

MedeA LAMMPS: A Powerful Gateway to a Powerful Molecular Dynamics Program

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1 Introduction

MedeA LAMMPS provides flexible calculation setup and analysis capabilities to unlock the power of the LAMMPS molecular dynamics (MD) code.

MedeA LAMMPS automates the details of properly formatting molecules, fluids, and solids into the required LAMMPS coordinate, connectivity, forcefield parameter, and command-line formats. It also provides access to the core capabilities of LAMMPS, including minimization, molecular dynamics simulations using the *NVE*, *NVT*, and *NPT* ensembles, and materials properties calculations.

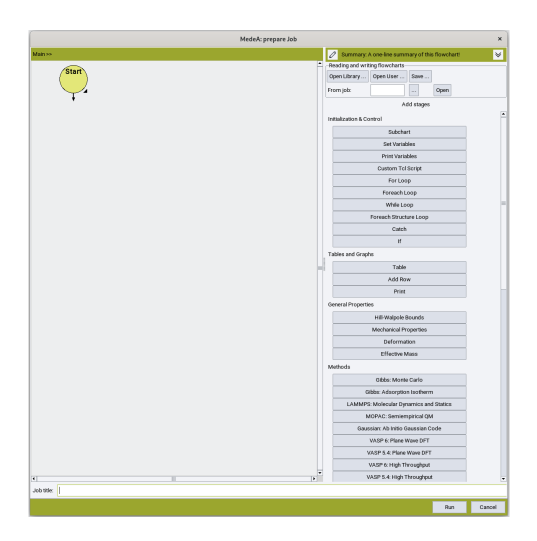
MedeA LAMMPS fully integrates with *MedeA Forcefield* for advanced forcefield handling and assignment, and any compatible custom forcefield can be used. There are also options for expert LAMMPS users to add any LAMMPS commands to existing protocols, or to prepare completely customized simulations.

After each calculation, *MedeA LAMMPS* automatically analyzes the block averages and fluctuations of temperature, pressure, density, cell parameters, total energy and all energy components (potential, kinetic, Coulomb and van der Waals), and stress tensor elements, reporting convergence statistics and uncertainties computed according to a method applicable to properties sampled using molecular dynamics or Monte Carlo methods. Additionally, visualization of trajectories of MD simulations and structure optimizations can be performed.

The user interface to LAMMPS is based on flowcharts and those from any previous *MedeA LAMMPS* calculation can be reused, edited, shared with colleagues, and rerun, even on different systems and compute servers.

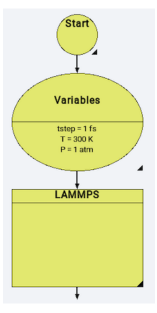
Bring up the Flowchart interface by selecting |Jobs| >> |New Job...|, and you should see the following:





2 Variables

A MedeA LAMMPS flowchart usually starts with a Variables stage:



With simulation variables defined in the following way:



	MedeA: prepare Job			×
Main >>	Edit stage: Variables			
Start	Variable	Value	Units	
()	tstep	1	fs	Delete
	т	300	к	Delete
	P	1	atm	Delete
Variables				Add
Valiables	Apply	Reset		
LAMMPS				
•		ок	Cancel	Help
Job title:				
			Run	Cancel

If no simulation variables are defined, the **LAMMPS** stages use the following default variables:

- Time step (tstep): 1 fs
- Temperature (T): 300 K
- Pressure (P): 1 atm

Note: A time step size of 1 fs works well for covalent forcefields with pre-defined bonds, angles, and dihedrals. For reactive potentials without pre-defined bonds, such as Tersoff, ReaxFF, and SNAP, time step sizes from 0.1 fs to 0.4 fs are recommended.

