

# VMD: Visual Molecular Dynamics

## Computational Microscope / Tool to Think

*amino acid tyrosine*

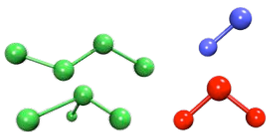
enzymatic control  
**BPTI**  
*VMD tutorial*

trafficking  
**Ubiquitin**  
*case study*

<http://www.ks.uiuc.edu/Training/CaseStudies/>  
<http://www.ks.uiuc.edu/Training/Tutorials/>

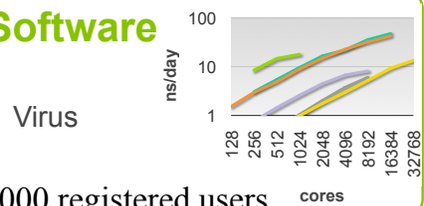
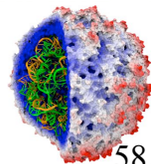
## Our Microscope is Made of...

### Chemistry



$$U(\vec{R}) = \underbrace{\sum_{\text{bonds}} k_i^{\text{bond}} (r_i - r_0)^2}_{U_{\text{bond}}} + \underbrace{\sum_{\text{angles}} k_i^{\text{angle}} (\theta_i - \theta_0)^2}_{U_{\text{angle}}} + \underbrace{\sum_{\text{dihedrals}} k_i^{\text{dih}} [1 + \cos(n_i \phi_i + \delta_i)]}_{U_{\text{dihedral}}} + \underbrace{\sum_i \sum_{j \neq i} 4 \epsilon_{ij} \left[ \left( \frac{\sigma_{ij}}{r_{ij}} \right)^{12} - \left( \frac{\sigma_{ij}}{r_{ij}} \right)^6 \right]}_{U_{\text{nonbond}}} + \sum_i \sum_{j \neq i} \frac{q_i q_j}{\epsilon r_{ij}}$$

### NAMD Software



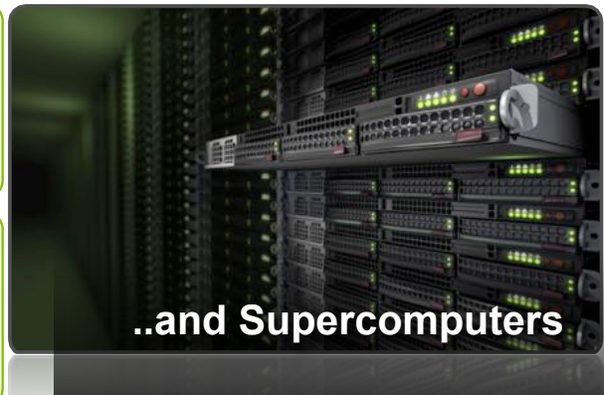
### Physics

$$m_i \frac{d^2 \vec{r}_i}{dt^2} = \vec{F}_i = -\vec{\nabla} U(\vec{R})$$

### Math

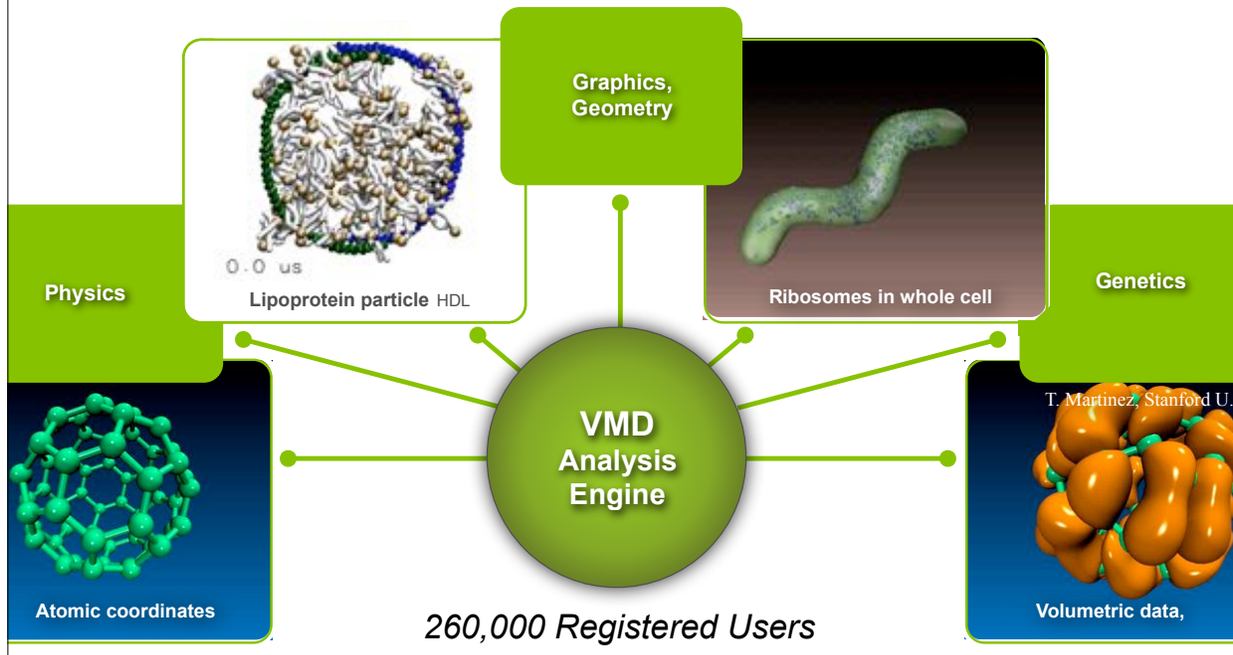
$$\vec{r}_i(t + \Delta t) = 2\vec{r}_i(t) - \vec{r}_i(t - \Delta t) + \frac{\Delta t^2}{m_i} \vec{F}_i(t)$$

(repeat **one billion times** = microsecond)



# Our Microscope is Made of...

and of...



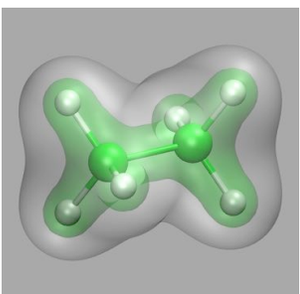
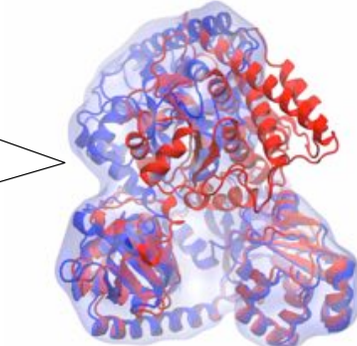
## VMD – A Tool to Think

**Volumetric Data:**  
Density maps,  
Electron orbitals,  
Electrostatic potential,  
Time-averaged occupancy, ...

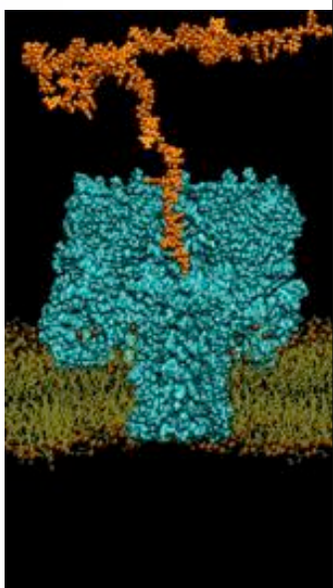
**Sequence Data:**  
Multiple Alignments,  
Phylogenetic Trees

Seq1	...
Seq2	...
Seq3	...
Seq4	...
Seq5	...
Seq6	...
Seq7	...
Seq8	...
Seq9	...
Seq10	...

