

# NeNa

## Untertitel

### Projektarbeit

im Fachbereich Physik



TECHNISCHE  
UNIVERSITÄT  
WIEN

vorgelegt von: Gabriel Sommer

Studiengang: Physik

Matrikelnummer: 11912404

Prüfer: Andreas Grüneis

August 28, 2025

# **Zusammenfassung**

Zugsamenfassung: Ziel, Methoden, Ergebnisse, Schlussfolgerungen

# Contents

<b>List of Figures</b>	<b>II</b>
<b>List of Tables</b>	<b>III</b>
<b>Listings</b>	<b>IV</b>
<b>List of Abbreviations</b>	<b>V</b>
<b>Symbolverzeichnis</b>	<b>VI</b>
<b>1 Einleitung</b>	<b>1</b>
<b>2 Aufgabenstellung</b>	<b>2</b>
<b>3 Theory</b>	<b>3</b>
3.1 From Statistical Mechanics to Vacancy configuration . . . . .	3
3.2 The explicit Formation Energy F . . . . .	6
3.3 Relaxation in LAMMPS . . . . .	6
3.4 Simulated Annealing . . . . .	6
3.5 Symmetry optimization . . . . .	7
3.6 DFT optical spectra . . . . .	7
<b>4 Finding Minima of Neon Replacements</b>	<b>9</b>
4.1 Pair Potential Interaction . . . . .	9
4.2 Geometric LAMMPS Setup . . . . .	12
4.3 Sodium Monomer . . . . .	13
4.4 Sodium Dimer . . . . .	14
<b>5 DFT optical spectra results</b>	<b>16</b>
<b>6 Conclusion</b>	<b>17</b>
<b>7 Ausblick</b>	<b>18</b>
<b>Bibliography</b>	<b>i</b>
<b>A Anhang</b>	<b>ii</b>
A.1 Inhalt des beigelegten Datenträgers . . . . .	ii

## List of Figures

3.1	state evolution ( $\vec{q}(t), \vec{p}(t)$ ) through 6N-dimensional phase space . . . . .	4
4.1	Plots of interpolations of pair potentials of Ne-Ne, Na-Na, Ne-Na calculated with Moeller Plesset 2 perturbation theory. . . . .	11
4.2	Initial setup for the Large-scale Atomic/Molecular Massively Parallel Simulator ( <a href="#">LAMMPS</a> ) relaxation. Only the inner blue sphere is allowed to relax, forces on the orange atom positions are overridden to 0. . . . .	13
4.3	Simulated Annealing for single sodium atom inserted, compared to a brute force minima search for a fixed number of removed atoms S. . . . .	14
4.4	Simulated annealing results and their relative occurrence after x sweeps	15

## List of Tables

## Listings

3.1	Simulated annealing algorithm . . . . .	7
3.2	lexicographically sorting point group symmetries . . . . .	8

## List of Abbreviations

<b>LAMMPS</b>	Large-scale Atomic/Molecular Massively Parallel Simulator
<b>HF</b>	Hartree-Fock
<b>BSSE</b>	Basis Set Superposition Error
<b>fcc</b>	face-centered-cubic
<b>MP2</b>	Second Order Møller-Plesset Perturbation Theory

## Symbolverzeichnis

Symbol	Bedeutung	Einheit
$B$	magnetische Flussdichte	T
$D$	Elektrische Flussdichte	$\text{A s m}^{-2}$



# **1 Einleitung**

Einleitung: Wieso sollte diese Arbeit gelesen werden?

## 2 Aufgabenstellung

We are here trying to calculate optical absorption spectra with vasp. The atomic configuration is a cavity inside a neon crystal that was proposed to be filled with either a single sodium atom or two sodium atoms (a sodium dimer). Now the first step was to find configurations of cavities, i.e. how many neon atoms will be replaced by an inserted sodium atom and subsequently by a sodium dimer. This substitution number will be denoted as  $S$  in the following thesis. For these specific configurations the optical absorption spectra will be calculated and compared with the experiment.

