Knowledgebase of Interatomic Models Application Programming Interface (KIM API)

Ryan S. Elliott, Ellad B. Tadmor, and Valeriu Smirichinski Dept. of Aerospace Engineering and Mechanics, University of Minnesota

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This document describes how KIM **Tests** and **Models** written in different languages work together. A unified interface, tuned for the specific needs of atomistic simulations, is presented. This interface is based on the concept of "descriptor files". A descriptor file specifies all variables and methods required for communication between a particular **Model** and a **Test**. A "KIM API object" is created, based on the descriptor files, that holds all arguments (variable/data and method pointers) needed for **Test/Model** interaction. A complete set of KIM API service routines are available for accessing the various pointers in the KIM API object.

openkim-api package 1

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- KIM repository: KIM data

KIM API concept and implementation:

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KIM TEAM























PIs

Ellad Tadmor (U. Minnesota) Ryan Elliott (U. Minnesota) James Sethna (Cornell)

Developers

Alex Alemi (Cornell) Matt Bierbaum (Cornell) Woo Song Choi (Cornell) Daniel Karls (U. Minnesota) John Crow (Silicon Life Sciences) Trevor Wenblom (Silicon Life Sciences)

Advisory Board

Graeme Ackland (U. Edinburgh) Michael Baskes (LANL) Chandler Becker (NIST) Noam Bernstein (NRL) Ioana Cozmuta (NASA) Karsten Jacobsen (Tech. U. Den.)

Ronald Miller (Carleton)

John Moriarty (LLNL)

Sadasivan Shankar (Intel)

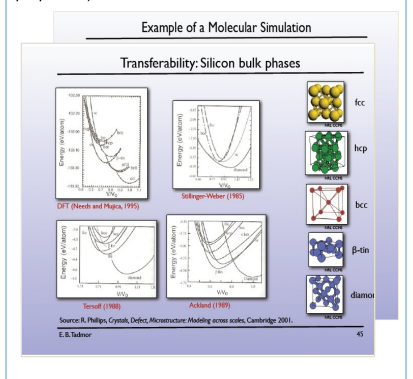
Adri van Duin (Penn State)

Gabriel Wainer (Carleton)

Molecular/atomistic simulations: tests and models

Tests

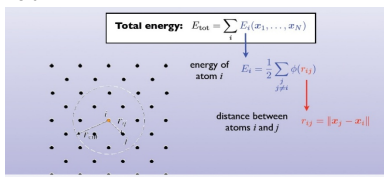
Test: a specific computer program which, when coupled with a suitable Model, calculates and returns a specific Prediction about a particular Configuration (or sequence of Configurations for dynamical properties).



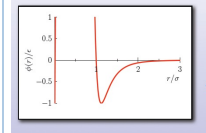


Models

Model: Computer implementation representing a specific interaction between atoms, e.g. an interatomic potential or force field



The Lennard-Jones potential is a simple pair potential, which described the interaction between two uncharged atoms:



$$\phi(r) = 4\epsilon \left[\left(\frac{\sigma}{r} \right)^{12} - \left(\frac{\sigma}{r} \right)^{6} \right]$$

repulsion due to overlapping electrons (Pauli principle) van der Waals attraction between transient dipoles

- Two fitting paramters (σ, ε)
- Designed for the nobles gasses (Ne, Ar, Kr, Xe).

Types of molecular modelers

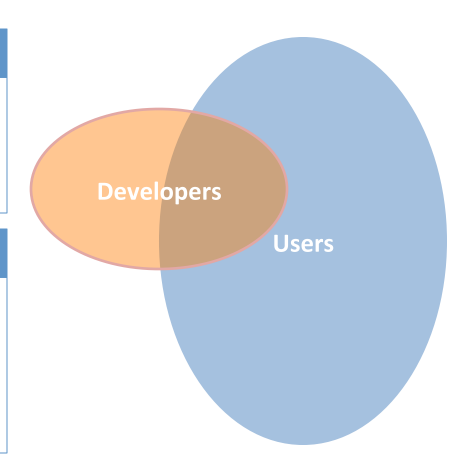
Very broadly speaking there are two types of *molecular modelers*:

Developers

- Create new models
- Study materials physics and applications
- Create new knowledge

Users

- Use models to study materials problems of scientific/technological importance
- Build sophisticated simulations to extract meaningful data
- Create new knowledge



Barriers faced by molecular modelers

The difficulties faced by developers and users of interatomic models include:

- No easy access to an extensive list of reliable reference data from experiments and first principles calculations for fitting.
- No easy access to implementations of existing models with known provenance and cross-language capability.
- No standardized tests for evaluating properties of molecular systems.
- 4. No framework for evaluating the *precision and transferability* of models and therefore no *rigorous guidelines* for choosing an appropriate model for a given application.

Knowledgebase of Interatomic Models (KIM) is proposed to overcome the barriers

The *Knowledgebase of Interatomic Models (KIM)* project is based on a four-year NSF cyber-enabled discovery and innovation (CDI) grant. The KIM project is designed to overcome the barriers mentioned on the previous page. KIM has the following main objectives:

- Development of an *online open resource* for standardized testing and long-term warehousing of interatomic models (potentials and force fields) and data.
- Development of an *application programming interface* (*API*) standard for atomistic simulations, which will allow any interatomic model to work seamlessly with any atomistic simulation code.
- Fostering the development of a quantitative theory of transferability of interatomic models to provide guidance for selecting application-appropriate models based on rigorous criteria, and error bounds on results.
- Striving for the permanence of the KIM project, including development of a sustainability plan, and establishment of a long-term home for its content.

