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## Administrivia

- Reminder: Homework 1 due today (Java part).

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## A Few Words About Design Patterns

- Title of our book includes the word "patterns".
- What do we mean? "Design patterns".

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### A Few (More) Words About Design Patterns

- Idea originated with architect Christopher Alexander (first book 1977). Briefly — look for problems that have to be solved over and over, and try to come up with “expert” solution, write it in a form accessible to others. Usually this means adopting “pattern format” to use for all patterns. Characteristics of a good pattern:
  - Neat balancing of competing “forces” (tradeoffs).
  - Name either tells you what it’s about, or is a good addition to vocabulary.
  - “Aha!” aspect.
- First used in CS in OOD/OOP, about 1987. Really started to take off in OO community with “Gang of Four” book (Gamma, Helms, Johnson, and Vlissides; 1995). Now can find people writing patterns in many, many areas.
- Simple low-level example — iterator.

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### “A Pattern Language for Parallel Programming”?

- Goal of our book (and preceding work) — apply this idea in parallel computing.
- We started out looking for patterns representing high-level structures for parallel programs, thinking there might be a dozen of them.
- At some point we realized we also wanted to talk about how you get from the original problem to one of these structures — i.e., how do expert parallel programmers think about how to decompose a problem, etc.? and also about commonly-occurring data structures and program structures, and how to map high-level designs/structures into real programming environments.
- After much thought and discussion . . .

### “A Pattern Language for Parallel Programming”, Continued

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- Eventually — four-layer “pattern language”. (Notice that “pattern language” connotes common vocabulary more than grammatical structure. Not a programming language!)
- Currently being revised/extended, primarily by Mattson and a group at UC Berkeley.

### Overall Organization of Our Pattern Language

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- Four “design spaces” corresponding to phases in design.
  - *Finding Concurrency* — how to decompose problems, analyze decomposition.
  - *Algorithm Structure* — high-level program structures.
  - *Supporting Structures* — program structures, data structures.
  - *Implementation Mechanisms* — generic discussion of programming environment “building blocks”.
- Idea is that you start at the top, work your way down, possibly with some backtracking.

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### *Finding Concurrency — Preview*

- Decomposition patterns (*Task Decomposition, Data Decomposition*): Break problem into tasks that maybe can execute concurrently.
- Dependency analysis patterns (*Group Tasks, Order Tasks, Data Sharing*): Organize tasks into groups, analyze dependencies among them.
- *Design Evaluation*: Review what you have so far, possibly backtrack.

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### *Algorithm Structure — Preview*

- *Task Parallelism* — decompose problem into lots of tasks, independent or nearly so. Example: numerical integration.
- *Divide and Conquer* — decompose recursively as in divide-and-conquer algorithms. Examples: quicksort, mergesort.
- *Geometric Decomposition* — decompose based on data (by rows, by columns, etc.). Example: Mesh-based computation.
- *Recursive Data* — rethink computation to expose unexpected concurrency. Ignore for now.
- *Pipeline* — decompose based on assembly-line analogy.
- *Event-Based Coordination* — decompose problem into entities interacting asynchronously.

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### *Supporting Structures — Preview*

- Program structure patterns:
  - *SPMD* (Single Program, Multiple Data) — “like an MPI program”.
  - *Loop Parallelism* — “like an OpenMP program”.
  - *Master/Worker* — like the name suggests.
  - *Fork/Join* — when none of the others fits.
- Data structure patterns:
  - *Shared Data* — generic advice for dealing with data dependencies.
  - *Shared Queue* — example of applying *Shared Data*.
  - *Distributed Array*.

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### *Implementation Mechanisms — Preview*

- Generic discussion of “building blocks” for parallel programming — analogous to assignment, if/then/else, loops in procedural programming languages. (Can think of this as “what basic questions do I ask about a new parallel programming environment?”)
- Three basic categories:
  - UE management.
  - Synchronization.
  - Communication.

### Example Applications

- Before starting on *Finding Concurrency* patterns — two example applications to be used as running examples.

