Visualization of Nano-Scale Structures

John E. Stone

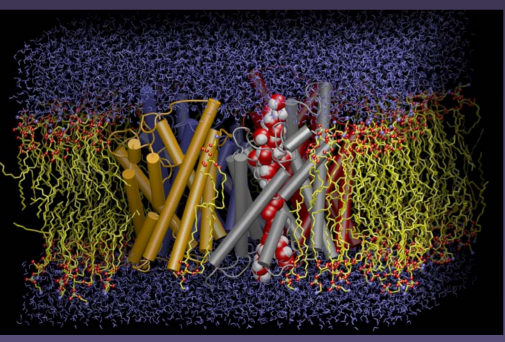
NIH Resource for Macromolecular Modeling and Bioinformatics

Beckman Institute
University of Illinois



VMD

- VMD "Visual Molecular Dynamics"
- Visualization of molecular dynamics simulations, sequence data, volumetric data
- User extensible with scripting and plugins
- http://www.ks.uiuc.edu/Research/vmd/







Overview

- Will be showing a lot of VMD images, feel free to ask questions
- General visualization concepts and methods
- Specific visualization examples for molecular dynamics trajectories, CryoEM maps, etc.
- Graphics technology driving molecular visualization capabilities and performance
- Challenges encountered exploiting these technologies for our purposes
- Where things are headed...



What is Visualization?

Visualize:

"to form a mental vision, image, or picture of (something not visible or present to sight, or of an abstraction); to make visible to the mind or imagination" [The Oxford English Dictionary]



http://www.ks.uiuc.edu/

Goals of Visualization

- Exploring data, making the invisible visible
- Gaining insight and understanding, interpret the meaning of the data
- Interactivity
- Communicating with others



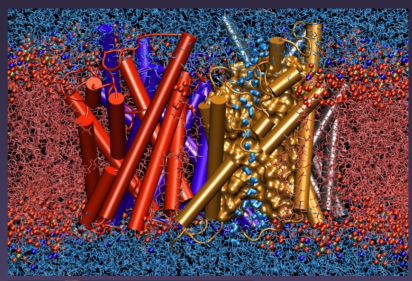
Attributes of the Data We're Interested in Visualizing

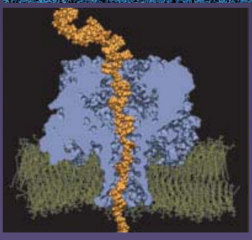
- Multiple types of data
 - Atomic structures
 - Sequence Data
 - Volumetric data
- Many attributes per-atom
- Millions of atoms, voxels
- Time varying (simulation trajectories)
- Multiple structures



Visualizing Data with Shape

- Direct rendering of geometry from physical data (e.g. atomic structures)
- Indirect rendering of data, feature extraction (e.g. density isosurfaces)
- Reduced detail representations of data (e.g. ribbons, cartoon)
- Use size for emphasis

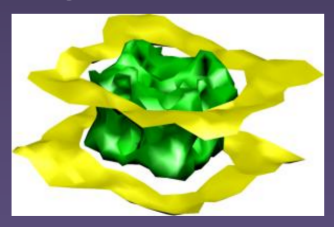


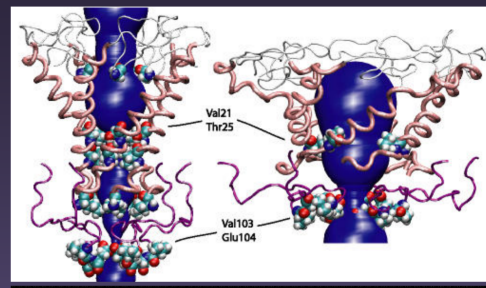


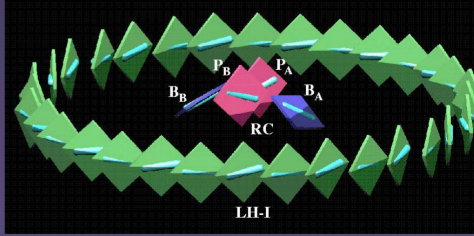


Schematic Representations

- Extract and render pores, cavities, indentations
- Simplified representations of large structures









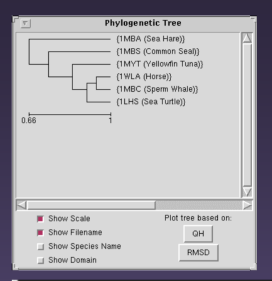
Visualizing Data with Texture and Color

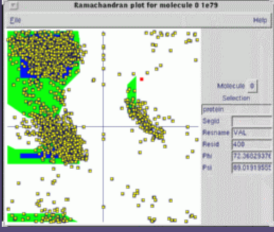
- Direct mapping of properties/values to colors (e.g. color by electrostatic potential)
- Indirect mapping via feature extraction (e.g. color by secondary structure)
- Use saturated colors to draw attention
- Use faded colors and transparency to deemphasize
- Use depth cueing/fog to de-emphasize background environment



