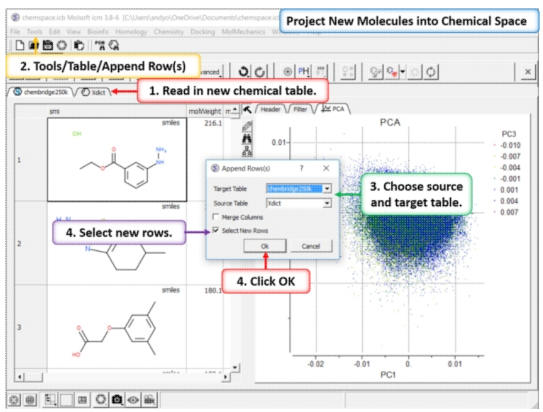


Step 7: Copy or append selected rows to a new chemical spreadsheet.

#### 14.4.4.5 Export Chemical Space to SDF File Format

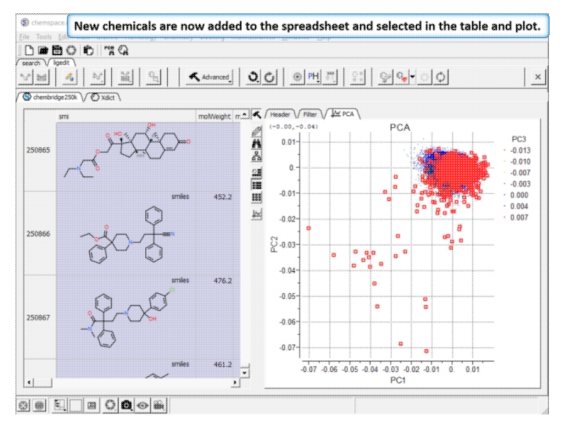


Step 8: Export as SDF, csv or Excel.



#### 14.4.4.6 Project New Molecules in the Chemical Space

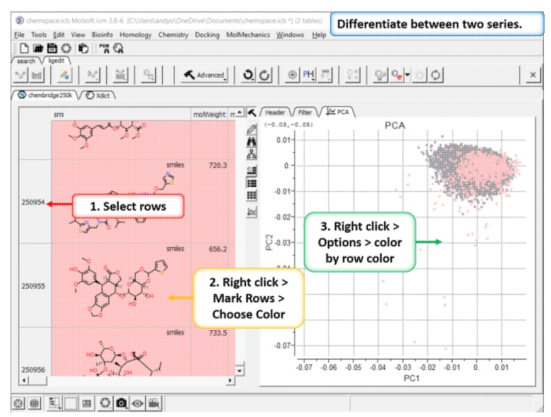
**Step 9:Project New Molecules into Chemical Space** Use Tools/Table/Append Row(s).



14.4.4 Working with Large Chemical Space

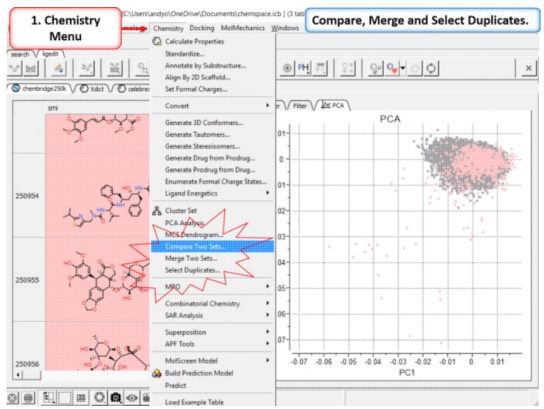
Step 10:New chemicals are now added to the spreadsheet and selected in the table and plot.

# 14.4.4.7 Distinguish between different sets of molecules in Chemical Space



**Step 11:Differentiate between two series.** You can differentiate between two series by coloring rows and then propagating that row color to the plot.

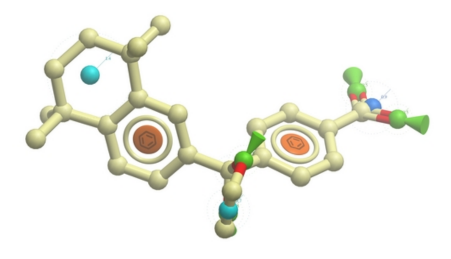
# 14.4.4.8 Compare, Merge and Select Duplicates in Chemical Space



Step 12: Compare, Merge and Select Duplicates.

### 14.4.5 3D Pharmacophore Searching

**Objective** Undertake a 3D pharmacophore search of a table containing 3D coordinates.



• File/Open **example\_ph4.icb** (this file is provided in the ICM distribution and therefore can be found in \$ICMHOME or in Windows

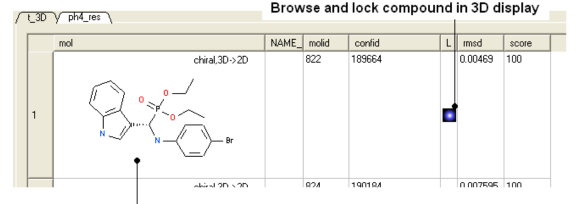
Program Files/MolSoft

- In this example the 3D pharmacophore has already been extracted from a ligand. To find out how to generate a 3D pharmacophore see the section entitled Pharmacophore Draw 3D.
- In this example a table containing 3D coordinates is already provided containing 3D coordinates. The table is called t\_3D.
- To run 3D pharmacophore searching right click on the name of the pharmacophore object in the ICM Workspace and select **Search Pharmacophore**.
- Select the table t\_3D from the drop down list and click OK.
- A table of search results will be displayed.

#### Right click here pharma Clear Default Display... 🕁 💽 рЗ 1 Edit Description... ? × ≶ Search Pharmacophore - o p5 Edit - 💽 p7 table To Search In t\_3D -Search Pharmacophore. 8a 💿 🖩 tables (1 it Advanced Ok ۶ ancel i…t 3D 9994 ۲ Center . ∃-script (1 item -run Connect to Object Enter name of table containing 3D coordinates

# • You can browse the results by clicking on the table and the ligand will be displayed in the graphical display.

• Remember you can use the check boxes in column L to lock compounds and overlay them.



### Double click to view in 3D

#### Manual References (Web Links)

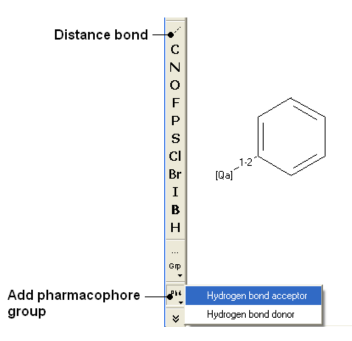
3D Pharmacophore Search

### 14.4.6 2D Pharmacophore Searching

Objective Undertake a 2D pharmacophore search of a chemical spreadsheet.

