All About Species, Atoms, and Iterators

David A. Kofke

Department of Chemical Engineering University at Buffalo, State University of New York



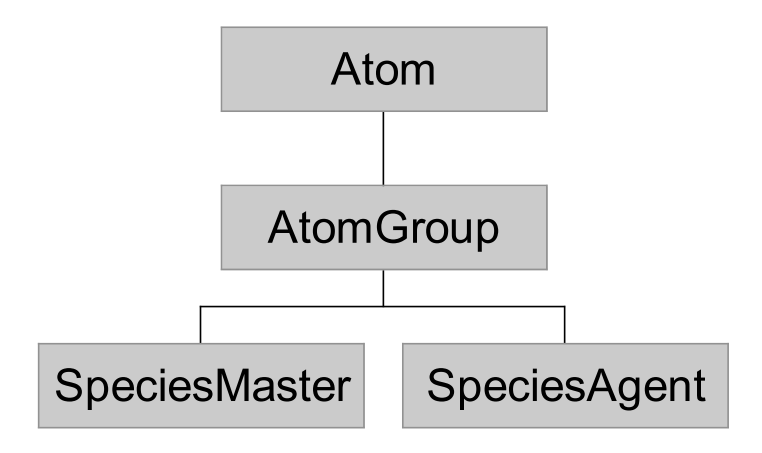


Species

- Top-level simulation element
 - Species, Potential, Integrator, Controller, Meter, Device, Display
- Defines methods and holds fields to specify, build, and arrange molecules/atoms
- Employs several lower-level classes
 - SpeciesMaster, SpeciesAgent
 - Atom, AtomGroup
 - AtomFactory
 - AtomType, Space.Coordinate, Parameter Bond, BondInitializer
 - Configuration