3 Initialization

Upon opening the LAMMPS stage, you will see the following interface:





The *Initialization* section in the right-side panel has one available stage: Initialize LAMMPS . The LAMMPS stage must start with the Initialize stage:

Med	eA: prepare Job				
fain >> LAMMPS		Edit stage: Initialize			
Start	-	Periodicity:	3-d period	lic	•
(•••••)		Nonbond method:	PPPM		•
\mathbf{v}		Nonbond cutoff:	9.5		
		Long range precision:	0.00001		
Initialize		✓ Add tail correctio	ns for Van de	r Waals interac	tions
		Run LAMMPS on:	CPU		-
3-d periodic CuoTri 9: 5 Logrange: PPM W/ tail correction	=				
	•		ок	Cancel	Help
III	()				

The parameters are:

• Periodicity: Choose from 3-d periodic for bulk systems and layer perpendicular to Z for slab models.

Hint: Even with 3-D periodic boundary condition, a slab model can effectively be created by having large enough vacuum size (> ~20 Å)



- *Nonbond method*: For long-range Coulombics, choose from Cutoff for a simple spherical cutoff, PPPM for the particle–particle–mesh long-range electrostatic solver, and Ewald for conventional Ewald. This is a forcefield-sensitive option, meaning this option does not appear for all forcefields.
- *Nonbond cutoff*: The cutoff value for spherical or real-space cutoff. The default value of 9.5 is usually good. This is a forcefield-sensitive option.
- Long range precision: Only used with PPPM and Ewald methods. Determines the reciprocal-space convergence. The default value of 0.00001 is usually a good value. This is a forcefield-sensitive option.
- Add tail corrections for Van der Waals interactions: The default is yes. This is a forcefield-sensitive option.
- Run LAMMPS on: Choose from CPU, OpenMP, and GPU.
 - CPU : Simulations will run on the CPU cores and parallelize over MPI ranks.
 - OpenMP : Simulations will run on the CPU cores and parallelize over OpenMP threads.
 - GPU : Simulations will run on the combination of CPU cores and GPU cards.

Hint: This option is only visible if the \Box **Enable running Job on GPU** checkbox from File >> Preferences... >> Miscellaneous is checked.

Note: *MedeA LAMMPS* supports NVIDIA GPU cards with compute capabilities (https://developer. nvidia.com/cuda-gpus) from 3.0 to 8.6. This includes the Kepler, Maxwell, Pascal, Volta, Turing, and Ampere series GPU cards. For HPC cluster compute nodes the Tesla Kepler, Pascal, Volta, and Ampere series are recommended. For workstations and desktops, GeForce Titan V, and Quadro GP100 and GV100 are recommended. Both Linux and Windows are supported.

4 Bias

The Bias section has one available stage: Orientation :

Edit stag	e: OrientationBias		
Subset:	Other		
Force:	5.0	kcal/mol/Ang	-
x:	1.0		
y:	0.0		
z:	0.0		

The parameters are:

- Subset: Add bias to this atom pair subset.
- Force: the magnitude of the added bias.
- *x*, *y*, and *z*: the direction of the added bias.

5 Single Point

The *Single Point* section has two available stages: Single Point Energy and Single Point Forces . Both of these stages have no adjustable parameters.



6 Minimization

The Minimization section has one available stage: Minimize .

The Minimize stage has the following parameters:

• *Relax cell*: Choose from No , Isotropically , With x=y , With y=z , With x=z , and Anisotropically . Depending on the choice, there are 0–6 cell parameters to fix (remaining unchanged during minimization), 0–6 stress values to relax each cell parameter to, and maximum volume change:

