

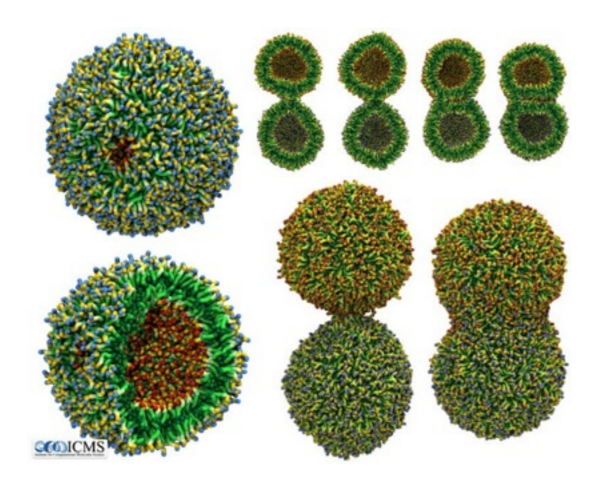
ictp/vmd-viz/

#### About VMD

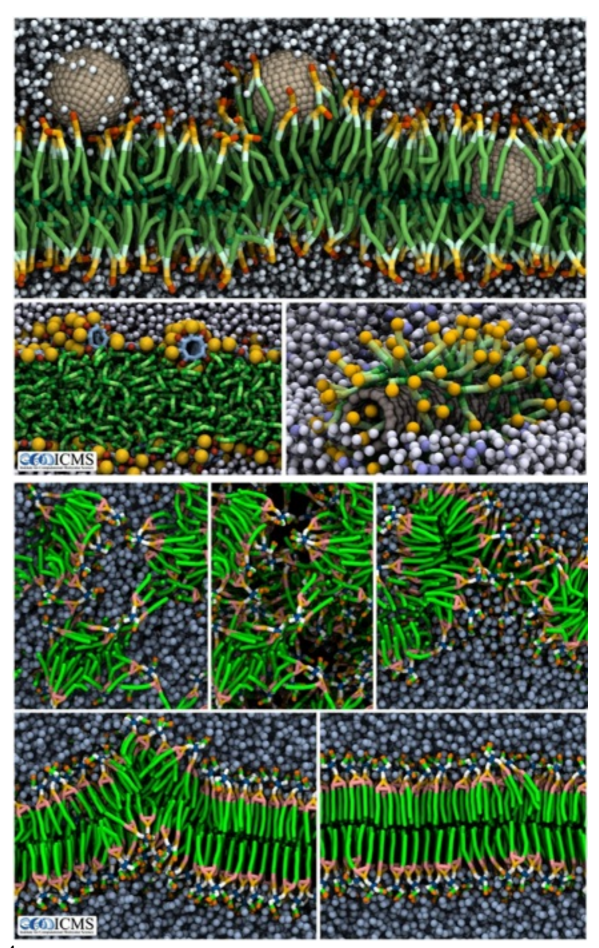
- Powerful tool for molecular visualization, modeling and analysis.
- Theoretical and Computational Biophysics Group at the University of Illinois Urbana-Champagne.
- John Stone, Senior Developer
- http://www.ks.uiuc.edu/Research/vmd/
- VMD Mailing List

### VMD Features

- Support for all major computer platforms
- Support for multicore processors
- Support for GPU accelerated computation
- No limits on the number of molecules, atoms, residues or number of trajectory frames, except available memory
- Many molecular rendering and coloring methods
- Extensive atom selection syntax for choosing subsets of atoms for display (includes boolean operators, regular expressions, and more)
- Support for over 60 molecular file formats and data types through an extensive library of built-in file reader/writer plugins and translators