- File/Open example\_ph4.icb (this file is provided in the ICM distribution and therefore can be found in \$ICMHOME or in Windows Program Files/MolSoft
- Chemistry/Chemical Search
- Draw the query as shown below using the

#### 14.4.6 2D Pharmacophore Searching



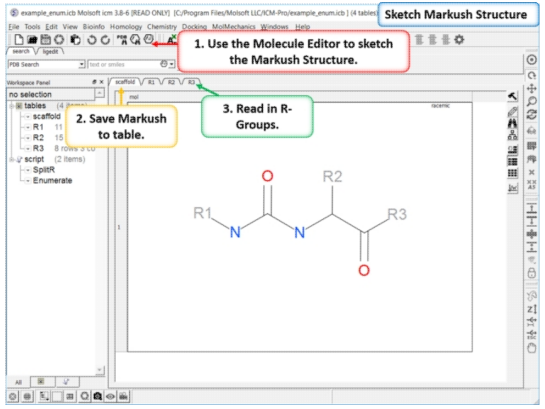
• Fill in the query and results option as shown below.

	Data Source
	📀 Local Tables 🛛 🔗 MolCart
	Query Options         Database         Table         Table         Search Type         Substructure         Maximal # of Hits         1000         Max Distance         0.4         Number of Matches         any         Match stereo         Ignore Salt         By Selection
	-Result
	sult_new 💌 🔽 Auto 🔽 Append
	New Table     C Select in Source
Relation Value	T Add to DB
	🔽 Highlight Match 🔲 Rotate by Match
	🦵 Display as Grid
	Search

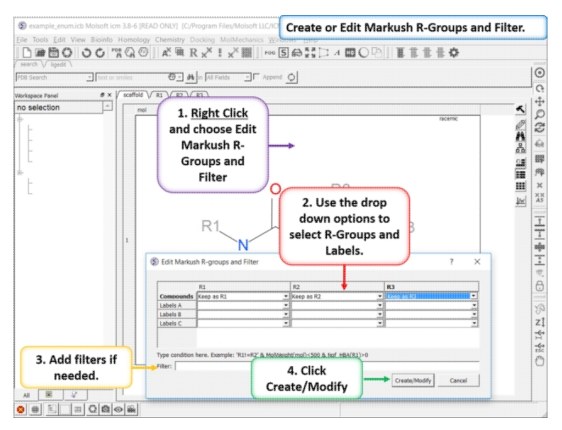
#### Manual References (Web Links)

2D Pharmacophore Search

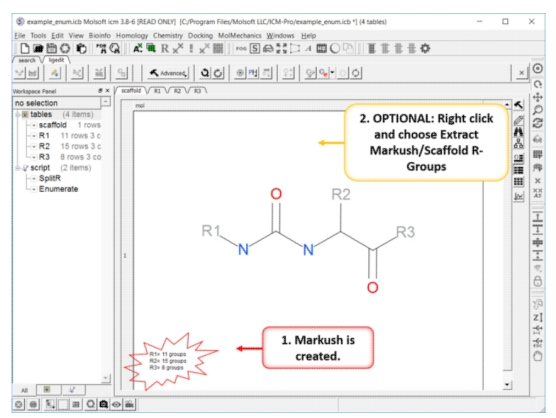
### 14.4.7 How to create a Markush structure.



**Setup** A pre-prepared example is available in File/Open Examples > example\_enum.icb. The markush structure has already been drawn and the R-groups have been read into ICM. The Markush structure is ready to be created.



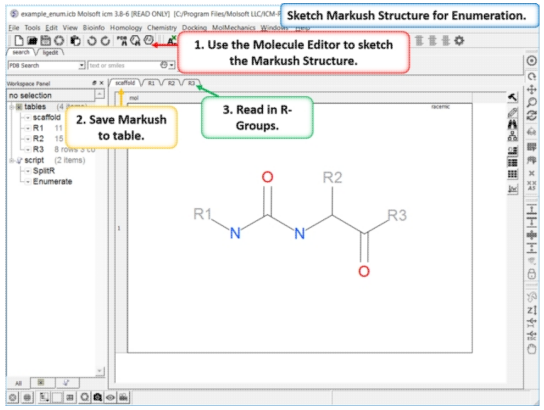
**Create or Edit Markush R-Groups and Filter.** Right Click and choose Edit Markush R-Groups and Filter. Add the R-groups, labels and filters and then click on Create/Modify.



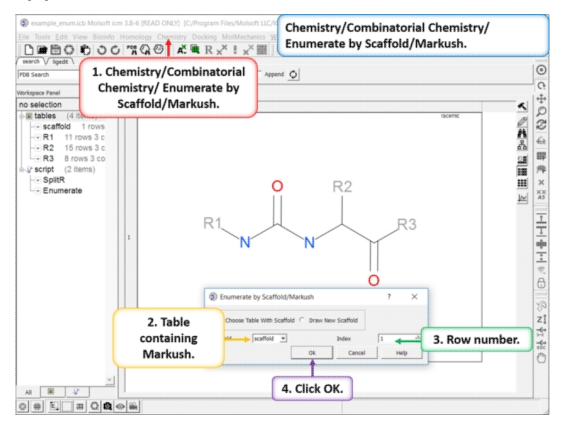
**The Markush Structure is Created.** If you want to see the R-Groups in a separate table you can extract them by right clicking on the Markush and choose Extract Markush/Scaffold R-Groups.

#### 14.4.8 How to enumerate a Markush library.

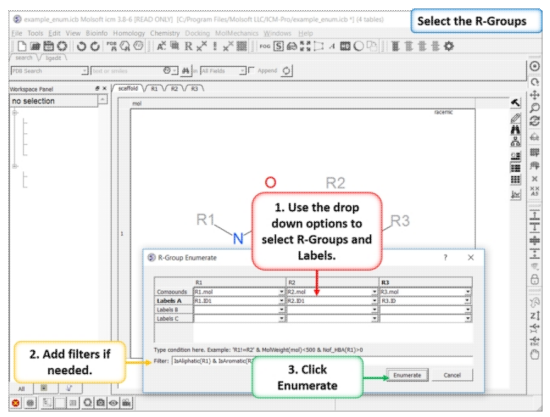
|Documentation|



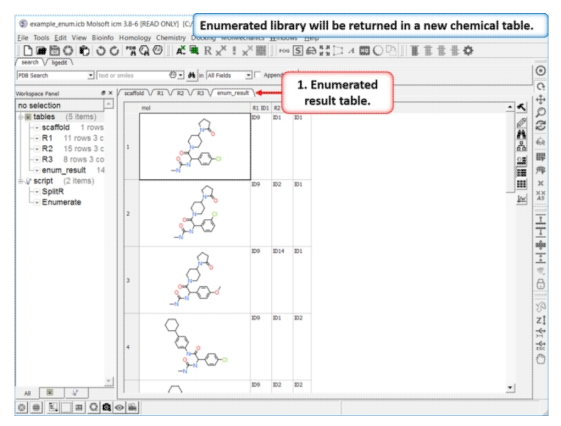
**Enumeration Setup** A pre-prepared example is available in File/Open Examples > example\_enum.icb. The markush structure has already been drawn and the R-groups have been read into ICM. The Markush structure is prepared for enumeration.