More information on KIM is available at the project website: http://openKIM.org9
University of Minnesota

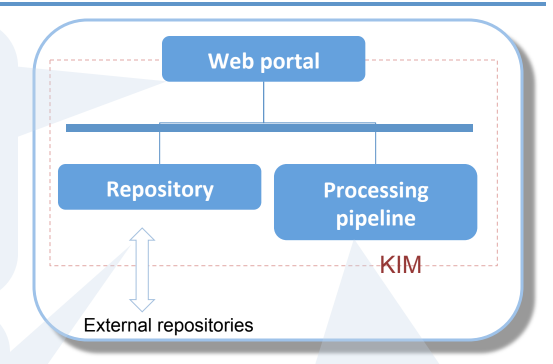
KIM framework

A web interface that will facilitate:

- user upload and download of Tests, Models and Reference Data
- searching and querying the repository
- comparing and visualizing Predictions and Reference Data
- recording user feedback (ranking and discussion forums)

A user-extendible database of

- interatomic Models
- standardized Tests (simulation codes)
- Predictions (results from Model-Test couplings)
- Reference Data (obtained from experiments and first principles calculations)



Processing Pipeline:

An automatic system for generating Predictions due to new Test or Model upload or changes:

- detect viable Test-Model couplings
- assign computational resources based on priority and dependencies
- store results in Repository
- requires an application programming interface (API) to be defined

KIM repository: Models

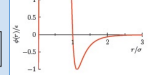
Models

Tests Predictions Reference Data KIM API

Model: Computer implementation representing a specific interaction between atoms, e.g. an interatomic potential or force field.

- Model Format
 - Stand-alone Model (black box)
 - Model Driver (e.g. Lennard-Jones)
 - + Parameter Set (e.g. ε_{Ar} =10.4 meV, σ_{Ar} =0.34 nm)

 $\phi(r) = 4\epsilon \left[\left(\frac{\sigma}{r} \right)^{12} - \left(\frac{\sigma}{r} \right)^{6} \right]^{\frac{5}{5}}$



Lennard-Jones (pair) • Ar parameterization •	Stillinger-Weber (3-bdy) • Si parameterization	:: CHARMM/AMBER ::	EDIP Brenner
 ⊞ Morse (pair)	• !!	EAM/Finnis-Sinclair/glue	Bond-order potentials
Cu parameterization	MGPT (4-body)	MEAM	ReaxFF
• • • • • • • • • • • • • • • • • • • •	Mo parameterizationTa parameterization	∺ Tersoff	⊪ GAP
Born-Mayer (ionic pair)	• [']	• 0 • 0 • 0	• 0 • 0 • 0

Every model will have a unique KIM ID for referencing in papers.

KIM repository: Tests

Models

Tests

Predictions Reference Data

KIM API

Test: a specific computer program which when coupled with a suitable Model, possible including additional input, calculates and returns a specific Prediction about a particular Configuration (or sequence of Configurations for dynamical properties).

Prediction of a Test will be a logical, scalar, tensor, graph, configuration or field, computed from a Test-Model coupling

Scalars

- lattice constants
- cohesive energy
- vacancy formation energy
- surface energy
- grain boundary energy
- vacancy migration barrier
- dislocation mobility
- peierls stress
- melting temperature

Tensors

- stress
- elastic constants

Configurations

- dislocation core structure
- surface structure
- grain boundary structure
- nanocluster structure

Graph

- phonon spectrum
- cohesive energy vs volume
- energy along transition path
- radial distribution functions

Fields

- simulated TEM hi-res image
- gamma surface

- Popular codes (ddcMD, DL POLY, GROMACS, GULP, iMD, LAMMPS, NAMD, SPaSM, etc.) can be included in a library of tools for writing *Tests*.
- Automatic test generation by linking to external repositories of first principles results.

KIM repository: KIM Data

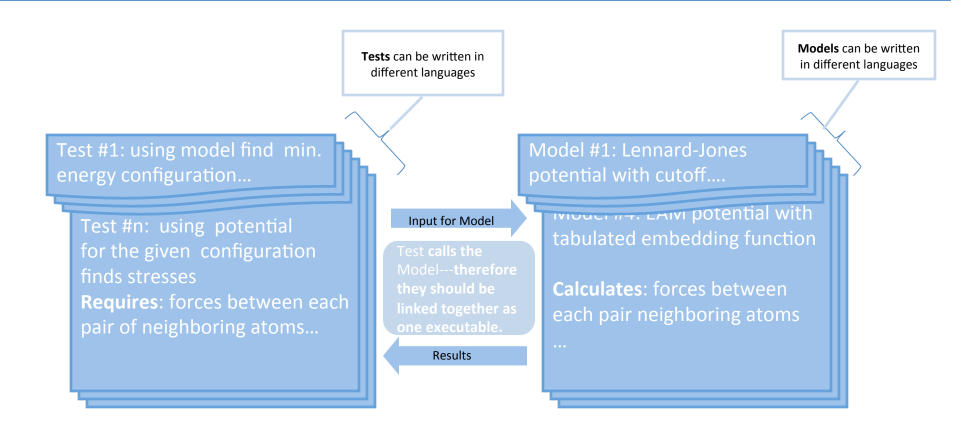
Models Tests Predictions Reference Data KIM API

Data in KIM can either be

- → a *Prediction* computed from a Test-Model coupling, or
- Reference Data computed by first principles or measured experimentally.
- Standardization of Data
 - Identified in terms of a set of "descriptors" drawn from a standardized "dictionary" (similar to that used in the Protein Data Bank project)
 - Descriptors will be automatically generated when possible (for example, the "Space Group" descriptor will be automatically generated for a given crystal structure).
- Data classes
 - Logical (true/false result for a test, e.g. a given crystal phase is stable)
 - Scalar or Tensor (lattice constant, cohesive energy, elastic constants...)
 - Graphs (transition pathway energy, phonon spectrum, ...)
 - Configurations (relaxed defect core, surface structure, ...)
 - Fields (simulated hires TEM image, ...)
- Quality assurance
 - Acceptance of only "publication quality" data enforced by KIM Editor
 - "Data Provenance"

KIM API concept and implementation

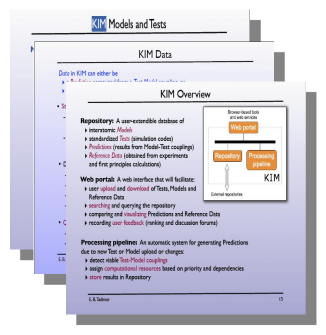
The KIM API facilitates communication between Models and Tests



Users and developers will be able to download **Tests** and **Models** (from openkim.org), then compile, link and run the resulting programs to produce new results.