			Edit stage: Minimize	
Edit stage: Minimize			Optimization parameters	
Optimization parameters			Method:	Conjugate gradients 👻
	jugate gradients 👻		Linesearch:	Fast -
Linesearch: Fast			Maximum step:	0.05
Maximum step: 0.05			Energy tolerance:	0.0
Energy tolerance: 0.0			Force tolerance:	1.0
Force tolerance: 1.0			Maximum iterations:	1000
Maximum iterations: 1000			Maximum force evaluations	
Maximum force evaluations: 1000	00			10000
Trajectory Write trajectory			Trajectory Write trajectory	
Every (steps): 100			Every (steps): 100	
Cell relaxation			Cell relaxation	
Relax cell:	No		Relax cell:	Isotropically -
			Maximum volume change:	0.01
			Pressure 1.0	par 🗸
			·	
			Edit stage: Minimize	
			Optimization parameters Method:	
				Conjugate gradients 🔻
			Linesearch:	Fast
			Maximum step:	0.05
			Energy tolerance:	0.0
Edit stage: Minimize			Force tolerance:	1.0
Optimization parameters			Maximum iterations:	1000
Method: Co	onjugate gradients 👻		Maximum force evaluations	^{3:} 10000
Linesearch: Fa	ast 🔻		Trajectory	<u> </u>
Maximum step: 0.0	05		Write trajectory	
Energy tolerance: 0.0	0		Every (steps): 100	
Force tolerance: 1.0	0		-Cell relaxation	
	000		Relax cell:	Anisotropically 🔽
Maximum force evaluations: 10			Maximum volume change:	0.01
-Trajectory			Fix the cell in the x direc	ction
Write trajectory	_		Fix the cell in the y direc	stion
Every (steps): 100			Fix the cell in the z direc	
Cell relaxation			Fix the cell in the yz dire	
Relax cell:	With x=y		Fix the cell in the xz dire	
Maximum volume change:	0.01		Px: 1.0 Py:	1.0 Pz: 1.0
Fix the cell in the z direction				
Px, Py: 1.0 Pz: 1.0) bar 👻		Pyz: 0.0 Pxz:	0.0 Pxy: 0.0 bar
Method: Choose	e from Conjugate Gradie	en	it (CG), Ste	epest Descent (SD), and

Hessian-free truncated Newton (HFTN). The default is CG.

The CG method is the Polak–Ribiere (PR) version. At each iteration, the force gradient is combined with the previous iteration information to compute a new search direction perpen-

v. 3.8



dicular (conjugate) to the previous search direction. The PR variant affects how the direction is chosen and how the CG method is restarted when it ceases to make progress. The PR variant is thought to be the most effective CG choice for most problems.

With the SD method, at each iteration, the search direction is set to the downhill direction corresponding to the force vector (negative gradient of energy). Typically, steepest descent will not converge as quickly as CG but may be more robust in some situations.

With the HFTN algorithm, at each iteration, a quadratic model of the energy potential is solved by a conjugate gradient inner iteration. The Hessian (second derivatives) of the energy is not formed directly but approximated in each conjugate search direction by a finite difference directional derivative. When close to an energy minimum, the algorithm behaves like a Newton method and exhibits a quadratic convergence rate to high accuracy. In most cases, the behavior of HFTN is similar to CG, but it offers an alternative if CG seems to perform poorly.

Warning

When changing cell parameters, only the conjugate gradient and steepest descent methods can be used.

- · Linesearch: Choose from Fast, Quadratic, or Backtrack. The default is Fast.
- Maximum step: Maximum distance (Å) to move atoms. The default value of 0.05 is usually good.
- Energy tolerance: The default is 0.0 eV or kcal/mol.
- Force tolerance: The default is 1.0 eV/Å for metallic/NIST and COMB3 forcefields and 1.0 kcal/mol-Å for all other forcefields. For numerical stability, the force tolerance value is limited to a minimum of 1e-12 eV/Å or kcal/mol-Å. If a user specifies a smaller value than 1e-12, this value is overwritten with 1e-12 when generating the input files for LAMMPS.
- Maximum iterations: Maximum iterations for energy evaluations. The default is 1000.
- Maximum force evaluations: Maximum iterations for force evaluations. The default is 10000.
- Write trajectory and the default value for Every (steps): is 100.

7 Building and Editing

This section has two available stages: Set Cell and Compress Layer .