#### **Enumerate Library - Step 1:** Choose the menu option Chemistry/Combinatorial Chemistry/ Enumerate by Scaffold/Markush.



Enumerate Library - Step 2: Select R-groups, labels and filters.



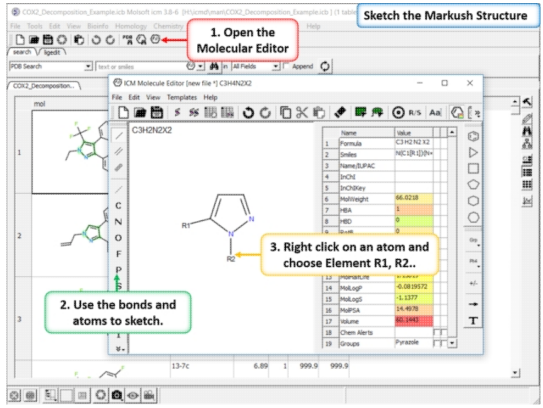
14.4.8 How to enumerate a Markush library.

**Results Table** The enumerated library will be displayed in a new chemical table.

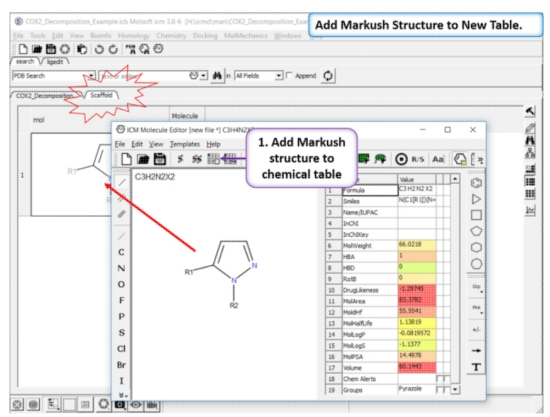
# 14.4.9 How to decompose a library based on a Markush structure.

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mol		Molecule Name	ACTIVITY	SET	Min dist	Avg dist	
1	to	13-10	7.48	2	999.9	999.9	
2	A.	13-6a	4	3	0.22	0.56	
3	HO.	13-7a	7.12	2	0.08	0.55	
2		13-7c	6.89	1	999.9	999.9	-

**Read in the SDF File to be decomposed.** A prepared ICM file containing COX-2 inhibitors with a common scaffold and activity data can be downloaded from here: ftp://ftp.molsoft.com/pub/workshop/ (1. cut and paste the address into a web browser. 2. Right click on COX2\_Decomposition\_Example.icb and choose "Save Link As". 3. Open the .icb in ICM File/Open).



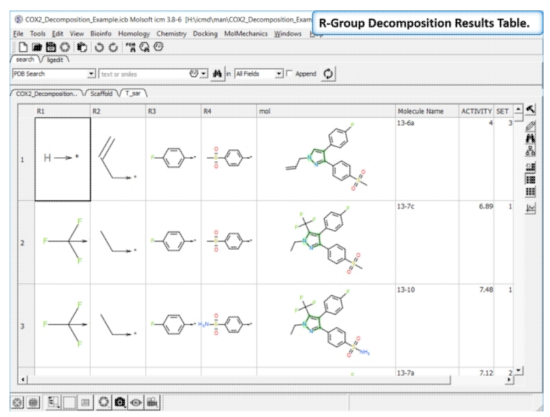
**Sketch the Markush Structure.** Sketch the Markush structure using the Molecule Editor.



**Create Markush Structure Table** Save the Markush structure to a new chemical table.

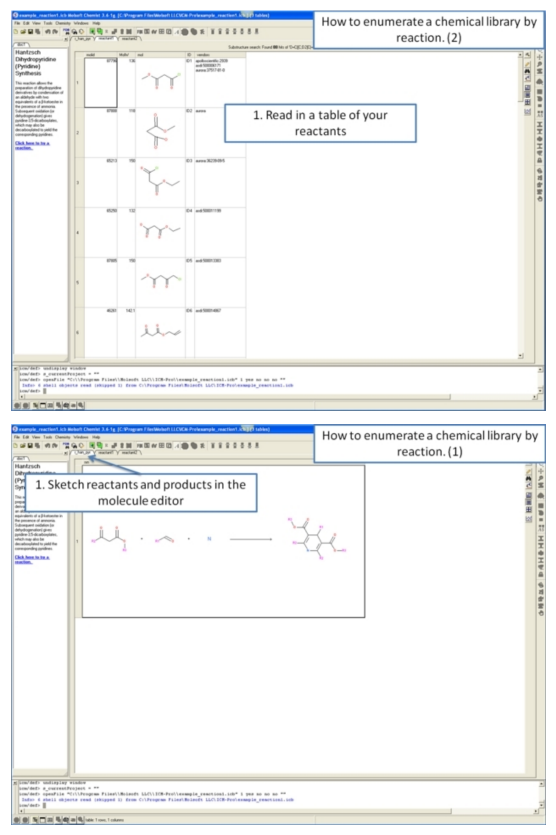
368001		/SAR Analysis composition.	105 ¥11 ADDE	- <u>0</u>		
mol	N R2	Molecule Name C3H2N2X2				
	(F Choose	Decomposition	Draw New Scaffold		?	×
2. Scaffold tabl	Table For Dec	Scaffold amposition and a scale Single SAR Table 3. Table f decompositi	or	Index ure Column mol to Add Missing R-Groups Ck Ck Ca		

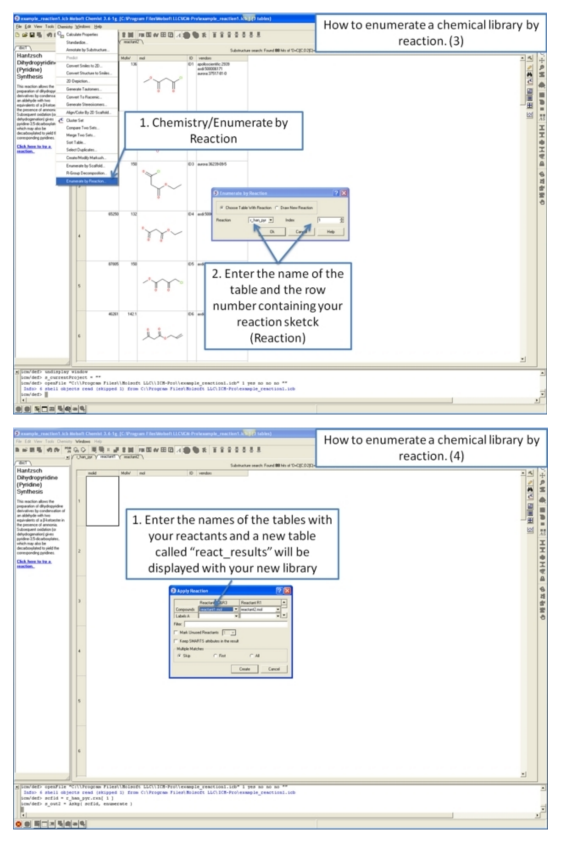
#### Chemistry/SAR Analysis/ R-Group Decomposition.