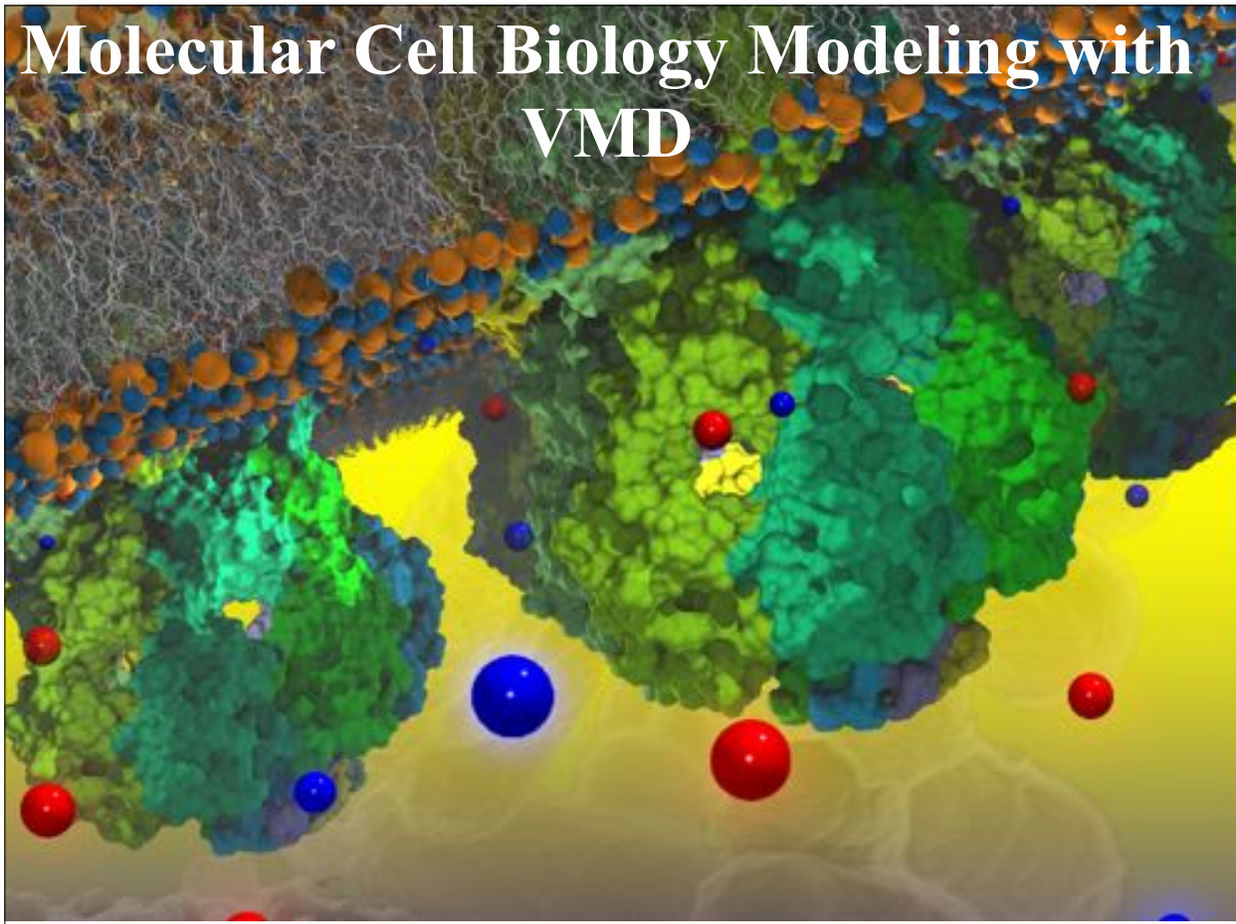
VMD



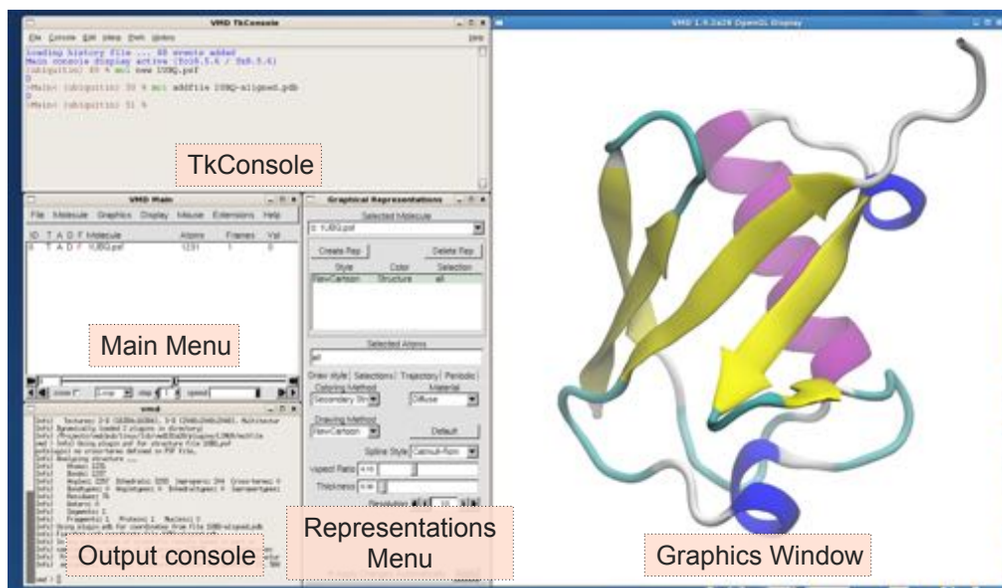
**Atomic Data:**  
Coordinates,  
Trajectories,  
Energies,  
Forces, ...