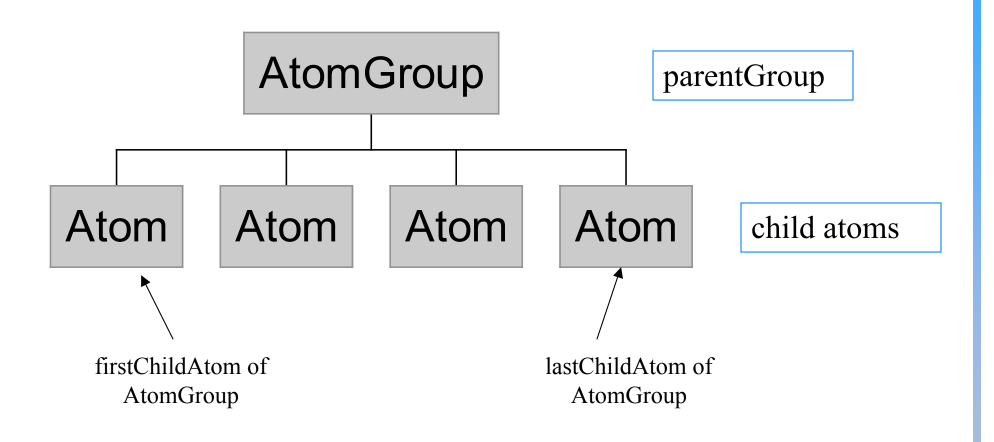


Inheritance Hierarchy



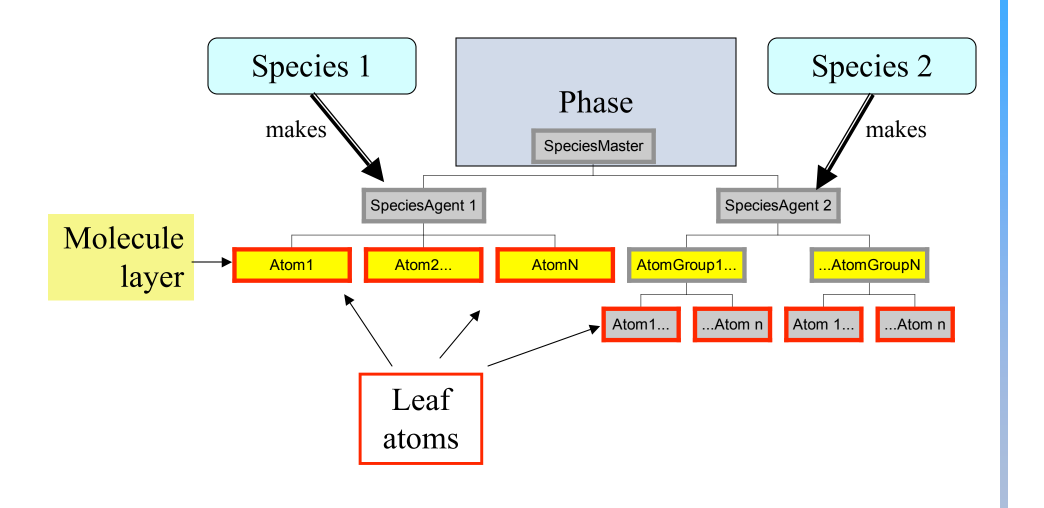


Containment



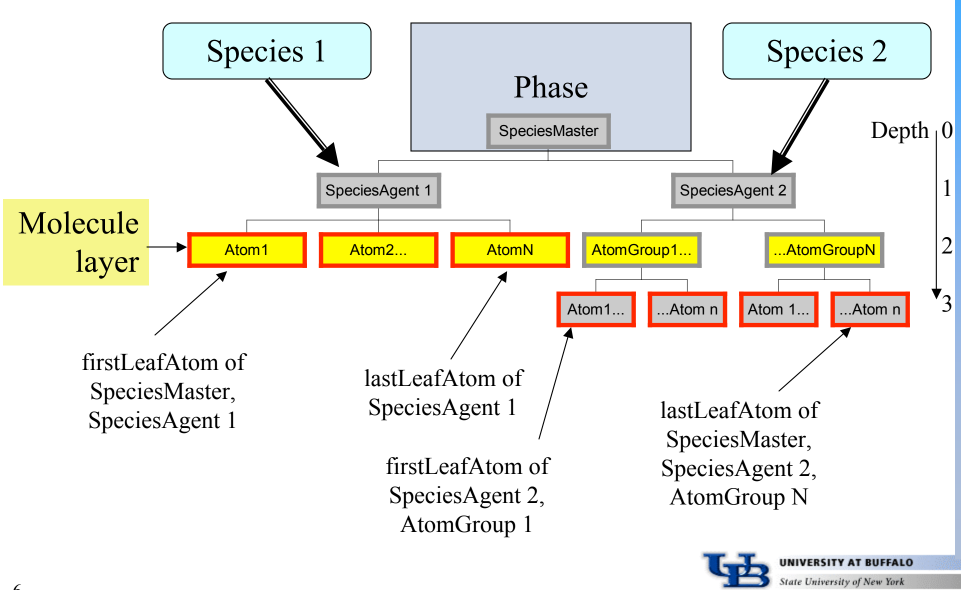


Containment Hierarchy



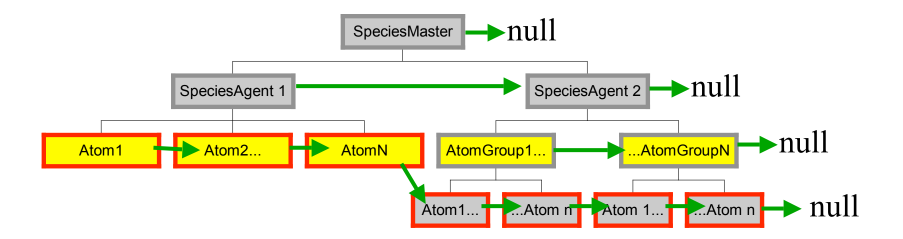


Containment Hierarchy



Sequencing

- Atoms are sequenced to facilitate looping through them
 - All leaf atoms in phase are sequenced together
 - Otherwise, only siblings are sequenced