## 3 Theory

### 3.1 From Statistical Mechanics to Vacancy configuration

In later chapters only discrete configurations of the system at hand will be examined. Going from a continuous picture of classical phase space to a discrete description is straight forward. First we consider the system of the crystal of Neon Atoms as a canonical ensemble, this means it will be described under the assumption of an infinitely large heat bath by Boltzmann statistics. The probability distribution of members of the ensemble will then be

$$p(\vec{\mathbf{q}}, \vec{\mathbf{p}}) = \frac{e^{-\beta H(\vec{\mathbf{q}}, \vec{\mathbf{p}})}}{\int \dots \int e^{-\beta H(\vec{\mathbf{q}}, \vec{\mathbf{p}})} d^{3N} \vec{\mathbf{q}} d^{3N} \vec{\mathbf{p}}}, \quad (3.1)$$

with  $N$  being the number total of particles of the system.

Now an ensemble is to be understood as many fictional copies of the same system representing the states that a system will explore when propagating in time according to Hamilton's equation of motion. The ensemble distribution represents the relative number of times that an ergodic system will come by a given state after an infinite amount of time has passed. Now for a crystal in principle this still holds. At low temperatures though the phase space that is being explored can be abstracted in such cases. The reason is that the system will for a given finite time usually explore only nearby points in the phase space. This is the crystal configuration but allowing for dynamics such as lattice vibrations and thermal movement of the atoms with respect to their lattice site. So for a given initial point in phase space the system will mostly explore its proximity until it probabilistically jumps to another volume of phase space whose proximity then will be explored for some further time. Now this means the systems lattice configuration has changed which at low but non zero temperature should be reasonable. We now consider these volumes of phase space (being approximately confined regions, i.e. configurations of e.g. a crystalline structure) to be discrete states that the system can be in. So our canonical distribution now is a discrete one, each state representing such a configuration volume. What's left is to figure out what state of the still continuously possible states inside will be used to represent each configuration. We in this work will use the state  $(\vec{\mathbf{q}}_{0i}, \vec{\mathbf{p}}_{0i})$  that locally minimizes  $H(\vec{\mathbf{q}}, \vec{\mathbf{p}})$ . The Boltzmann distribution now follows to be discrete:

$$p(\vec{\mathbf{q}}_{0i}, \vec{\mathbf{p}}_{0i}) = \frac{e^{-\beta H(\vec{\mathbf{q}}_{0i}, \vec{\mathbf{p}}_{0i})}}{\sum_n e^{-\beta H(\vec{\mathbf{q}}_{0n}, \vec{\mathbf{p}}_{0n})}}. \quad (3.2)$$

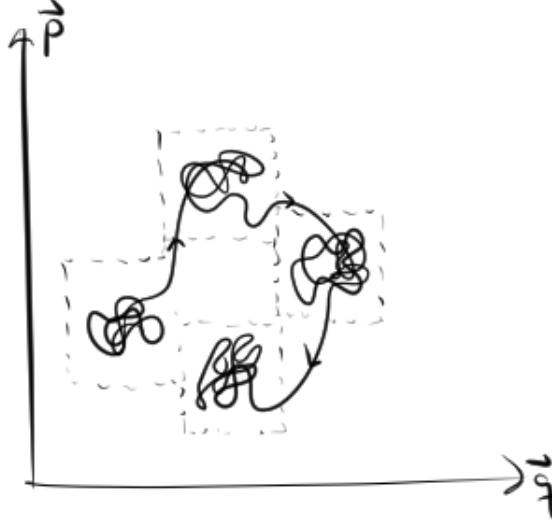


Figure 3.1: state evolution  $(\vec{q}(t), \vec{p}(t))$  through  $6N$ -dimensional phase space

Since we know that the momenta  $\vec{p}$  only show up in the kinetic term of the Hamiltonian, we can immediately say that the momenta  $\vec{p}_{0i}$  representing the minimal hamiltonian will be 0 for all possible configurations  $(\vec{q}_{0i}, \vec{p}_{0i})$ , so we could just drop the  $\vec{p}$  dependency from here on, but for now we'll continue to write them for completeness and only later on will remember this argument.

