Simulating irradiation stability of 2D materials

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1 Introduction

In this experiment, we use different interatomic potentials to simulate the behaviour of a single-layer graphene lattice after a transfer of kinetic energy to an atom in the lattice, as would be the case in an electron microscope. We test this in the static-lattice approximation as well as for lattices undergoing thermal vibrations corresponding to different temperatures. In the latter case, we obtain statistical distributions of threshold energies for different temperatures that we analyse. Furthermore, we discuss the behaviour and compare the performance of the different potentials employed.

2 Theoretical background

A solid with a crystalline structure that only consists of a single layer of atoms is referred to as a 2D material. Today, various 2D materials with varying properties are studied, the most famous one being graphene, which consists of carbon atoms and has a hexagonal lattice structure with basis vectors

$$\vec{a}_1 = a \cdot \begin{pmatrix} \frac{\sqrt{3}}{2} \\ \frac{1}{2} \end{pmatrix} \qquad \vec{a}_2 = a \cdot \begin{pmatrix} -\frac{\sqrt{3}}{2} \\ \frac{1}{2} \end{pmatrix}$$

and lattice constant a = 2.46 Å [1].

In recent years, Scanning Transmission Electron Microscopy (STEM) has become a valuable tool for studying 2D materials. In contrast to (traditional) light microscopes, STEM devices use a focussed beam of electrons that is scanned over a sample in order to form an image. Since the de Broglie wavelength of electrons at relativistic velocities typically found in electron microscopes is orders of magnitude smaller than the wavelength of visible light, far better resolutions are possible through this technique. With the continuing development and improvement of aberration correctors, even higher resolutions are possible while at the same time reducing the necessary electron beam energy. [2]

2.1 Irradiation damage and displacement threshold energy

Relativistic electrons in electron microscopes carry high kinetic energies. Therefore, when scattered with a sample, they are sometimes able to cause defects to form in said lattice through different

mechanisms such as elastic and inelastic scattering or chemical etching. [2]

In this experiment, we study pristine graphene, which is most dominantly affected by elastic scattering. When an electron scatters elastically with an atom, it transfers some of its kinetic energy to the atom. In this process, kinetic energy and momentum are conserved. If the transferred energy was high enough, the atom is ejected from the lattice. The minimum energy required for this to happen is called the displacement threshold energy E_{KO} . [2]

2.2 Thermal vibrations

In the simplified case where the target atoms are at rest (static-lattice approximation), no damage is observed below E_{KO} , since the energy is then not high enough to cause the breaking of bonds and therefore the ejection of an atom. Above E_{KO} , every scattering process then leads to the ejection of an atom.

In reality however, the target atoms are not at rest due to thermal and quantum mechanical zeropoint vibrations. Depending on the out-of-plane velocity of the target atom during the scattering process, it is both possible for the transferred energy to be above E_{KO} but the atom to remain bound and for it to be below the threshold energy and the atom to be ejected [2]. Therefore, instead of a sharp increase in the ejection probability like in the static-lattice approximation, we expect the probability to rise smoothly with increasing electron beam energy. This results in a probability distribution for damage to occur that is sampled each time an electron is scattered on the sample.

2.3 Types of potentials

In atomistic simulations, interatomic potentials are necessary for computing energies and forces. The optimization process of traditional potentials relies on parameters fitted to energies calculated via Density Functional Theory (DFT), but mainly to experimental data. Such potentials consist of a rather small number of fitting parameters. Since they scale linearly with the number of atoms, computations are much faster compared to ab-initio-simulations. However, traditional potentials only provide limited accuracy.

In contrast, Machine learning potentials feature a much higher number of parameters which are fitted using ML approaches. Usually, they rely on a reference database for regression; interpolation of DFT-calculated energies is applied. In this process, local structural parameters for capturing the local atomic environment are learnt; this fitting process is not grounded in physical insights: in contrast to traditional interatomic potentials, the ML-generated potentials are directly fitted to the potential energy surface and not to physical properties, i.e., the ML architecture learns the Potential Energy Surface (PES) as a function of atomic configurations. The approximate PES will differ from exact DFT results; nevertheless, within the range of the training data, DFT-like accuracy can be achieved.

Furthermore, one can distinguish so-called physically informed potentials in which additional physical knowledge has been incorporated.[3]

2.4 ML potentials

Several Machine learning techniques can be used to fit the potentials, e.g. Kernel ridge regression. In the last few years, neural networks became very popular. In the context of interatomic potential learning, usually feedforward neural network architectures are used. They consist of an input layer, hidden layers and a final output layer. The input is passed through the layers which consist of

trainable weights and a non-linear (e.g. sigmoid or ReLU) activation function. For ML potentials, various descriptors (e.g. Zernike or Gaussian descriptors or Atomic cluster expansion) are used to describe the local environment; they provide processable data for the input layer. Usually, the descriptors encompass information about the atomic configuration of the neighbourhood of a specific atom up to a cut-off.

The optimization process of neural networks works via the minimization of a loss function (which is usually done via backpropagation). Due to its many trainable parameters (depending on the number of hidden layers), i.e. the weights of the network, they are highly flexible. This, however, causes the danger of overfitting, i.e. the network parameters get perfectly fit to the training data, but it generalises poorly.

In particular, the NN architecture has no physical foundation (i.e. it is a ,black box`) and in principle, highly non-physical behaviour could emerge. In general, one can expect DFT-like accuracy, as long as similarity with training configurations is given, but the behaviour outside is unpredictable. In any case, since the training process basically consists of interpolating the DFT energies but does not directly encompass physical knowledge, there is no guarantee that it is still physical. In contrast, traditional potentials which are fitted to physical properties should generalise comparatively well. [3]

2.5 Tested potentials

In the following, the performance of four carbon potentials has been tested:

The first one is a Tersoff-style three-body potential, in the following referred to as tersoff. This is a classical inter-atomic carbon potential which was proposed by Tersoff in 1988. It was the first multi-body potential which uses a bond order formalism to encompass the bonding state of an atom in the calculations.[4]

Another potential [5] uses the Gaussian Approximate Potential, referred to as GAP. GAP uses non-parametric kernel regression to express the non-linear function of the atomic positions as a linear function in the kernel [6]. It should be applicable to a broad range of configurations. It was trained on optB8-vdW DFT data; this method augments a semilocal exchange-correlation functional with a nonlocal correlation functional which takes dispersion interactions into account. For long range interactions an additional term is applied.

A hybrid neural network potential (`hNN-Grx') for multilayer graphene is presented in [7]. `Hybrid' refers to the combination of neural network output and theoretical physical knowledge (similar to the GAP potential described above): For long-range dispersion attraction, a Lennard-Jones-term is used, whereas the short-range interactions are computed via a neural network. The neural network was trained on a dataset consisting of 14500 mono-, bilayer graphene and graphite DFT calculations performed via VASP. It will be referred to as nn in this report.

Finally, a potential was created using the PYACE-package which allows fitting of interatomic potentials [8]. It relies on atomic cluster expansion (ACE) [9]; in the training step, neural networks are applied. We will refer to it as pyace.

3 Methods and Results

In the following, the process of testing the different potentials on both static-lattice and thermalized systems is shown in a step-by-step way, each step briefly described and immediately followed by

the results it produced.

3.1 Packages used

For the computations performed in the following, we used the ASE (Atomic Simulation Environment) [10]. Its interface allows us to run different atomistic simulation codes and import various potentials.