**Results Table.** The decomposition data will be displayed in a new table. You can sort the columns by Actitiy or R-Group. The R-group decomposition can also be used to build more complex SAR tables as described here.

# 14.4.10 How to enumerate a chemical library by reaction.

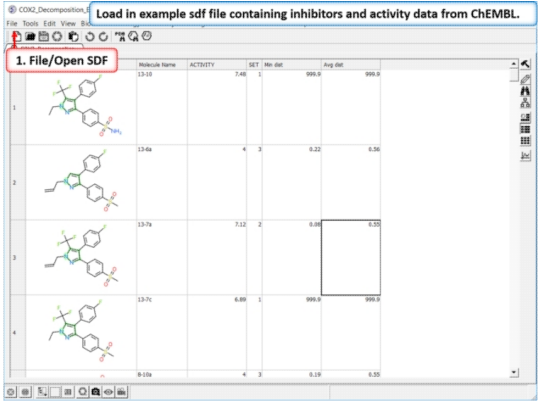




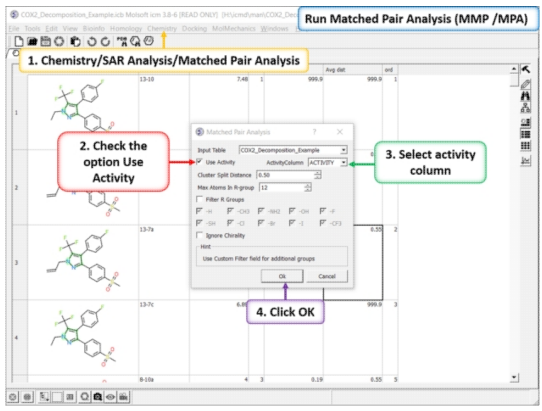
## 14.4.11 Detecting Activity Cliffs

There are two methods to detect activity cliffs Matched Pair and Structure--activity landscape index(SALI).

The Matched Pair Analysis performs Maximum Common Substructure search on various pairs to find one-point variations in R-group. It returns a chemical spreadsheet with the delta difference in Activity and a score which is a combination of delta activity and size of R-group therefore looking for smaller R-groups with bigger activity difference.

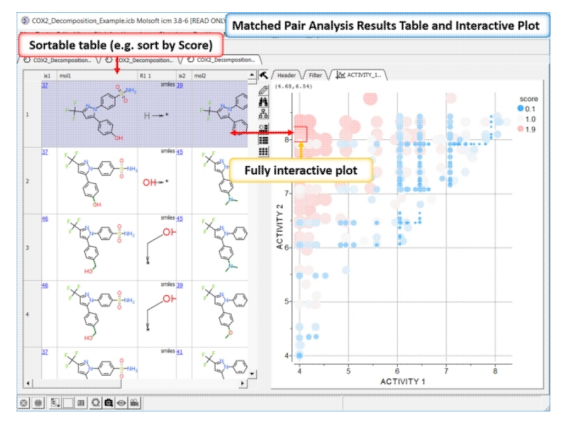


**Read in Activity Data** A prepared ICM file containing COX-2 inhibitors with a common scaffold and activity data can be downloaded from here: ftp://ftp.molsoft.com/pub/workshop/ (1. cut and paste the address into a web browser. 2. Right click on COX2\_Decomposition\_Example.icb and choose "Save Link As". 3. Open the .icb in ICM File/Open).



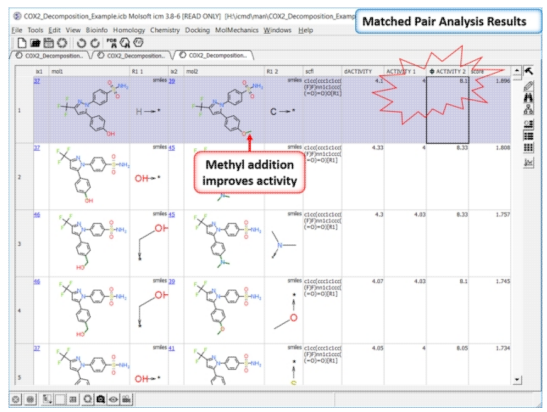
#### **Run Matched Pair Analysis**

- Chemistry/SAR Analysis/Matched Pair Analysis
- Enter the Input table name.
- Check the option "Use Activity".
- Click OK.



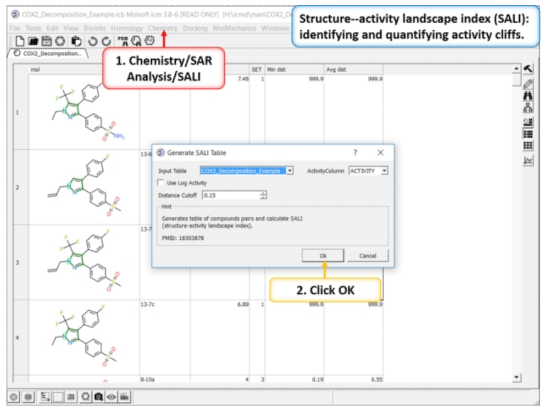
14.4.11 Detecting Activity Cliffs

\*{Matched Pair Analysis Results} A table of the results will be displayed - you can sort by score. The plot is displayed activity 1 versus activity 2 colored and sized by score (red is better). The plot and the table are linked so a selection in one is propagated to the other.

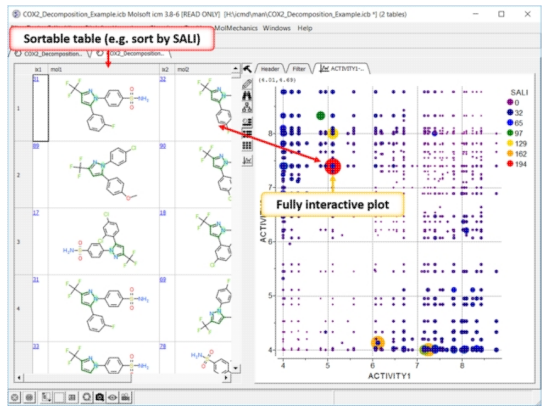


**Results Analysis** The top scoring comparison shows that the addition of a methyl group significantly improves activity. You can also add filtering e.g. R1\_2 filter and select methyl groups - right click on column header and choose filter.