Now for the problem at hand we will switch to the grand canonical sum, since we need to introduce two more degrees of freedom, i.e. the particle exchange with the heat and particle bath, which here in general involves two types of particles numbers  $N_1, N_2$ . They naturally introduce only discrete degrees of freedom so we can quickly write the grand canonical distribution for the system:

$$\rho(N_1, N_2, \vec{q}_{0i}, \vec{p}_{0i}) = \frac{e^{-\beta(H_{N_1 N_2}(\vec{q}_{0i}, \vec{p}_{0i}) - \sum_{i=1}^2 \mu_i N_i)}}{\frac{1}{h^{3N} \prod_{j=1}^2 N_j!} \sum_{N_1} \sum_{N_2} \sum_n \left[ H_{N_1 N_2}(\vec{q}_{0n}, \vec{p}_{0n}) - \sum_{i=1}^2 \mu_i N_i \right]} = \quad (3.3)$$

$$= \frac{e^{-\beta(H_{N_1 N_2}(\vec{q}_{0i}, \vec{p}_{0i}) - \sum_{i=1}^2 \mu_i N_i)}}{\Xi_{\mu_1 \mu_2}(T, V)}, \quad (3.4)$$

with

$$\mu_i = -T \frac{\partial S}{\partial N_i} = \frac{\partial E}{\partial N_i}. \quad (3.5)$$

the  $i$ -th chemical potential. The most probable states  $(N_1, N_2, \vec{\mathbf{q}}_i, \vec{\mathbf{p}}_i)$ <sup>1</sup> in the grand canonical distribution are the ones for which the term  $F = H_{N_1 N_2}(\vec{\mathbf{q}}_i, \vec{\mathbf{p}}_i) - \sum_{i=1}^2 \mu_i N_i$  is minimal. We here in this work call  $F$  the formation energy. In the following chapters we will fix the number of sodium atoms  $N_2$ . The term  $\mu_2 N_2$  (because of the exponential), just factors out, giving us the conditional probability:

$$\rho(N_2|N_1, \vec{\mathbf{q}}_i, \vec{\mathbf{p}}_i) = \frac{e^{-\beta(H_{N_1 N_2}(\vec{\mathbf{q}}_i, \vec{\mathbf{p}}_i) - \mu_1 N_1)}}{\Xi_{\mu_1 \mu_2}(T, V)} \quad (3.6)$$

leading us to the effective quantity that will be minimized in the following chapters:

$$F_m(N_1, \vec{\mathbf{q}}_i, \vec{\mathbf{p}}_i) = H_{m N_1}(\vec{\mathbf{q}}_i, \vec{\mathbf{p}}_i) - \mu_1 N_1 \quad (3.7)$$

for  $m$ , being a fixed sodium insertion, so  $m \in \{1, 2\}$ . We remember that we already figured out that the minimum of that term needs to have the momenta  $\vec{\mathbf{p}}_i$  set to zero, since they only contribute the positive kinetic terms to  $H_{m N_1}(\vec{\mathbf{q}}_i, \vec{\mathbf{p}}_i)$  meaning we can just set them to zero and drop the dependency, finally leading us to the final form of the formation energy:

$$F_m(N_1, \vec{\mathbf{q}}_i) = H_{m N_1}(\vec{\mathbf{q}}_i) - \mu_1 N_1 \quad (3.8)$$