The Wen/Tadmor-potentials [3,4] are available via openKIM [11]. For Python, the kimpy-package provides an interface to the KIM-API.

```
[]: import ase.units as units
     from ase import Atoms
     from ase.build import graphene
     from ase.io import Trajectory
     from ase.md.verlet import VelocityVerlet
     from ase.visualize import view
     from ase.calculators import calculator
     from ase.md.velocitydistribution import MaxwellBoltzmannDistribution, Stationary
     from ase.md.langevin import Langevin
     from quippy.potential import Potential
     from ase.calculators.kim.kim import KIM
     from pyace import PyACECalculator
     import time
     from matplotlib import pyplot as plt
     import numpy as np
     import scipy as sc
     import re
```

3.2 Preparation of the system

For preparing the systems that will be simulated, we implement two functions: - get_middle() for retrieving the index of the atom in the approximate center of the grid produced by ASE - get_atoms() for actually generating a grid of given size with some vacuum above and below. Here, ASE provides a built-in function to automatically generate the lattice structure of graphene with the correct parameters for its atoms.

It is also important to note that we impose periodic boundary conditions for which we will perform tests in order to avoid finite-size effects.

```
[]: def get_middle(atoms : Atoms) -> int:
"""

For a given atoms list, returns the index of the center atom.
```

```
Parameters
------
atoms: ase.Atoms
The atoms array to get the dimensions from

Returns
-----
middle_atom: int
Index of the center atom
"""

middle_atoms = {7: 49, 6: 42, 5: 25, 4: 20, 3: 9, 2: 6, 1: 1}
return middle_atoms[int(np.sqrt(len(atoms)/2))]
```

```
[]: def get_atoms(size : tuple) -> Atoms:
         Returns a graphene lattice of dimensions size and with a vacuum layer of 10 \Box
      \hookrightarrow on
         each side of the lattice
         Parameters
         size : tuple
              The dimensions of the lattice to create (in units of unit cells)
         Returns
         _____
         qra : ase.Atoms
              The generated lattice
         11 11 11
         a = 2.45
         gra = graphene(a=a, size=size, vacuum=10)
         gra.center()
         gra.pbc = (True, True, True)
         return gra
```

3.3 Simulation code for static-lattice ejection simulations

ASE equips us with some very powerful tools to perform MD simulations. The main ideas are that we provide it with a propagation algorithm, a file name to which the trajectory of the simulation will be saved and a timestep and it then allows us to perform an arbitrary amount of simulation steps.

The function run_simulation() below is a wrapper for the ASE propagator that not only propagates the system but also gives the middle atom an initial momentum kick corresponding to the kinetic energy ekin specified. While propagating the system, the function then periodically checks whether

it thinks that the kicked atom was ejected or whether it returned to the lattice. This is done through a simple algorithm: If the atom's momentum is reversed, we know that the attracting force was great enough (i.e., the kinetic energy was small enough) to cause the atom to not be ejected. This part is fairly straightforward and there is no chance of incorrectly detecting an atom as not ejected.

The tricky part however is to detect whether the atom was ejected or not, since, as long as there is a force, we do not know whether it will be great enough to cause the atom to eventually return to the lattice. The way this was implemented is that we periodically check the relative change of momentum of the atom from one timestep to the other. If this change drops below a certain threshold, we label the atom as ejected.

This algorithm is not particularly trustworthy, since the threshold value must be found empirically and changes with parameters such as cell size and timestep. It is therefore always better to let the simulation run a certain number of timesteps until we are sure that the atom either returns or not. Choosing a threshold that is too small results in non-ejected atoms to be labelled as ejected and prematurely ending the simulation. However, for doing many simulations in order to optimise parameters, it is necessary to impose this restriction, since we cannot afford to let every simulation run until the end.

For the static-lattice approximation, we use the velocity verlet algorithm to propagate the system in an NVE (constant number of particles N, volume V and total energy E) ensemble.

```
[]: def run simulation(atoms : Atoms, calc : calculator, ekin : float, n_timesteps :
      → int, timestep : float, traj_file : str, check_ejected=True, printing=True, 
      →ejection_threshold=0.0001) -> str:
         Performs n timesteps of simulation steps using the specified calculator
      \hookrightarrow calc and writes
         trajectories to the file traj file. Uses timestep in fs and kicks the center
         atom of the atoms array with a kinetic energy ekin. If check_ejected is \Box
      ⇔true, the function
         will automatically try to determine whether the atom has been ejected or u
      ⇔whether it returns to
         the lattice, the latter with perfect accuracy, the former based on the rate_
      ⇔of change of
         momentum, which, for small timesteps, must be modified appropriately.
         Parameters
         _____
         atoms : ase.Atoms
             The lattice to use for the simulation
         calc : ase.calculators.calculator
             The calculator to use
         ekin: float
             The kinetic energy in eV to kick the center atom with
```

```
n_timesteps: int
       The number of simulation timesteps to perform (might be more or less_{\sqcup}
→depending on check_ejected)
  timestep : float
       The simulation timestep in fs
  traj file : str
       The name of the simulation file to write the trajectory to
  check_ejected : bool
       Whether or not to check for atom ejection. If an atom is detected as \sqcup
⇒either ejected or remained in the
       lattice, the simulation will stop and return the respective result
  printing : bool
       If True, prints status into console
  ejection_threshold : float
       Threshold percentage/100 value of momentum differential between two \Box
→runs for labeling atom as ejected
  Returns
  result: str
      If check\_ejected is True: ejected if the atom was ejected, not ejected\sqcup
⇔if the atom was not ejected or
      unknown if neither condition was met
  atoms.calc = calc
  # Integrator for the equations of motion, timestep depends on system
  dyn = VelocityVerlet(atoms, timestep * units.fs)
  momenta = []
  middle = get middle(atoms)
  # Saving the positions of all atoms after every time step
  with Trajectory(traj_file, 'w', atoms) as traj:
      dyn.attach(traj.write, interval=1)
       # Running one timestep before impact
      dyn.run(1)
       # Giving the target atom a kinetic energy of ene in the -z direction
       atoms[middle].momentum[2] = -(2 * ekin * atoms[-1].mass)**0.5
```

```
counter = 0 # Introduce a counter. If, after n_timesteps, the status is_{\sqcup}
⇔not determined, repeat the simulation a maximum of 2 times
       if not check_ejected:
           counter = 2
      while counter < 3:
           times = []
           for step in range(n_timesteps):
               start = time.time() # Measure time
               dyn.run(1) # Perform one simulation step
               times.append(time.time() - start) # Measure time
               momenta.append(atoms[middle].momentum[2]) # Save momentum for
⇔ejection check
               if not check_ejected:
                   # Continue loop if we do not check for ejection
                   continue
               if step > 10 and momenta[-1] > 0:
                   # Check for ejection by looking at the momentum of the atom.
→ If it has inverted, we know it is returning to the lattice
                   # and was therefore not ejected
                   if printing:
                       print(f"[ekin={ekin:0.1f}] Atom definitely not ejected_
→({np.mean(times)*1000} ms/step).")
                   return "not ejected"
               if (step+1) \% 10 == 0:
                   # Check for ejection. This part is tricky and one needs tou
⇒be careful with the percentage in the if statement below
                   if momenta[-1] <= 0 and np.abs((momenta[-1] - momenta[-2])/
→momenta[-2]) < ejection_threshold:</pre>
                       if printing:
                           print(f"[ekin={ekin:0.1f}] Atom ejected ({np.
→mean(times)*1000} ms/step).")
                       return "ejected"
           counter += 1
           if check ejected:
               if printing:
                   print(f"[ekin={ekin:0.1f}] No result, continuing simulation

¬for another {n_timesteps} timesteps ({counter}/3)")

  return "unknown"
```