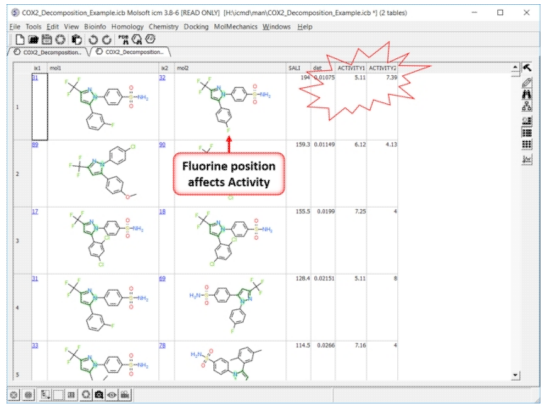
The SALI method developed by John Van Drie is calculated using chemical distances and activity and can identify different changes compared to MPA.



Structure--activity landscape index (SALI): identifying and quantifying activity cliffs. Use the menu option Chemistry/SAR Analysis/SALI.



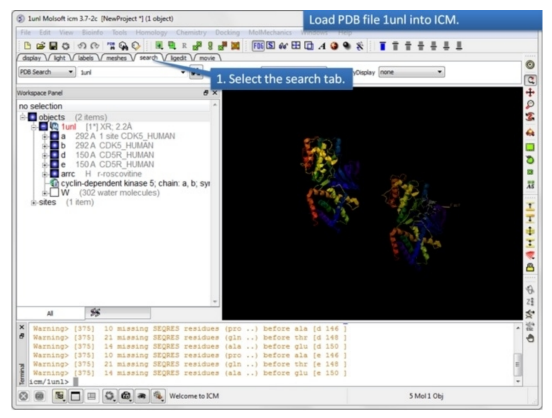
**SALI Results** A fully interactive table and plot is returned containing the SALI index.



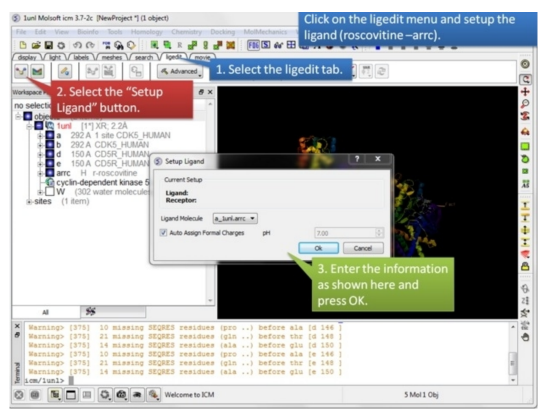
**Results** In this example the fluorine movement around the ring affects activity.

# 14.5 3D Ligand Editor Tutorial

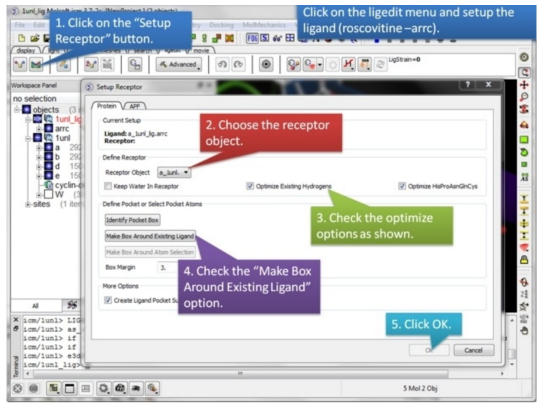
In this example we will use the 3D interactive editor to modify a cyclin-dependent kinase 5 inhibitor (PDB Code 1UNL). A hydrophobic cavity not occupied by the crystal structure ligand (roscovitine) can be exploited to improve the interaction between the ligand and the receptor. Roscovitine is an experimental drug candidate in the family of cyclin-dependent kinase (CDK) which preferentially inhibits multiple enzyme targets including CDK2, CDK7 and CDK9. It alters the growth phase or state within the cell cycle of treated cells and is in clinical trials for treatment of non-small cell lung cancer (NSCLC), Cushing's Disease, leukemia, HIV infection, herpes simplex infection, cystic fibrosis



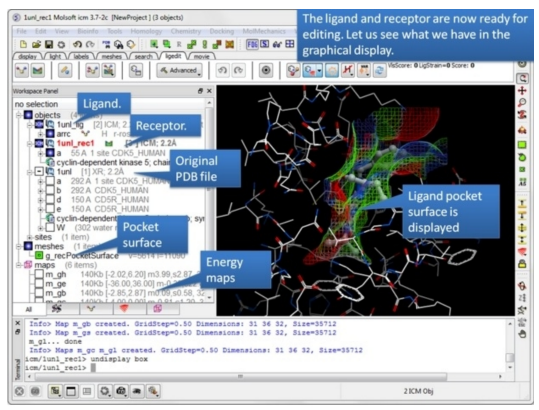
Step 1: Read in the PDB file 1UNL.



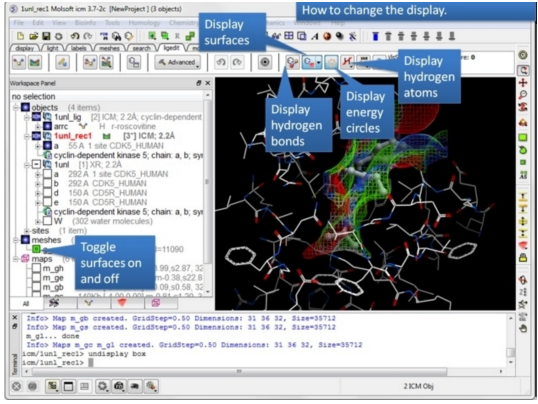
Step 2: Setup the ligand in the 3D ligand editor.



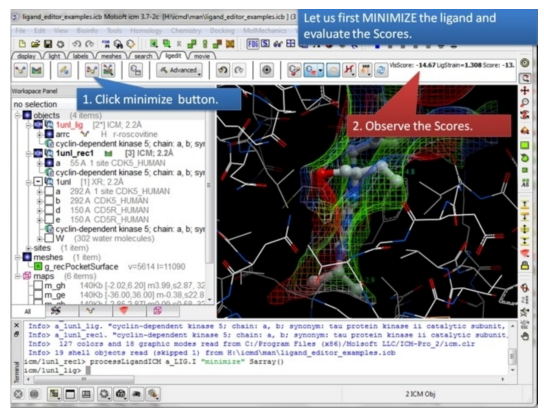
Step 3: Setup the receptor in the 3D ligand editor.