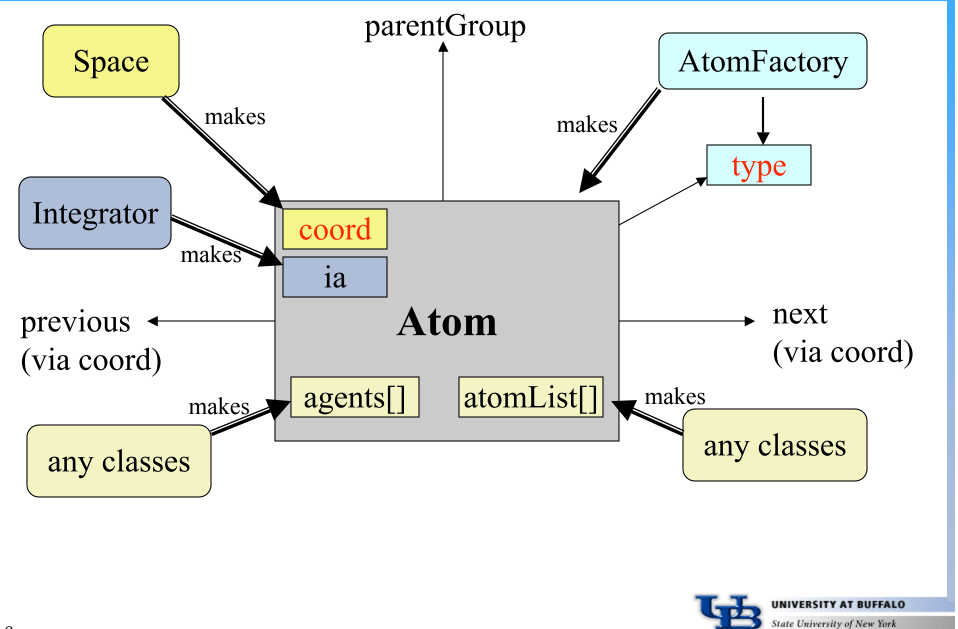


Atom

- Corresponds to a physical atom (in simplest case)
- Holds data relating to
 - Spatial coordinate (position and/or momentum)
 - Atom type
 - Position in atom hierarchy
 - Position in atom sequence
- Can also hold auxiliary data
 - Agents from other classes
 - Parameters specified by other classes (in type field)
 - Lists of other atoms for use by other classes



Atom Fields



Atom Methods

- Sequence
 - nextAtom; previousAtom
 - preceeds; index
 - clearPreviousAtom, setNextAtom
- Hierarchy
 - parentGroup, parentMolecule, parentSpeciesAgent; isDescendedFrom
 - firstChildAtom; lastChildAtom; firstLeafAtom; lastLeafAtom
 - All return this if not an AtomGroup
 - depth, leafAtomCount; signature
 - setParentGroup
- Type
 - type, creator
- Simulation
 - parentPhase; parentSimulation; parentSpecies; requestAtomListIndex