The next step is to load the different potentials we will later use to propagate the system and to determine quantities like potential energy or temperature.

```
nn = KIM("hNN_WenTadmor_2019Grx_C__MO_421038499185_001")
tersoff = KIM("Tersoff_LAMMPS_Tersoff_1988_C__MO_579868029681_004")
pyace = PyACECalculator("PyACE/output_potential.yaml")
pyace.set_active_set("PyACE/output_potential.asi")
```

We are now ready to perform our simulations. There are two main parameters that we need to determine and that are different for each potential: The size of the system (measured in units of unit cells) and the timestep. Of course, we would like to choose values for those parameters that minimize computational effort whilst maintaining accuracy and convergence and avoiding finite-size effects.

For that, we employ the following procedure: 1. Fix the timestep to a low value we can be reasonably sure will produce stable results 2. Create a list of different cell sizes we'd like to test (we started at 7×7) 3. Starting with the largest cell size, perform ejection simulations for each cell size and find the energy at which the first ejection is observed respectively

We expect the energy of the first ejections to start changing with decreasing cell size, as finite size effects become more prevalent. Therefore, we choose the cell size to be the smallest one that produced the same ejection energy as the larger ones.

The next step is to determine the largest possible timestep and the procedure here is analogous to the one for the cell size. We perform multiple ejection simulations at increasing time steps and fixed cell size (to the size previously determined) and stop when we see the threshold energy change. Then, we simply choose the largest timestep at which no change was observed.

These optimizations, of course, have to be done for each potential, since we expect them to behave very differently in simulations.

```
[]: energies = np.arange(19, 19.5, 0.1)
     timestep = 1.4
     n_{timesteps} = 120
     potential = "gap"
     calc = eval(potential)
     sizes = [(5, 5, 1)]
     abort_on_ejected = False # Should the simulation for a specific cell size abort_
      →if ejection was detected for one of the energies
     for size in sizes: # Loop through all sizes of grids
         print(f"\nDoing size {size}")
         ejected = False # If ejection was detected, cancel simulation for the
      →current size (if abort_on_ejected is True)
         for ekin in energies:
             if ejected and abort_on_ejected:
                 continue
             traj_file = f'trajectories_final_1/{potential}_{ekin:0.1f}.traj' # Load_
      \hookrightarrow trajectory
```

```
atoms = get_atoms(size) # Prepare grid with current size
if run_simulation(atoms, calc, ekin, n_timesteps, timestep, traj_file,_
check_ejected=True) == "ejected":
ejected = True
```

Along with the optimizations, we performed some rudimentary benchmarking for the different potentials, i.e., for a fixed cell size of 7×7 , we measured the average time it took the propagator to perform one timestep. The results of the optimizations and the benchmarking are the following:

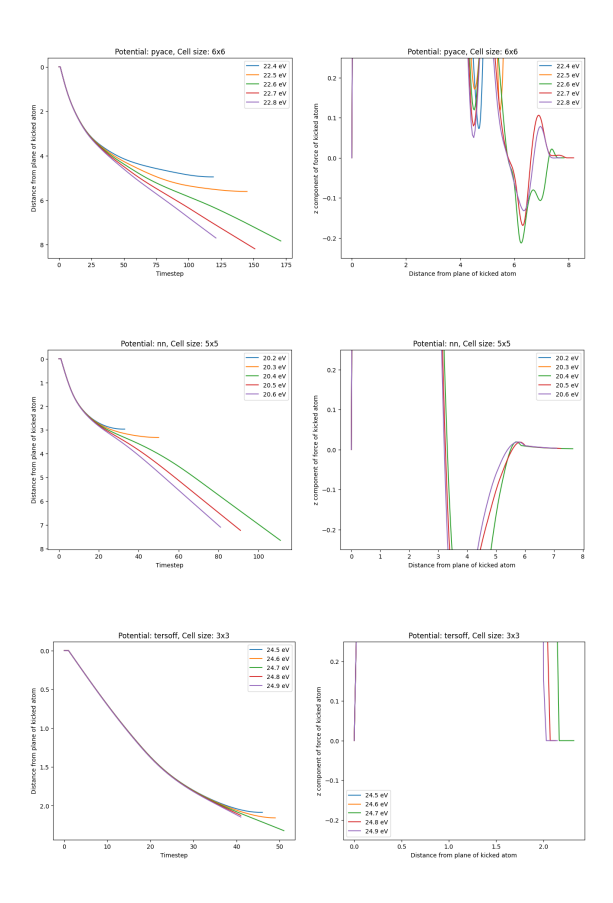
Potential	Minimum cell size	Maximum timestep	Energy of first ejection	Average simulation time/timestep
GAP	5×5	1,4 fs	19.2 eV	8444 ms
pyace	6×6	1,2 fs	22,6 eV	133 ms
nn	5×5	1,6 fs	20,4 eV	52 ms
tersoff	3×3	0.4 fs	24.7 eV	3.5 ms

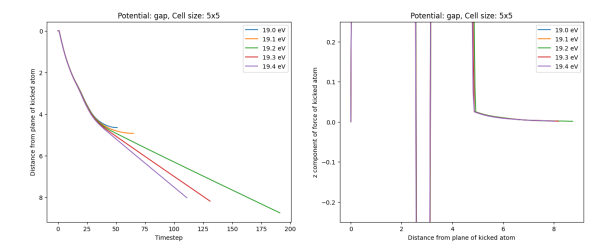
The first ejection energies are not only a metric for how well the simulation performs at different parameter values but are actually the threshold energies E_{KO} in the static-lattice approximation.

In the following, we are going to visualize a simulation for each potential performed using the parameters we just obtained. Furthermore, we will investigate the behaviour of the force acting on the ejected atom as a function of distance from the plane. We do this in order to determine whether the different potentials implement a sharp cut-off at a certain distance or whether the force decays smoothly, for example $\propto r^{-2}$ with r being the distance from the plane.

```
[]: # Dictionary for plots. The keys are the potential names for which to look in
     →the file system under trajectories_final_1/
    # and the values are the energy ranges to look for.
    →0.1), "tersoff": np.arange(24.5, 25.0, 0.1), "gap": np.arange(19, 19.5, 0.1)}
    for i, item in enumerate(potentials.items()):
        fig, ax = plt.subplots(1, 2, figsize=(16, 6))
        potential, energies = item
        positions = []
        forces = []
        for ekin in energies:
           traj = Trajectory(f"trajectories_final_1/{potential}_{ekin:0.1f}.traj")__
     →# Load trajectory for potential and kinetic energy
           middle = get_middle(traj[-1]) # Determine middle atom to know which_
     ⇒atom to plot for
           cell_size = int(np.sqrt(len(traj[-1])/2)) # For plot title
```

```
position = []
      force = []
      for j in range(len(traj)):
          position.append(traj[j][middle].position[2])
          force.append(traj[j].get_forces()[middle][2])
      positions.append(position)
      forces.append(force)
  ax[0].set_xlabel("Timestep")
  ax[0].set_ylabel("Distance from plane of kicked atom")
  ax[0].set_title(f"Potential: {potential}, Cell size:
ax[1].set_xlabel("Distance from plane of kicked atom")
  ax[1].set_ylabel("z component of force of kicked atom")
  ax[1].set_title(f"Potential: {potential}, Cell size:
for j, position in enumerate(positions):
      ax[0].plot(np.abs(np.array(position)-10), label=f"{energies[j]:0.1f}_u
→eV") # Plot distance from plane vs. timestep in left plot
  for j in range(len(forces)):
      ax[1].plot(np.abs(np.array(positions[j])-10), forces[j],__
□label=f"{energies[j]:0.1f} eV") # Plot force on kicked atom vs. distance
→ from plane in right plot
  ax[0].legend()
  ax[1].legend()
  ax[0].invert_yaxis()
  ax[1].set_ylim(-0.25, 0.25)
  plt.show()
```