The most challenging technical requirement is the need for multi-language support

openKIM.org framework



Processing pipeline: an automatic system for generating predictions when Tests or Models are uploaded or changed.

Requirements:

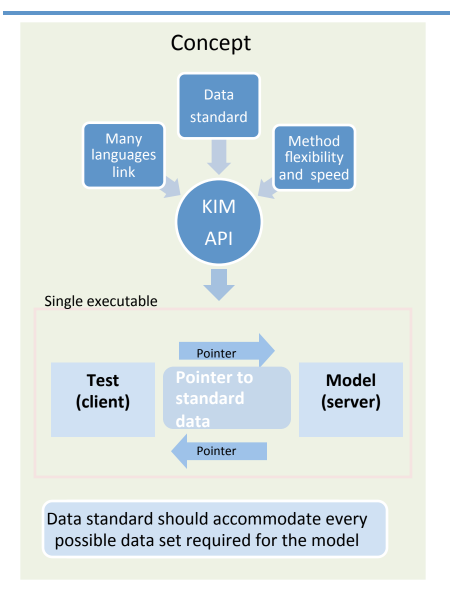
- •Multilanguage support (C, C++, F77, FORTRAN 90, Python ...)
- A variety of data structures need to be accommodated: scalars, multidimensional arrays, variable size arrays, etc..
- Speed & performance are very important
- Standardized API, version tracking, etc...

Processing pipeline: sequence of actions

- detect a viable Model/Test coupling
- build (compile and link)
 Tests against Model
- run probe-tests
- assign computational resources
- run full-scale Test against
 Model
- analyze results ...
- store results in the repository

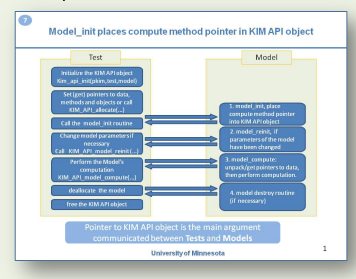
Need a simple interface: ideally just one argument per call

The KIM API is based on exchanging pointers to data and methods



Schematic of implementation

- Data and method pointers are packed in one object. The Interface consists of exchanging one pointer to the KIM API object between a **Test** and a **Model**
- 2. All languages naturally support pointers:
 - •FORTRAN (cray or 2003 standard)
 - •C/C++
 - Java
 - Python



Using C-style pointers in Fortran

In order to implement the KIM API concept in a cross-language environment, all languages have to work with C-style pointers.

FORTRAN 77 and Fortran 90/95 do not support C-style pointers directly, however essentially all compilers support the `cray pointers' extension which provides this capability.

A cray pointer is an integer that can store a memory address. An example below shows the general syntax and usage of a cray pointer in Fortran compared with C.

C code FORTRAN code Keyword pointer, followed by two arguments double precision :: y=10.0d0 double y=10.0; double precision :: x px - is a pointer (analog pointer (px,x) double *x; double *x in C) x - is a pointee px = loc(v)x = &v; print*,"x=",x printf("*x=%f \n", *x); As soon as px holds an address, access to that address is done by pointee x

How can a Test know what type of input/output data is required by a Model? We have solved this problem by introducing the KIM API descriptor file

ex_model_Ne_P_MLJ_NEIGH_PURE_H.kim

```
MODEL NAME := ex model Ne P LJ NEIGH PURE H
Unit Handling := fixed
SUPPORTED ATOM/PARTICLES TYPES:
# Symbol/name
               Type
                               code
                               1
Ne
               spec
MODEL INPUT:
# Name
                        Unit.
                                    Shape
                                               Requirements
                Type
numberOfParticles
                integer
                                    []
                        none
numberParticleTypes
                integer
                                    []
                        none
particleTypes
                integer
                                    [numberOfParticles]
                        none
```

KIM API descriptor file defines all arguments that the model needs for computation including input and output arguments. Also on the test side, the .kim file defines what the Test can provide as input for the Model and what it expects from the Model as a result.

Tests and Models expose the required input/output arguments that will be communicated using the KIM API

Structure of descriptor file

Model/Test name and system of units lines

MODEL_NAME := ex_model_Ar_P_Morse

Unit_Handling := flexible

Section lines

SUPPORTED ATOM/PARTICLES TYPES:

CONVENTIONS:

MODEL_INPUT:

MODEL_OUTPUT:

MODEL_PARAMETERS:

Data lines

- * Species Data lines
- * Flag Data lines
- * Argument Data lines

Brief description of Section lines

These lines identify logically distinct sections within the KIM descriptor file.

All lines following a Section line, up to the next Section line or end of the file, will be assigned to the indicated section.

These sections may occur in any order within a KIM descriptor file, however the order given here is recommended. A section line may only occur once within a KIM descriptor file.

Brief description of Data lines

These lines are used to specify the information that a Model (Test) will provide to and require from a Test (Model), as well as the conventions that the Model (Test) uses.

- * Species Data lines allow for the definition of atomic species by providing a symbol and an integer code. These lines are located in section SUPPORTED ATOM/PARTICLES TYPES.
- * Flag Data lines this line type defines a convention that can be used to ensure that Models and Tests are able to work together, and should only be used within the CONVENTIONS section of the KIM descriptor file.
- * Argument Data lines the main KIM descriptor file line format, used within the MODEL_INPUT, MODEL_OUTPUT, and MODEL PARAMETERS sections.



