

### **GPU Teaching Kit**

**Accelerated Computing** 



Module 16 - Application Case Study - Electrostatic Potential Calculation

Lecture 16.1 - Electrostatic Potential Calculation - Part 1

# Objective

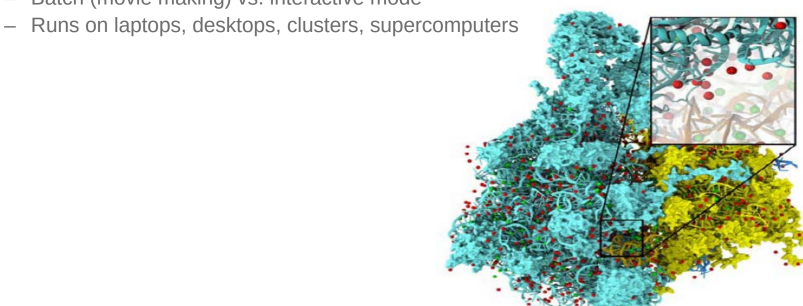
- To learn how to apply parallel programming techniques to an application
  - Thread coarsening for more work efficiency
  - Data structure padding for reduced divergence
  - Memory access locality and pre-computation techniques

## **VMD**

#### Visual Molecular Dynamics

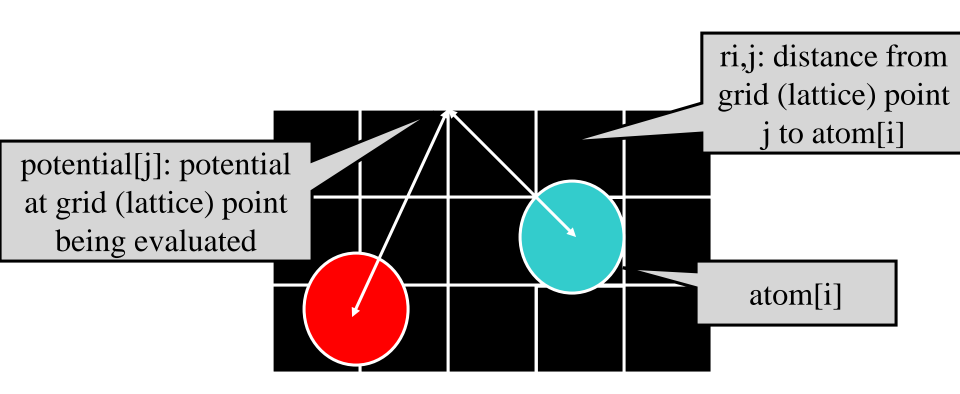
- Visualizing, animating, and analyzing bio-molecular systems
- More than 200,000 users worldwide

Batch (movie making) vs. interactive mode



# Electrostatic Potential Map

- Calculate initial electrostatic potential map around the simulated structure considering the contributions of all atoms
  - Most time consuming, focus of our example.



## **Electrostatic Potential Calculation**

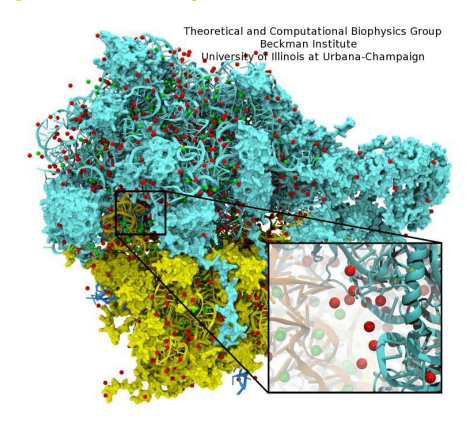
- The contribution of atom[i] to the electrostatic potential at lattice[j] is potential[j] = atom[i].charge / rij.
- In the Direct Coulomb Summation method, the total potential at lattice point j is the sum of contributions from all atoms in the system.