As one can see from the left-hand side plots above, which show the distance of the kicked atom from the plane, we observe clear ejections for energies larger than the previously determined E_{KO} . On the other hand, we can see the atoms starting to return for energies below the threshold. In the latter cases, the simulation automatically stopped due to the criterion discussed above.

The plots on the right-hand side show the out-of-plane component of the force acting on the kicked atom as a function of distance from the plane. We can make different observations for the four potentials: - pyace: Here, we see that the force does not suddenly drop to zero but rather that it decays smoothly. Also, it is interesting to note that this decay does not seem to start at a fixed distance from the plane as is the case for the other potentials. - nn: With this potential, we observe a rather sharp drop of the force to a small value which then quickly approaches zero. This initial drop happens at a distance of 6 Å from the plane, indicating a fixed cut-off distance in the potential. - tersoff: In this case, the sharp drop to zero is most noticeable. Although not exactly at the same distance for each trajectory, at about 2 Å from the plane, the force suddenly drops to zero with no smooth decay whatsoever. - GAP: For this potential, the same behaviour as in the case of the nn potential albeit at a distance of 5 Å from the plane can be observed: A sudden drop of the force to a small value followed by a smooth but quick decay to zero.

To conclude the static-lattice simulations, we store the obtained values for the timesteps and the cell sizes in dictionaries for later use.

```
[]: TIMESTEPS = {tersoff: 0.4, pyace: 1.2, nn: 1.6, gap: 1.4}
SIZES = {tersoff: (3, 3, 1), pyace: (6, 6, 1), nn: (5, 5, 1), gap: (5, 5, 1)}
```

3.4 Simulations for a thermalized system

In this section, we investigate the effects of a thermalized system, i.e., lattice atoms that are undergoing thermal vibrations corresponding to different temperatures. As discussed in the theoretical background, we now expect the ejection procedure to be a stochastic process. Therefore, we will need to perform many simulations for each potential and at different temperatures in order to obtain probability distributions.

The static-lattice simulations have shown that the GAP potential has a simulation time per timestep

that is orders of magnitude higher than those of the other potentials. Since, for a thermalized system, we will need to perform a very large number of simulations that already take a long time even for the faster potentials, it is not possible for us to continue investigating the GAP potential. We will therefore restrict our analysis on the three remaining potentials.

In order to being able to perform ejection simulations on a thermalized system, we first need to produce such a system and equilibrate it. A factor that has not been taken into account before is internal stress: Although we prepare the systems in the <code>get_atoms()</code> function with the equilibrium lattice constant for graphene, the different interatomic potentials result in the system being prepared with internal stresses. In order to get rid of those, we need to re-scale the cell with a certain factor, which, in the following, will be referred to as the scaling factor.

Determining the scaling factors for the different potentials is relatively simple. Thermodynamically speaking, our goal is to equilibrate the system with respect to its volume, i.e., we want to minimize the equation of state

$$-p = \left(\frac{\partial U}{\partial V}\right)_{S, N}$$

where U is the energy, V is the volume, p is the pressure (in our case: stress), S is the entropy and N the number of particles. The subscripted variables will be held constant.

The function determine_scaling_factors() implements the procedure of first generating a system that is not in equilibrium through the get_atoms() function, re-scaling it with a range of scaling factors and then finding the scaling factor for which the total potential energy is minimal. This is equivalent to setting the pressure (stress) to zero and finding the corresponding system volume. This is done for all the potentials given to the function. We immediately run the function for the three potentials that we will be performing simulations on and store the scaling factors in a dictionary.

```
for calc in calcs:
    energies = np.zeros(factors.size)

for i, factor in enumerate(factors):
    atoms2 = atoms.copy()
    atoms2.set_cell(atoms.cell*factor, scale_atoms=True)
    atoms2.calc = calc
    energies[i] = atoms2.get_potential_energy()

scaling_factors[calc] = factors[np.argmin(energies)]

return scaling_factors

scaling_factors = determine_scaling_factors([tersoff, pyace, nn])
```

With the static system now being in equilibrium, we can start thermalizing and equilibrating it. The following function <code>generate_equilibrated_system()</code> does exactly that and supports thermalization and quilibration in two different ensembles: The NVE ensemble and the NVT ensemble.

Thermalization in the NVE ensemble is achieved by assigning each atom in the static lattice a random momentum drawn from a Maxwell Boltzmann (MB) distribution. The temperature we use for the MB distribution is double the temperature we want our system to have after equilibration, since, due to the equipartition theorem, the initial average kinetic energy of $\langle E_{kin}^{init} \rangle = k_B 2T$ will, in equilibrium, be equally partitioned on the kinetic and potential energy, i.e., $\langle E_{kin}^{eq} \rangle = \langle E_{pot}^{eq} \rangle = k_B T$.

On the other hand, the approach for thermalizing in the NVT ensemble is to use Langevin dynamics, which implements a thermostat that couples the system to a heat bath. After equilibration, the system will have the desired temperature, at which point we can switch to an NVE ensemble and equilibrate for a couple more time steps.