# Molecular Cell Biology Modeling with VMD



## Molecular Graphics with VMD



typical VMD session

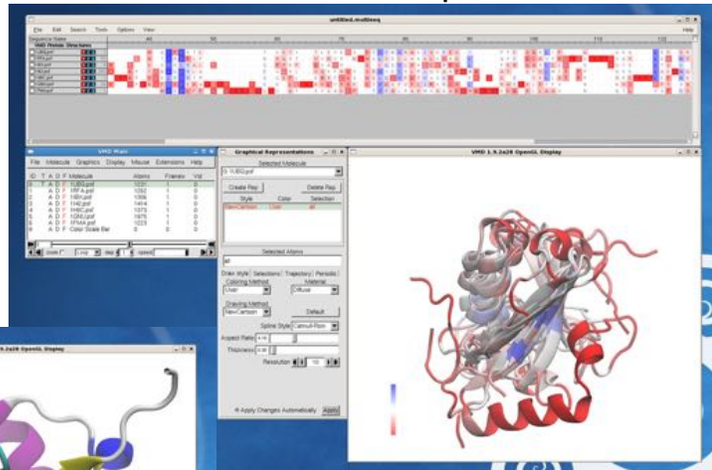
# Bioinformatics

## MultiSeq

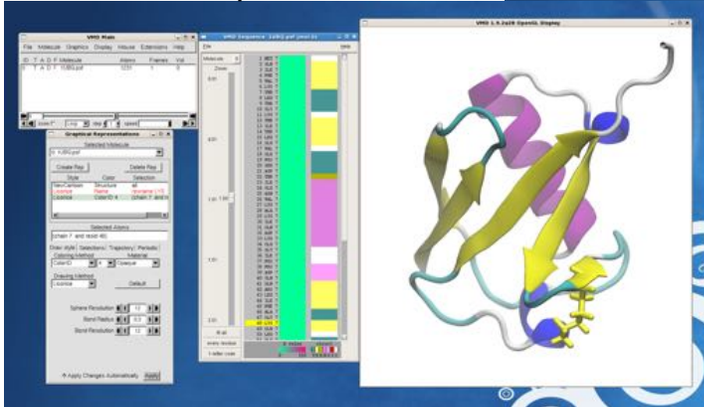
Link sequence to structure

Multiple sequence/  
structure alignments

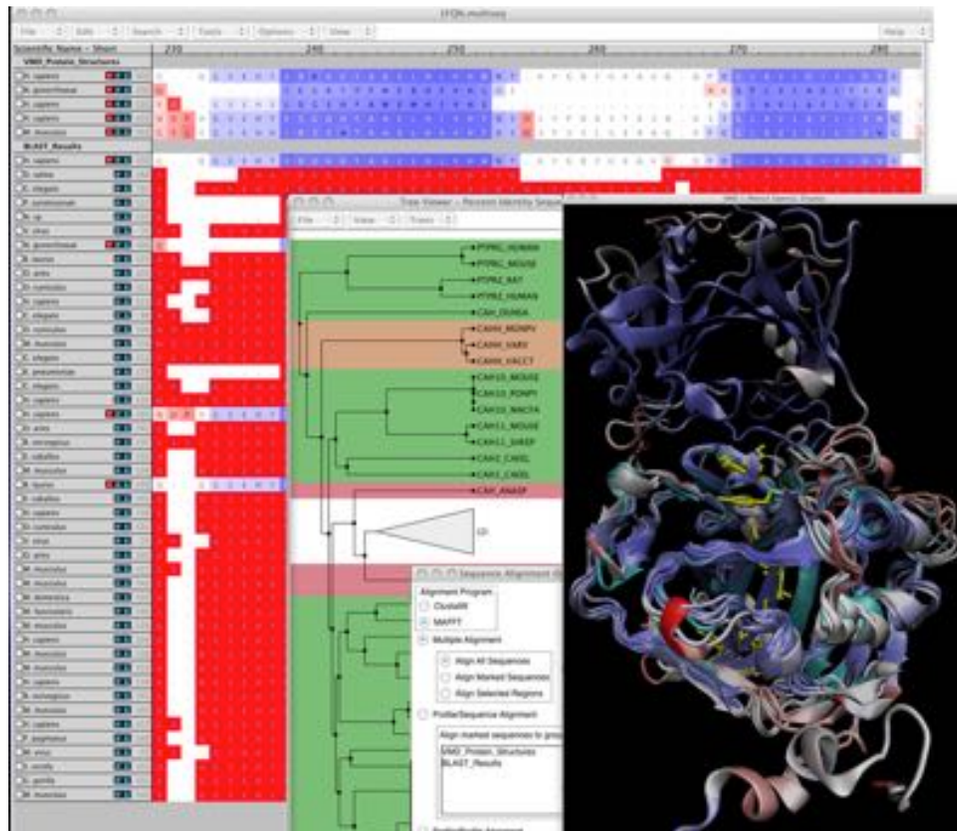
Phylogenetic trees



## Sequence Viewer



## Structure, Sequence and Phylogenetic Analysis with VMD

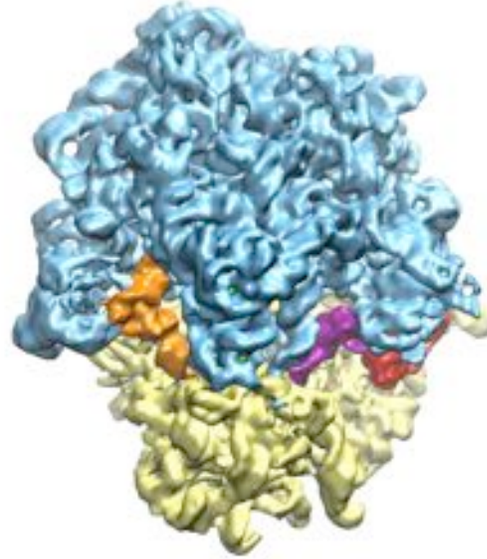


# VMD Handles Volumetric Data

Cryo-EM map of the *E. coli* ribosome at 6.7-Å resolution

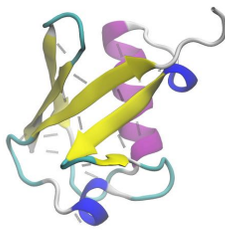


electron density  
from QM/MM  
calculation  
Martinez, Stanford

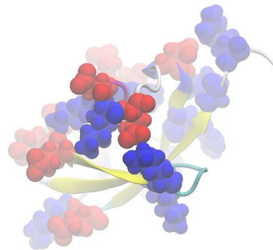


cryo-EM density

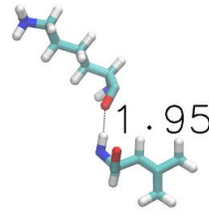
# VMD Determines Physical Properties



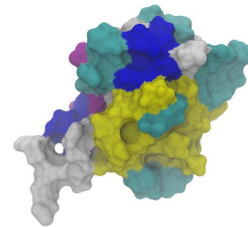
hydrogen bonds



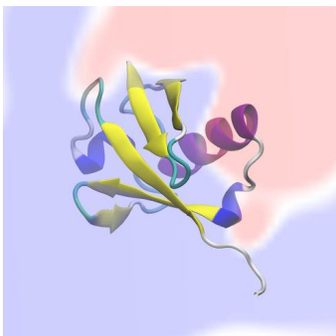
salt bridges



distances



SASA



electrostatics

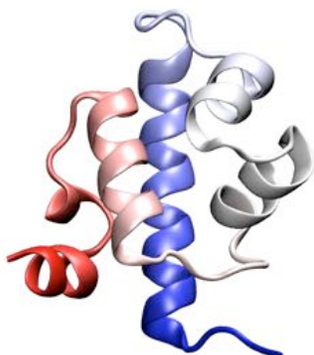
- SASA
- secondary structure
- interaction surfaces
- hydrogen bonds
- salt bridges
- electrostatics
- distances
- angles
- dihedrals
- RMSD
- RMSF
- interaction energies
- forces
- free energy profiles
- normal modes