Visualizing Data Topologically

- Data relationships indicated by grouping (e.g. phylogenetic trees)
- Abstract or schematic representation (e.g. Ramachandran plot)

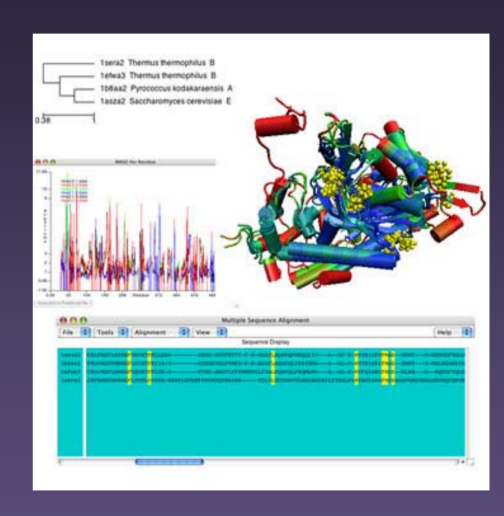






Bringing it all together...

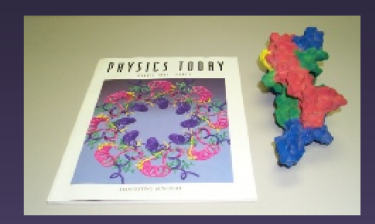
- Aligned sequences and structures, phylogeny
- Simultaneous use of shape, color, topology, and interactivity
- Multiple simultaneous representations
- Multiple data display modalities
- Selections in one modality can be used to highlight or select in others





What else can we do?

- Enhance visual perception of shape
 - Motion, interactive rotation
 - Stereoscopic display
 - High quality surface shading and lighting
- Enhance tactile perception of shape
 - Print 3-D solid models
 - Interactive exploration using haptic feedback

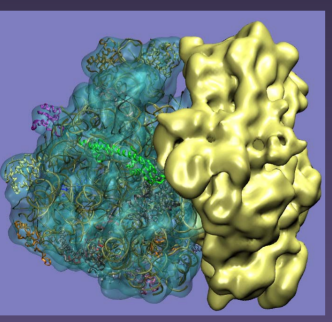


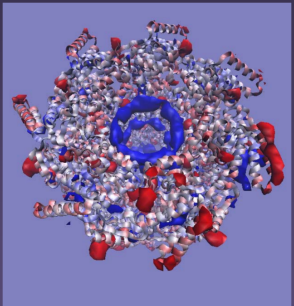


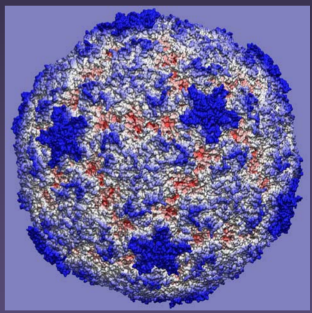


VMD Representation Examples

- Draw atomic structure, protein backbone, secondary structure, solvent-accessible surface, window-averaged trajectory positions, isosurfaces of volumetric data, much more...
- Color by per-atom or per-residue info, position, time, electrostatic potential, density, user-defined properties, etc...







Ribosome, J. Frank

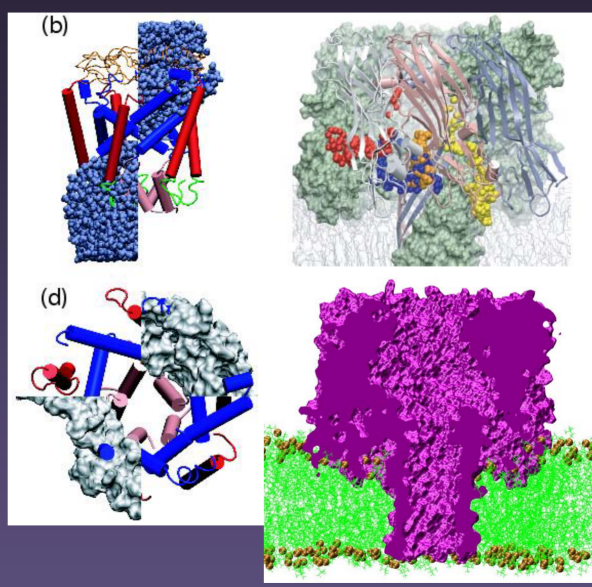
GroEL /w Situs

4HRV, 400K atoms



Multiple Representations, Cut-away Views

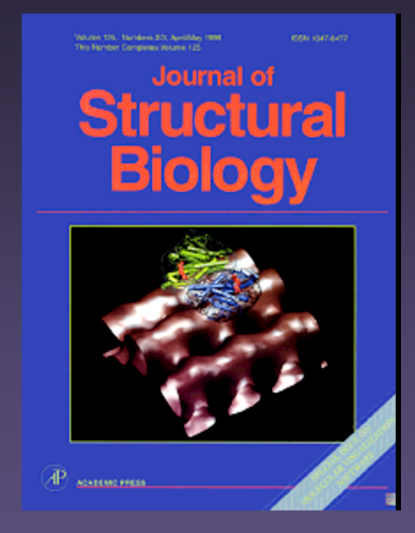
- Multiple reps are often used concurrently
 - Show selected regions in full atomic detail
 - Simplified cartoonlike or schematic form
- Clipping planes can slice away structure obscuring interesting features