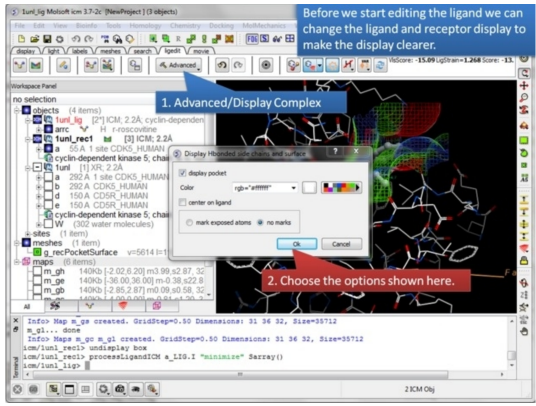
**Step 4:** Observe the new objects, meshes and maps that have been created.



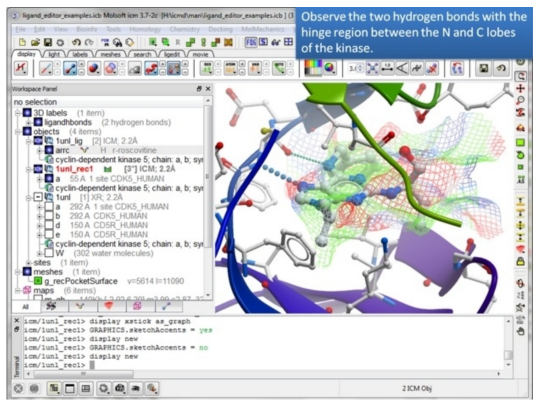
**Step 5:** You can display and undisplay, hydrogen atoms, surfaces, and hydrogen bonds.



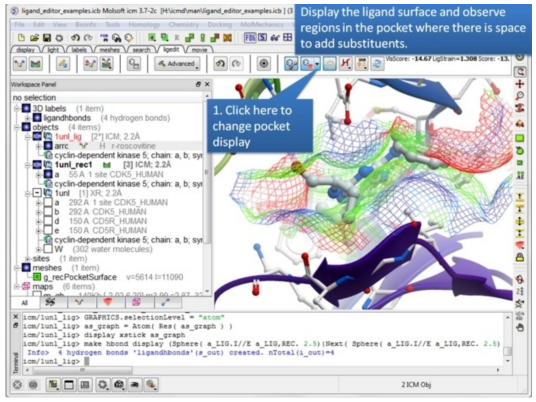
Step 6: First minimize the ligand and observe the Scores.



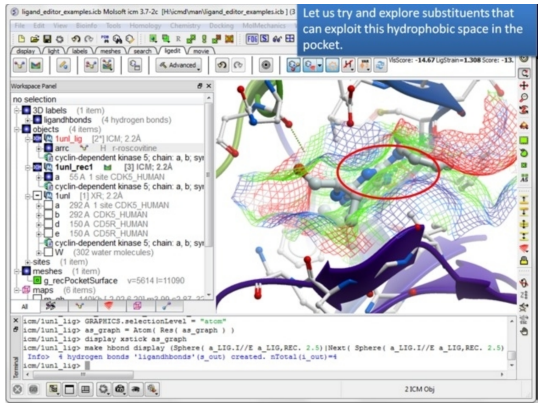
**Step 7:** In the Advanced menu there is an option to make a clearer view of the complex.



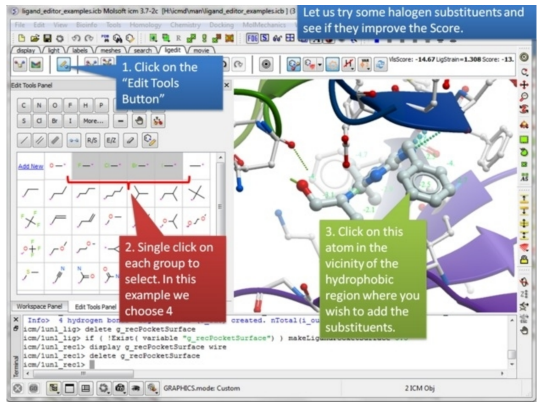
**Step 8:** Observe that the ligand makes key hydrogen bond interactions with the hinge region of the kinase. It is important to conserve these interactions when making modifications.



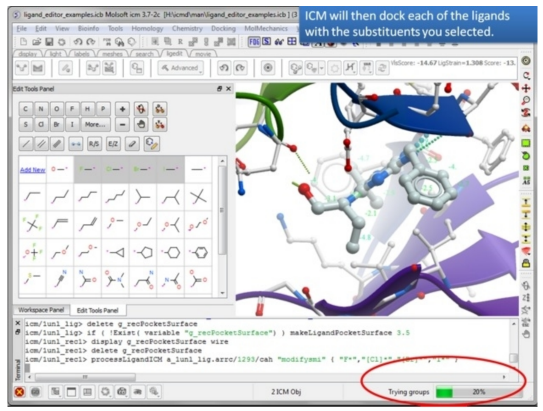
**Step 9:** The ligand surface representation is a useful way of identifying space in the pocket where you can add groups to the ligand. The surface is colored: White=aromatic lipophilic, Green=non-aromatic other (mostly aliphatic) lipophilic surface, Red=hydrogen bonding acceptor potential, Blue=hydrogen bond donor potential. The coloring of the surface helps you determine which modifications to make, for example if the surface is colored red you ideally would add a hydrogen bond acceptor in that region.



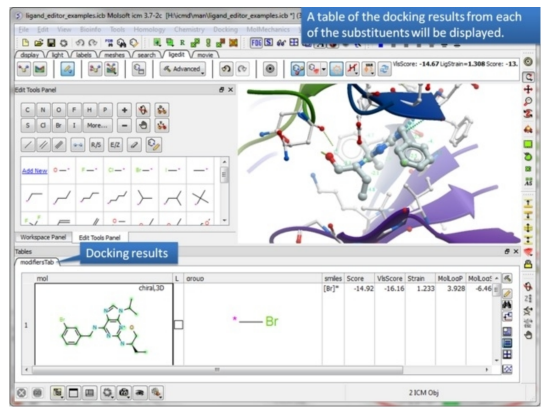
**Step 10:** There seems to be a hydrophobic region in this pocket that we might try to explore.



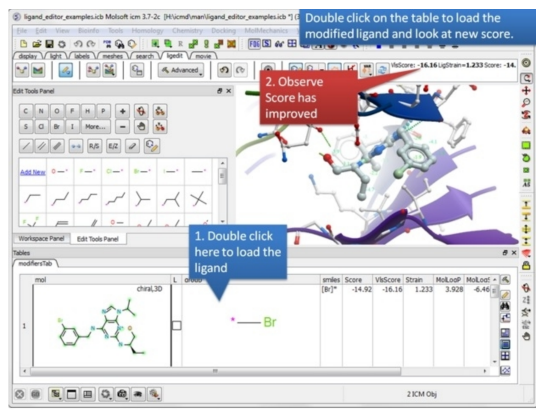
**Step 11:** You can try any modification group you wish from the "edit tools panel". We will try some halogen substituents and see if they improve the score.