# Trajectory Graphics/Analysis with VMD

**VMD Plugin:  
Timeline**

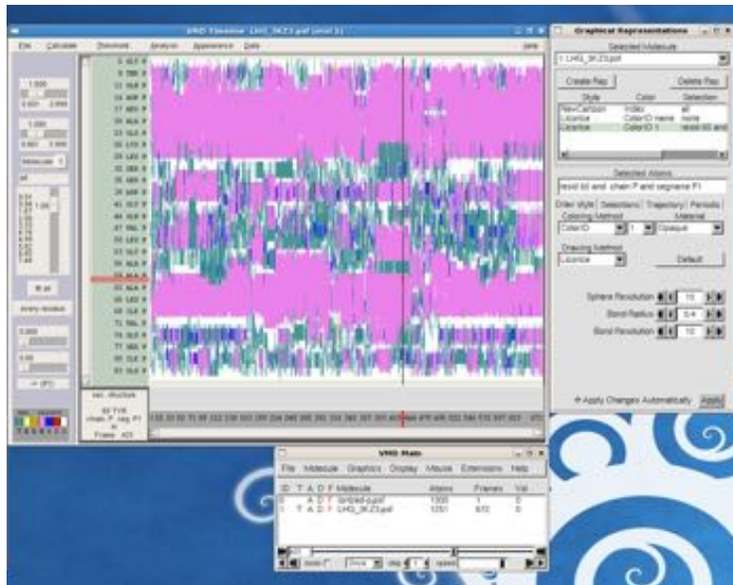
plugin to analyze MD trajectories for events  
plot properties, e.g. RMSD, secondary  
structure, hydrogen bonds, for each residue  
across a trajectory

Example: protein folding



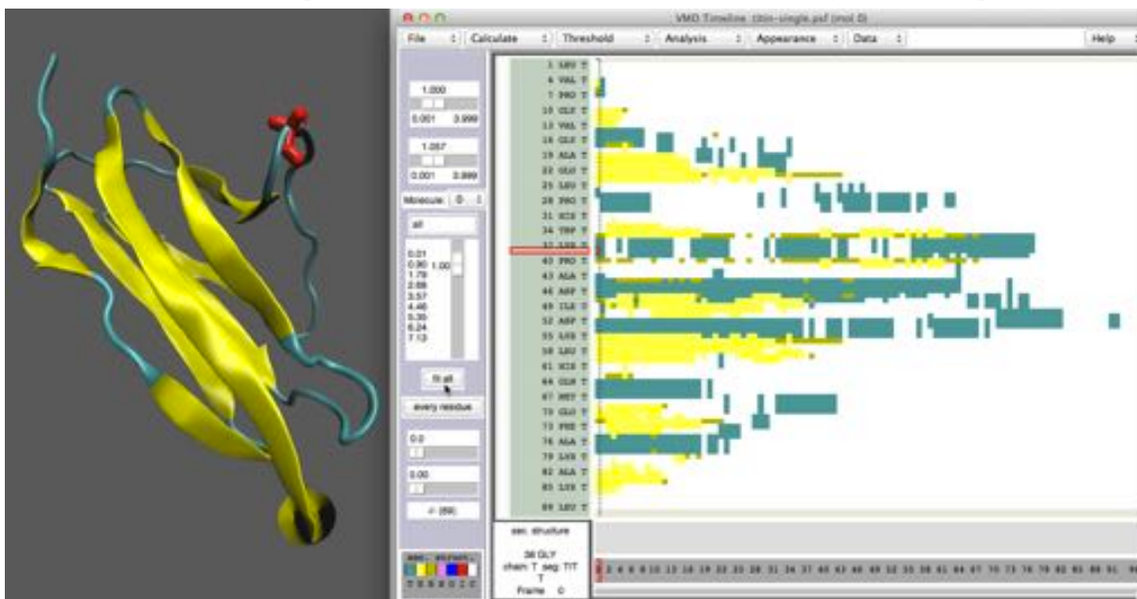
lambda repressor (2012)  
80 amino acids, 100  $\mu$ s

*Journal of Physical Chemistry Letters*, 3:1117-1123,  
2012. *Proceedings of the National Academy of  
Sciences, USA*, In press, 2013.



**VMD Session 2:**  
*trajectory of villin head piece*

# Timeline: a graphing and analysis tool to identify events in an MD trajectory

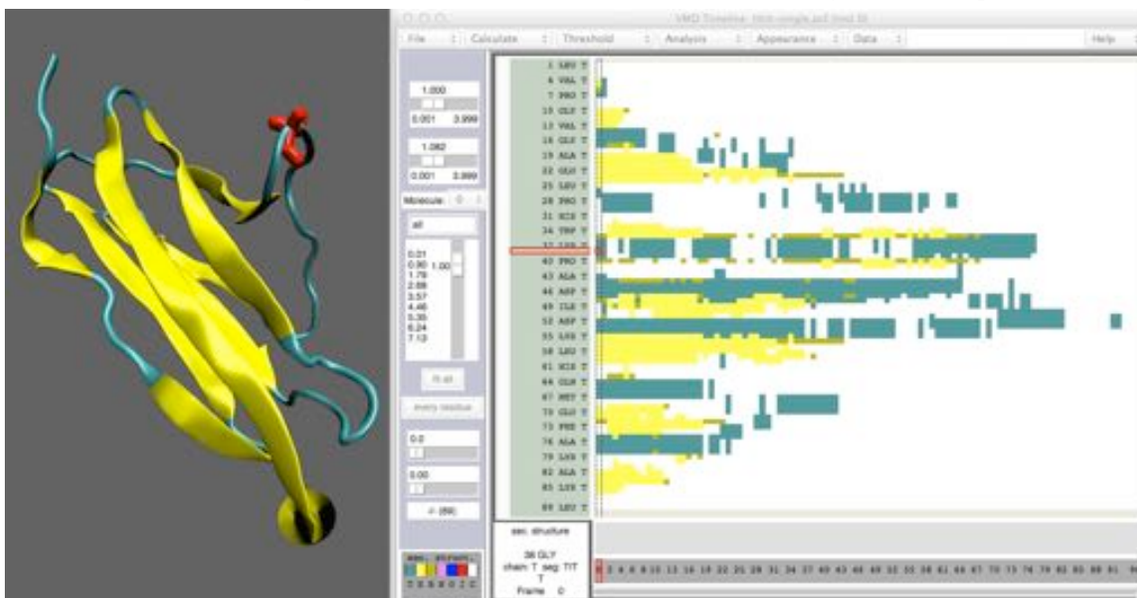


■ Beta turn   ■ Extended beta   ■ Isolated bridge   □ None (coil)