Display of User-Defined Graphics

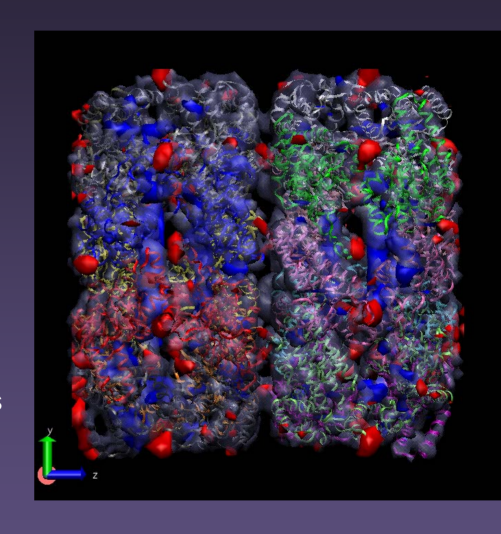
- Means for third-party tools to generate graphics for display in VMD
- SITUS did isosurface extraction for itself, which was then loaded into VMD
- Good way to prototype new visualization features and develop interfaces to other software





GroEL: Docked Map and Structure

- SITUS + VMD since 1999
- SITUS:
 - Dock map+structure
 - Synthesize map from PDB
 - Calculate difference between EM map and PDB
- VMD:
 - Load SITUS maps or meshes
 - Display isosurfaces
 - Display map/structure alignment error as isosurfaces
 - Texture reps by density or map/structure alignment error

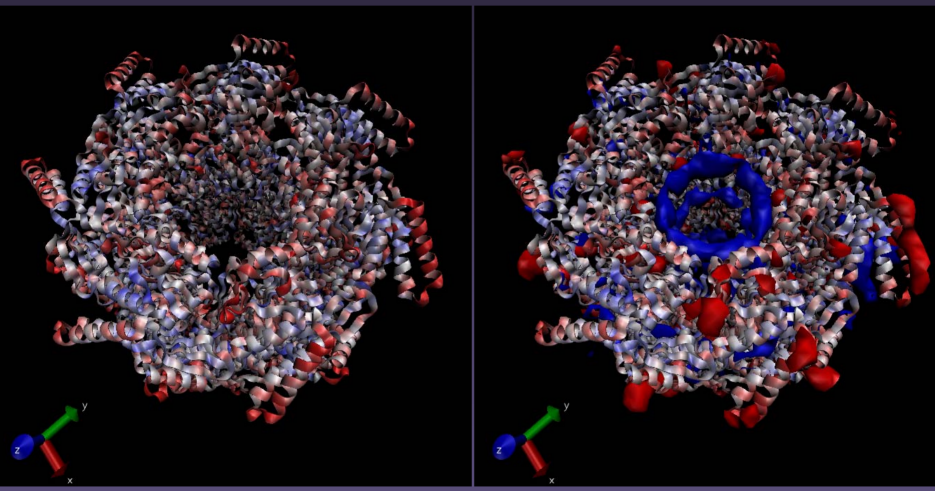




GroEL: Display of Difference, Error

Ribbons textured by difference map

...with difference isosurfaces

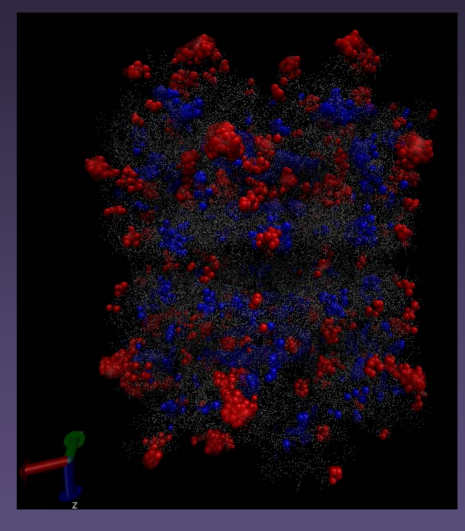


http://www.ks.uiuc.edu/



GroEL: Select Atoms by Map Values

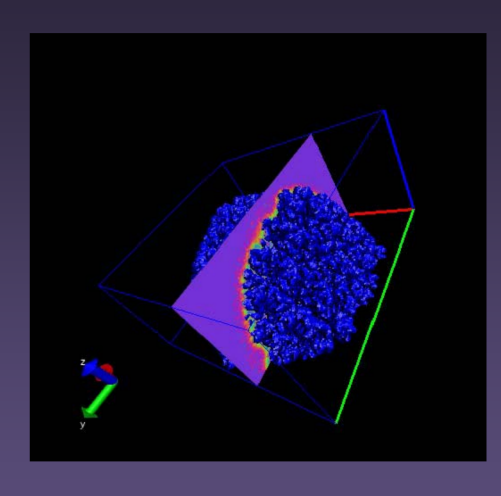
- Superimpose the difference map isosurface with VDW rep of atoms in difference areas
- Atoms can be selected by map values:
 - Nearest voxel
 - Interpolated voxel value
 - Selections can be used for purposes other than visualization, scripting, etc.