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### Example — Molecular Dynamics

- Goal is to simulate what happens to large molecule. Of interest, e.g., in modeling how a drug interacts with a protein.
- Approach is to treat molecule as a collection of balls (atoms) connected by springs (chemical bonds). Then do “standard time-stepping” — divide time into discrete steps, and at each step use classical mechanics to figure out new positions for atoms based on current positions and forces among them.  
In more details . . .

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## Molecular Dynamics — Computation

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- At each time step:
  - Compute forces (vibrational and rotational) on atoms caused by chemical bonds between them. Short-range interaction, so not too much computation here.
  - Compute forces on atoms caused by their electrical charges. Potentially must consider all pairs of atoms, so lots of computation here.
  - Use forces to update atoms' positions and velocities.
  - Compute other physical properties of the system — e.g., energies.
- To reduce the computational load, can limit computation of electrical-charge-induced forces to atoms that are “close”. To do this, calculate for each atom a list of “neighbors”. If time steps are short, atoms don't move much, and we don't have to do this every step.

## Molecular Dynamics Pseudocode

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```

Int const N    // number of atoms
Array of Real :: atoms   (3,N) //3D coordinates
Array of Real :: velocities (3,N) //velocity vector
Array of Real :: forces  (3,N) //force in each dimension
Array of List  :: neighbors(N) //atoms in cutoff volume

loop over time steps
  vibrational_forces (N, atoms, forces)
  rotational_forces (N, atoms, forces)
  neighbor_list (N, atoms, neighbors)
  non_bonded_forces (N, atoms, neighbors, forces)
  update_atom_positions_and_velocities
    (N, atoms, velocities, forces)
  physical_properties ( ... Lots of stuff ... )
end loop

```

### Pseudocode for Non-Bonded Force Computation

```
function non_bonded_forces (N, Atoms, neighbors, Forces)
  Int const N // number of atoms
  Array of Real :: atoms (3,N) //3D coordinates
  Array of Real :: forces (3,N) //force in each dimension
  Array of List :: neighbors(N) //atoms in cutoff volume
  Real :: forceX, forceY, forceZ

  loop [i] over atoms
    loop [j] over neighbors(i)
      forceX = non_bond_force(atoms(1,i), atoms(1,j))
      forceY = non_bond_force(atoms(2,i), atoms(2,j))
      forceZ = non_bond_force(atoms(3,i), atoms(3,j))
      force(1,i) += forceX;   force(1,j) -= forceX;
      force(2,i) += forceY;   force(2,j) -= forceY;
      force(3,i) += forceZ;   force(3,j) -= forceZ;
    end loop [j]
  end loop [i]
end function non_bonded_forces
```

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### Example — Heat Diffusion

- A simple example, representative of a big class of scientific-computing applications — “heat distribution problem”.
- Goal is to simulate what happens when two ends of a pipe are put in contact with things at different (constant) temperatures — pipe conducts heat, its temperature changes over time, eventually converging on a smooth gradient.
- Can model mathematically how temperature in pipe changes over time using partial differential equations.
- Can approximate solution by “discretizing” — spatially and with regard to time.

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## Heat Diffusion Code

```
double *uk = malloc(sizeof(double) * NX);
double *ukpl = malloc(sizeof(double) * NX);
double *temp;
double dx = 1.0/NX; double dt = 0.5*dx*dx;
double maxdiff, diff;

initialize(uk, ukpl);

for (int k = 0; (k < NSTEPS) && (maxdiff >= threshold); ++k) {

    /* compute new values */
    for (int i = 1; i < NX-1; ++i) {
        ukpl[i]=uk[i]+ (dt/(dx*dx))*(uk[i+1]-2*uk[i]+uk[i-1]);
    }

    /* check for convergence */
    maxdiff = 0.0;
    for (int i = 1; i < NX-1; ++i) {
        diff = fabs(uk[i] - ukpl[i]);
        if (diff > maxdiff) maxdiff = diff;
    }

    /* "copy" ukpl to uk by swapping pointers */
    temp = ukpl; ukpl = uk; uk = temp;

    printValues(uk, k);
}
```

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## Minute Essay

- What has been most interesting about Homework 1? most difficult?

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