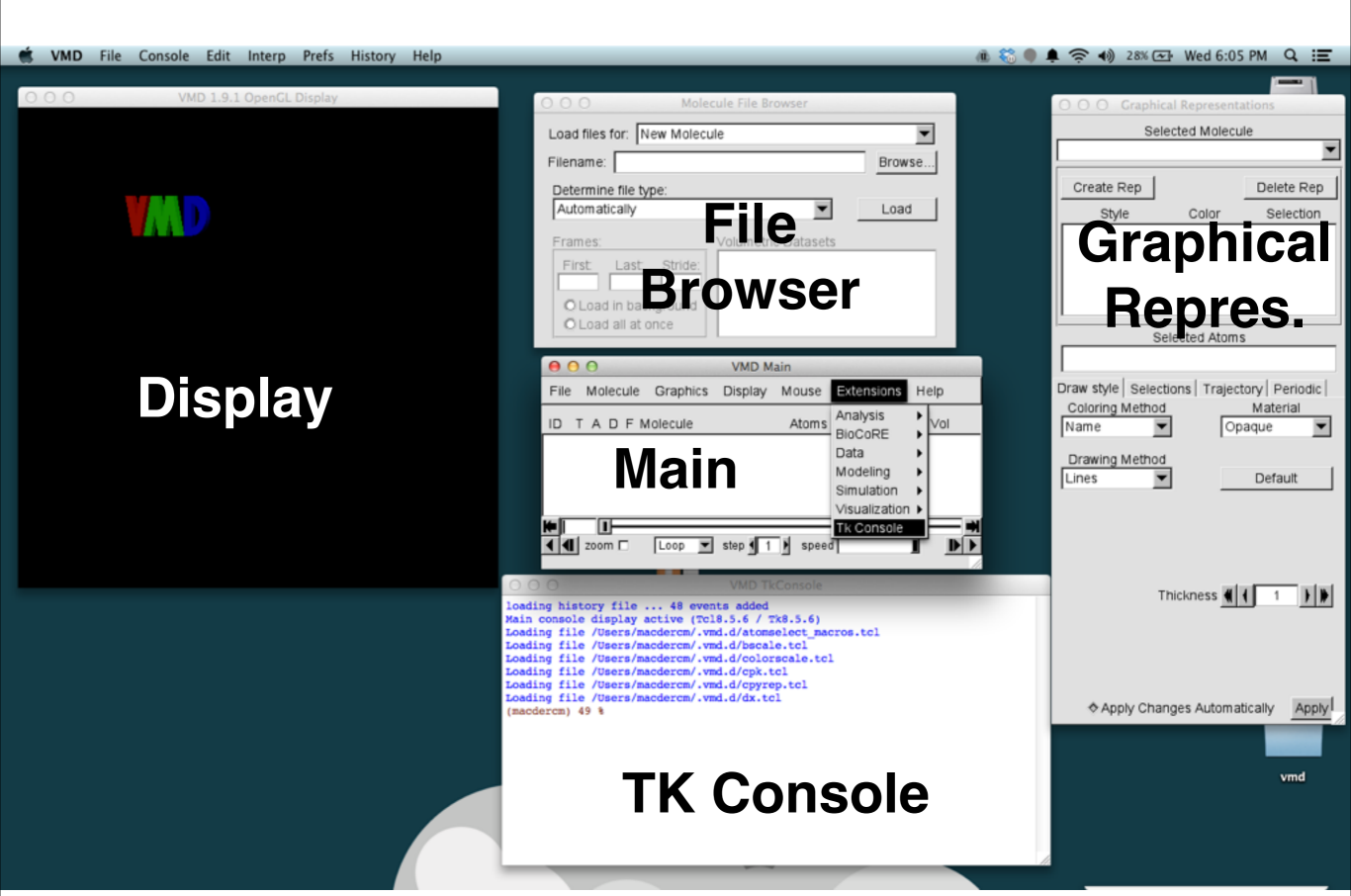


Some nice images/movies courtesy of Axel



# Installing VMD

- On windows/OSX: Install like anything else.
- Linux: (install to root-fs)
  - 1. Unpack the archive, change to created directory
  - 2.> ./configure; cd src; make install
- 2.or Linux: (install to \${HOME})
  - 1. Unpack the archive, change to created directory
  - 2. Edit the configure file: nano ./configure
    \$install\_bin\_dir="\\${HOME}/pkg/bin";
    \$install\_library\_dir="\\${HOME}/pkg/lib/\$install\_name";
  - 3.> mkdir -p \${HOME}/pkg/lib \${HOME}/pkg/bin; cd src; make install; export PATH=\${HOME}/pkg/bin:\${HOME}



# 10 minute introduction to VMD with Myoglobin

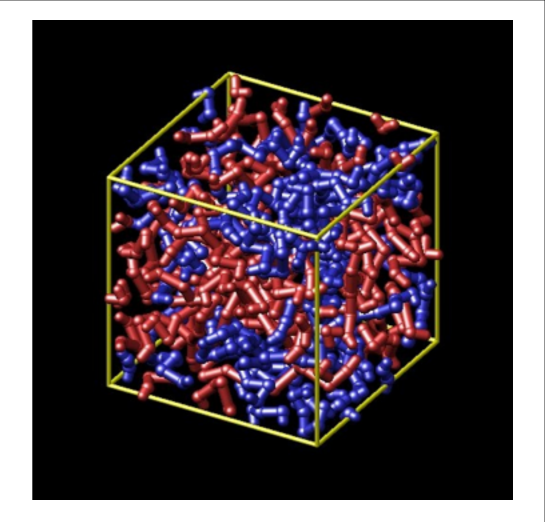
You should become familiar with:

- VMD's Main window
- Loading molecules
- Changing Representations
  - Coloring Methods, Drawing methods, Materials
- Highlighting particular groups of atoms
   myoglobin

# Image/Move Creation with LAMMPS

#### · <u>dump image</u>

write an image every 200 steps



```
dump_modify d1 pad 5 element N O
```

Enlarge the image and zoom in a bit. Color by type.

```
dump d1 all image 200 file.*.jpg type element size 1024 1024 zoom 1.2 dump_modify d1 pad 5 element N O
```

```
# in.03
# 3d random polymer
                                   03/in.03
atom_style bond
units lj
read_data data.ab
special_bonds fene
# Step 1: soft potential push-off to remove overlaps
pair_style <u>soft</u> 1.12246
pair_coeff * * 0.0 1.12246
# these have to match the data in def.chain.ab
bond_style harmonic
bond_coeff 1 50.0 0.97
bond_coeff 2 50.0 1.17
# make type 2 particles a bit heavier
mass 2 1.5
velocity all create 0.45 2349852
fix
        1 all nve
# use a crude implicit solvent model
        2 all langevin 0.45 0.45 0.02 8742
fix
```

# LAMMPS input for random polymer

 $Imp_g++ < in.03$ 

```
# use this to ramp up the height of the soft core repulsion
variable prefactor equal ramp(1.0,20.0)
        3 all adapt 1 pair soft a * * v_prefactor
fix
thermo
            50
                                            03/in.03
            10000
run
unfix
unfix
#undump d1
neighbor 0.5 bin
# Main run
           lj/cut 5.0
pair_style
pair_coeff 1 1 1.0 1.0
pair_coeff
            1 2 1.1 1.122462
pair_coeff
            2 2 1.2 1.2
thermo
            1000
reset_timestep
dump d1 all image 200 snap-03.*.jpg element element
dump_modify d1 pad 5 element N O
        10000
run
```

# LAMMPS input for random polymer

Can you adjust this line to output a movie? Coordinates of the atoms?