7.1 Set Cell

Edit stage: \$	Set Cell
Set cell	using density 🗸
De	ensity: \$rho_calc
Edit stage: S	Set Cell
Set cell	using volume 💌
Vo	lume: \$V_calc



Edit stage: S	et Cel	I		
Set cell	scali	ng by a factor	•	
Fa	ctor:	0.9]

- Set Cell: Changes the cell dimension by one of the following:
 - using density and its Density
 - using volume and its Volume
 - scaling by a factor and its Factor

All of the above options can be a value or a variable.

7.2 Compress Layer

The **Compress Layer** stage performs combined equilibration and compression of layered materials in the simulation cell, ensuring that none of the atoms in the system are outside the desired layer. First, a minimization is performed, then a compression with a set of "indenters" under an *NVT* ensemble, followed by one last minimization.

Edit stage: Compre	ss Layer					
-Layer						
Layer thickness:	Explicit			-		
Thickness:	15				Ang	-
Optimization para	meters					
Method:		Conju	gate grad	lients		•
Linesearch:		Fast		•		
Maximum step:		0.1]		
Energy tolerance:		0.0]		
Force tolerance:		1.0]		
Maximum iteration	ns:	5000]		
Maximum force e	valuations:	10000				
Equilibration/anne	ealing paran	neters-				
Time:	100		ps	•		
Time step:	\$tstep			•		
Initial Temperature	e: \$T			•		
Final Temperature	: \$т			•		

The **Compress Layer** stage has some similar options to the **Minimize** stage with the following exceptions:

- *Layer thickness*: Set the desired layer thickness with options of Explicit, From target density, or Use initial c dimension :
 - Explicit : Enter the desired layer thickness explicitly and the **Compress Layer** stage adjusts the c dimension accordingly.
 - From target density : Enter the desired layer density (excluding vacuum area) and the **Compress** Layer stage adjusts the c dimension accordingly.
 - Use initial c dimension : The final c dimension does not change.
- *Time*: Simulation time.



- *Time step*: Time step size for *NVT* integration.
- Initial Temperature and Final Temperature: Initial and final temperature for NVT integration.

Warning

A **Compress Layer** stage does not apply periodic boundaries in all 3 directions and must be used with layer perpendicular to Z periodicity and the Cutoff *Nonbond method*. It should therefore only be used by itself or with other **Compress Layer** stages. Results may be unpredictable if combined with other stages, such as those found in the *Minimize* or *Dynamics* sections that are normally used with 3-dimensional systems.

Warning

The **Compress Layer** stage may not work correctly when there are frozen atoms in the system. We highly recommend to un-freeze all of the atoms before utilizing a **Compress Layer** stage.

8 Dynamics

The *Dynamics* section has ten stages and the following stages are included in the standard *MedeA LAMMPS* license:

- Initialize Velocities
- NVE Ensemble
- NVT Ensemble
- NPT Ensemble

While the following stages (modules) require additional licenses:

- Cohesive Energy Density
- Thermal Conductivity
- Viscosity
- Diffusion
- Surface Tension
- Deposition

8.1 Initialize Velocities

The Initialize Velocities stage thermalizes the cell to a user-defined temperature:

Edit stage: velocities	
Initial Temperature:	\$T 🗸
✓ No net translation	
No net rotation	
Random Seed (positive integer):	72489
Assign a random seed autom	natically

• Initial Temperature: Thermalize the cell to this temperature.



- *No net translation*: Whether to ensure no net translational velocity is present in the system.
- *No net rotation*: Whether to ensure no net rotational velocity is present in the system.
- *Random Seed (positive integer):* A random seed for the random number generator. The seed can be assigned manually.
- Assign a random seed automatically: randomly generates a random seed.

8.2 NVE Ensemble

The NVE Ensemble stage performs time integration without any alterations to the equations of motion.