1/8

secondary structure during titin domain extension

# Timeline: a graphing and analysis tool to identify events in an MD trajectory

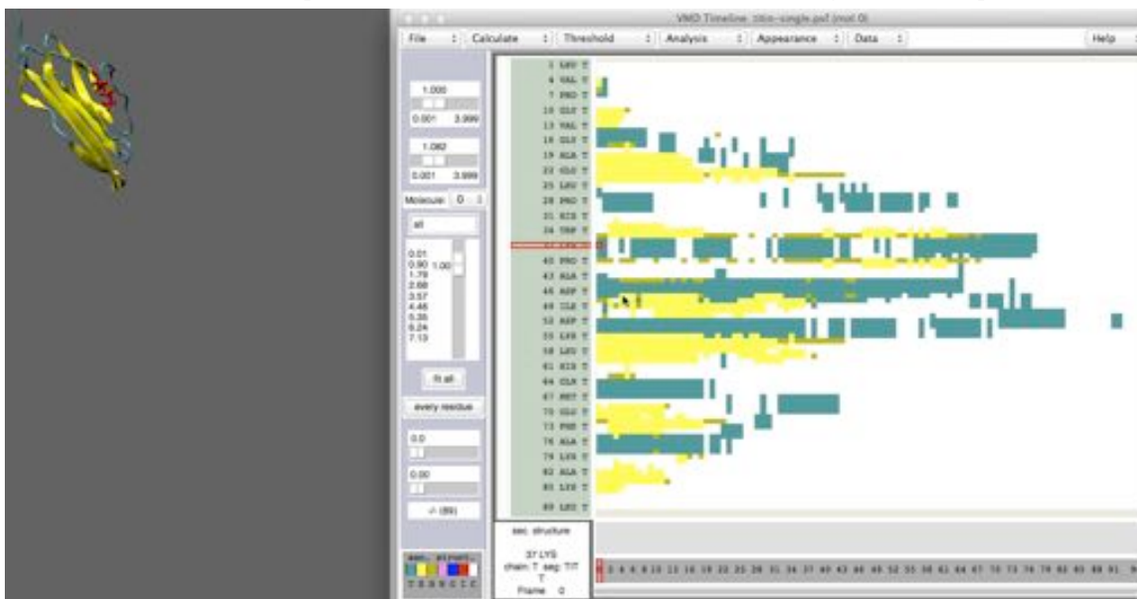


■ Beta turn   ■ Extended beta   ■ Isolated bridge   □ None (coil)

2/8

secondary structure during titin domain extension

# Timeline: a graphing and analysis tool to identify events in an MD trajectory

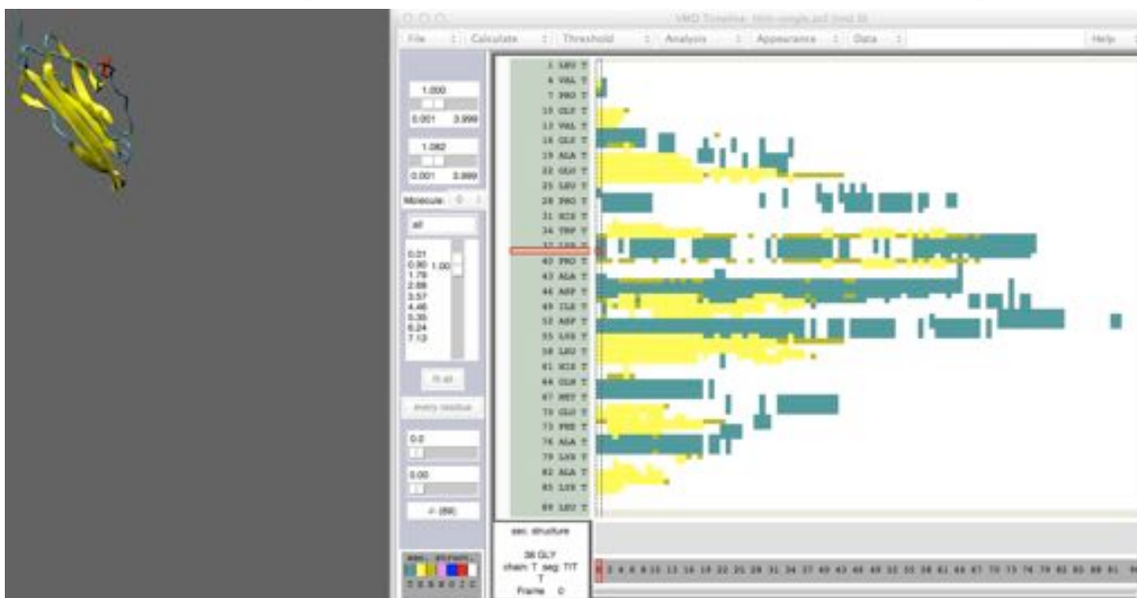


Beta turn  
  Extended beta  
  Isolated bridge  
  None (coil)