# Image/**Move** Creation with LAMMPS

 Piece together individual images to make a movie using ffmpeg

```
> ffmpeg -i file.%05d.jpg movie.mp4
```

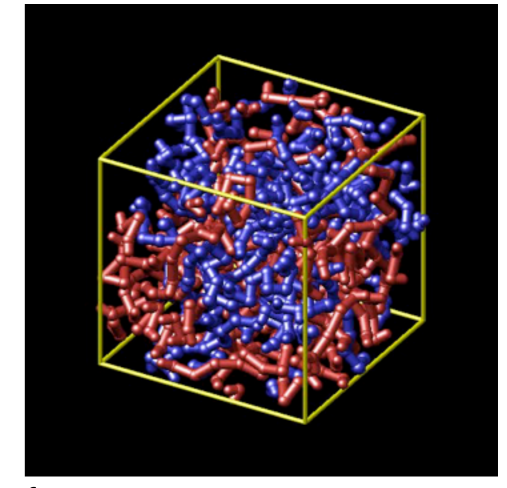
Filenames must be sequential:

```
file.00001.jpg
file.00002.jpg
file.00003.jpg
```

```
sh -c 't=0 ; for s in file.?????.jpg; do mv $s file.$t.jpg ; t=`expr $t + 1`; done'
```

# Image/**Move** Creation with LAMMPS

#### · dump movie



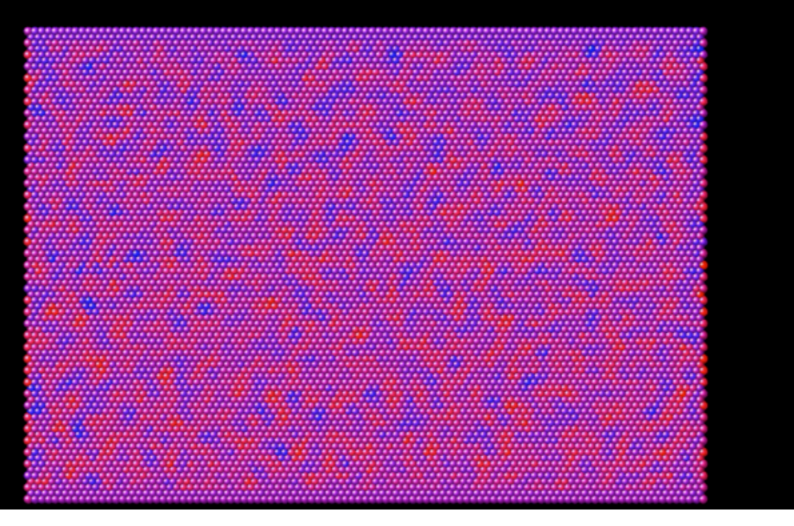
create a movie with 200 steps between frames

dump\_modify d1 element N O

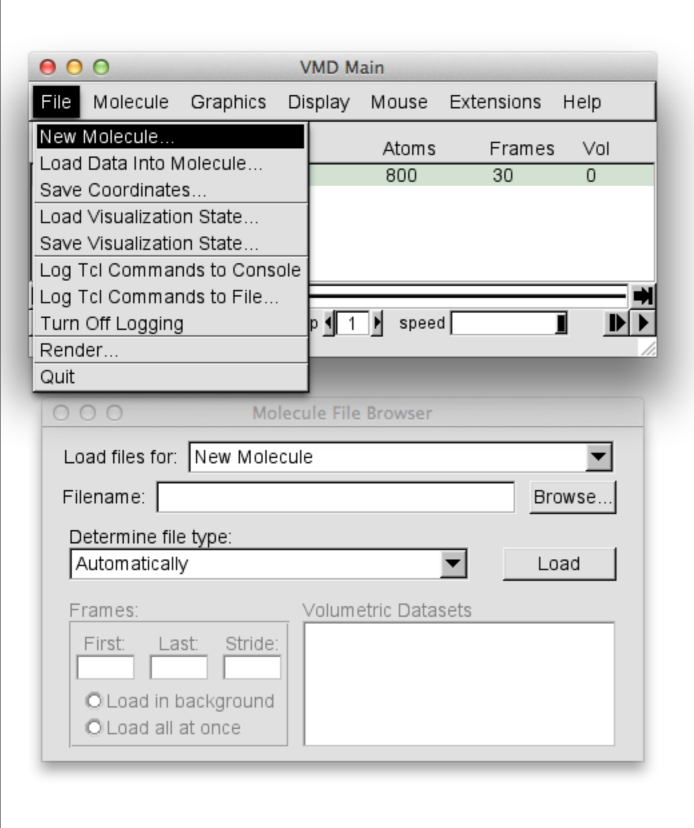
Enlarge the movie and zoom in a bit

dump d1 all movie 200 file.mpg element element size 1024 1024 zoom 1.2 dump\_modify d1 element N O

#### Axel Kohlmeyer



shell sh -c 't=0; for s in snap-movie.?????.ppm; do mv \$s snap-movie.\$t.ppm; t=`expr \$t + 1`; done' shell ffmpeg -y -an -i:v snap-movie.%d.ppm -r 24 -b:v 2400k -c:v libx264 crack-strain.mp4



#### Loading a molecule/ trajectory into VMD

- Use the GUI: "New Molecule"
- Use the 'mol' command:

```
mol new myfile.xyz
mol addfile mytraj0.dcd
mol addfile mytraj1.dcd
mol addfile mytraj2.dcd
```