E	dit stage: NVE	E		
	Control A	nalysis		
	Time:	100	ps	•
	Time Step:	\$tstep		•
	Sampling:	10000	samples	-
	Trajectory:	0	frames	•

The parameters in the Control tab are:

- Time: Duration of the simulation run (defaults to 100 ps).
- Time Step: Time step size employed in solving the equations of motion.
- *Sampling:* Number of samples employed in performing averaging. This parameter does not affect dynamics.
- *Trajectory:* Number of trajectory frames saved during the molecular dynamics simulation. This parameter does not affect dynamics.

The Analysis tab allows you to add analyses to the dynamics stage. See The Analysis Tab for more details.

Warning

Using an **NVE** stage does not automatically guarantee an *NVE* ensemble. You should examine *Job.out* and the energy profile to ensure that the total energy is conserved so that an *NVE* ensemble is achieved.

8.3 NVT Ensemble

The NVT Ensemble stage performs time integration with a thermostat added to the equations of motion.



t stage: NVT	
Control Analys	s
nitial Temperatu	e: \$T
Final Temperatur	: \$т 🗸
Time:	100 ps 🔻
Time Step:	\$tstep 🗸
Controls	
Control:	Nose-Hoover
Damping (fs):	100.0
Drag (0.0 -> 2'ish	: 0.0
Trajectory	
Sampling: 100	0 samples 🔻
Trajectory: 0	frames 👻

The parameters, in addition to those in the NVE Ensemble stage, are:

- Initial Temperature and Final Temperature: These two parameters can be the same or different (for establishing cooling or heating). These parameters can be values or variables.
- · Control: Choose a thermostat algorithm from one of the following:

rescaling : interval (steps), window, amount of rescaling

Langevin : Damping (fs), Random Seed (integer)

Berendsen : Damping (fs)

Nose-Hoover : Damping (fs) and Drag

Hint: The default options of Nose-Hoover and Langevin are the good choices for a thermostat. The *Damping* parameter is recommended to be 100 times the time step size. For example, for a 1 fs time step, *Damping* is recommended to be 100 while for a 0.2 fs time step, the recommended value is 20.

Warning

Using an **NVT** stage does not automatically guarantee an *NVT* ensemble. You should examine *Job.out* and the temperature profile to ensure that temperature is maintained at the defined value so that an *NVT* ensemble is achieved.

8.4 NPT Ensemble

The NPT Ensemble stage performs time integration with both a thermostat and a barostat added to the equations of motion.



it stage: NP	т								
Control /	Analysis								
Initial Temp	erature:	\$т					•		
Final Tempe	erature:	\$т					•		
Initial Press	ure:	\$P					•		
Final Pressu	ure:	\$P					•		
Time:		100			ps		•		
Time Step:		\$tstep	D				-		
Cell								_	
Restrict cel	ll motion:	isotr	ор	ic				•	
Controls								_	_
Control:				Nose	-Hoo	ver	Т&	Р	•
Temperatu	re Dampi	ng (fs)	:[100.0)				
Pressure D	amping (fs):		100.0)				
Drag (integ	er):			0					
Trajectory									
Sampling:	10000		sa	mple	s	•			
Trajectory:	0		fra	ames		•			

The parameters, in addition to those in the **NVT** stage are:

- *Initial Pressure* and *Final Pressure*: These two parameters can be the same or different (for pressurization or depressurization). These parameters can be values or variables.
- Restrict cell motion: Controls how the cell volume and shape are equilibrated and relaxed:
 - *isotropic*: Only a, b, and c cell parameters (x, y, and z dimensions) are relaxed and the three components relax to one averaged value.
 - fixed angles: Also known as anisotropic with a, b, and c cell parameters relaxed independently.
 - *constrained*: Allows users to choose to relax any of the six cell parameters (a, b, c, alpha, beta, and gamma) independently.
 - unconstrained: All six cell parameters relax independently.

Warning

To relax the cell with the *constrained* or *unconstrained* options, the cell must be non-orthorhombic or you need to check the box *Allow orthorhombic cell angles to relax* in the **Initialize** stage.