In addition to the <code>generate_equilibrated_system()</code> function, we also implement a function <code>plot_temperature()</code> that takes a simulation trajectory and plots the system's temperature as a function of time. We use this function later to determine whether the systems we prepared actually equilibrated to the correct temperatures.

```
T: float
       The temperature in Kelvin used for the MB distribution
  n_{equilibration}: int
      Number of equilibration timesteps to perform
   timestep : float
       Timestep to use for equilibration in fs
   calc : ase.calculators.calculator
       Calculator to use for equilibration
  scaling_factors : dict
      Dictionary with scaling factors for cell that contains the calculator ...
\neg used as a key
  out_trajectory : bool
       Whether or not to return the equilibration trajectory
  NVT : bool
       Equilibrate in NVT rather than in NVE ensemble
  filename : str
       Name of the trajectory file to write to. If none, use temporary file_
⇔and return trajectory instead
  Returns
  gra : Atoms
       The equilibrated graphene lattice
   traj : Trajectory
       The equilibration trajectory, only returned if out_trajectory is True_
⇔and no filename is given
  11 11 11
  gra = get_atoms(size)
  gra.set_cell(gra.cell*scaling_factors[calc], scale_atoms=True)
  gra.calc = calc
  if not NVT:
      MaxwellBoltzmannDistribution(gra, temperature_K=2*T) # Thermalize atoms.
→ Use 2*T to get desired kinetic energies due to equipartition theorem
      Stationary(gra) # Move into center of mass reference frame
  if filename == None:
      filename = f"/tmp/traj_equil_{time.time()}.traj"
```

```
traj = Trajectory(filename, "w", gra) if out_trajectory else None
    if NVT:
        Langevin(gra, timestep=timestep*units.fs, trajectory=traj,__
 otemperature_K=T, friction=0.01/units.fs).run(n_equilibration) # Equilibrate⊔
 ⇔system in NVT ensemble
    else:
        VelocityVerlet(gra, timestep=timestep*units.fs, trajectory=traj).
 →run(n_equilibration) # Equilibrate system in NVE ensemble
    if not traj == None:
        traj.close()
    if out_trajectory and filename.startswith("/tmp"):
        return gra, Trajectory(filename)
    return gra
def plot_temperature(traj : Trajectory, const=None) -> None:
    Plot the temperature of a trajectory
    Parameters
    _____
    traj : Trajectory
        The trajectory to plot the temperature from
    const : float
        Temperature to draw hline at
    fig, ax = plt.subplots(1, 1, figsize=(10, 7))
    temp = np.zeros(len(traj))
    for i in range(len(traj)):
        temp[i] = traj[i].get_temperature()
    if const != None:
        ax.hlines(const, 0, len(traj), ls="--", color="gray", label=f"T={const}_\( \)
 ⇔K")
    ax.plot(temp, label="Simulation temperature")
    if const != None:
        ax.legend()
    ax.set_xlabel("Timestep")
    ax.set_ylabel("Temperature [K]")
    plt.show()
```

The next function we implement (propagate_system()) is for propagating a thermalized system in the NVE ensemble for a given number of time steps and writing the trajectory to a file.

```
[]: def propagate_system(atoms : Atoms, n_timestep : int, timestep : float,__

¬filename=None) → None:
         .....
         Propagate a system of atoms using Velocity Verlet
         Parameters
         _____
         atoms : Atoms
             System to propagate
         n_timestep : int
             Number of timesteps to propagate for
         timestep : float
             Timestep in fs
         filename : str
             If not None, save the trajectory to a file
         traj = Trajectory(filename, "w", atoms) if filename != None else None
         VelocityVerlet(atoms, timestep=timestep*units.fs, trajectory=traj).
      →run(n_timestep) # Propagate system in NVE ensemble
         if not traj == None:
             traj.close()
```

The idea for the simulations that we will do is that we first generate and equilibrate systems in different potentials and at different temperatures and subsequently propagate them for a large number of timesteps. This way, we will end up with very long trajectories that we can use for ejection simulations. Then, we use multiple frames of each trajectory to perform ejection simulations and determine the threshold energy, resulting in a large sample of threshold energies for each trajectory, i.e., for each temperature and potential.

In order to avoid correlations between the sampled threshold energies in a trajectory, we need to carefully choose the number of timesteps to wait between the frames that we use for ejection simulations. For this, we can use the time autocorrelation function of the momentum of the kicked atom, which should ideally approach zero after some time, when correlations have died away. We then know how long we have to wait between simulation frames in order to have uncorrelated initial configurations.

The function autocorrelate_momentum() calculates the time autocorrelation function of the kicked atom's momentum using NumPys correlate() function.

```
[]: def autocorrelate_momentum(traj : Trajectory) -> np.ndarray:
         Compute the autocorrelation function for the z component of the momentum of \Box
      ⇒the middle atom of the
         given trajectory
         Parameters
         traj : Trajectory
             The trajectory to use
         Returns
         autocorrelation : np.ndarray
             The autocorrelation function values for the trajectory
         middle = get_middle(traj[-1])
         M = len(traj)
         momenta = np.zeros(M)
         for i in range(M):
             momenta[i] = np.linalg.norm(traj[i][middle].momentum) # Store magnitude_
      →of the target atom's momentum
         momenta -= np.mean(momenta) # Remove com motion
         autocorrelation = np.correlate(momenta, momenta, mode="full")[-M:] / np.
      \rightarrowarange(M, 0, -1)
         autocorrelation /= np.std(momenta)**2 # normalization
         return autocorrelation
```

For each potential, we now generate an equilibrated thermalized system.

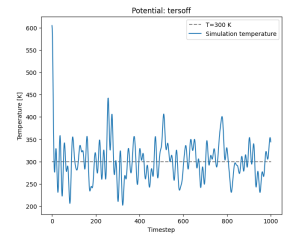
```
# Determining n_steps to simulate in order to avoid correlations, using large_
system size

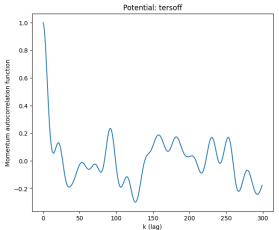
autocorrelation_test_potentials = {"tersoff": tersoff, "pyace": pyace, "nn": nn}

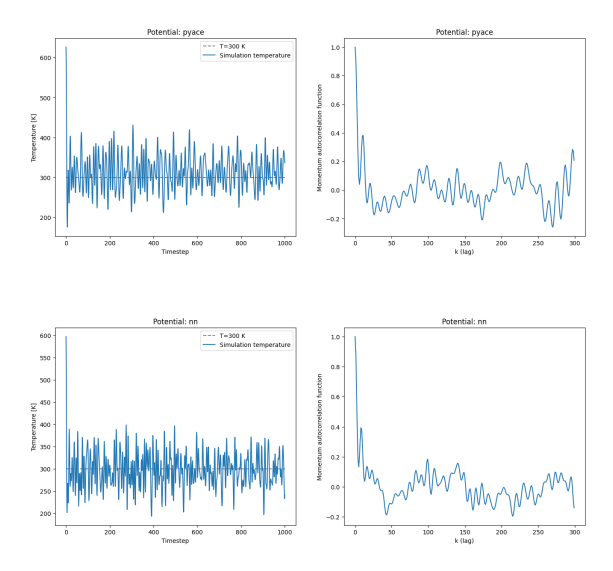
for name, calc in autocorrelation_test_potentials.items():
    equil_test = generate_equilibrated_system(size=(6, 6, 1), T=300,__
n_equilibration=1000, timestep=TIMESTEPS[calc], calc=calc,__
scaling_factors=scaling_factors, out_trajectory=True, NVT=False,__
filename=f"trajectories_correlations/{name}.traj")
```

We then apply the autocorrelate_momentum() function to those systems and plot the results.

```
[]: # Plot the results
     for name, calc in autocorrelation_test_potentials.items():
         traj_equil_test = Trajectory(f"trajectories_correlations/{name}.traj")
         fig, ax = plt.subplots(1, 2, figsize=(16, 6))
         temp = np.zeros(len(traj_equil_test))
         for i in range(len(traj equil test)):
             temp[i] = traj_equil_test[i].get_temperature()
         ax[0].hlines(300, 0, len(traj_equil_test), ls="--", color="gray", __
      ⇔label="T=300 K")
         ax[0].plot(temp, label="Simulation temperature")
         ax[0].legend()
         ax[0].set_xlabel("Timestep")
         ax[0].set_ylabel("Temperature [K]")
         ax[1].plot(autocorrelate_momentum(traj_equil_test)[:300]) # Plot_
      \rightarrowautocorrelation function up to k = 100 to avoid finite size effects
         ax[1].set_xlabel("k (lag)")
         ax[1].set_ylabel("Momentum autocorrelation function")
         ax[0].set_title(f"Potential: {name}")
         ax[1].set_title(f"Potential: {name}")
         plt.show()
```