# Overview of Direct Coulomb Summation (DCS) Algorithm

- One way to compute the electrostatic potentials on a grid, ideally suited for the GPU
  - All atoms affect all map lattice points, most accurate
- For each lattice point, sum potential contributions for all atoms in the simulated structure:
  - potential += charge[i] / (distance to atom[i])
- Approximation-based methods such as cut-off summation can achieve much higher performance at the cost of some numerical accuracy and flexibility

# Irregular Input vs. Regular Output

- Atoms come from modeled molecular structures, solvent (water) and ions
  - Irregular by necessity
- Energy grid models the electrostatic potential value at regularly spaced points
  - Regular by design



# Summary of Sequential C Version

- Algorithm is input oriented
  - For each input atom, calculate its contribution to all grid points in an x-y slice
- Output (energy grid) is regular
  - Simple linear mapping between grid point indices and modeled physical coordinates
- Input (atom) is irregular
  - Modeled x,y,z coordinate of each atom needs to be stored in the atom array
- The algorithm is efficient in performing minimal number of calculations on distances, coordinates, etc.

# A Sequential C Version

```
void cenergy(float *energygrid, dim3 grid, float gridspacing, float z, const float
*atoms, int numatoms)
```

```
int atomarrdim = numatoms * 4; //x,y,z, and charge info for each atom
  int grid_slice_offset = (grid.x*grid.y*z) / gridspacing;
for (int n=0; n<atomarrdim; n+=4) { // calculate potential contribution of each atom
    float dz = z - atoms[n+2]; // all grid points in a slice have the same z value
    float dz^2 = dz^*dz^*
   float charge = atoms[n+3];
    for (int j=0; j<grid.y; j++) {</pre>
     float y = gridspacing * (float) j;
     float dy = y - atoms[n+1]; // all grid points in a row have the same y value
     float dy2 = dy*dy;
     int grid_row_offset = grid_slice_offset+ grid.x*j;
     for (int i=0; i<grid.x; i++) {
        float x = gridspacing * (float) i;
        float dx = x - atoms[n];
        energygrid[grid_row_offset + i] += charge / sqrtf(dx*dx + dy2+ dz2);
                                        The grid parameter gives the
```

number of grid points in each dimension of the lattice.

# A Sequential C Version

```
void cenergy(float *energygrid, dim3 grid, float gridspacing, float z, const float
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 int atomarrdim = numatoms * 4; //x,y,z, and charge info for each atom
  int grid_slice_offset = (grid.x*grid.y*z) / gridspacing;
for (int n=0; n<atomarrdim; n+=4) { // calculate potential contribution of each atom
   float dz = z - atoms[n+2]; // all grid points in a slice have the same z value
   float dz^2 = dz^*dz^*
   float charge = atoms[n+3];
   for (int j=0; j<grid.y; j++) {</pre>
     float y = gridspacing * (float) j;
     float dy = y - atoms[n+1]; // all grid points in a row have the same y value
     float dy2 = dy*dy;
     int grid_row_offset = grid_slice_offset+ grid.x*j;
     for (int i=0; i<grid.x; i++) {
        float x = gridspacing * (float) i;
        float dx = x - atoms[n];
        energygrid[grid_row_offset + i] += charge / sqrtf(dx*dx + dy2+ dz2);
                                        The grid parameter gives the
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                                        dimension of the lattice.
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# A Sequential C Version