• Control: Choose a combination of thermostat and barostat from the following options:

Rescaling/Berendsen : Interval (steps), Window, Amount of rescaling, Pressure Damping (fs), Bulk Modulus

Langevin/Berendsen : Damping (fs), Random Seed (integer), Pressure Damping (fs), Bulk Modulus

Berendsen/Berendsen : Damping (fs), Pressure Damping (fs), Bulk Modulus

Nose-Hoover T/Berendsen : Damping (fs), Drag (integer), Pressure Damping (fs), Bulk Modulus

Nose-Hoover T & P : Temperature Damping (fs), Pressure Damping (fs) and Drag (0 to 2'ish)

Hint: The default option of Nose-Hoover T & P is the recommended combination of thermostat and barostat for most systems.



Warning

Using an **NPT** stage does not automatically guarantee an *NPT* ensemble. You should examine *Job.out* and the temperature, stresses, and pressure profiles to ensure that temperature, stresses, and pressure are maintained at the defined value so that an *NPT* ensemble is achieved.

8.5 Cohesive Energy Density

The Cohesive Energy Density stage is a separate module available from Materials Design. It performs time integration under an *NVT* ensemble while making automated extractions and calculations to compute the cohesive energy density.

Edit stage: CED)		
Temperature:	\$T	•	
Time:	100	ps 🝷	
Time Step:	\$tstep	•	
Control:	Nose-Hoover	-	
	Damping(fs):	100.0	
	Drag (0.0 -> 2'isl	h): 0.0	
Sampling:	10000	samples 🔻	
Trajectory:	0	frames 💌	
	CED Sampling:	200	samples 🔻
	CED Cutoff:	12.0	

The parameters, in addition to those in the **NVT** stage are:

- *CED Sampling*: Calculate cohesive energy density every this many samples , steps , or amount of time.
- CED Cutoff: The cutoff value to calculate the energy.

Results are written in Job.out.

8.6 Thermal Conductivity

The **Thermal Conducitivity** stage is a separate module available from Materials Design and described in section Thermal Conductivity.

8.7 Viscosity

The Viscosity stage is a separate module available from Materials Design and described in section Viscosity.

8.8 Diffusion

The **Diffusion** stage is a separate module available from Materials Design and described in section Diffusion.

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8.9 Surface Tension

The **Surface Tension** stage is a separate module available from Materials Design. It performs time integration under an *NVT* ensemble while calculating the surface tension. It has the same parameters as the **NVT** stage.

8.10 Deposition

The **Deposition** stage is a separate module available from Materials Design and described in section MedeA Deposition.

9 Custom

The **Custom Code** stage is available with the standard *MedeA LAMMPS* license. It enables the addition of any custom LAMMPS commands not accessible from the above stages. Variables defined in any previous stages are passed into the **Custom Code** stage.

Edit stage: Custom			
Description:	Custom code		

Note: Characters in *MedeA LAMMPS* **Custom Code** stages with special meaning in Tcl (such as square brackets) will be interpreted in the usual Tcl manner (which can be convenient as this enables access to Flowchart variables in your LAMMPS custom stage). If you wish to avoid such interpretation, you can escape such characters with a preceeding backslash.

10 The Analysis Tab

The *Analysis* tab is available in the following stages: **NVE**, **NVT**, **NPT**, **Surface Tension**, and **Deposition**. You can add one or more of the following analyses to these stages:

• Distances : The Distance Analysis analyzes the interatomic distance between pairs of atoms defined by a pair subset. Therefore, the subset must be defined for the structure and entered in the *Pair subset:* box. The pair distances are analyzed every *Frequency* and plotted using the defined *Number of histogram bins.* Please see the tutorial Distance Analysis with LAMMPS [1] for more details.