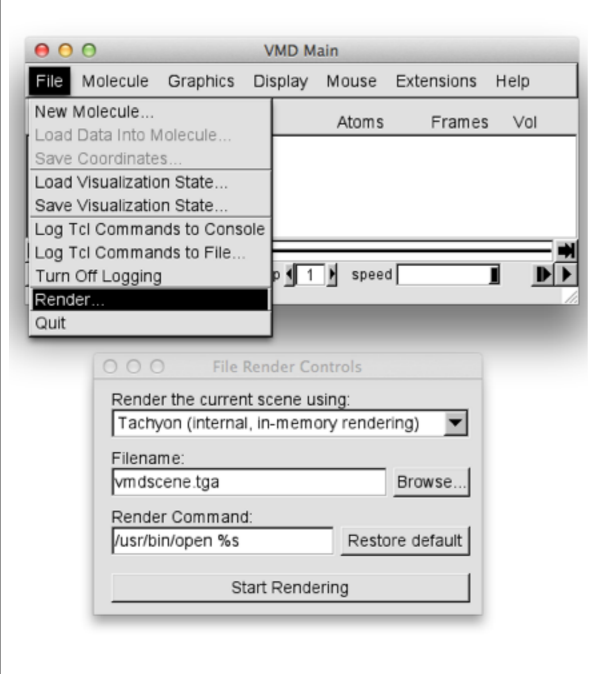
. . . . .

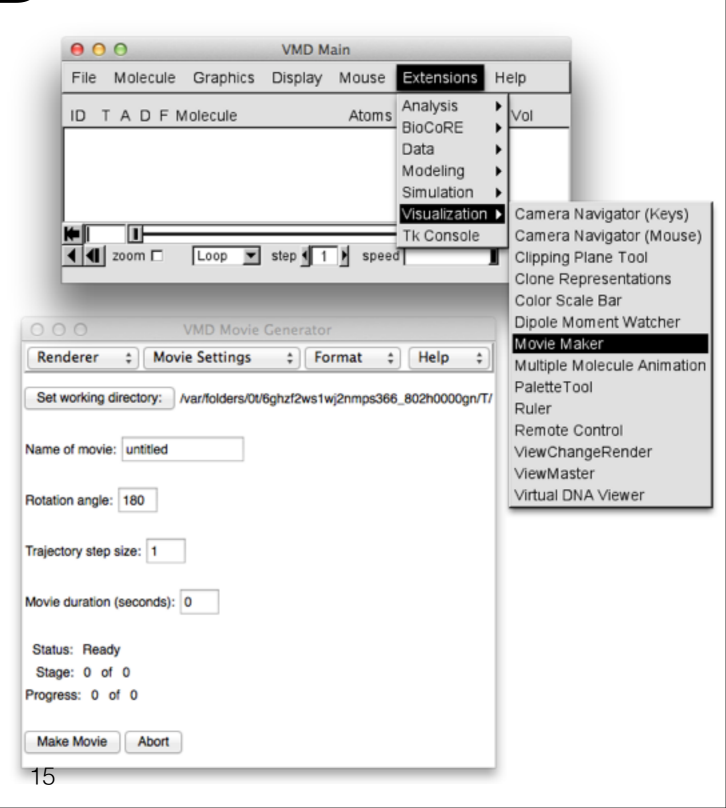
•Lammps data:

topo readlammpsdata file.data

#### Image/Move Creation with VMD

 The GUI provides both a render menu and moviemaker tool.





# Image/**Move**Creation with VMD

- Use GUI. (What fun is this??)
- Use TCL.
  - Load trajectory and LAMMPS topology (.data) files.
     topo readlammpsdata my.data Requires TopoTools mol addfile my.dcd
  - Adjust scene to your liking: orientation, colors, materials, drawing mode, display size. Save a state file for easy reloading.
  - Movie: Simple 'for' loop over vmd frames, calling your preferred rendering engine in the body of the loop.
    - 'render engine filename options'

```
# movie03.tcl
# To run, in the vmd tckon type: 'source movie03.tcl'
# Load a LAMMPS data file, a trajectory,
# set some representations to our liking,
# adjust the display size and rotate/zoom,
# loop over frames and render
# Script requires TopoTools and pbctools
package require topotools
package require pbctools
# Load the data file and trajectory dcd
set molid [topo readlammpsdata data.ab]
mol addfile 03.dcd type dcd waitfor all
# Set the representation, delete the initial representation
# created by vmd
mol delrep 0 top
mol representation CPK 2.0 0.30 10.0 10.0
mol color Name
mol selection {all}
mol material AOEdgy
mol addrep top
```

## Load Necessary Packages

#### **Load Files**

# **Create Representation**

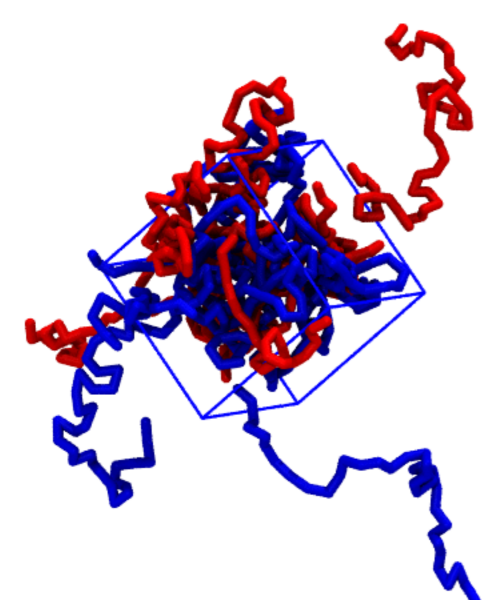
```
# Set the scene: adjust display height
# and rotate axes a bit, set the background
# to white
display height 15
rotate x by -45; rotate y by 45; rotate z by 45
color Display Background white
# Draw the periodic box centered at the origin
pbc box_draw -center origin
# Get the number of frames
set nframes [molinfo $molid get numframes]
# Loop over frames, calling render.
# Format the output filename as snap-03.00001.tga
# Skip the first "junk" frame from lammpsdata
for {set i 1} {$i < $nframes} {incr i} {
      render TachyonInternal\
        [format "snap-03.%05d.tga" $i]
## Call external script to encode movie
exec ./03-encode-vmd.sh
```