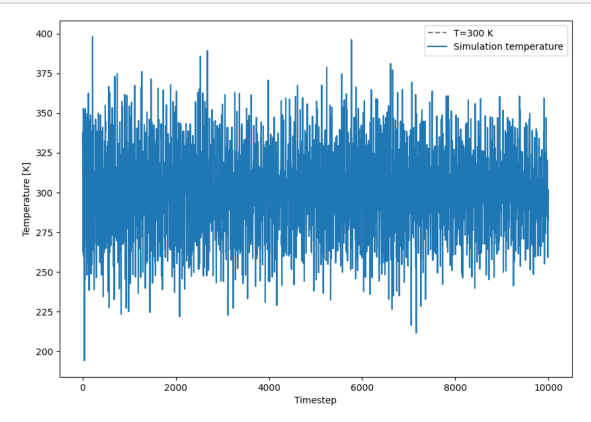
In the left-hand side plots above, one can see the temperature curve of the propagated systems for each potential. On the right-hand side, the corresponding autocorrelation functions are plotted. Although they are by far not ideal, probably due to the small system size of 6×6 , we can be reasonably sure that correlations have mostly died away after 100 time steps, so we will use this value in the following simulations.

With all the preparations done, we can now start to actually generate the before-mentioned long trajectories. For each potential, we will generate four trajectories, one for each temperature in $\{100\mathrm{K},\,200\mathrm{K},\,300\mathrm{K},\,350\mathrm{K}\}$. Each trajectory has a length of 10.000 time steps after equilibration (with 2.500 equilibration times steps). In order to minimize finite-size effects, we choose a system size of 6×6 for all potentials, which is the largest system size that still takes a reasonable time to simulate. For time steps, we use the values we previously determined through optimization.

This leaves us with a total number of 101 frames per trajectory (i.e., potential and temperature) that we can use to perform ejection simulations and subsequently produce a probability distribution

for the threshold energies.

The following plot shows the temperature curve of one of those long trajectories (pyace at 300K).



The last step of simulations is to perform ejection simulations on all the trajectories we just generated. For each frame in each trajectory, we simulate until we reach the first ejection, record the energy at which this first ejection was achieved and save it in a corresponding .csv file for later analysis, such that we are left with one results file per trajectory containing all the recorded threshold energies.

It is important to note that we do not use the algorithm that tries to detect ejection described above, since it has proven to be too unstable for thermalized systems. So, we let every simulation run until the end, unless a reversal of momentum was detected, i.e., the atom was not ejected. This is slower but produces considerably better results.

```
[]: temps = [100, 200, 300, 350] # Temperatures to test
     potentials = {"tersoff": tersoff, "nn": nn, "pyace": pyace} # Potentials to test
     #potentials = {"nn": nn}
     energies = {tersoff: np.arange(22.0, 27.5, 0.1), pyace: np.arange(20.0, 25.5, 0.
      \rightarrow 1), nn: np.arange(18.0, 24.0, 0.1)}
     thresholds = {} # Dict for the results (keys are potentials and temperatures,
      →and values are lists of threshold energies)
     # Prepare dictionary with empty lists
     for name in potentials.keys():
         for temp in temps:
             thresholds[f"{name}@{temp}"] = []
     for temp in temps:
         for name, calc in potentials.items():
             traj = Trajectory(f"trajectories_vibrations/{name} {temp}.traj") # Loadu
      ⇔the prepared trajectory
             key = f"{name}@{temp}"
             for frame in range(0, len(traj), 100):
                 atoms = traj[frame] # Load the frame (we go through the whole_
      →trajectory file and always skip 100 frames due to correlations)
                 for ekin in energies[calc]:
                     prefix = f''[{name}_{0}[{temp}_{K}] [{ekin:0.1f}] "
                     filename = f"/tmp/trajectories_vibration_ejection.traj" #_
      → Temporary file to write trajectories to (we don't need them)
                     # We simulate until one simulation ran through completely_
      without the atom having returned (momentum having reversed) before
                     # We set timesteps to 80 but are really simulating 80*3=240
      ⇔timesteps, so we should be fine
                     if run_simulation(atoms.copy(), calc, ekin, 80, ⊔
      →TIMESTEPS[calc], filename, check_ejected=True, printing=False,
      →ejection_threshold=0) != "not ejected":
```

3.5 Analysis of the thermalized system simulations

With the simulations done, we can now start to reconstruct the sampled ejection probability distributions. For that, we first define two functions that allow us to load the .csv files produced in the previous simulation. load_results() loads only a specific sample with given potential and temperature, while load_all_results() returns a dictionary of all samples it could find, organized by potential and temperature.

```
[]: def load_results(potential : str, temp : float) -> np.ndarray:
        Loads a specific result for the simulation of the threshold energy of a
         thermalized system.
        Parameters
         _____
        potential : str
             The name of the potential used in the simulation
         temp : float
             The temperature used in the simulation
        Returns
         _____
         result : np.ndarray
             The first ejection energies obtained in the simulation
        path = f"results_vibrations/{potential}@{temp}.csv"
        assert os.pyth.isfile(path), f"Result {path} does not exist."
        return np.loadtxt(path, delimiter=",")
     def load_all_results() -> dict:
        Load all the results found in results_vibrations/ and return it as a dict
        Returns
         loaded_results : dict
```

```
A dictionary containing all the results, like the one produced in the
\hookrightarrow simulation that
       generated the .csv files
       Keys: potential@temperature
       Values: np.ndarray of the obtained threshold energies
  potentials : list
       The names of the potentials that were loaded
   temps : list
       The temperatures that were loaded
  pattern = re.compile(r"^[a-zA-Z0-9\s]+0[0-9]+(?:\.[0-9]+)?\.csv$", re.
→IGNORECASE)
  files = [f for f in os.listdir("results_vibrations") if os.path.isfile(os.
→path.join("results_vibrations", f)) and pattern.match(f)]
  loaded_results = {}
  for file in files:
       loaded_results[file[:-4]] = np.loadtxt(os.path.

→join("results_vibrations", file), delimiter=",")
  potentials = list(set([name.split("0")[0] for name in loaded_results.
⇒keys()])) # Find all unique potential names
  temps = list(set([float(name.split("@")[1]) if "." in name.split("@")[1]__
⇒else int(name.split("@")[1]) for name in loaded_results.keys()])) # Find all_
→unique temperatures
  return loaded_results, potentials, temps
```

Since we want to fit a normal (gaussian) distribution to the data, we need to implement the corresponding probability density function

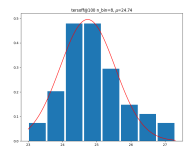
$$f(x) = \frac{1}{\sigma\sqrt{2\pi}} \cdot \exp\left(-\frac{(x-\mu)^2}{2\sigma^2}\right)$$

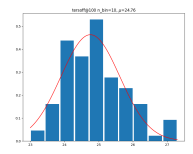
with the mean μ and the standard deviation σ .