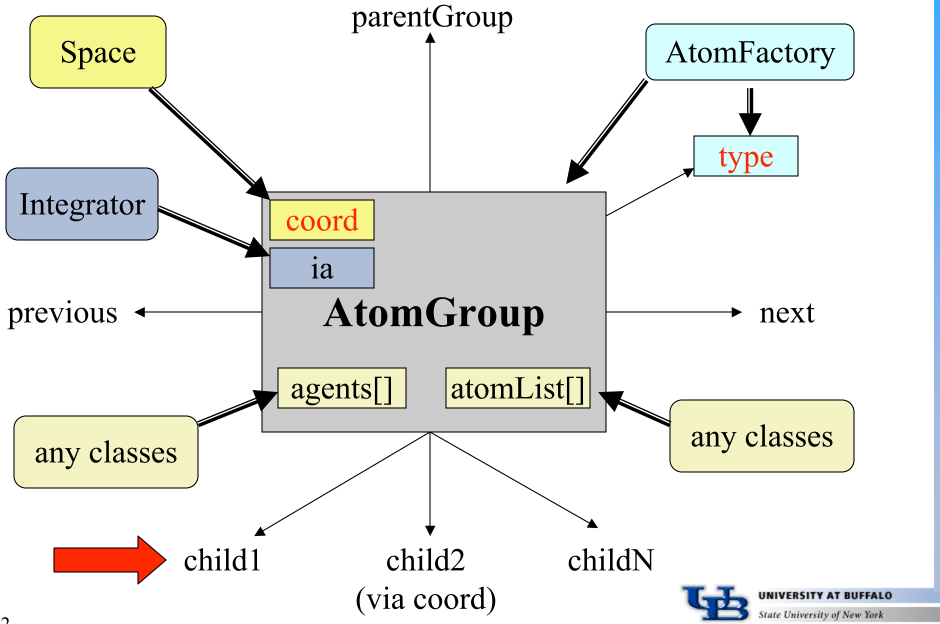


AtomGroup

- Extends Atom
 - Inherits all fields given to an atom
 - Uses CoordinateGroup for Coordinate
 - e.g. translateTo method relays command to all child atoms' coordinates
- Defines a collection of atoms or other atom groups
 - Molecule
 - Atom subgroup, e.g., CH₂, amino acid, polymer segment, etc.
- Has methods to add, remove, count contained atoms
 - addAtom; removeAtom; removeAll
 - addAtomNotify; removeAtomNotify
 - childAtomArray; getAtom; randomAtom
 - firstChildAtom; lastChildAtom
 - firstLeafAtom; lastLeafAtom
 - childAtomCount; leafAtomCount



Fields in an AtomGroup



AtomFactory

- Constructs an Atom (or AtomGroup)
 - Refers to other atom factories to build group from subgroups
- Uses a BondInitializer to attach "bonds" to atom pairs
 - Bonds may be used by potentials or displays
- Uses a Configuration to put the atoms in some arrangement
- Reservoir can accept atoms and hold them for reuse to avoid repeated construction
 - Grand-canonical simulations
- Important methods
 - makeAtom() takes atom from reservoir, or builds new one if empty
 - build() defines how to make a new atom
 - set/get methods for bondInitializer and configuration
 - requestAgentIndex() gives integer for index of placement of an agent



Basic AtomFactory Classes

AtomFactoryMono

- Produces a single atom
- setType method specifies AtomType instance to be referenced by all Atoms made by factory

AtomFactoryHomo

- Produces AtomGroup composed of arbitrary number of identical atoms
- Constructor:
 - AtomFactoryHomo(Space, AtomFactory, int (nAtoms), BondInitializer,Configuration)
- AtomFactoryHetero
 - Produces AtomGroup composed of non-identical atoms
 - AtomFactoryHetero(Space, AtomFactory[], Configuration)



More Complex Atom Factories

• Other atom factories can be built up from the basic ones

```
//red atoms
AtomFactoryMono atomFactoryMono0 = new AtomFactoryMono(space);
AtomType type0 = new AtomType.Sphere(atomFactoryMono0, Default.ATOM MASS, Default.ATOM SIZE);
atomFactoryMono0.setType(type0);
    //black atoms
AtomFactoryMono atomFactoryMono1 = new AtomFactoryMono(space);
AtomType type1 = new AtomType.Sphere(atomFactoryMono1, Default.ATOM MASS, Default.ATOM SIZE);
atomFactoryMono1.setType(type1);
    //red segment
atomFactoryHomo0 = new AtomFactoryHomo(space,atomFactoryMono0,3,new BondInitializerChain(),
                                        new ConfigurationLinear(space));
    //black segment
atomFactoryHomo1 = new AtomFactoryHomo(space,atomFactoryMono1,3,new BondInitializerChain(),
                                        new ConfigurationLinear(space));
    //molecule
AtomFactoryHetero atomFactoryHetero = new AtomFactoryHetero(space,
                              new AtomFactory[] {atomFactoryHomo0,atomFactoryHomo1},
                              new MyBondInitializer(),
                              new ConfigurationLinear(space,1.25*Default.ATOM SIZE));
speciesSpheres2 = new Species(this,atomFactoryHetero);
```