- > ./03-encode-vmd.sh
- -or-
- > ffmpeg -i snap-03.%05d.tga 03.movie.mp4

#### **Set Scene**

#### **Render Frames**

### Combine frames to make a movie

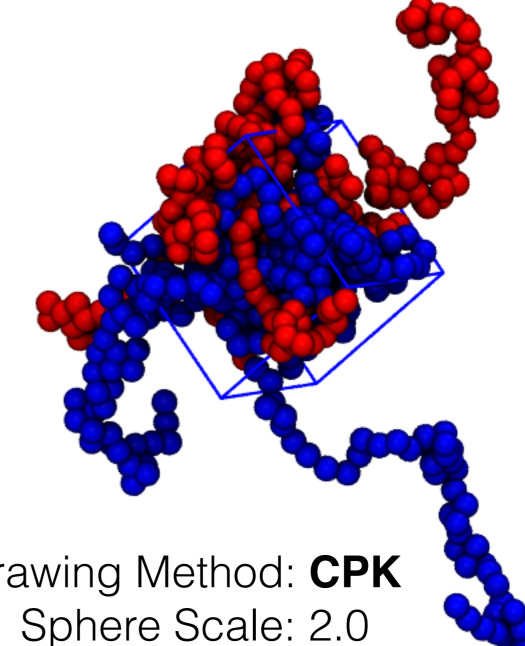


Drawing Method: tube

Color Method: name

Material: ao shiny

Ambient Occlusion: yes



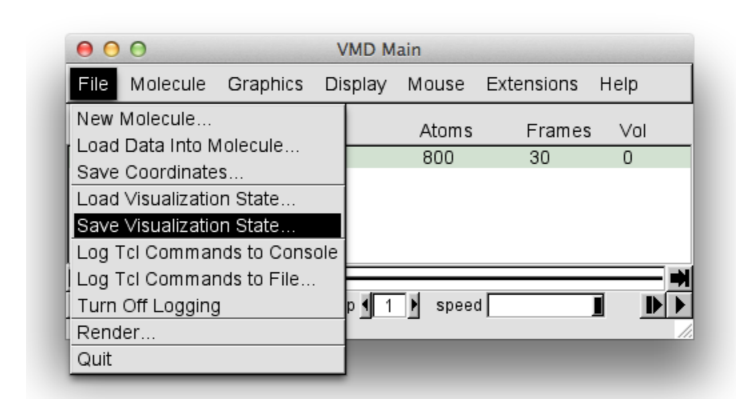
Drawing Method: **CPK** 

Color Method: name

Material: ao edgy

Ambient Occlusion: yes

# Image/**Move**Creation with VMD



Once you're happy with your scene, make sure you save its state using the 'save visualization state' option!

# VMDs true power comes from the TK (TCL) console

```
loading history file ... 48 events added
Main console display active (Tcl8.5.6 / Tk8.5.6)
Loading file /Users/macdercm/.vmd.d/atomselect_macros.tcl
Loading file /Users/macdercm/.vmd.d/bscale.tcl
Loading file /Users/macdercm/.vmd.d/colorscale.tcl
Loading file /Users/macdercm/.vmd.d/cpk.tcl
Loading file /Users/macdercm/.vmd.d/cpyrep.tcl
Loading file /Users/macdercm/.vmd.d/dx.tcl
(macdercm) 49 %
```

Interact with VMD by issuing commands in the TCL scripting language!

# TCL (tickle)

- A "high level" interpreted scripting language particularly adept at text processing and interfacing with C, C++ codes. Has a closely related GUI construction language "tk".
- All data types are manipulated as strings.
- VMD has a built in TCL interpreter for interacting with its internal data structures including molecules, atoms or groups of atoms, visualization, rendering, vector/ matrix operations, topologies and data analysis.

# TCL Syntax

- command -flag -flag -flag <args>
- Arguments are typically \$variables containing strings or one of two data structures: the "list {}" or the "array(), aka, hash-table"
- Example: Search a list of elements, return the index of the first matching element:
  - lsearch {a b c d e c} "c" => 2
  - lsearch -index 1 { {a b c d e} {f g h i j} }
    "g" => {f g h i j}

- Define and print a list:
  - set mylist {3 5 1 4 2} or [list 3 5 1 4 2]
  - puts \$mylist => 3 5 1 4 2
- Access a list element:
  - puts [lindex \$mylist 2] => 1
- Sort the list:
  - puts [lsort -increasing \$mylist] => 1 2 3 4 5

- Change a list element
  - lset mylist  $2 9 \Rightarrow 3 5 9 4 2$
- Iterate over each element in a list and print it
  - foreach x \$mylist {puts \$x} =>
    3
    5
    1

- Loop over two elements at a time and print them
  - foreach {x y} \$mylist {puts "\$x \$y"} =>
    3 5
    1 4
    2
- Append an element to a list
  - lappend mylist  $a \Rightarrow 35142a$

- Assign each element in a list to a variable:
  - (tcl8.5) lassign {1 2 3} a b c
    puts "\$a \$b \$c" => 1 2 3
  - (tcl 8.4) foreach {a b c} {1 2 3} {break} puts "\$a \$b \$c" => 1 2 3
- Unset a variable or list: unset mylist

Variables contained within {\$a \$b \$c} are not substituted. How then can we define a list of variables?

```
set a 1; set b 2; set c 3
set mylist [list $a $b $c]
```

#### TCL Resources

- http://www.tcl.tk/man/tcl8.5/tutorial/tcltutorial.html
- http://www.tcl.tk/man/tcl8.5/
- http://www.tcl.tk/man/tcl8.5/tutorial/tcltutorial.html