^[1] https://download.materialsdesign.com/downloads/tutorial/Tutorial-Geometrical-analysis-in-LAMMPS.pdf



Edit stage: NVE					
x	Introl Analysis Distances 1 Title Distance analysis Pair subset:				
	OK Cancel Help				

Distributions: The Distribution Analysis analyzes the spatial distribution of the given Subset along the defined Direction, using spatial bins of the specified width. The Profile can be Number or Density/mass. The Number value means the number is computed for each chunk, i.e., number/chunk. The Density/mass value means the mass density is computed for each chunk, i.e. total-mass/volume/chunk. The Number of intervals defines how many analyses are performed throughout the simulation. Please see the tutorial Deposition of O2 on a Si Surface with Reactive Potentials [2] for more details.

Edit stage: NVE			
Control Analysis Distributions 1 Title: Distribution analysis Subset: OXY Vnumber of intervals: 10 Thickness of spatial bins (Ang Direction: z Profile: Density/mass Number add analysis Density/mass): 1.0		
	ОК	Cancel	Help

• PairCorrelations : The Pair Correlation Analysis plots the time-averaged pair correlation function (also known as the radial distribution function) between Subset 1 and Subset 2. These subsets must be created for the structure prior to setting up the flowchart, and these subsets must contain only one atom type each. The Number of intervals defines how many analyses are performed throughout the simulation using the defined Number of histogram bins.

Edit stage: NVE				
	X	trol Analysis Pair Correlations 3 Title: Pair Correlation analysis Subset 1: Subset 2: Number of intervals: 10 Number add analysis	of hi:	
		OK Cancel He	lp	

GroupInteractions: The Group Interaction Analysis computes the non-bond interactions between Subset 1 and Subset 2 and the interactions are time-averaged every Frequency. You can choose to include the pair interactions (van der Waals + Coulomb) and kspace contributions (long-range summation) or not. Additional keywords such as boundary and molecule can be added in the Additional Keywords

^[2] https://download.materialsdesign.com/downloads/tutorial/Tutorial-Deposition-of-O2-on-Si-surface-with-reactive-forcefields.pdf

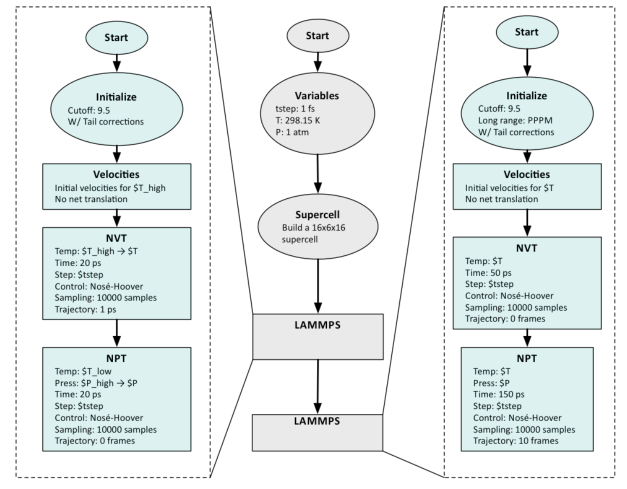
box. See compute group/group [3] for detailed usage of these keywords.

Edit stage: NVE					
	Imported Analysis Group Interactions 4 Title: Group Interactions analysis Subset 1: Import Prequency: 100 Import Prepuestor Import Prepuesto				

Note: GroupInteractions analysis only supports valence force fields (such as PCFF+).

11 Example

This flowchart illustrates the efficient use of MedeA LAMMPS flowcharts:



The Variables stage defines the overall parameters for temperature, pressure, and basic time step.

A larger supercell is constructed from the provided molecule.

[3] https://lammps.sandia.gov/doc/compute_group_group.html



Two different **LAMMPS** stages can be used. For instance, an equilibration stage that makes use of a computationally less demanding approach for non-bond interactions (e.g. *Cutoff*) can be used before a simulation with more computationally intense settings.