RDV: Fast, Coarse Map Display

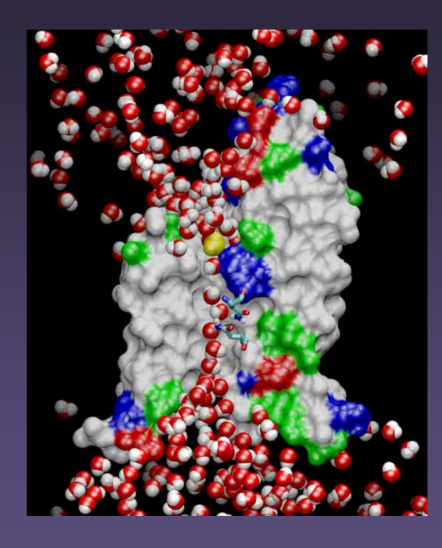
- Shaded points can be rendered very efficiently
- Normals are retrieved from a volume gradient map that VMD generates when maps are loaded
- Effective for dense surfaces
- Even a 3 yr. old laptop can interactively rotate a shaded points isosurface of the 230x230x140 RDV map





Trajectory Animation

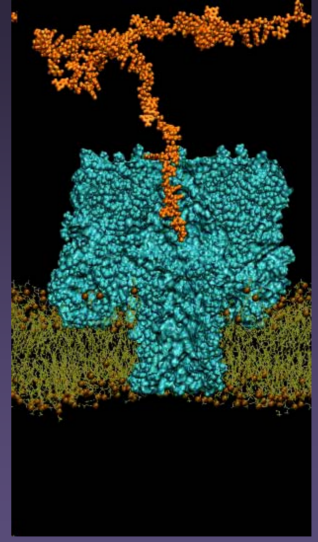
- Motion aids perception of shape, understanding of dynamic processes
- Animate entire model, or just the parts where motion provides insight
- Window-average positions onthe-fly to focus on significant motions
- Selected atoms updated onthe-fly (distance constraints, etc)





Visualization of Large All Atom Molecular Dynamics Simulations (1)

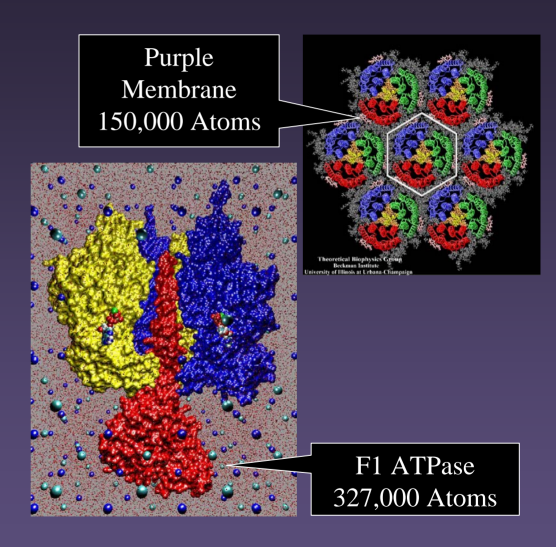
- All-atom models of proteins, membranes, DNA, in water solution
- 100K to 2M atoms
- 512 CPU jobs run on remote supercomputers for weeks at a time for a 10ns simulation
- Visualization and analysis require workstations with 4-32 GB of RAM, 1-4 CPUs, high-end graphics accelerators





Visualization of Large All Atom Molecular Dynamics Simulations (2)

- Multiple representations show areas in appropriate detail
- Large models: 1,00,000 atoms and up
- Long trajectories: thousands of timesteps
- A 10 ns simulation of 100K atoms produces a 12GB trajectory
- Multi-gigabyte data sets break 32-bit addressing barriers



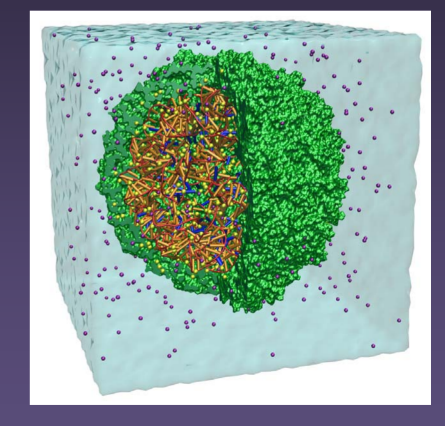


Visualization of Large All Atom Molecular Dynamics Simulations (3)