### VMD's '<u>Atomselect</u>' Command

- set sel [atomselect molid "selection text"] => atomselect0
- Primary method for interacting with atoms
- Returns a handle to an internally created proc that references the atoms satisfying the selection text criteria.
  - e.g. "all", "resname popc", "chain A", "residue 5", "resid 10 to 20", "resid 10 20"

### VMD's 'Atomselect' Command

- set sel [atomselect molid "selection text"]=> atomselect0
- Combine selection criteria using logical operators:
  - "and", "or", "not", >, <, =
  - "protein and chain A and x > 5"
     "not (water or protein)"
     "name "0.\*" and charge < 0.5"</li>

# Special "phrases"

Select all water atoms within 5 angstrom of protein

```
"water within 5 or protein" vs. "water exwithin 5 of protein"
```

Make sure we get complete water molecules:

same residue as (water exwithin 5 of protein)

### VMD's '<u>Atomselect</u>' Command

- List all keywords: "atomselect keywords"
- name type index serial atomicnumber element residue resname resid chain segname segid x y z radius mass charge beta occupancy all none
- fragment numbonds backbone sidechain protein nucleic water waters helix alpha\_helix helix\_3\_10 pi\_helix sheet betasheet beta\_sheet turn coil structure phi psi sequence
- user user2 user3 user4
- sqr sqrt abs floor ceil sin cos tan atan asin acos sinh cosh tanh exp log
- acidic cyclic acyclic aliphatic alpha amino aromatic basic bonded buried cg charged hetero hydrophobic small medium large neutral polar purine pyrimidine surface lipid ion sugar solvent carbon hydrogen nitrogen oxygen sulfur noh heme

# Selection Querying

- set sel [atomselect molid "selection text"] => atomselect0
- Querying a selection:set idx [\$sel get index]=>{0 1 2 3 4 5}
- returns the value of the keyword requested for each atom as a TCL list
- Atoms are returned in the order in which they were loaded/created in VMD. NOT the order you specify:
  - set sel [atomselect molid "name CB CA"] and set sel [atomselect molid "name CA CB"] return identical lists!

# Selection Querying

Number of atoms selected
 \$sel num

The selection text:\$sel text

The associated molecule (molid)
 \$sel molid

# Selection Querying

- Querying multiple keywords (performance considerations)
  - Two separate selections:

```
set id [$sel get id]
set an [$sel get name]
```

Vs. Aggregated list (expensive, but has its uses)

```
set id_an [$sel get {index name}] =>
{{0 N} {1 CA} {2 CB} {3 CG}}
```

### Modifying Atom Properties

- Change selected atoms' name to CA
   \$sel set name CA
- Change selected atoms' chain to "A"
   \$sel set chain A
- Change the selected atoms' residue name to "POPC"
   \$sel set resname POPC
- Set the coordinates of selected atoms to {1 2 3}
   \$sel set {x y z} {1 2 3}

### Coordinate Manipulation

Translate the selected atoms -5 angstroms along z-axis

```
$sel moveby {0.0 0.0 -5.0}
```

Rotate the selected atoms 180 degrees about z-axis

```
$sel move [transaxis z 180.0 deg]
```

 Rotate selected atoms 104 degrees about bond having coordinates r1, r2

```
$sel move [trans bond r1 r2 104 deg]
```

- How could you get the total number of different atom names and their corresponding values for a given structure?
- How could you renumber all the residues in all chains from a mis-formatted <u>PDB</u> file?
- How could you select all atoms within 5A of the origin?
- How about all atoms within a cube with edge length 3A centered at the origin? What if I wanted entire residues/ fragments not just atoms?

```
## Get unique atom names

set sel [atomselect top "all"]

set names [lsort -unique [$sel get name]]

puts [llength $names]

$sel delete
```

```
#Renumber residues in each chain from 1 to N
set sel [atomselect top "all"]
foreach c [lsort -unique [$sel get chain]] {
  set i 1
  foreach id [lsort -unique [$sel get residue]] {
     set sel2 [atomselect top "chain $c and resid $id"]
     $sel2 set resid $i
     incr i
     $sel2 delete
$sel delete
```

```
# Select all atoms within 5 angstroms of the origin sqrt(x*x + y*y + z*z) < 25
```

```
# Select all atoms within a cube with # edge length 3 angstroms x > -3 and x < 3 and y > -3 and z > -3 and z < 3
```

 How could you construct a <u>lookup (hash) table</u> that uses the atom index as a key, and the value being the atom type?

```
idx(0) type0idx(1) type1idx(2) type2
```

```
# Build a lookup table associating
# atom index to atom type

set sel [atomselect top "all"]
foreach idx [$sel get index] type [$sel get type] {
    set idx($idx) $type
}
```

 How would you write a procedure to output a <u>PDB</u> formatted file using VMD's atomselection functionalities?

```
• proc writePDB {sel} {
    ...
}
```

Note: there is already the command '\$sel writepdb' file.pdb' command, this is just an exercise!

```
#Write out a PDB formatted file for a selection
proc writePDB {sel} {
# Loop over each atom property in parallel
foreach charge [$sel get charge]\
     type [$sel get type]\
      atomname [$sel get name]\
      resname [$sel get resname]\
      resid [$sel get resid]\
      X [$sel get x]\
      Y [$sel get y]\
      Z [$sel get z]\
      chain [$sel get chain]\
      occ [$sel get occupancy]\
      beta [$sel get beta]\
      segid [$sel get segid]\
      ele [$sel get element]\
      serial [$sel get serial] {
    # Display it using the PDB format
    puts [format "%-6s%5d %4s %3s %s%4d %8.3f%8.3f%8.3f%6.2f%6.2f
                                                                     %-4s%2s"\
                  "ATOM " $serial $atomname $resname $chain\
                  $resid $X $Y $Z $occ $beta $segid $ele]
```