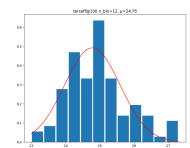
```
[]: def normal(x, mu, sigma):
    return 1/(sigma*np.sqrt(2*np.pi)) * np.exp(-0.5*np.power((x - mu)/sigma, 2))
```

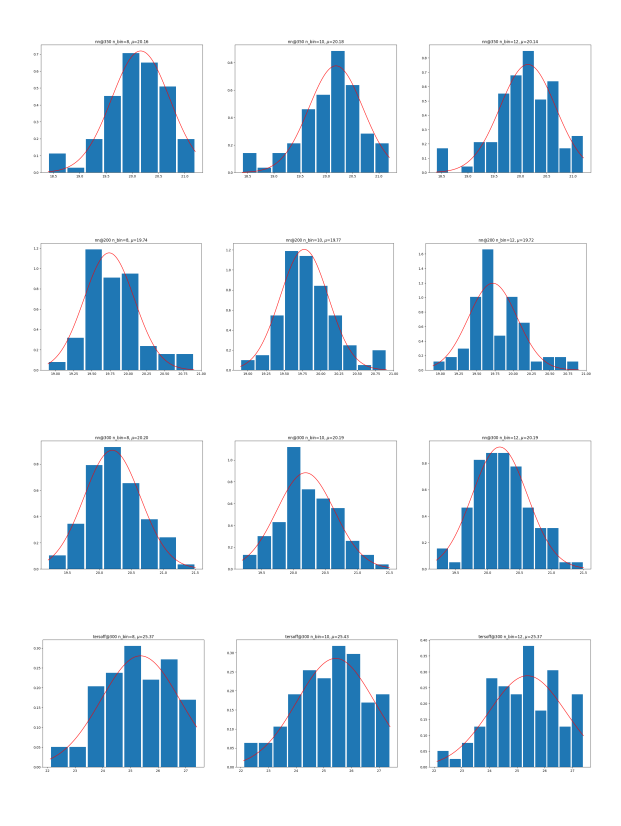
We will be fitting the normal distribution to a histogram with bins of equal width. To determine the ideal number of bins, we plot the histograms of all the trajectories with the corresponding fit for different bin counts and visually inspect the results.

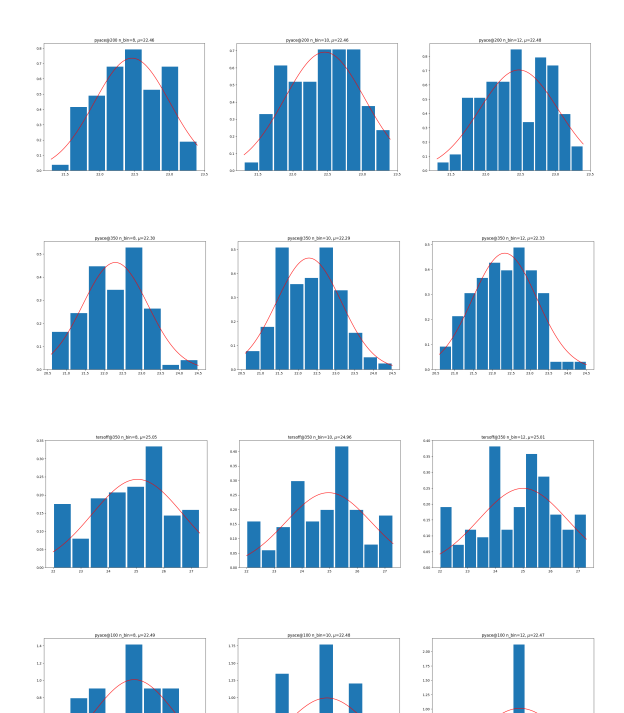
```
[]: # Analyzation code for the results of the thermalized threshold energies, u
      ⇔deciding on a number of bins to use
     loaded_results, potentials, temps = load_all_results()
     n_bins = [8, 10, 12]
     for key, result in loaded_results.items():
         fig, ax = plt.subplots(1, len(n_bins), figsize=(10*len(n_bins), 7))
         for i, n_bin in enumerate(n_bins):
             pdf, bins = np.histogram(result, density=True, bins=n_bin)
             bin_centers = (bins[1:] + bins[:-1])/2 # Calculate bin centers
             bin_width = (bins[1:] - bins[:-1])[0]
             p0 = [np.mean(result), np.std(result)]
             popt, pcov = sc.optimize.curve_fit(normal, bin_centers, pdf, p0=p0)
             mu, sigma = popt[0], popt[1]
             x = np.arange(bins[0], bins[-1], 0.01)
             ax[i].bar(bin_centers, pdf, align="center", width=bin_width/1.1)
             ax[i].plot(x, normal(x, *popt), color="red")
             ax[i].set_title(f"\{key\} n_bin=\{n_bin\}, \mus=\{mu:0.2f\}")
         plt.show()
```

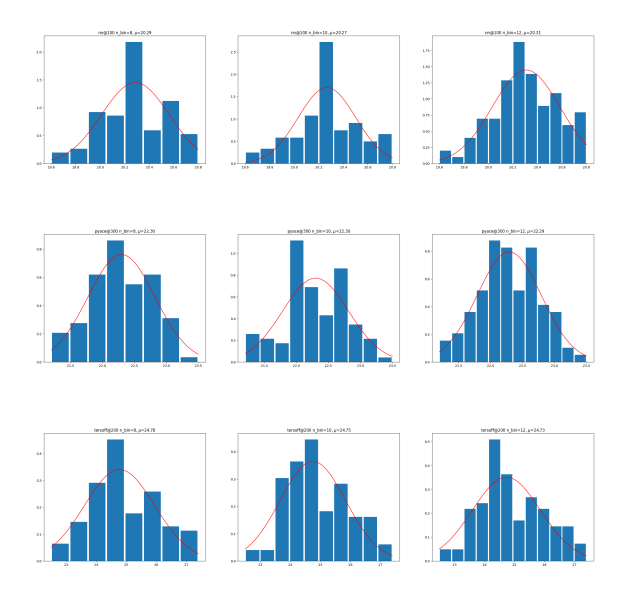












This process could be improved with advanced binning strategies, but this goes beyond the scope of this experiment. For now, we decide on a bin count of 8.

The last step we do in our analysis is to fit the normal distribution to the histogram of each trajectory. The plots containing the histograms are sorted by temperature. The plots below those are then comparing the probability density functions at different temperatures of each potential.