3/8

secondary structure during titin domain extension

# Timeline: a graphing and analysis tool to identify events in an MD trajectory

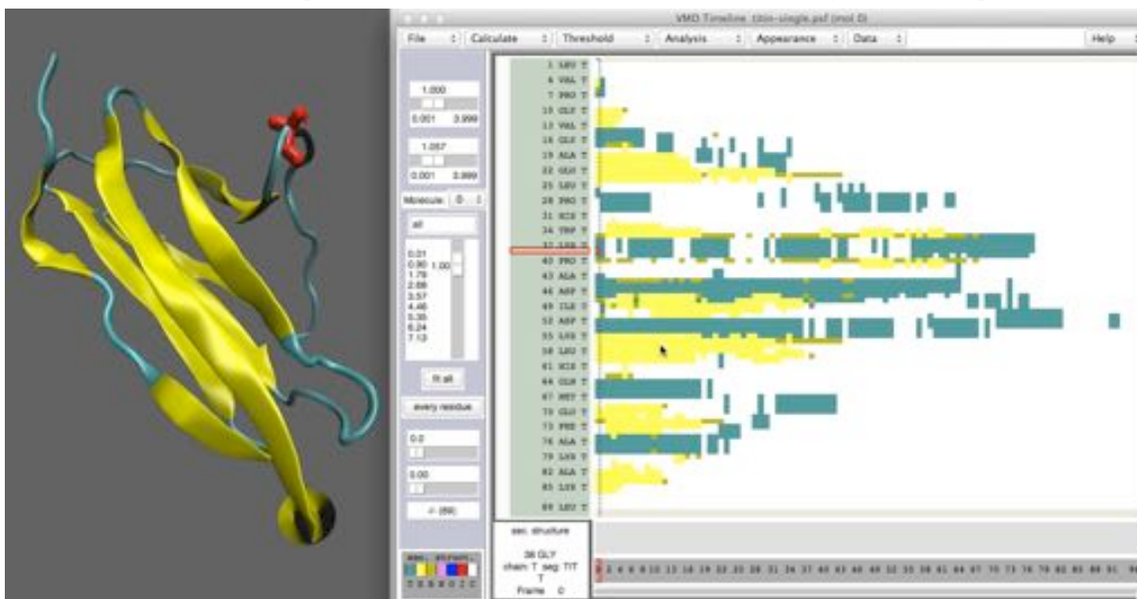


Beta turn  
  Extended beta  
  Isolated bridge  
  None (coil)

4/8

secondary structure during titin domain extension

# Timeline: a graphing and analysis tool to identify events in an MD trajectory

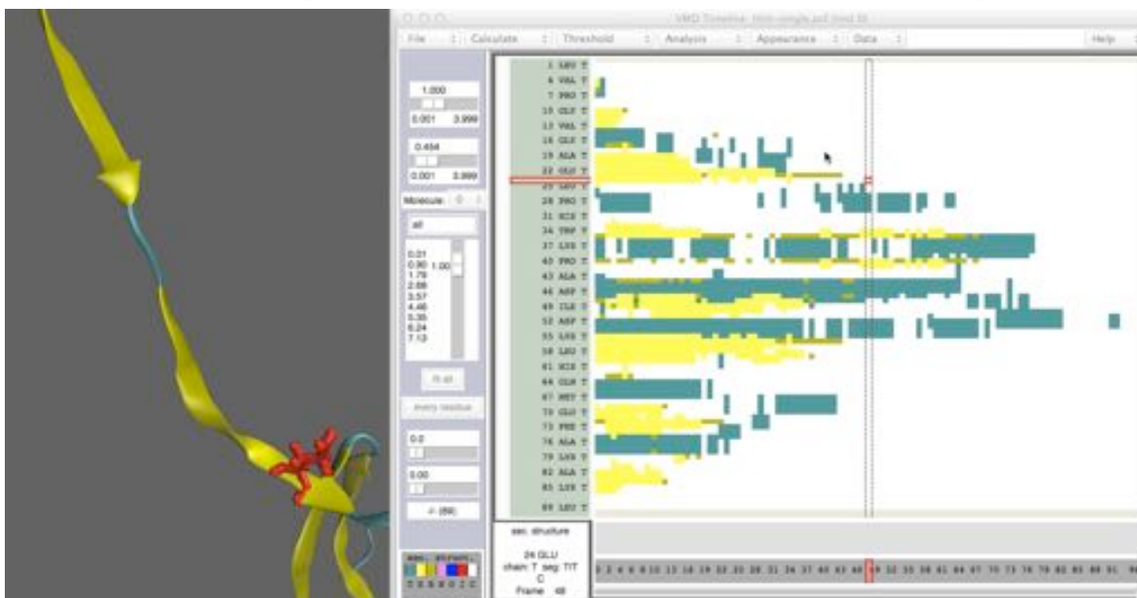


Beta turn   
  Extended beta   
  Isolated bridge   
  None (coil)

5/8

secondary structure during titin domain extension

# Timeline: a graphing and analysis tool to identify events in an MD trajectory

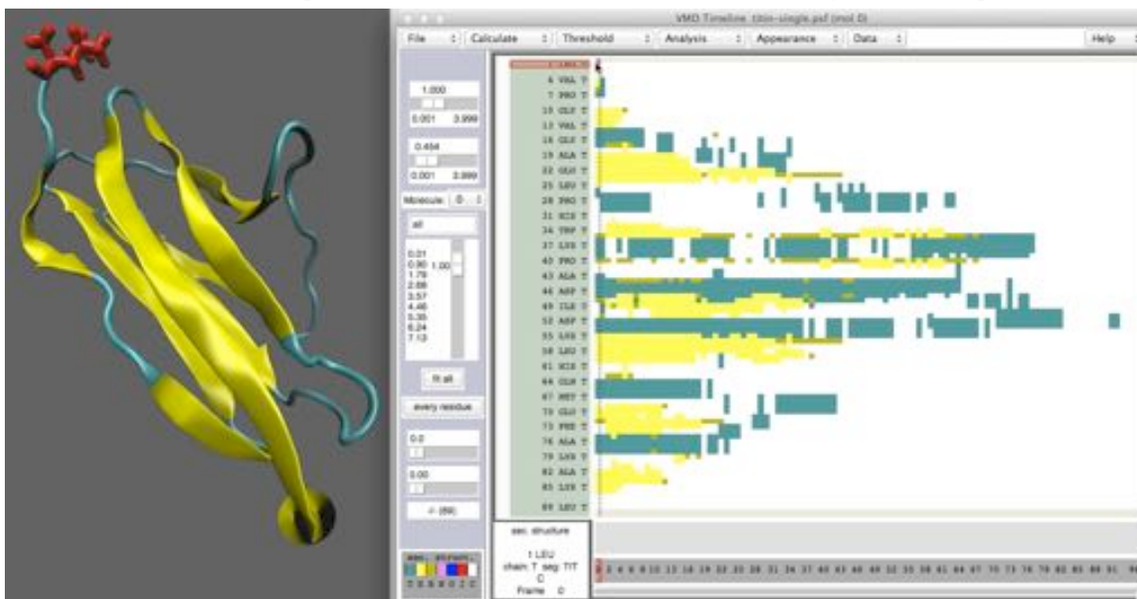


Beta turn   
  Extended beta   
  Isolated bridge   
  None (coil)

6/8

secondary structure during titin domain extension

# Timeline: a graphing and analysis tool to identify events in an MD trajectory



■ Beta turn   
 ■ Extended beta   
 ■ Isolated bridge   
  None (coil)

7/8

secondary structure during titin domain extension

## VMD Plugins: extensible analysis

### Modeling Plugins

- Autonize
- AutoPSF
- Chirality
- Cispeptide
- CGTools
- Dowser
- FFTK
- Inorganic Builder
- MDFF
- Membrane Builder
- MergeStructs
- Molefacture
- Mutator
- Nanotube
- Paratool
- Psfgen
- Solvate
- SSRestrains
- Topotools