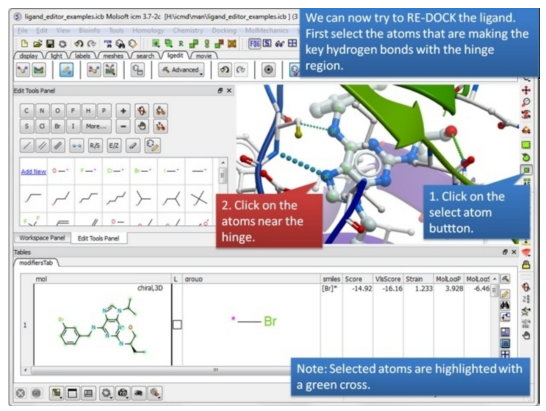
**Step 12:** The editor will add each substituent and evaluate the score. This may take some time depending on the number of groups you chose but you can monitor the progress in the bottom right hand corner.



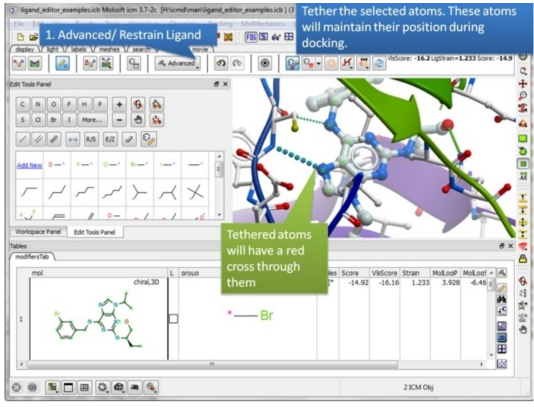
**Step 13:** A table of ranked hits will be displayed. In this case the bromine substituent scored best.



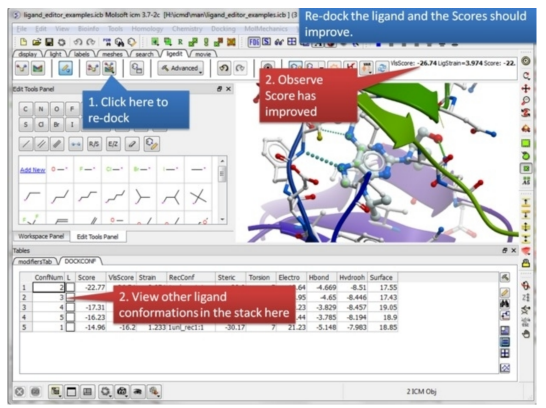
**Step 14:** Double click on the table row to display the complex and observe the improved score.



**Step 15:** We can now re-dock this ligand to make sure the interactions are optimal. We must remember to tether the atoms making a hydrogen bond with the hinge and so first we must select these atoms.



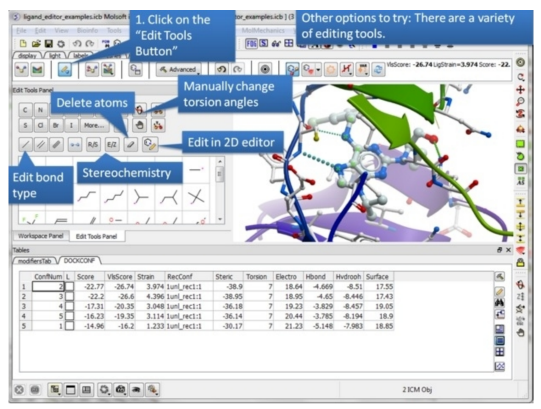
Step 16: Tether the atoms.



**Step 17:** Click on the re-dock button and a stack of the best conformations will be displayed in a table called DOCKCONF. Observe the score has improved significantly for this modified ligand.

isplay		Bioinfo	Tools H	search V	Chemistry Chemistry R g <sup>2</sup> 9 Igedit <u>movie</u> Advanced	Docking I	FD6 S 0	nics 1) Ar EB	modif	icatio	n. You on to a	o and redo any ligand can also save any table. core: -26.74 LigStrain=3.974 Score: -22.
t Tools F	N O Br	Add ci confo to a ta	rmati			Undo/ I	Redo		-			
Add Net	• ••	•• - مر	• •	· ,	 - ×	•		Å		2		
oles	stab V DO	Edit Tools P	o —	~ / _	. /	•	*					ð ×
	onfNum L	Score	VIsScore	Chroin	RecConf	Steric	Torsion	Electro	Hbond	Hvdroph	Suchas	4
	2	-22.77	-26.74		1unl_rec1:1	-38.9	7		-4.669	-8.51	17.55	
	-	-22.2	-26.6		1unl_rec1:1	-38.95	7		-4.65		17.43	
1	3	-17.31	-20.35		1unl_rec1:1	-36.18	7				19.05	24
	3				1unl_rec1:1	-36.14	7		-3.785		18.9	
1 Co		-16.23				-30.17	7				18.85	
Co 1 2 3	4	-16.23 -14.96	-16.2	1.233	1unl_rec1:1	-30.17	/	61.23				
Co 1 2 3 4	4		-16.2	1.233	lun_rec1:1	-30.17	,	21,23				

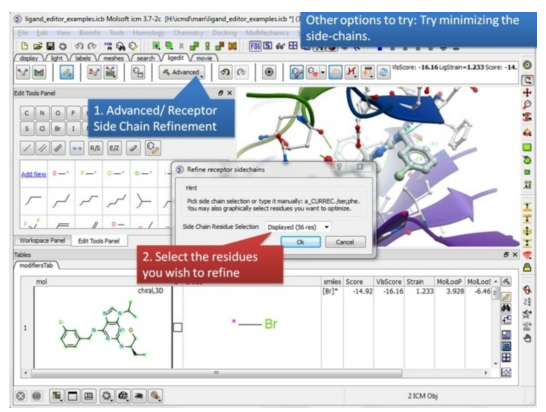
**Step 18:** Never worry about mofifying a ligand because everything is saved. Try anything you want because you can always undo and redo. There is also a convenient way of storing a modification you like in a a table.



Step 19: Explore the "Edit Tools Panel".

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Step 20: Perform a virtual screen of a substituent database.



**Step 21:** You can also minimize side-chains inside the ligand editor. This may remove unwanted clashes between your receptor and ligand.