Satellite Tobacco Mosaic Virus 932,508 atoms



Coarse Representation

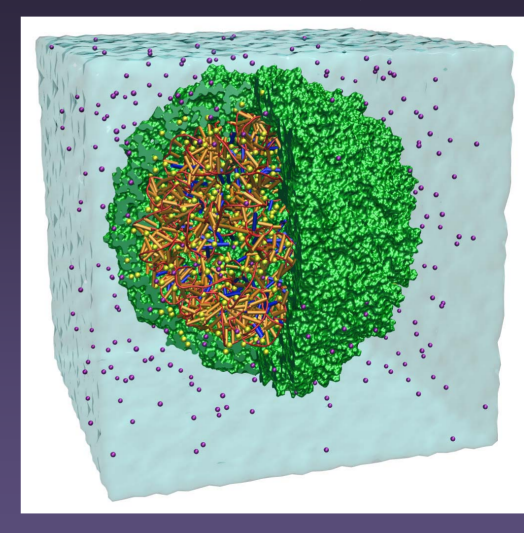




Visualizing Coarse-Grain Simulations

Satellite Tobacco Mosaic Virus, CG Model

- Visualization techniques can be used for both allatom and CG models
- Groups of atoms replaced with "beads", surface reps, or other geometry
- Display 1/20th the data
- No standard file formats for CG simulation trajectories yet, done with scripting currently





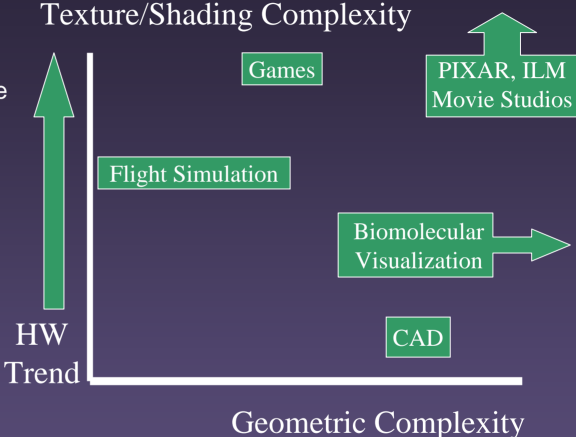
User Interface Issues

- Ease of use is important
- Graphical picking and text-based selection languages need higher level selection keywords to work well with huge complexes
- Viewing huge structures involves more clutter, even with coarse reps, software must do more to help you see what you want to see automatically
- Software needs to know what's "important" at a higher level, much of this information must come from the structure/map files themselves



Comparison of Molecular Visualization with Other Graphics Intensive Applications

- Geometric complexity limits molecular visualization performance
- All atoms move every simulation timestep, thwarts many simplification techniques
- Commodity graphics hardware is tuned for requirements of games
- Solution: Use sophisticated shading instead of geometry where possible





Timeline: Graphics Hardware Used for Molecular Visualization

60's and 70's:

Mainframe-based vector graphics on Tektronix terminals

Evans & Sutherland graphics machines

80's:

Transition to raster graphics on Unix workstations, Mac, PC

Space-filling molecular representations

Stereoscopic rendering

90's - 2002:

3rd-generation raster graphics systems

Depth-cueing

Texture mapping: coloring by potential, density, etc

Full-scene antialiasing



Programmable Graphics Hardware

Groundbreaking research systems:

AT&T Pixel Machine (1989):

82 x DSP32 processors

UNC PixelFlow (1992-98):

64 x (PA-8000 +

8,192 bit-serial SIMD)

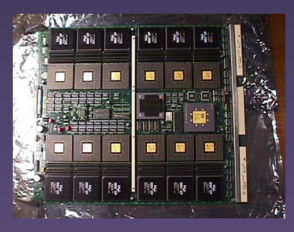
SGI RealityEngine (1990s):

Up to 12 i860-XP processors perform vertex operations (ucode), fixed-func fragment hardware

Most graphics boards now incorporate programmable processors at some level



UNC PixelFlow Rack

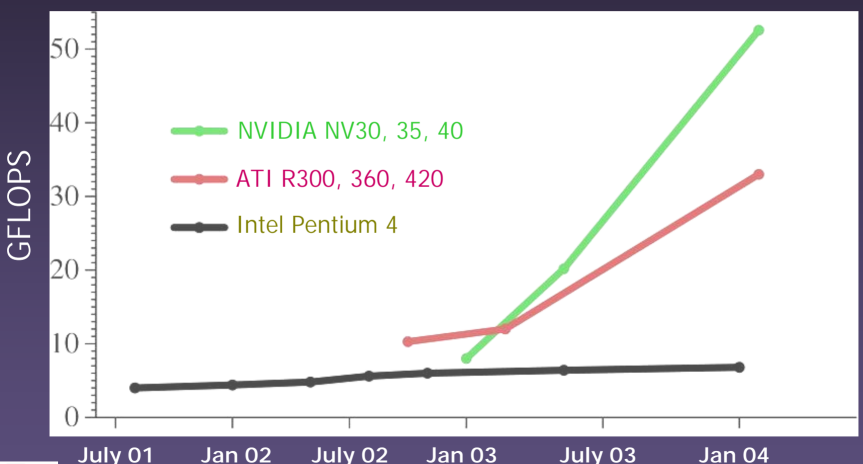


Reality Engine Vertex Processors



GPUs Already Outperformed CPUs for Raw Arithmetic In 2004. The Performance Gap Continues to Widen.....