Each argument line in the descriptor file describes an argument and its properties

EXAMPLEs/MODELs/ex_model_Ar_P_MLJ_F90.kim

All characters after a '#' are ignored (a comment field)

MODEL_NAME := ex Unit_Handling		MLJ_F90			
 compute	method	none	[]	Method means a subroutine or function	
MODEL_OUTPUT: # Name	Type	Unit	Shape	pointer Requirements	
energy	real*8	energy	[]	optional	
force	r∋al*8	force	[numberOf?articles	opt; nal	

The name of an argument is its "key word". By using key words, the KIM service routines can pack/unpack data pointers from the KIM API object. Key words are standardized as part of the KIM API.

Type of data in computer representation

Physical dimensions

The shape of an argument describes its array properties. It specifies the number (rank) and size (range or extent) of indices. For example, [] means a scalar (zero-dimensional array), [numberOfParticles] means a one-dimensional array and [numberOfParticles,3] means a two-dimensional array of size numberOfParticles x 3.

The "requirements" field is only used in **Model** descriptor files. An empty field indicates that the argument is required. A value of "optional" indicates that the associated data will be computed only if the argument is in the **Test**'s descriptor file and if the **Test** explicitly requests it.

Specifying particle types – species data lines

EXAMPLEs/MODELs/ex_model_Ar_P_MLJ_F90.kim

 Species data lines define the atom/particle types supported by the Test/Model and should only be used within the SUPPORTED_ATOM/PARTICLES_TYPES section of the KIM descriptor file. Each line consists of three white-space separated (case sensitive) strings The three strings are as follows:

code: This is the integer that the Model uses internally to identify the atom/particle type. The value specified by a Test is ignored.

Type: This must be `spec'.

Name: This string gives a unique name to the atom/ particle type. This name is checked against the standard list in `standard.kim'.

The **KIM_API_get_partcl_types()** service routine allows one to obtain a list of all particle types used by the model during runtime. Also the **KIM_API_get_partcl_type_code()** service routine allows one to get the particle type integer code (see DOCs/KIM_API_Description.txt).

In order to define "conventions" of test/model behavior, flag data lines are reserved

EXAMPLEs/MODELs/ex_model_Ar_P_MLJ_F90.kim

######################################				
# Name	Туре			
OneBasedLists	flag			
Neigh_IterAccess	flag			
Neigh_LocaAccess	flag			
NEIGH_RVEC_H	flag			
NEIGH_PURE_H	flag			
NEIGH_RVEC_F	flag			

A flag data line defines a convention, that can be used to ensure that **Models** and **Tests** are able to work together, and should only be used within the CONVENTIONS section of the KIM descriptor file. The line consists of two white-space separated (case sensitive) strings. The two strings, in order, are as follows:

Name: This string gives a unique name to the convention. This name is checked against the standard list in 'standard.kim'

Type: This must be 'flag'

KIM_API_allocate() has no effect on "flag" type arguments, because they are not "data pointer holders".

For a detailed description of all flag lines see the file DOCs/standard.kim. Also see the files in DOCs/.



Parameter arguments are used to publish/access internal parameters of a Model

EXAMPLEs/MODELs/ex_model_Ar_P_MLJ_CLUSTER_F90/ex_model_Ar_P_MLJ_CLUSTER_F90.kim

MODEL_PARAMETERS:				
# Name	Туре	Unit	Shape	Requirements
PARAM_FREE_sigma	real*8	length	[]	
PARAM_FREE_epsilon	real*8	energy	[]	
PARAM_FREE_cutoff	real*8	length	[]	
•••				

The format for parameter arguments in a KIM descriptor file is the same as that for argument data types.

Two types of model parameters are allowed

- 1) PARAM_FIXED_XXXXXXX these should not be changed by the Test
- 2) PARAM_FREE_XXXXXXX these may be changed by the Test (which must then call the Model's reinit() function to inform the model that its parameters have changed)

KIM_API_get_params() service routine will return a list of all parameters in the object during runtime (as an array of text strings).

KIM API get free params() service routine will return a list of FREE parameters and

KIM_API_get_fixed_params() will return a list of FIXED parameters (see KIM_API_Description.txt)

Names of parameter arguments are not checked against standard.kim

Specifying units that model can handle: Units Handling and base units

EXAMPLEs/MODELs/ex model Ar P MLJ F90/ex model Ar P MLJ F90.kim

For Models, a variable `Unit_Handling' specifies whether the Model can adjust its input and output to match a Test (`flexible') or can only work with one set of units (`fixed'). This information is ignored for Tests.

Base unit lines:

Five lines that describe a set of five base units from which all other units are derived in a consistent way:

Unit_charge := `C' | `e' | `statC'

Unit temperature := `K'

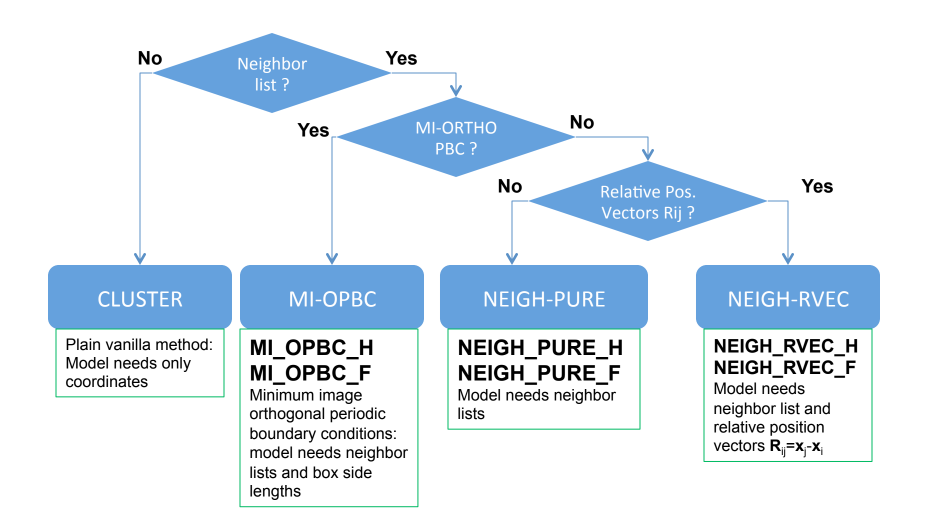
Unit_time := `fs' | `ps' | `ns' | `s'

The list of recognized units above may be extended in the future.