Since we expect the spread of threshold energies to increase with temperature, as lower as well as higher energies then become more likely to cause ejections, we also calculate the full width half maximum (FWHM) of each probability density function and visualize them in the last plot, which shows the FWHMs as a function of temperature for each potential.

```
[]: # Doing the plots separated by temperature for 8 bins per potential with 

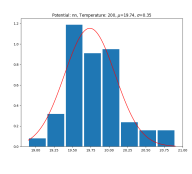
→histogram
```

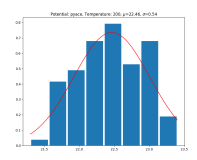
```
loaded_results, potentials, temps = load_all_results()
# Organize the results by temperature
temp_results = {temp: {key.split("@")[0]: loaded_results[key] for key in_
 →loaded_results.keys() if int(key.split("@")[1]) == temp} for temp in temps}
n bin = 8
fit_params = {potential: {} for potential in potentials} # Store the fit_
 ⇒parameters found for later use
for temp in temp results.keys():
   n_potentials = len(temp_results[temp].keys())
   fig, ax = plt.subplots(1, n_potentials, figsize=(10*n_potentials, 7))
   for i, potential in enumerate(temp_results[temp]):
       result = temp_results[temp][potential]
       pdf, bins = np.histogram(result, density=True, bins=n_bin)
       bin_centers = (bins[1:] + bins[:-1])/2
       bin_width = (bins[1:] - bins[:-1])[0]
       p0 = [np.mean(result), np.std(result)]
       popt, pcov = sc.optimize.curve_fit(normal, bin_centers, pdf, p0=p0)
       mu, sigma = popt[0], popt[1]
       fit params[potential][temp] = (popt, bins[0], bins[-1])
       x = np.arange(bins[0], bins[-1], 0.01)
        ax[i].bar(bin_centers, pdf, align="center", width=bin_width/1.1)
        ax[i].plot(x, normal(x, *popt), color="red")
        ax[i].set_title(f"Potential: {potential}, Temperature: {temp},__

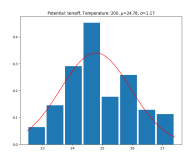
→$\mu$={mu:0.2f}, $\sigma$={sigma:0.2f}")

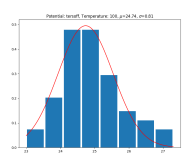
   plt.show()
# PDFs for different temperatures per potential
n_potentials = len(fit_params.keys())
fig, ax = plt.subplots(1, n_potentials, figsize=(10*n_potentials, 7))
for i, potential in enumerate(fit_params.keys()):
   x_{min}, x_{max} = 1000, 0
```

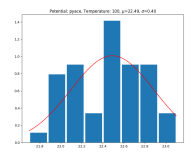
```
for temp in fit_params[potential].keys(): # Find the plotting range needed_
 →to plot all PDFs of the same potential in the same plot
        params = fit_params[potential][temp]
        if params[1] < x min:</pre>
            x_min = params[1]
        if params[2] > x max:
            x_max = params[2]
    for temp in fit_params[potential].keys():
        params = fit_params[potential][temp]
        FWHM = 2.355*params[0][1] # Calculate FWHM
        x = np.arange(0.95*x_min, 1.05*x_max, 0.01)
        ax[i].plot(x, normal(x, *params[0]), label=f"T = {temp} K, $\mu = 0
 \Rightarrow{params[0][0]:0.2f}, FWHM = {FWHM:0.2f}")
    ax[i].set_title(f"PDF of threshold energy for {potential}")
    ax[i].legend()
    ax[i].set_xlabel("Threshold energy [eV]")
    ax[i].set_ylabel("PDF for ejection")
plt.show()
# FWHMs per potential as a function of temperature
fig, ax = plt.subplots(1, n_potentials, figsize=(10*n_potentials, 7))
for i, potential in enumerate(fit_params.keys()):
    FWHMs = []
    for temp in fit_params[potential].keys():
        params = fit_params[potential][temp]
        FWHMs.append(2.355*params[0][1])
    ax[i].scatter(list(fit_params[potential].keys()), FWHMs)
    ax[i].set_title(f"FWHMs of PDF of threshold energies for {potential}")
    ax[i].set_xlabel("Temperature [K]")
    ax[i].set_ylabel("FWHM [eV]")
plt.show()
```

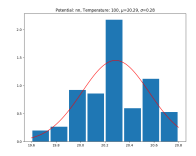


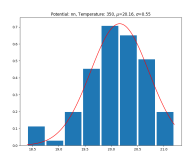


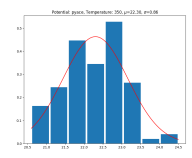


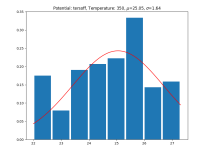


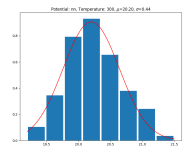


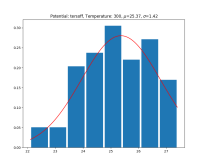


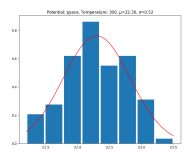


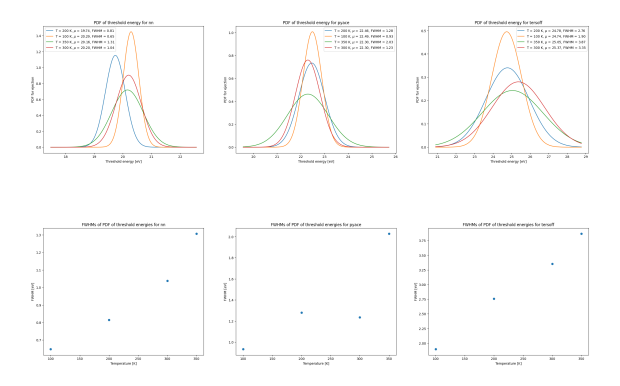












As one can see, the FWHMs in the last plot grow with increasing temperature, except for one data point for the pyace potential at 300 K. The average displacement threshold energies obtained from the fits are the following:

Potential	Average displacement threshold energy with standard error		
pyace	$(22.4 \pm 0.1) \text{ eV}$		
nn	$(20.1 \pm 0.2) \text{ eV}$		
tersoff	$(25.0 \pm 0.2) \text{ eV}$		

4 Discussion

In this experiment, we investigated the behaviour of graphene after momentum transfer to one of the lattice atoms using different inter-atomic potentials, both for the static-lattice approximation as well as for thermalized systems.

4.1 Comparison of results to previous studies

In a previous study [12], Susi et al. used DFT molecular dynamics to determine the displacement threshold energy for thermalized graphene systems and obtained results in an energy range of [21,25 eV; 21,375 eV], which they found to overestimate the result of 21,14 eV obtained from experiments. Nevertheless, this gives us a basis to compare our results to.

The potential that comes closest to the experimental value is the nn potential, followed by pyace. Tersoff generally yielded the worst results when compared to previous findings. Concerning the DFT-MD results, pyace is closest to the range mentioned above, with nn in second place.

Another study done by Chirita Mihaila et al. [13] used DFT tight-binding based MD to investigate the displacement threshold of thermalized graphene. In their article, they produced a similar plot to the FWHM plot we obtained at the end of our experiment. Comparing both plots, none of our potentials managed to reproduce these results, with pyace and nn coming closest. It has to be pointed out that this could partially be because our sample size was more than 4 times smaller than the one in [13] and that pyace, apart from the FWHM value at 350 K, comes somewhat near the study's results. Also, it is worth mentioning that the small cell size of 6×6 we used for the thermal simulations yields high variability in the system's temperature, which, combined with the rather small sample size, could negatively impact the accuracy of the results.

4.2 Possible improvements

As mentioned above, a larger sample and/or cell size would surely yield better results. For this, modifications to the simulation code would be necessary to increase performance. For example, a full simulation in the pyace potential for a single temperature (we used four of those) and a cell size of 6×6 took roughly 2,5 to 3 hours. The obvious improvement would be to use multiprocessing, which would be rather easy to implement, since each of the 101 ejection simulations per potential per temperature runs completely independent of the others. In case of our numeric example of pyace with a single-core runtime of 3 hours and assuming 8 cores working in parallel, this would reduce the simulation time to about 20 minutes. Further improvements could be made by running the simulations on a modern desktop processor instead of on a notebook.

5 References

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