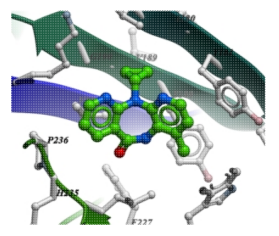
## **14.6 Atomic Property Field Tutorials**

## **APF Background**

Atomic Property Fields (APF) is a 3D pharmacophoric potential implemented on a grid. APF can be generated from one or multiple ligands and seven properties are assigned from empiric physico-chemical components (hydrogen bond donors, acceptors, Sp2 hybridization, lipophilicity, size, electropositive/negative and charge).Here we describe template APF superposition whereby the APF is generated from a single or multiple template and is then globally optimized with the internal force-field energy of the ligand. You can read more about the APF method here.

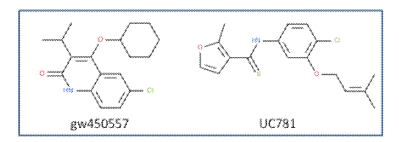
**About this Example** In this example we will use Atomic Property Fields to superimpose, screen and cluster a set of Non-nucleoside reverse-transcriptase inhibitors (NNRTIs) (antiretroviral drugs) used in the treatment of human immunodeficiency virus (HIV). We will use a non-trivial superposition example of two NNRTIs which have similar pharmacophoric properties but are significantly different substructure. We will then read in additional PDBs which contain first generation NNRTIs approved by the FDA (Nevirapine, Delavirdine, Efavirenz) we will then use the combined fields to screen an sdf library to the APF fields

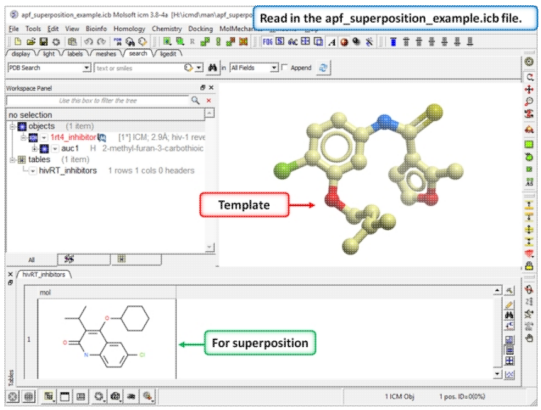
NNRTIs are chemically diverse but bind in the same allosteric pocket in the palm domain of the p66 subunit and part of the p51 subunit. Many NNRTIs bind in a "butterfly" conformation (see image below of Nevirapine) making pi-pi interaction with the rings in the pocket.



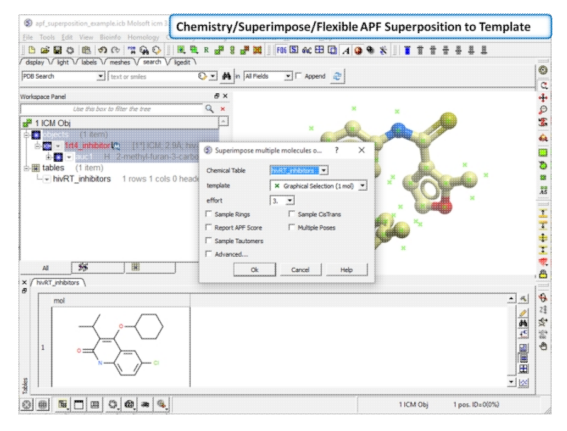
## 14.6.1 APF Superposition

This is an example of a non-trivial chemical superpostion, we will superimpose inhibitor UC781 to gw450557. A prepared ICM file containing the two inhibitors can be downloaded here ftp://ftp.molsoft.com/pub/workshop/ (1. cut and paste the address into a web browser. 2. Right click on apf\_superposition\_example.icb and choose "Save Link As". 3. Open the .icb in ICM File/Open).

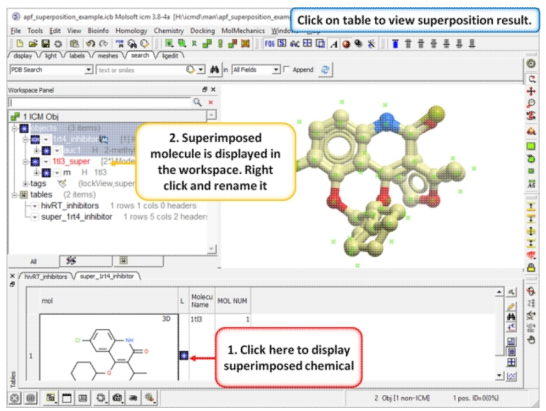




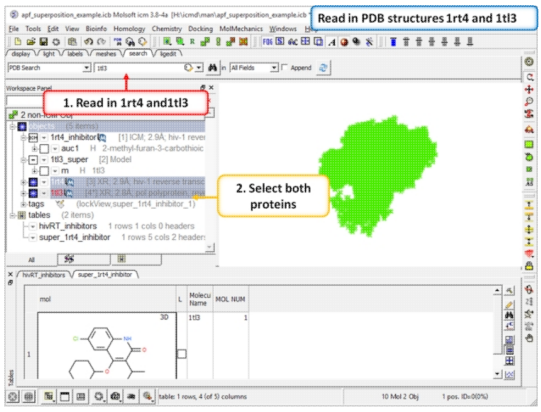
**Step 1:** Download and open the prepared ICM file. The file can be downloaded here ftp://ftp.molsoft.com/pub/workshop/(1.cut and paste the address into a web browser. 2. Right click on apf\_superposition\_example.icb and choose "Save Link As". 3. Open the .icb in ICM File/Open).



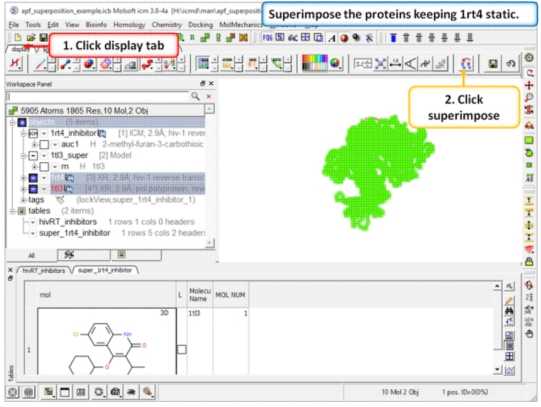
**Step 2:** Superimpose the chemical from the table onto the 3D structure using Chemistry/Superimpose/Flexible APF Superposition to Template. Enter the parameters as shown in the screenshot.



**Step 3:** When the superposition has finished a results table will be displayed. You can toggle the superimposed molecule display on or off in the tbale.



**Step 4:** Now lets compare the docked pose of UC781 with the crystal structure pose. First we need to read in the PDB structures containing UC781 (1tl3) and gw450557 (1rt4).



**Step 5:** Superimpose the crystal structures keeping 1rt4 static. You need to select the two pdbs and use the superimpose button in the

## display tab.

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**Step 6:** Compare the APF superimposed structure with the crystal structure. The RMSD should be