There are several service routines related to units and units handling in KIM API:

KIM_API_get_unit_handling(), KIM_API_convert_to_act_unit(), KIM_API_get_unit_length(), KIM_API_get_unit_energy(), etc...(see DOCs/KIM_API_Description.txt).

Handling of Neighbor lists and Boundary Conditions – NBC methods



Descriptions of the NBC methods

CLUSTER:

In the CLUSTER method, the Model receives the number of particles and coordinates without additional information (such as neighbor lists or other boundary condition specifiers) and computes requested quantities under the assumption that the particles form an isolated cluster. For example, if energy and forces are requested, it will compute the total energy of all the particles based on the supplied particle coordinates and the derivative of the total energy with respect to the positions of the particles.

NEIGH PURE [H|F]:

In the NEIGH_PURE methods (NEIGH_PURE_H and NEIGH_PURE_F), the Model receives the number of particles, coordinates and a full or half neighbor list. The neighbor list defines the environment of each particle, from which the particles's energy is defined. In the case of a half list, the value of the argument `numberContributingParticles' indicates that the first `numberContributingParticles' contribute their energy to the total and the remaining particles do not contribute to the energy (they are "ghost" particles). When `numberContributingParticles' is equal to `numberParticles' the half list is called "symmetric", otherwise it is called "unsymmetric." In the case of a full list, any particle that has one or more neighbors contributes its energy to the total and those particles with zero neighbors do not contribute to the total energy. The model computes the requested quantities using the supplied information. For example, if energy and forces are requested, it will compute the total energy of all the particles based on their neighbor lists and the derivative of the total energy with respect to the positions of the particles. This method can be used with codes that use ghost particles to apply boundary conditions. The ghost particles are treated as regular particles by the Model, and it is up to the calling code to discard some information such as the forces on the ghost particles and to compute the appropriate total energy from per-particle energies of the physical particles, or to use a modified neighbor list to obtain the desired values.

NEIGH PURE H:

This is the Pure Half neighbor list method. The model needs `coordinates', a half neighbor list (with data stored in the `neighObject' argument), the `numberContributingParticles', and the `get_neigh' method supplied by the Test. NEIGH PURE F:

This is the Pure Full neighbor list method. The model needs `coordinates', a full neighbor list (with data stored in the `neighObject' argument), and the `get_neigh' method supplied by the Test.

Descriptions of the NBC methods (2)

NEIGH RVEC [HIF]:

In the NEIGH_RVEC methods (NEIGH_RVEC_H and NEIGH_RVEC_F), the Model receives the number of particles, coordinates, a half or full neighbor list, and the relative position vectors r_ij (r_ij = x_i-x_i). The neighbor list and r_ij vectors define the environment of each particle, from which the particles's energy is defined. In the case of a half list, a neighbor pair i and j (where i < j) with relative position vector (RVEC) r_ij defines two pieces of information: (1) j is a neighbor of i with RVEC r_ij and (2) i is a neighbor of j with RVEC r_ji = -r_ij. Additionally, the value of the argument 'numberContributingParticles' indicates that the first 'numberContributingParticles' contribute their energy to the total and the remaining particles do not contribute to the energy (they are "ghost" particles). When 'numberContributingParticles' is equal to 'numberParticles' the half list is called "symmetric", otherwise it is called "unsymmetric." In the case of a full list, any particle that has one or more neighbors contributes its energy to the total and those particles with zero neighbors do not contribute to the total energy. The model computes the requested quantities using the supplied information. For example, if energy and forces are requested, it will compute the total energy of all the particles based on their neighbor lists and the derivative of the total energy with respect to the positions of the particles. These methods enable the application of general periodic boundary conditions, including multiple images. A possible future extension to these methods is to allow the Test to provide a ForceTransformation() function for each neighbor, which would enable the application of complex boundary conditions such as torsion and objective boundary conditions.

NEIGH RVEC H:

This is the Relative Vector Boundary Condition Full neighbor list method. The Model needs `coordinates', a full neighbor list (with data stored in the `neighObject' argument), and the `get_neigh' method supplied by the Test. The `neighObject' argument must also contain the relative position vectors (RVEC) (which are returned by the `get_neigh' function). NEIGH RVEC F:

This is the Relative Vector Boundary Condition Full neighbor list method. The Model needs `coordinates', a full neighbor list (with data stored in the `neighObject' argument), and the `get_neigh' method supplied by the Test. The `neighObject' argument must also contain the relative position vectors (RVEC) (which are returned by the `get_neigh' function).



Descriptions of the NBC methods (3)

MI_OPBC_[H|F]:

In the MI_OPBC methods (MI_OPBC_H and MI_OPBC_F), the Model receives the number of particles and coordinates, the side lengths for the periodic orthogonal box and a neighbor list. It assumes all particles lie inside the periodic box. Side lengths of the box must be at least twice the cutoff range. This method computes the requested quantities under the assumption that the particles are subjected to the minimum image, orthogonal, periodic boundary conditions. MI_OPBC_H:

This is the Minimum Image Orthogonal Periodic Boundary Condition Half neighbor list method. The Model needs 'coordinates', a half neighbor list (with data stored in the 'neighObject' argument), 'numberContributingParticles', the 'get_neigh' method supplied by the Test, and the 'boxSideLengths' argument (which specifies the three side-lengths of the orthogonal simulation box).