Floating point multiply-add performance (Data courtesey Ian Buck)

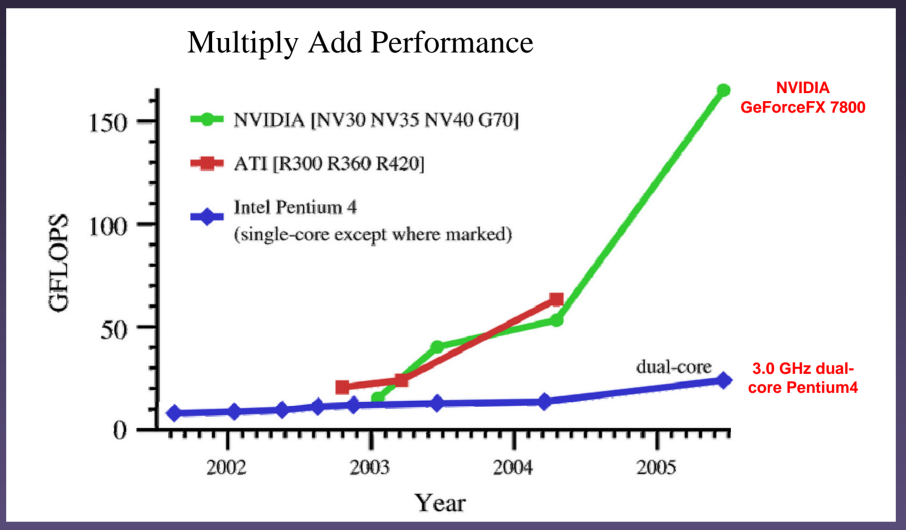




July 01 Jan 02 July 02 Jan 03 July NIH Resource for Macromolecular Modeling and Bioinformatics http://www.ks.uiuc.edu/

Beckman Institute, UIUC

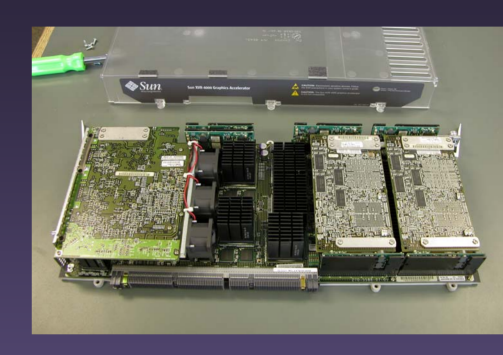
Programmable Shading: Computational Power Enables New Visualization and Analysis Techniques





Early Experiments with Programmable Graphics Hardware in VMD

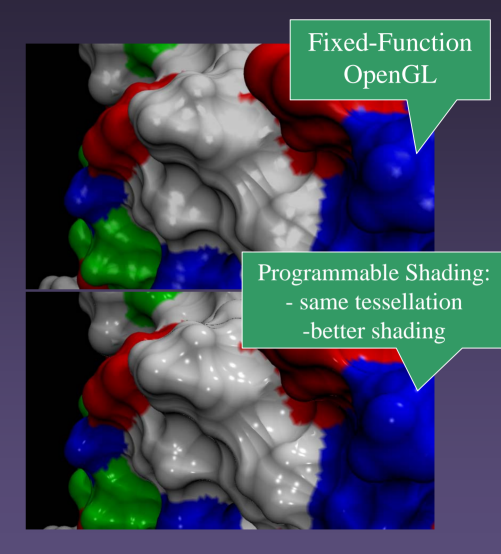
- Sun XVR-1000/4000 (2002)
 - 4xMAJC-5200 CPUs
 - 1GB Texture RAM
 - 32MB ucode RAM
 - 1 Teraflop Antialiasing Filter
 Pipeline
- Custom ucode and OpenGL extension for rendering spheres
 - Draw only half-spheres, with solid side facing the viewer
 - 1-sided lighting
 - Host CPU only sends arrays of radii, positions, colors
 - fast DMA engines copy arrays from system memory to GPU
 - Overall performance twice as fast, host CPU load significantly decreased





Benefits of Programmable Shading (1)

- Potential for superior image quality with better shading algorithms
- Direct rendering of:
 - Quadric surfaces
 - Density map data,
 solvent surfaces
- Offload work from host CPU to GPU