### Analysis Plugins

- APBSRun
- CatDCD
- Contact Map
- GofRGUI
- HBonds
- ILSTools
- IRSpecGUI
- MultiSeq
- NAMDEnergy
- NAMDPLOT
- NetworkView
- NMWiz
- ParseFEP
- PropKaGUI
- RamaPlot
- RMSD Tool
- RMSD Trajectory Tool
- RMSD Visualizer Tool
- Salt Bridges
- Sequence Viewer
- Symmetry Tool
- Timeline
- VolMap

### Visualization Plugins

- Clipping Plane Tool
- Clone Rep
- Dipole Watcher
- Intersurf
- Navigate
- NavFly
- MultiMolAnim
- Color Scale Bar
- Remote
- Palette Tool
- ViewChangeRender
- Viewmaster
- Virtual DNA Viewer
- VMDMovie

### Simulation Plugins

- AutoIMD
- IMDMenu
- NAMD GUI
- NAMD Server
- QMTool

### Data Plugins

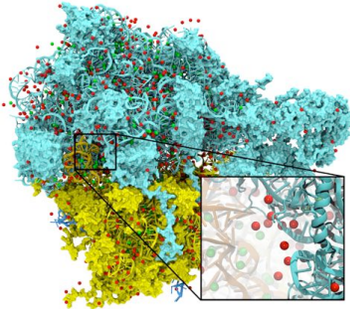
- Data Import
- Multiplot
- PDBtool
- MultiText

### Other Plugins

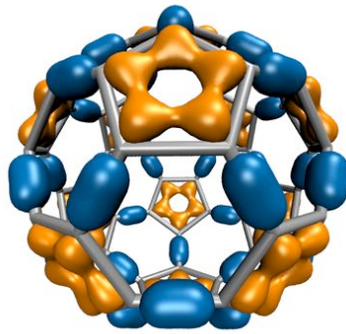
- AtomEdit
- DemoMaster
- ExecTool
- Hesstrans
- Optimization
- PBCTools
- RESPTool
- RNAview
- SignalProc
- TkCon

<http://www.ks.uiuc.edu/Research/vmd/plugins/>

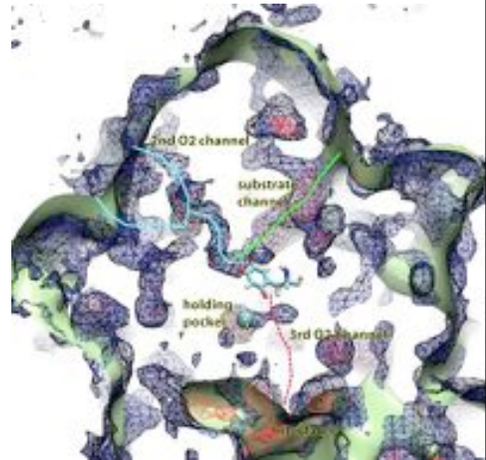
# VMD the Compute Engine



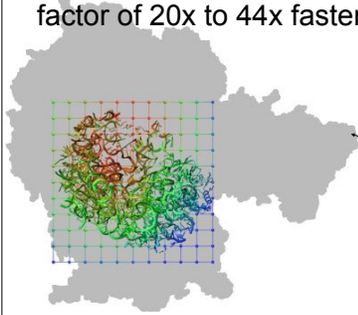
Electrostatic field calculation, ion placement: factor of 20x to 44x faster



Molecular orbital calculation and display: factor of 120x faster



Imaging of gas migration pathways in proteins with implicit ligand sampling: factor of 20x to 30x faster

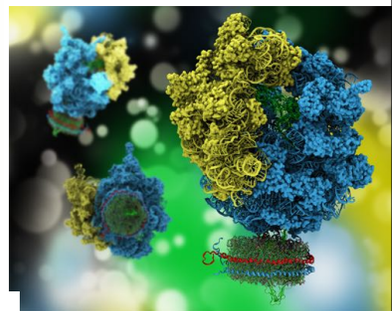
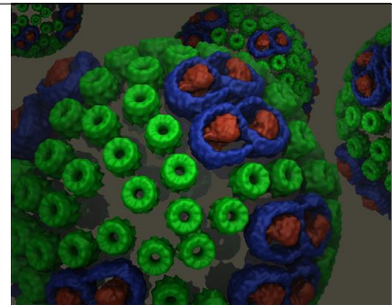


Parallel analysis on GPUs

VMD is first bio-software that is optimized for power consumption!

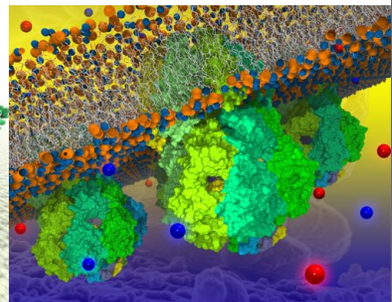
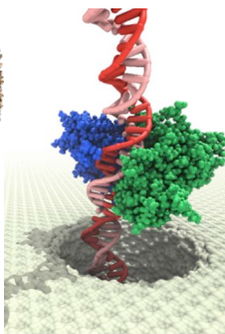
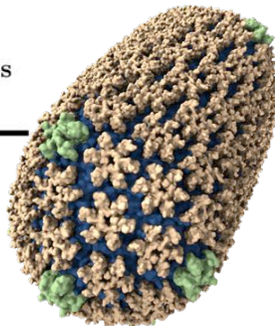
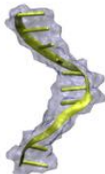
# VMD the Artist

Publication-quality images and movies



University of Illinois at Urbana-Champaign  
Beckman Institute for Advanced Science and Technology  
Theoretical and Computational Biophysics Group  
Computational Biophysics Workshop

VMD Images and Movies Tutorial



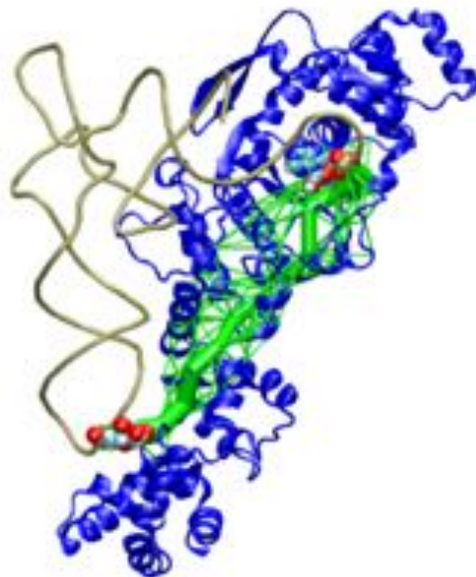
# VMD Session 3: *exchanging .vmd file of photosynthetic chromatophore for joint viewing*