More Complex Atom Factories

• Other atom factories can be built up from the basic ones

```
//red atoms
AtomFactoryMono atomFactoryMono0 = new AtomFactoryMono(space);
AtomType type0 = new AtomType.Sphere(atomFactoryMono0, Default.ATOM MASS, Default.ATOM SIZE);
atomFactoryMono0.setType(type0);
    //black atoms
AtomFactoryMono atomFactoryMono1 = new AtomFactoryMono(space);
AtomType type1 = new AtomType.Sphere(atomFactoryMono1, Default.ATOM MASS, Default.ATOM SIZE);
atomFactoryMono1.setType(type1);
    //red segment
atomFactoryHomo0 = new AtomFactoryHomo(space,atomFactoryMono0,3,new BondInitializerChain(),
                                        new ConfigurationLinear(space));
atomFactoryHomo1 = new AtomFactoryHomo(space,atomFactoryMono1,3,new BondInitializerChain(),
                                        new ConfigurationLinear(space));
AtomFactoryHetero atomFactoryHetero = new AtomFactoryHetero(space,
                              new AtomFactory[] {atomFactoryHomo0,atomFactoryHomo1},
                              new MyBondInitializer(),
                              new ConfigurationLinear(space,1.25*Default.ATOM SIZE));
speciesSpheres2 = new Species(this,atomFactoryHetero);
   //another approach
AtomFactoryHetero atomFactoryMolecule = new AtomFactoryHetero(space,
          new AtomFactory[] {atomFactoryMono0, atomFactoryMono0, atomFactoryMono0,
                              atomFactoryMono1, atomFactoryMono1, atomFactoryMono1} ...);
```



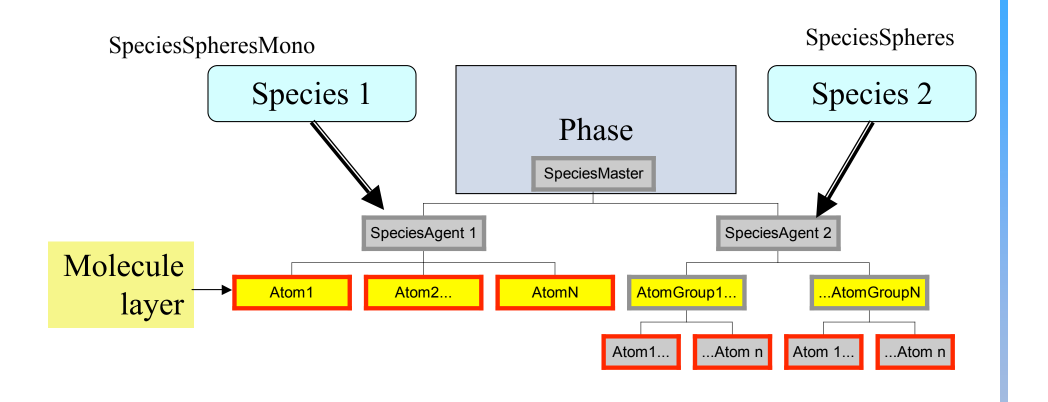


Species Definition

- Species differ in the AtomFactory they use
 - Species constructor (the only one)
 - Species(Space, AtomFactory)
- A few species are pre-defined
 - SpeciesSpheres
 - Multiatomic molecules
 - SpeciesSpheres(Simulation, int nM, int nA, BondInitializer, Configuration)
 - SpeciesSpheresMono
 - Molecules are monatomic (no AtomGroup layer)
 - SpeciesSpheresMono(Simulation, int nM)
 - SpeciesWalls, SpeciesSphereWells, SpeciesSpheresRotating, etc.
- Extensions should define particular chemical compounds via AtomFactory
 - new Species(space, atomFactoryWater) instead of new SpeciesWater



Spheres vs. Spheres Mono





Species Methods

- public void allAgents(AtomAction action);
 - Performs given action on all agents of this species in all phases
- public void allAtoms(AtomAction action);
 - Performs given action on all leaf atoms of this species in all phases
- public void allMolecules(AtomAction action);
 - Performs given action on all molecules of this species in all phases
- public SpeciesAgent getAgent(Phase p);
 - Returns agent of this species in the given phase
- public void setNMolecules(int nM);
 - Performs setNMolecules on all agents of this species in all phases
- makeAgent(SpeciesMaster); makeMolecule();



Iteration

- Common simulation activities involve looping through atoms and atom pairs, calculating properties based on their positions and separations
- Iterators are used to facilitate the looping



AtomIterator Interface

- public boolean hasNext();
 - true if iterator has another atom to give
- public boolean contains(Atom atom);
 - true if given atom or its parents would among this iterator's iterates
- public Atom next();
 - returns the next iterate
- public Atom reset();
 - resets iterator to be ready to loop over all its iterates
- public Atom reset(IteratorDirective id);
 - resets iterator to be ready to loop over iterates consistent with the directive
- public void setBasis(Atom atom);
 - defines the basic set of atoms given by this iterate
- public int size();
 - number of iterates given by this iterator
- public void setAsNeighbor(boolean b);
 - more on this later