MI OPBC H:

This is the Minimum Image Orthogonal Periodic Boundary Condition Full neighbor list method. The Model needs `coordinates', a full neighbor list (with data stored in the `neighObject' argument), the `get_neigh' method supplied by the Test, and the `boxSideLengths' argument (which specifies the three side-lengths of the orthogonal simulation box).

Example of using NBC methods in KIM file

EXAMPLEs/MODELs/ex_model_Ne_P_LJ/ ex_model_Ne_P_LJ.kim

CONVENTIONS: # Name OneBasedLists	Type flag
Neigh_IterAccess	flag
Neigh_LocaAccess	flag
NEIGH_RVEC_H	flag
NEIGH_PURE_H	flag
NEIGH_RVEC_F	flag
NEIGH_PURE_F	flag
MI_OPBC_H	flag
MI_OPBC_F	flag
CLUSTER 	flag

The example in ex_model_Ne_P_LJ.kim is designed to work with six different NBC methods.

If the Test can also work with multiple NBC methods and there are several matches, the first matched method listed in the Model's KIM file will have precedence.

The KIM_API_init () routine will check that all needed data lines for the chosen method are in the KIM descriptor file.

NBC Methods

Neighbor list access methods: all related lines in the KIM descriptor files

DOCs/standard.kim (only related to Neighbor list access are shown here)

```
CONVENTIONS:
# Name
                         Type
ZeroBasedLists
                         flag
                                  # presence of this line indicates that indexes
                                  # for particles are from 0 to numberOfParticles-1 (C-style)
                                  # presence of this line indicates that indexes for
OneBasedLists
                         flag
                                  # atoms are from 1 to numberOfParticles
                                                                               (Fortran-style)
                                  # works with iterator mode
Neigh IterAccess
                         flag
                                  # works with locator mode
Neigh LocaAccess
                         flag
Neigh BothAccess
                                  # needs both locator and iterator modes
                         flag
                                             neighObject stores completely encapsulated neighbor list object
NEIGH PURE H
                         flag
NEIGH PURE F
                         flag
                                             Access to the object is done through method get_neigh. The
NEIGH RVEC H
                         flag
                                             neighbor list object and the method to access it are supplied by
NEIGH RVEC F
                         flag
                                             the Test.
MI OPBC H
                         flag
MI OPBC F
                         flag
MODEL INPUT:
# Name
                                      Unit
                                                    Shape
                                                                 requirements
                         Type
get neigh
                         method
                                      none
neighObject
                         pointer
                                                    []
                                      none
                         real*8
                                                    [3]
boxSideLengths
                                      length
```

rij

Interface to get_neigh method

```
integer function get neigh(pkim,mode,request,particle,numnei,pnei1particle,prij)
get neigh function for access to the neighbor
                                                        implicit none
list object
                                                       integer(kind=kim intptr),
                                                                                                 :: pkim
                                                                                    intent(in)
here:
                                                                                                                    FORTRAN style
                                                                                                 :: mode
                                                        integer.
                                                                                    intent(in)
          operate in iterator or locator mode
mode -
                                                                                                 :: request
                                                                                    intent(in)
                                                        integer,
           mode = 0 : iterator mode
                                                       integer,
                                                                                                :: particle
                                                                                    intent(out)
           mode = 1 : locator mode
                                                                                    intent(out)
                                                        integer,
                                                                                                 :: numnei
                                                        integer,
                                                                                    intent(out)
                                                                                                :: pnei1particle
request - Requested operation
                                                       integer,
                                                                                                 :: nei1particle(1);
           If mode = 0
                                                                                                    pointer(pnei1particle,nei1particle)
              request = 0 : reset iterator
                                                        double precision,
                                                                                   intent(out)
                                                                                                 :: prij
              request = 1 : increment iterator
                                                        double precision,
                                                                                                 :: rij(3,*);
                                                                                                              pointer(prij,rij)
           If mode = 1
                                                    end function get neigh
              request = #: number of the
                            particle whose
                                                    int get neigh(void ** pkim, int * mode, int * request, int * particle,
                                                                                                                                C style
                            neighbor list
                                                                        int * numnei, int ** pnei1particle, double ** prij);
                            is requested
```

```
particle - the number of the particle whose neighbor list is returned
```

numnei - number of neighbors returned

nei1particle - integer array of neighbors of an particle which will point to the list of neighbors on exit.

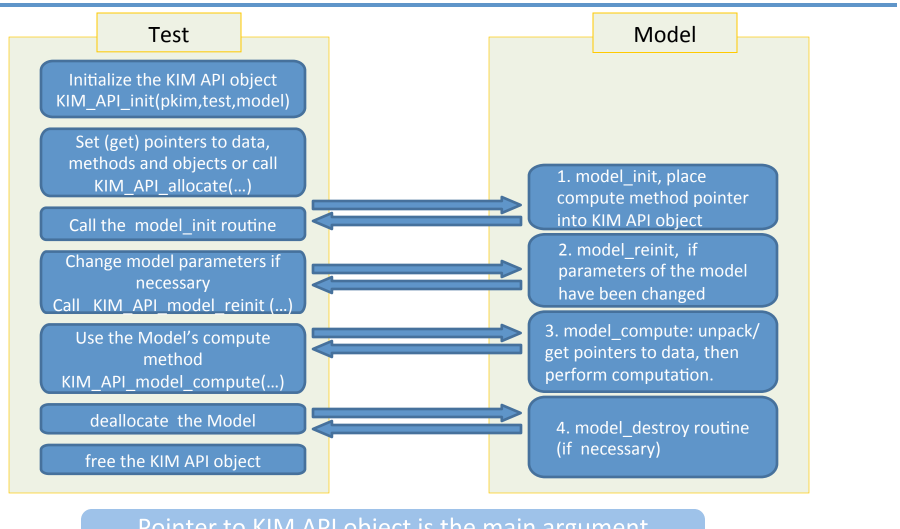
 array of relative position vectors of the neighbors of a particle (including boundary conditions if applied) if they have been computed (NBC scenario NEIGH_RVEC_[H|F] only). Has NULL value otherwise (all other NBC

scenarios).