>Main< (writepdb) 246 % writePDB \$sel								
ATOM	1190	CHA HEM A 154	-5.248	39.769	-0.250	1.00	7.67	C
ATOM	1191	CHB HEM A 154	-3.774	36.790	3.280	1.00	7.05	C
ATOM	1192	CHC HEM A 154	-2.879	33.328	0.013	1.00	7.69	C
ATOM	1193	CHD HEM A 154	-4.342	36.262	-3.536	1.00	8.00	C
ATOM	1194	C1A HEM A 154	-4.960	39.256	0.996	1.00	7.60	C
ATOM	1195	C2A HEM A 154	-5.177	39.925	2.270	1.00	7.96	C
ATOM	1196	C3A HEM A 154	-4.762	39.099	3.235	1.00	7.45	C
ATOM	1197	C4A HEM A 154	-4.279	37.882	2.615	1.00	7.44	C
ATOM	1198	CMA HEM A 154	-4.792	39.352	4.760	1.00	8.07	C
ATOM	1199	CAA HEM A 154	-5.789	41.338	2.464	1.00	7.92	C
ATOM	1200	CBA HEM A 154	-7.283	41.203	2.814	1.00	9.61	C
ATOM	1201	CGA HEM A 154	-8.000	40.335	1.803	1.00	10.04	C
ATOM	1202	01A HEM A 154	-8.314	39.154	2.116	1.00	11.60	0
ATOM	1203	02A HEM A 154	-8.252	40.816	0.681	1.00	11.56	0
ATOM	1204	C1B HEM A 154	-3.405	35.579	2.720	1.00	7.15	C

#### NICE!

Created by Axel Kohlmeyer

- Tools for getting/setting/guessing molecular topologies using VMDs atomselection paradigm
- Support for reading/writing LAMMPS data files
- Load TopoTools into tcl interpreter:
   package require topotools
- Use the 'topo' command topo somecommand someargument [...] [-sel ] [-mol | top]

- TopoTools primary function is to provide an interface for manipulating molecular structure, including:
  - adding and deleting bonds, angles, dihedrals
  - Associating atom types with bonds, angles, dihedrals
  - Guessing missing bonds, angles and dihedrals
  - Reading/Writing LAMMPS data files.

Fix stuff like this!



- Get a list of bonded connectivities based on atom index topo getbondlist => {{0 1} {1 2} {2 3} {3 4}}
- Guess angles based on bonded connectivity
   topo -sel \$sel guessangles
   topo -sel \$sel getanglelist =>
  - {{C-CA-N 0 1 2} {CA-C-0 1 2 3} {N-C-0 3 2 4} {CA-C-N 1 2 4} {C-N-CA 2 4 5}}
- Write out a LAMMPS data file
   topo writelammpsdata file.data full

# Putting it all together: Building a Water box using just VMD

- You can use VMD to construct a water box which can then be simulated in LAMMPS. Here's how you can do it yourself:
  - 1.Use VMDs <u>vector and matrix</u> routines to construct a single water molecule having <u>TIP3P</u> geometry.
  - 2.Use VMDs selection routines to set the water molecule's properties: name,type,mass,charge,radius
  - 3.Use topotools::replicatemol command to create a box of these water molecules.
  - 4.Use TopoTools to assign bonds/angles to all molecules.
  - 5.Use TopoTools to output a LAMMPS data file

```
#!/usr/bin/tclsh
                                  waterbox/waterbox.0.tcl
proc build_tip3 {} {
   # Create a new molecule that holds three "empty" atoms
   # return the molecule id and store in molid
    set molid [mol new atoms 3]
   # Add a frame to hold atomic data
    animate dup $molid
   # Define O-H bond length in angstroms and H-O-H angle in degrees
    set r 0.9572; set theta 104.52
   # Use VMDs vector/matrix routines to move the atoms into their
   # equilibrium positions and assign their coordinates
    set o \{0.0 \ 0.0 \ 0.0\}; # coordinate of Oxygen atom
    set h1 [list $r 0.0 0.0]; # coordinate of first hydrogen atom
   # We need to calculate the position of the second hydrogen atom
   # by rotating the v = \{h1-o\} vector a value $theta in the xy-plane
    set v [vecsub $h1 $o]
   # Create rotation matrix corresponding to a rotation about the
   # z axis a value $theta
    set R [transabout {0.0 0.0 1.0} $theta deg]
   # Multiply the vector v with R, this is the new position of atom h2
    set h2 [vectrans $R $v]
```

# Example: VMD Water box

## 1. Build a TIP3P Water





#### waterbox/waterbox.0.tcl

```
# Create a selection of the three "empty" atoms so we can
   # set their coordinates and their molecular properties
    set sel [atomselect $molid "all"]
                                       1. Build a
    set xyz [list $o $h1 $h2]
                                     TIP3P Water
    $sel set {x y z} $xyz
   # Now we need to set the atomic mass, charge radius, and name of each
   # of the atoms. We create a list-of-sublists, where each sublist
    # contains the properties of each atom. The order of the properties
   # in each sublist must be consistent across the sublists
   # { {o_mass o_charge o_radius o_name o_type}\
     {h1_mass h1_charge h1_radius h1_name h1_type}\
      {h2_mass h2_charge h2_radius h2_name h2_type}}
      set props { {15.9994 -0.830 1.299 OH OT}\
                   {1.008 0.415 1.00 H1 HT}\
                   {1.008 0.415 1.00 H2 HT} }
    $sel set {mass charge radius name type} $props
   # reanalyze the mol to let vmd know the
   # atoms have their properties set
   mol reanalyze $molid
   # Delete our selection (always a good idea)
    $sel delete
   # Have the procedure return our new molecules ID
    return $molid
```