Using an Atom Iterator

Perform actions on iterates (pass atoms to action)

```
iterator.reset();
while(iterator.hasNext()) {
    Atom atom = iterator.next();
    //do something with atom
}
```

• Also could pass action to iterator, but not fully implemented

```
AtomAction action = ...//define action
iterator.allAtoms(action);
   //performs action on all atoms
```

Iterator Directive

- IteratorDirective passed to *reset* instructs how to iterate
- Key attributes
 - direction
 - UP, DOWN, BOTH, NEITHER
 - atom
 - iteration performed in reference to atom, if not null
- Directive is conditioned before passing to iterator
 - set()
 - sets atom to null
 - direction unchanged
 - set(Atom a)
 - sets atom to a
 - direction unchanged
 - set(Direction d)
 - atom unchanged
 - sets direction to d



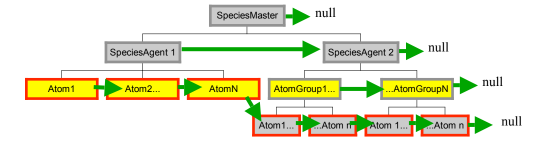
Setting the Basis

- Basis defines the source of atoms for the iterator
 - setBasis(Atom);
- Within a basis, iterator may be set to loop over atoms in different ways
 - Depends on iteratorDirective
 - Depends on design of iterator
- Setting basis does not reset the iterator
 - Need to call reset() before beginning iteration



AtomIteratorSequential 1.

- Most commonly used iterator
- Presents atoms in sequence defined with heirarchy



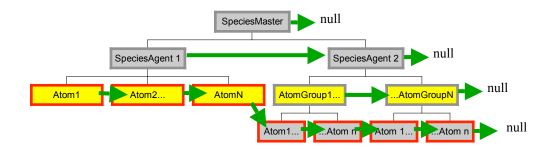
- setBasis indicates the parent atom of the iterates
 - isLeafIterator flag indicates if children of basis are iterated, or if all leaf atoms below basis are iterated



AtomIteratorSequential 2.

Directives

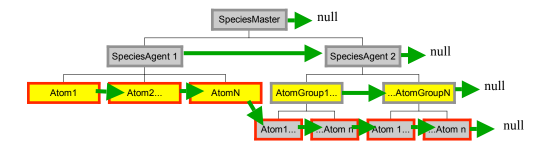
- atom == null indicates to do all atoms
- atom != null indicates to begin iteration with atom
- direction indicates which way to go in sequence
 - UP: natural order, beginning with specified atom
 - DOWN: opposite direction, beginning with specified atom
 - BOTH: *up* beginning with atom, then *down* beginning after it
 - NEITHER: do just specified atom





AtomIteratorSequential 3.

- Examples (set as leaf iterator)
 - setBasis(speciesAgent1);
 - iterator.reset(iteratorDirective.set().set(UP))
 - Loop through all molecules of species 1, from first to last
 - iterator.reset(iteratorDirective.set(atom2).set(UP));
 - Loop from atom2 up through last
 - setBasis(speciesMaster);
 - iterator.reset(iteratorDirective.set(atomN).set(BOTH));
 - Loop from atomN up through last atom in phase, then from N-1 down to first atom in phase





Other Atom Iterators

- AtomIteratorCompound
 - Forms a single iterator from a collection of atom iterators
 - Can also form from an atom group
- AtomIteratorBonds
 - Loops over atoms Bonded to Basis atom
- AtomIteratorNonbonded
 - Loops over atoms in molecule not bound to given atom
- AtomIteratorList
 - Loops over atoms in an AtomList
- AtomIteratorSinglet
 - Gives one atom and then expires
- AtomIteratorNeighbor
 - Loops over predefined "neighbors" of a given atom



AtomPairIterator

- Iterates over pairs of atoms, returning each as an AtomPair
- Formed from two AtomIterator instances
- IteratorDirective interpretation is slightly different
 - Details for another discussion!