VMD 1.9.1 Released Feb. 4, 2012

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## Key Features

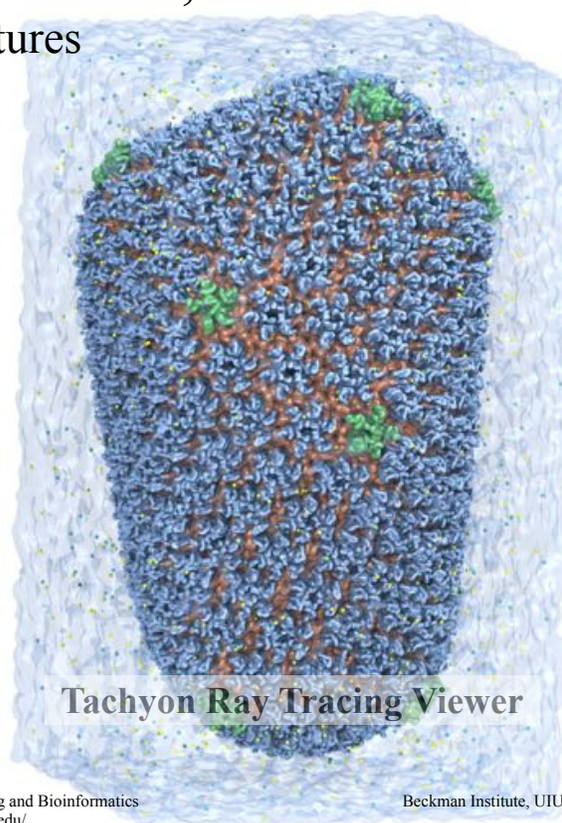
- New **NetworkView** plugin for study of allostery, signaling networks
- New **Force Field Toolkit (ffTK)** assists with CHARMM parameter development
- New “**QuickSurf**” surface representation
- New **user-contributed plugins**:
  - Normal Mode Wizard
  - PropKa interface
  - RMSD Trajectory Tool
  - “Heat map” plotting
  - Many others...



NetworkView: tRNA anti-codon w/  
ligand in GluRS active site

## Key Features

- New **Tachyon Ray Tracing** plugin for extremely realistic molecular views
- Vastly improved **Force Field Toolkit (ffTK)**: faster charge optimization routine; optimization of bond and angle parameters; projection of missing parameters onto molecular structure; interactive display for analyzing complex dihedral potential energy surfaces within the context of the molecular structure
- Updates **Molecular Dynamics Flexible Fitting (MDFF) Method**: options for MDFF with implicit solvent, xMDFF for low-resolution x-ray crystallography, multi-core CPU and GPU-accelerated analysis
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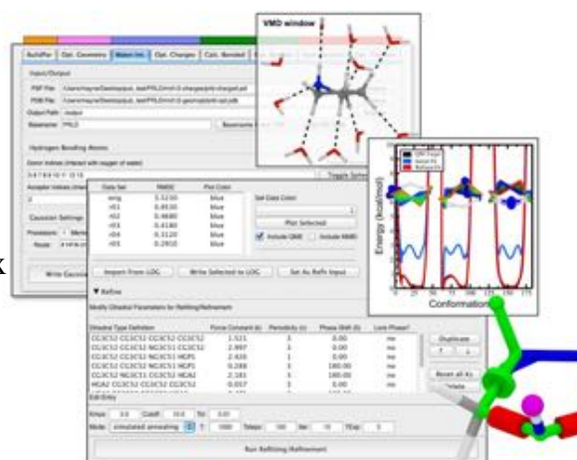
National Center for Research Fluoroscopes

BTRC for Macromolecular Modeling and Bioinformatics  
<http://www.ks.uiuc.edu/>

Beckman Institute, UIUC

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## Improved Force Field Toolkit



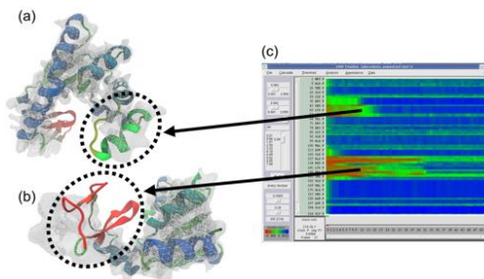
National Center for Research Fluoroscopes

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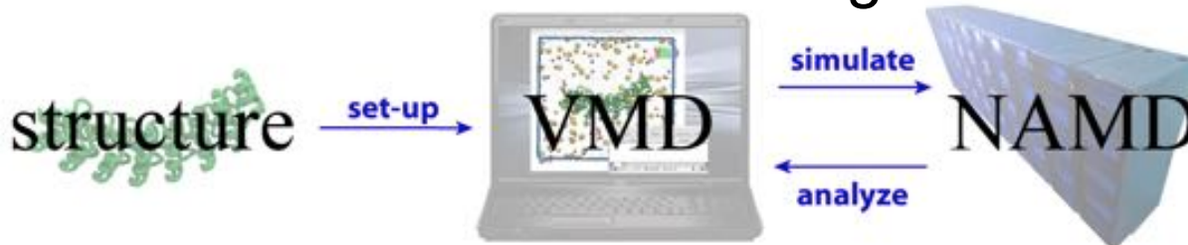
**Improved MDFF Analysis**



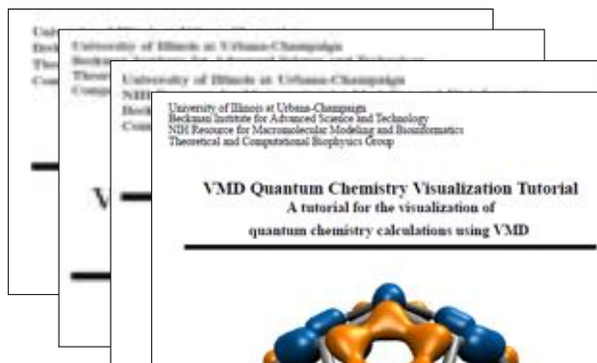
BTRC for Macromolecular Modeling and Bioinformatics  
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Beckman Institute, UIUC

## VMD and NAMD Work Together



- Over 212,000 registered VMD users
  - 18% (39,000) are NIH-funded
  - Over 49,000 have downloaded multiple VMD releases
- Over 8,000 citations
- User community runs VMD on:
  - MacOS X, Unix, Windows operating systems
  - Laptops, desktop workstations
  - Clusters, supercomputers
- VMD user support efforts:
  - 20,000 emails, 2007-2011
  - Develop and maintain VMD tutorials and topical mini-tutorials; 11 in total
  - Periodic user surveys



# The Biomedical Technology Research Center (BTRC) for Macromolecular Achievements Built on People

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- 5 faculty members (2 physics, 1 chemistry, 1 biochemistry, 1 computer science);
- 8 developers; 1 system admin.; 17 post docs; 46 graduate students; 3 administrative staff (assistant director, grants coordinator, training and dissemination)
- Funding/support from NIH and National Science Foundation



*Beckman  
Institute,  
Urbana,  
Illinois*

1-2