The return value depends on the results of execution. (see DOCs/KIM_API_Description.txt for details)

Test must supply the get_neigh method and store a pointer to it in the KIM API object

Model_init places compute method pointer in KIM API object



Pointer to KIM API object is the main argument communicated between **Tests** and **Models**

Initialization of KIM API object, setting and getting data-pointers can be done through the KIM service routines

KIM API/KIM API C.h

```
#include <stdint.h>
#ifdef cplusplus
extern "C" {
#endif
//global methods
int KIM API init(void * kimmdl, char * testname, char *mdlname);
void KIM_API_allocate(void *kimmdl, intptr t natoms, int ntypes);
void KIM API free(void *kimmdl, int * kimerror);
void KIM API print(void *kimmdl, int *kimerror);
int KIM_API_model_compute(void * kimmdl);
//element access methods
int KIM API set data(void *kimmdl,char *nm, intptr t size, void *dt);
void * KIM API get data(void *kimmdl,char *nm, int * kimerror);
//multiple data set/get methods
void KIM_API_setm_data(void *kimmdl, int *error, int numargs, ... );
void KIM API getm data(void *kimmdl, int *error, int numargs, ... );
```

Initialization is done by analyzing test and model descriptor files

One can use optional KIM service routine to allocate standard arguments and data

Call model_compute routine by address stored in KIM API object

Directly place data pointer into the KIM API object

"Multiple" version of set/get data

Description of all KIM API service routines are located in the file: **DOCs/KIM API Description.txt**

Examples of using KIM_API_init and KIM_API_allocate service routines

.../ex_test_Ar_free_cluster_CLUSTER_F90/ex_test_Ar_free_cluster_CLUSTER_F90.F90

.../ex_test_Ar_multiple_models/ex_test_Ar_multiple_models.c

```
status = KIM_API_init(&pkim_periodic_model_0, testname,
modelname0);
   if (KIM_STATUS_OK > status)
        KIM_API_report_error(_LINE__, __FILE__, "KIM_API_init())
for MODEL_ZERO for period
...
```

KIM API init will check the consistency of KIM descriptor file (Test and Model) against standard.kim, after that will check if Test and Model match: NBC methods, particle species (if any), conventions and argument data lines

If the match is successful, then the KIM API object is created. This object conforms to the Model descriptor KIM file and can store all described data as pointers

KIM_API_allocate will allocate memory for all arguments with, fully specified shape, stored in the KIM API object

It is not mandatory to use KIM_ API_allocate. A Test can use its own memory and set address of the data in the KIM API object.

Examples of using KIM API getm/setm data

("multiple" version of get/set data)

.../ex_test_Ar_free_cluster_CLUSTER_F90/ex_test_Ar_free_cluster_CLUSTER_F90.F90

```
! Unpack data from KIM object
 call kim api getm data f(pkim, ier, &
      "numberOfParticles", pnAtoms,
                                          1, &
      "numberParticleTypes", pnparticleTypes, 1, &
      "particleTypes", pparticleTypesdum, 1, &
      "coordinates", pcoor, 1, &
     "cutoff",
                                        1, &
                pcutoff,
      "energy",
                penergy, 1, & pvirialglob, 1, &
      "virial",
      "forces",
                        pforces,
                                           1)
 if (ier.lt.KIM STATUS OK) then
    idum = kim api report error f( LINE , THIS FILE NAME,&
                          "kim api getm data f", ier)
    goto 42
 endif
```

KIM_API_getm_data (or kim_api_getm_data_f) will return pointers stored in KIM_API object. "Multiple" version of get data routines allows to get several variable pointers from the KIM API object s at once.

KIM_API_setm_data (or kim_api_setm_data_f) allows to place (pack) several data pointers into KIM API objects See DOCs/KIM_API_Description.txt for the details

KIM_API_model_init will call model initialize routine that, in turn, will place model compute into KIM object

.../ex_test_Ar_multiple_models/ex_test_Ar_multiple_models.c

```
/* call model init routines */
status = KIM_API_model_init(pkim_periodic_model_0);
if (KIM_STATUS_OK > status) KIM_API_report_error
(_LINE__,_FILE__,"KIM_API_model_i
...
/* call compute functions */
status = KIM_API_model_compute(pkim_periodic_model_0);
if (KIM_STATUS_OK > status) KIM_API_report_error
(_LINE__,_FILE__,"compute", status);
...
```

KIM_API_model_init will call the model_init routine . KIM_API_model_init utilizes the KIM standard naming convention in order to make the call. In C the name of the model init routine must have all lower case letters in the following format modelname_init_, for example:

model_ar_p_mlj_cluster_init__

model_name

KIM_API_model_compute
calls the address of the
model compute subroutine
stored in KIM API object.
By the time
KIM_API_model_compute is
called the address is placed in
KIM API object by
model_init_routine

.../ex_model_Ar_P_MLJ_C/ex_model_Ar_P_MLJ_C.c

Place address of actual compute routine into the KIM API object

An example of using get_neigh method through KIM API service routines

.../ex_model_Ar_P_MLJ_NEIGH_PURE_H_F/ex_model_Ar_P_MLJ_NEIGH_PURE_H_F.F90

```
do i = 1, numberOfParticles
   ! Get neighbors for atom i
   atom = i ! request neighbors for atom i
   ier= kim api get neigh f(pkim, 1, atom, atom ret, numnei, pneilatom, &
                            pRij dummy)
   if (ier.lt.KIM STATUS OK) then
          idum = kim api report error f( LINE , THIS FILE NAME,&
                                        "kim api get neigh", ier)
          return
   endif
    ! Loop over the neighbors of atom i
   do jj = 1, numnei
          j = neilatom(jj)
          Rij(:) = coor(:,j) - coor(:,i) ! distance vector between i j
          Rsqij = dot product(Rij,Rij) ! compute square distance
         if ( Rsqij < model cutsq ) then ! particles are interacting?</pre>
             r = sqrt(Rsqij)
                                                         ! compute distance
             call pair (model epsilon, model sigma, model A, model B, &
                     model C, r,phi,dphi,d2phi) ! compute pair potential
```

Locator mode -- get neighbors of a particle using half or full neighbor lists as requested.