```
void cenergy(float *energygrid, dim3 grid, float gridspacing, float z, const float
*atoms, int numatoms) {
  int atomarrdim = numatoms * 4; //x,y,z, and charge info for each atom
                                                                       Input oriented
  int grid_slice_offset = (grid.x*grid.y*z) / gridspacing;
  for (int n=0; n<atomarrdim; n+=4) { // calculate potential contribution of each atom
    float dz = z - atoms[n+2]; // all grid points in a slice have the same z value
   float dz2 = dz*dz;
   float charge = atoms[n+3];
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     float y = gridspacing * (float) j;
     float dy = y - atoms[n+1]; // all grid points in a row have the same y value
     float dy2 = dy*dy;
     int grid_row_offset = grid_slice_offset+ grid.x*j;
     for (int i=0; i<grid.x; i++) {
        float x = gridspacing * (float) i;
        float dx = x - atoms[n];
        energygrid[grid_row_offset + i] += charge / sqrtf(dx*dx + dy2+ dz2);
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                                        dimension of the lattice.
```

## An Sequential C Version

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void cenergy(float *energygrid, dim3 grid, float gridspacing, float z, const float
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    float dz = z - atoms[n+2]; // all grid points in a slice have the same z value
    float dz2 = dz*dz;
    float charge = atoms[n+3];
    for (int j=0; j<grid.y; j++) {</pre>
     float y = gridspacing * (float) j;
     float dy = y - atoms[n+1]; // all grid points in a row have the same y value
     float dy2 = dy*dy;
      int grid_row_offset = grid_slice_offset+ grid.x*j;
      for (int i=0; i<grid.x; i++) {
         float x = gridspacing * (float) i;
         float dx = x - atoms[n]
         energygrid[grid_row_offset + i] += charge / sqrtf(dx*dx + dy2+ dz2);
```

# A More Optimized Sequential C Version

```
void cenergy(float *energygrid, dim3 grid, float gridspacing, float z, const float
   *atoms, int numatoms) {
 int grid_slice_offset = (grid.x*grid.y*z) / gridspacing;
  int atomarrdim = numatoms * 4; //x,y,z, and charge info for each atom
  for (int n=0; n<atomarrdim; n+=4) { // calculate potential contribution of each
   at.om
   float dz = z - atoms[n+2]; // all grid points in a slice have the same z value
    float dz2 = dz*dz;
   float charge = atoms[n+3];
   for (int j=0; j < grid.y; j++)
      float y = gridspacing * (float) j;
      float dy = y - atoms[n+1]; // all grid points in a row have the same y value
      float dy2 = dy*dy;
      int grid row offset = grid slice offset+ grid.x*j;
      for (int i=0; i<grid.x; i++) {
         float x = gridspacing * (float) i;
         float dx = x - atoms[n] ];
         energygrid[grid_row_offset + i] += charge / sqrtf(dx*dx + dy2+ dz2);
```

# An Intuitive Sequential C Version

```
void cenergy(float *energygrid, dim3 grid, float gridspacing, float z, const float
   *atoms, int numatoms) {
   int grid slice offset = (grid.x*grid.y*z) / gridspacing;
   int atomarrdim = numatoms * 4; //x,y,z, and charge info for each atom
  for (int n=0; n<atomarrdim; n+=4) { // calculate potential contribution of each
   atom
   float dz = z - atoms[n+2]; // all grid points in a slice have the same z value
    float dz2 = dz*dz;
   float charge = atoms[n+3];
    for (int j=0; j < grid.y; j++) {
     float y = gridspacing * (float) j;
      float dy = y - atoms[n+1]; // all grid points in a row have the same y value
     float dy2 = dy*dy;
      int grid row offset = grid slice offset+ grid.x*j;
      for (int i=0; i<grid.x; i++) {</pre>
         float x = gridspacing * (float) i;
        float dx = x - atoms[n];
        energygrid[grid row offset + i] += charge / sgrtf(dx*dx + dy2+ dz2);
```

# **CUDA DCS Implementation Overview**

- Allocate and initialize potential map memory on host CPU
- Allocate potential map slice buffer on GPU
- Preprocess atom coordinates and charges
- Loop over potential map slices:
  - Copy potential map slice from host to GPU
  - Loop over groups of atoms:
    - Copy atom data to GPU
    - Run CUDA Kernel on atoms and potential map slice on GPU
  - Copy potential map slice from GPU to host
- Free resources



# Straightforward CUDA Parallelization

- Use each thread to compute the contribution of an atom to all grid points in the current slice
  - Scatter parallelization
- Kernel code largely correspond to intuitive CPU version with outer loop stripped
  - Each thread corresponds to an outer loop iteration of CPU version
  - numatoms used in kernel launch configuration of the host code