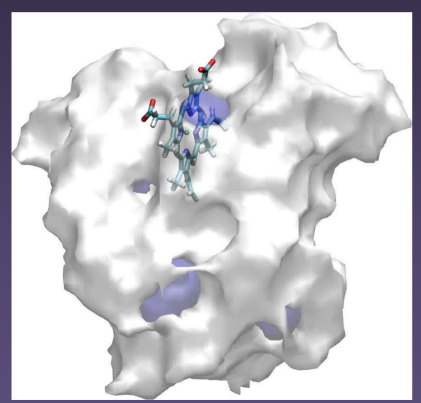


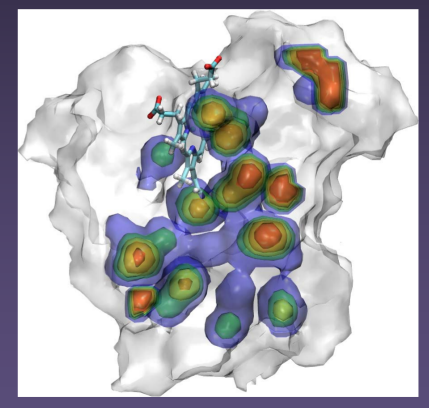
Benefits of Programmable Shading (2)

Myoglobin cavity "openness" (time averaged spatial occupancy)

Single-level OpenGL screen-door transparency obscures internal surfaces

Programmable shading shows transparent nested probability density surfaces with similar performance



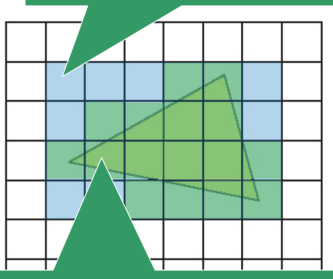




Rendering Non-polygonal Data with Present-day Programmable Shading

- Algorithms mapped to vertex/fragment shading model available in current hardware
- Render by drawing bounding box or a viewer-directed quad containing shape/data
- Vertex shader sets up
- Fragment shader performs all the work

Fragment shader is evaluated for all pixels rasterized by bounding box.

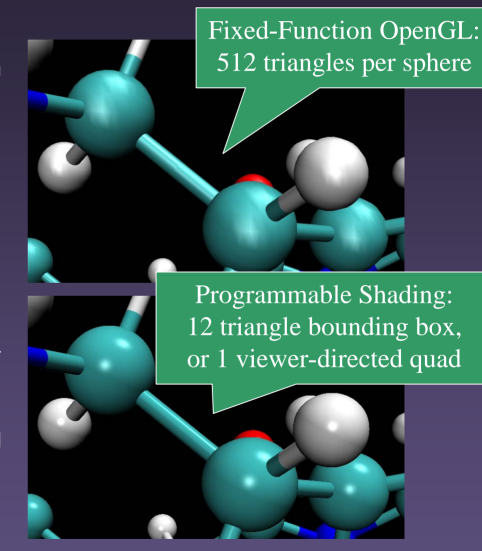


Contained object could be anything one can render in a point-sampled manner (e.g. scanline rendering or ray tracing of voxels, triangles, spheres, cylinders, tori, general quadric surfaces, etc...)



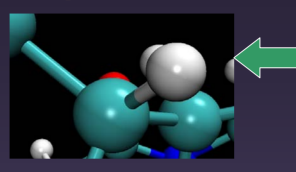
Ray Traced Sphere Rendering with Programmable Shading

- Fixed-function OpenGL requires curved surfaces to be tessellated with triangles, lines, or points
- Fine tessellation required for good results with Gouraud shading; performance suffers
- Static tessellations look bad when one zooms in
- Dynamic tessellation too costly when animating huge trajectories
- Programmable shading solution:
 - Ray trace spheres in fragment shader
 - GPU does all the work
 - Spheres look good at all zoom levels
 - Rendering time is proportional to pixel area covered by sphere
 - Overdraw is a bigger penalty than for triangulated spheres