KIM_API_get_neigh will call the method using the address stored in the KIM API object. These methods are supplied by the Test.

KIM_API_get_neigh will check if the arguments are set correctly. It will also convert the result from OneBaseLists to ZeroBaseLists (or vice versa) if necessary.

Details on the interface and a description of error codes are in DOCs/KIM_API_Description.txt

Computing quantities from the first derivative

MODELs/ ex model Ne P fastLJ/ex model Ne P fastLJ.c

```
while (KIM_STATUS_OK == *ier)

i = currentAtom + model_index_shift;;
    zi=i*DIM;

/* loop over the neighbors of currentAtom */
    for (jj = 0; jj < numOfAtomNeigh; ++ jj)

{
        ...

        /* process dEdr */
        if (comp_process_dEdr)
        {
            R = sqrt(Rsqij);
            double DE = fac*R;
            ier = KIM_API_process_dEdr(km, &DE, &R, &odx, &i, &j):
        }
        ...

This routine contribution, dEw with respect to to particle 'j'. The Totalin-rule, many and other therm automatic index
OneBasedList flate process_dEdr further proc
```

This routine can be called by a Model to provide the Test with a contribution, dEdr, to the first derivative of the Model's energy with respect to the (scalar) distance r_ij between particle `i' and particle `j'. The Test can use this information to compute, via the chain-rule, many properties. Examples include forces, the virial, and other thermodynamic tensions. The KIM API performs automatic index conversion (based on ZeroBasedList and OneBasedList flag settings) before calling the Test's supplied process_dEdr function. If the Test does not provide its own process_dEdr routine, then the KIM API standard process_dEdr routine is used. If the standard process_dEdr routine is used, the KIM API ensures that any appropriate memory initializations are performed. This routine and currently supports the computation of `virial' and `particleVirial'.

void **km -- pointer to KIM_API_model object

double *dE -- pointer to the contribution to the first derivative of the energy with respect to the pair-distance r_ij

double *r -- pointer to r_ij
-- the distance between
particles i and j

int * I,j $\mathrel{\mathsf{J}}$ -- pointers to particle index I and j.

double **pdx -- pointer to the relative position vector of particle j relative to particle i (i.e., r_ij = x_j - x_i).

On details of interface using process_dEdr see documentation in KIM_API_Description.txt and standard.kim

Appendix

Every argument that needs to be communicated between Tests and Models must be in the descriptor file

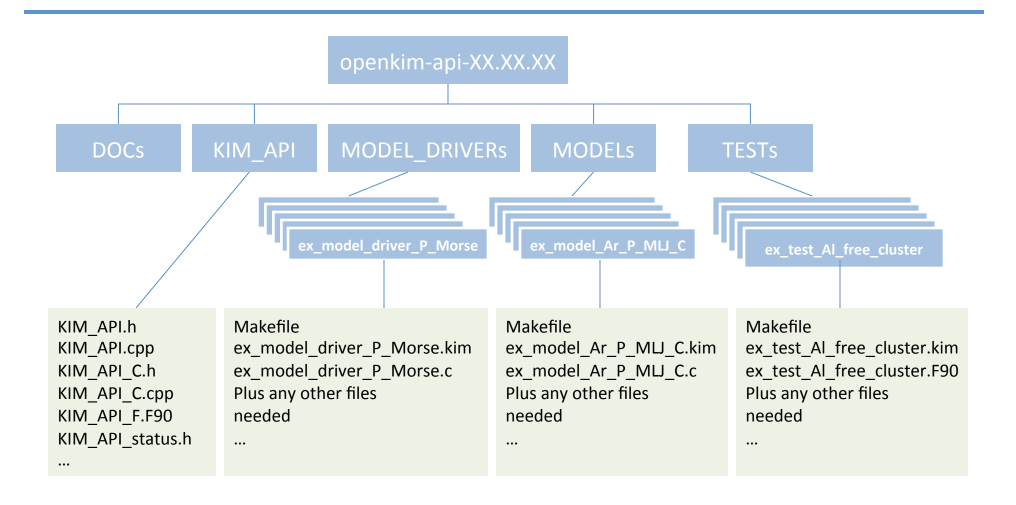
Each **Test** has its own descriptor file that describes the data it can supply to the **Model** and what data it expects the **Model** to compute. There are no optional arguments in a **Test**'s descriptor file (because the Test knows, a priori, what it will need to compute).

Each **Model** has its own descriptor file that describes the data it needs to perform its computations and what results it can compute. Some of the arguments/ methods can be identified as optional. Optional arguments/methods are ones that the **Test** does not have to provide or are results that the **Model** will only compute if the **Test** explicitly requests it.

KIM service routines (such as KIM_API_init) use both **Test** and **Model** descriptor files to:

- Check if the **Model** and **Test** match, also check if their descriptor files conform to the KIM API standard
- If they do -- create a KIM API object to store all arguments described in the Model's descriptor file
- Mark each optional argument that is not used by the **Test** "do not compute" (i.e., compute = false)
 The flag here is an integer: KIM_COMPUTE_TRUE compute, KIM_COMPUTE_FALSE do not compute
 Other service routines are used to:
- Set (get) argument or method pointers into (from) the KIM API object (e.g., KIM_API_set_data, KIM_API_get_data, etc.)
- Check if the "compute flag" is set to "compute" for an argument in the KIM API object (KIM_API_get_compute).
- Execute the Model's compute method (KIM API model compute)
- etc...

KIM API directory structure



Each **Test** and **Model** has its own descriptor file

The end