# Example: VMD Water box

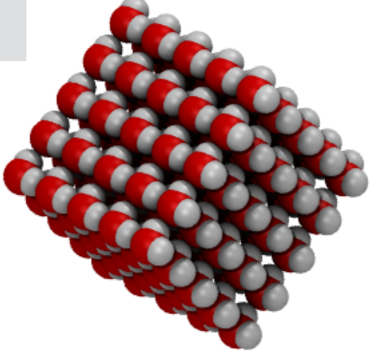
2. Apply
Atomic
Properties

#### waterbox/waterbox.1.tcl

```
proc makebox {molid} {
   # We require the package TopoTools to replicate our
   # water molecule, set our bonding and angle topologies,
   # and write out our input lammps data file.
    package require topotools
   # Set the edge length of our periodic box to
   # the 2*sigma_0-0 bond (3.1507) length.
   molinfo $molid set {a b c} {3.1507 3.1507 3.1507}
   # Replicate our water molecule in x,y,z 5x5x5 using topotools'
   # replicatemol command.
    set newmol [::TopoTools::replicatemol $molid 5 5 5]
```

# Example: VMD Water box

3. Replicate waters to make box

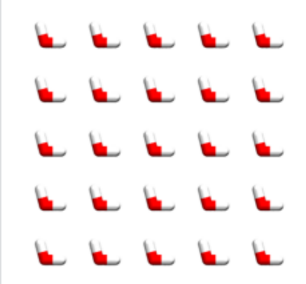


#### waterbox/waterbox.1.tcl

```
#Let's set our bonds and angles using
 # topotools, using the 'topo addbond', 'topo addangle' commands.
 # Make a selection of all {0 H*} atoms, and retrieve their indices
 # (required by addbond/addangle command).
 set sel [atomselect $newmol "name OH H1 H2"]
# Get our list of indices, they are returned as
 # a list of n atom indices in the order in which
 # they were loaded into VMD:
 # { idx_01 idx_H11 idx_H21\
     idx_02 idx_H12 idx_H22\ ...
     idx_On idx_H1n idx_H2n}
 set ids [$sel get index]
# Loop over tuples of atom indices and add bonds and angles
 foreach {id1 id2 id3} $ids {
     topo addbond $id1 $id2; # Add bond between O-H1
     topo addbond $id1 $id3; # Add bond between O-H2
     topo addangle $id2 $id1 $id3; # Add angle H1-O-H2
 ## Update VMDs and internal data structures
 mol reanalyze $newmol
 topo -molid $newmol retypebonds
 topo -molid $newmol retypeangles
```

# Example: VMD Water box

4. Apply bonding and angle topologies



#### waterbox/waterbox.1.tcl

```
# write out a protein structure file (topology), pdb (coordinates)
# and a lammps data file.
set fname tip3_125
animate write pdb $fname\.pdb
animate write psf $fname\.psf
topo -molid $newmol writelammpsdata $fname\.data full

# Delete our selection
$sel delete

# Return our new waterbox
return $newmol
```

# Example: VMD Water box

## 5. Write out input files

```
375 atoms
                                                   Masses
                                Pair Coeffs
250 bonds
                                ##
125 angles
                                                    1 1.008000 # HT
                                     HT
0 dihedrals
                                                    2 15.999400 # OT
                                # 2
                                     OT
0 impropers
2 atom types
                                # Bond Coeffs
1 bond types
                                                   Atoms
1 angle types
                                # 1 HT-0T
0 dihedral types
                                                   1 1 2 -0.830000 x y z # OT UNK
0 improper types
                                                   2 1 1 0.415000 x y z # HT UNK
                                # Angle Coeffs
                                                          0.415000 x y z # HT UNK
-7.547650 8.205850
                    xlo xhi
                                                   3 1 1
                   ylo yhi
-7.562936 8.190564
                                # 1 HT-0T-HT
                    zlo zhi
-7.876750 7.876750
```

#### waterbox/tip3\_125.param

```
lj/cut/coul/long 10.0 8.0
pair_style
               harmonic
bond_style
angle_style
               harmonic
#Masses
mass 1 1.008000 # HT
mass 2 15.999400 # OT
               1 1 0.0460 0.4000 #HT-HT
pair_coeff
             1 2 0.0836 1.7753 #HT-OT
pair_coeff
             2 2 0.1521 3.1507 #OT-OT
pair_coeff
                    450.0 0.9572 # HT-0T
bond_coeff
                    55.0 104.52 # HT-OT-HT
angle_coeff
```

# Example: VMD Water box

# Waterbox echo both waterbox/tip3\_125.in 2.5 bin neighbor neigh\_modify every 2 delay 0 check yes units real atom\_style full read\_data tip3\_opt.data include tip3\_125.param kspace\_style pppm 1.0e-6 minimize 1.0e-4 1.0e-6 300 1000 <u>velocity</u> all create 310.0 12345 mom yes rot yes dist gaussian mynpt all npt temp 310.0 310.0 400.0 iso 1.0 1.0 1000.0 fix d1 all dcd 10 tip3\_125.eq.dcd dump dump\_modify d1 unwrap yes timestep 0.5 thermo 10 1000 run

#### $lmp_g++ < tip3_125.in$

Example:

**VMD** 

Water box

#### Your turn!

- Visualize the output from the water box simulation. Can you make a movie?
- Can you calculate the O-O <u>radial distribution function</u>?
   Check VMD's "extensions->analysis" menu.
- There are two other water box scripts provided, waterbox.
   3.tcl and waterbox.4.tcl. Try running them and examining their output. What do you think they are doing?
- Axel has two very nice <u>TopoTools Tutorials</u> for creating topologies for various molecules. Companion files are in 03/ and start at 05.tcl, in.05