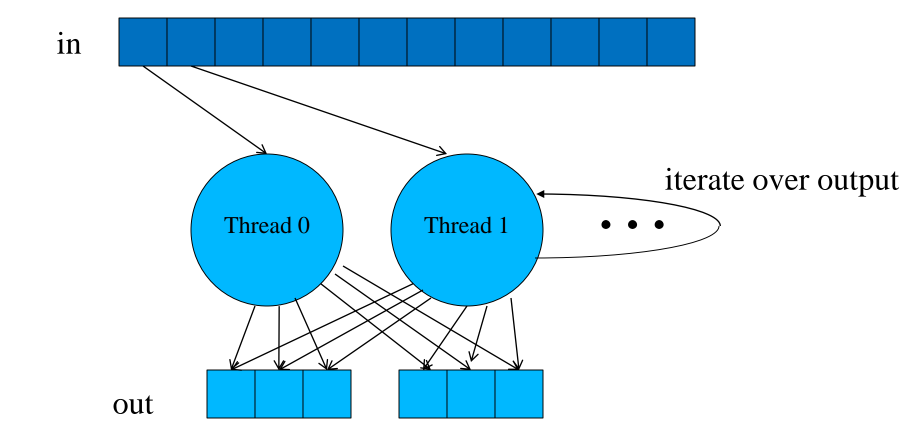
# A Very Slow DCS Scatter Kernel!

```
void global cenergy(float *energygrid, float *atoms, dim3 grid, float gridspacing,
float z) {
   int n = (blockIdx.x * blockDim .x + threadIdx.x) * 4;
   float dz = z - atoms[n+2]; // all grid points in a slice have the same z
   float dz2 = dz*dz;
   int grid_slice_offset = (grid.x*grid.y*z) / gridspacing;
   float charge = atoms[n+3];
                                                          Needs to be calculated
   for (int j=0; j<grid.y; j++) {
                                                          redundantly by every thread
     float y = gridspacing * (float) j;
     float dy = y - atoms[n+1]; // all grid points in a row have the same y value
     float dy2 = dy*dy;
     int grid_row_offset = grid_slice_offset+ grid.x*j;
     for (int i=0; i<grid.x; i++) {
        float x = gridspacing * (float) i;
        float dx = x - atoms[n];
        energygrid[grid_row_offset + i] += charge / sqrtf(dx*dx + dy2+ dz2));
```

# A Very Slow DCS Scatter Kernel!

```
void global _ cenergy(float *energygrid, float *atoms, dim3
grid, float gridspacing, float z) {
    int n = (blockIdx.x * blockDim .x + threadIdx.x) *4;
    float dz = z - atoms[n+2]; // all grid points in a slice have
the same z value
    float dz^2 = dz^*dz^*
    int grid slice offset = (grid.x*grid.y*z) / gridspacing;
    float charge = atoms[n+3];
    for (int j=0; j<qrid.y; j++) {
      float y = gridspacing * (float) j;
      float dy = y - atoms[n+1]; // all grid points in a row have
the same y value
      float dy2 = dy*dy;
      int grid_row_offset = grid_slice_offset+ grid.x*j;
      for (int i=0; i<grid.x; i++) {
                                               Needs to be done as
         float x = gridspacing * (float) i;
                                               an atomic operation
         float dx = x - atoms[n];
         energygrid[grid row offset + i] += charge / sgrtf(dx*dx
+ dy2 + dz2));
                                                        18
```

## **Scatter Parallelization**



#### Why is input oriented algorithm often used?

- In practice, each in element does not have significant effect on all out elements
- Output tends to be much more regular than input
  - Input usually comes as sparse data structure, where coordinates are part of the data
  - One needs to look at the input data to see if an input is relevant to an output value
  - Output is usually a regular, grid
  - Given an input value, one can easily find output via index calculation

## Pros and Cons of the Scatter Kernel

#### - Pros

- Follows closely the simple CPU version
- Good for software engineering and code maintenance
- Preserves computation efficiency (coordinates, distances, offsets) of sequential code

#### Cons

- The atomic add serializes the execution, very slow!
- Not even worth trying this.



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