Sphere Fragment Shader



- Written in OpenGL Shading Language
- High-level C-like language with vector types and operations
- Compiled dynamically by the graphics driver at runtime
- Compiled machine code executes on GPU

```
// VMD Sphere Fragment Shader (not for normal geometry)
void main(void) {
  vec3 raudir = normalize(V):
  vec3 spheredir = spherepos - rayorigin:
  // Perform ray-sphere intersection tests based on the code in Tachyon
  float b = dot(raydir, spheredir);
  float temp = dot(spheredir, spheredir):
  float disc = b*b + sphereradsq - temp;
  // only calculate the meanest intersection, for speed
  if (disc <= 0.0)
    discard; // ray missed sphere entirely, discard fragment
  // calculate closest intersection
  float thear = b - sqrt(disc);
  if (tnear < 0.0)
    discard:
  // calculate hit point and resulting surface normal
  vec3 pnt = rayorigin + thear * raydin;
  vec3 N = normalize(pnt - spherepos);
// Output the ray-sphere intersection point as the fragment depth
  // rather than the depth of the bounding box polygons.
  // The eye coordinate Z value must be transformed to normalized device
  // coordinates before being assigned as the final fragment depth.
  if (vmdprojectionmode == 1) {
    // perspective projection = 0.5 + (hfpn + (f * n / pnt.z)) / diff
    gl_FragDepth = 0.5 + (vmdprojparms[2] + (vmdprojparms[1] * vmdprojparms[0]
31:
  } else {
    // orthographic projection = 0.5 + (-hfpn - pnt.z) / diff
   gl_FragDepth = 0.5 + (-vmdprojparms[2] - pnt.z) / vmdprojparms[3];
#ifdef TEXTURE
  // perform texturing operations for volumetric data
  // The only texturing mode that applies to the sphere shader
```



Efficient 3-D Texturing of Large Datasets

- MIP mapping, compressed map data
- Non-power-of-two 3-D texture dimensions
 - Reduce texture size by a factor of 8 for worst-case (e.g. 2^N-1 dimensions on 3-D potential map)
- Perform volumetric color transfer functions on GPU rather than on the host CPU
 - perform all range clamping and density-to-color mapping on GPU
 - update color transfer function without redownloading large texture maps



Strategies for Working Within Current Hardware Constraints

- GPUs <= 512MB RAM currently
- Use bricked data, multi-level grids, view-dependent map resolution
- Use occlusion culling to prevent rendering of bricks that aren't visible, thus avoiding texture download/access
- Use reduced precision FP types for surface normal / gradient maps



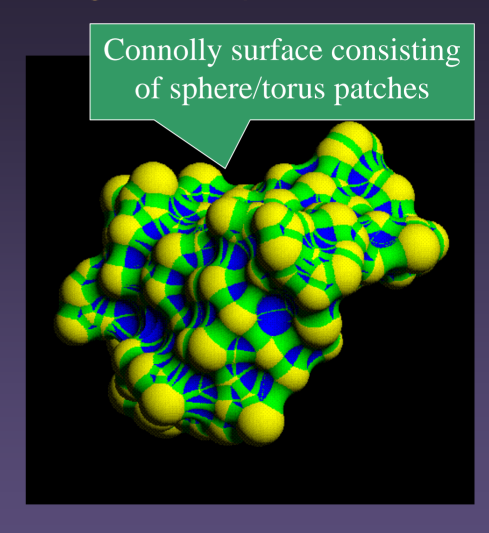
Near Term Possibilities with More Flexible / Powerful GPUs

- Atomic representation tessellation and spline calculations done entirely on GPU
- Direct rendering of isosurfaces from volumetric data via ray casting (e.g. electron density surfaces, demo codes exist already)
- Direct rendering of metaball ("Blob")
 approximation of molecular surfaces via ray casting (demo codes exist already)



The Wheel of Reincarnation: Revival of Old Rendering Techniques?

- Graphics hardware is making another trip around Myer and Sutherland's wheel (CACM '68)
- Visualization techniques that weren't triangle-friendly lost favor in the 90's may return
- Some algorithms that mapped poorly to the OpenGL pipeline are trivial to implement with programmable shading
- Non-polygonal methods get their first shot at running on graphics accelerator hardware rather than the host CPU
 - increased parallelism
 - higher memory bandwidth





Data Structures for Display of 10M Atom Complexes

- Uncompressed atom coordinates120MB (float)
- Avoid traversing per-atom data, hierarchical data structure traversal is a must
- Caching, lazy evaluation, multithreading, overlapped rendering with computation
- Geometry caching, symmetry/instancing accelerate static structure display
- Representation geometry may be 10-50x size of atom coordinate data
- GPU must generate geometry itself, not enough CPU->GPU bandwidth otherwise, particularly for trajectory animation



Next-Gen Graphics Architectures

Short Term:

- "Unlimited" shader instruction count
- Full IEEE floating point pipelines, textures, render targets
- Virtualized texture / render target RAM

Later:

- New programmable pipeline stages: geometry shader, pre-tessellation vertex shader
- Predicated rendering commands, conditions evaluated in hardware (culling operations, etc)



Next-Gen GPUs

- Increased parallelism in GPUs
 - Fragment processors: <u>48-way now</u> (ATI x1900), what next???
 - Multiple boards (NVIDIA "SLI", ATI "Crossfire", etc)
- Double (64-bit) and quad-precision (128-bit) floating point on GPUs
- Improved flexibility in on-GPU data structures, algorithms



Acknowledgements

- NIH NCRR
- UIUC Theoretical and Computational Biophysics Group Members
- Willy Wriggers: SITUS docked GroEL
- J. Frank, E. Villa: docked Ribosome
- Authors of HOLE, MSMS, SITUS, SURF, STAMP, STRIDE, VRPN, and many other freely available packages used in concert with VMD