Now we start with an initial number of neon atoms  $N_1^{[0]}$  determined in the simulation setup, then remove atoms  $S$  from  $N_1^{[0]}$ . The number of removals is denoted as  $S$ . We rewrite the formation energy with  $N_1 = N_1^{[0]} - S$  and get:

$$F_m(S, \vec{\mathbf{q}}_i) = H_{m N_1}(\vec{\mathbf{q}}_i) - \mu_1 (N_1^{[0]} - S). \quad (3.9)$$

The constant term  $-\mu_1 N_1^{[0]}$  drops obviously when minimizing, leading us further to

$$F_m(S, \vec{\mathbf{q}}_i) = H_{m N_1}(\vec{\mathbf{q}}_i) + \mu_1 S. \quad (3.10)$$

The chemical potential  $\mu_1$  is the change in energy when adding one atom, or vice versa the change in energy of the whole system when removing one atom, i.e.<sup>2</sup>:

$$\mu_1 = \frac{\partial E}{\partial N_1} = -\frac{\partial E}{\partial S} \quad (3.11)$$

which we identify as the negative cohesive energy  $-E_{\text{coh}}$ , so negative the energy that is required to remove one atom from the lattice.

---

<sup>1</sup>We from now on drop the superscript  $(\vec{\mathbf{q}}_i^{[0]}, \vec{\mathbf{p}}_i^{[0]}) \rightarrow (\vec{\mathbf{q}}_i, \vec{\mathbf{p}}_i)$ .

<sup>2</sup> $S$  here is not to be confused with the Entropy.

We remember that we are in the grand canonical ensemble, so removing one atom from the microscoping system means adding it to the heat and particle bath. This is intuitive, because when we look at the crystal, removing one atom to create a vacancy essentially means we are placing it some other part of the crystal, that acts as a heatbath, we do not send it to the gasphase, so we must add second chemical term in  $F_m(S, \vec{q}_i)$ , or remove  $S$  times the cohesive energy  $E_{\text{coh}}$ . The final form of the formation energy thus is:

$$F_m(S, \vec{q}_i) = H_{mN_1}(\vec{q}_i) - E_{\text{coh}}S. \quad (3.12)$$

It is the formula that is used in literature.

## 3.2 The explicit Formation Energy F

Now we can write out the Hamiltonian, that only includes the pair interaction of neon and sodium (kinetic energy already being set to zero, or alternatively being ignored).

## 3.3 Relaxation in LAMMPS

relaxation theory from lammmps described in [1] Bitzek, Koskinen, Gahler, Moseler, Gumbusch, Phys Rev Lett, 97, 170201 (2006). Testcite Griffiths[2]

## 3.4 Simulated Annealing

- proves to be effective for heisenberg model, spin systems
- analogy of these state vectors with the state vector in this case
- add gaussian or boltzman pick distribution in state vector

The object that is being annealed is a state vector containing binary information for every lattice site. It refers to the ideal static lattice of the neon structure. A positive value or 1 refers to an atom being present in the initial structure and a negative value or 0 means the neon atom is vacant at this site. So every digit of the binary state refers to a certain specific lattice site. The energy functional of the annealing process will be the energy after minimization (i.e. relaxation) of the above described state vector. This state vector therefore refers to an initial pre-relaxation configuration where sites are simply removed with [LAMMPS](#).

Energy functional contains cohesive Energy/chemical potential, not just hamiltonian!!

Listing 3.1: Simulated annealing algorithm

```
1 stateVector = [1,1,1,1,1,1,1,1,1,1,1,1,1,1]
2 alpha = (Tend / Tstart) ** (1 / N)
3
4 for i in N:
5     T = Tstart * (alpha**i)
6
7     n = randomInt(1,len(stateVector))
8     stateVector[n] = 1 - stateVector[n]
9
10    stateVectorOld = stateVector
11
12    EOld = minimizeEnergy(stateVectorOld)
13    E = minimizeEnergy(stateVector)
14
15    if E >= EOld:
16        stateVector = stateVectorOld
17    elif random(0,1)<=exp(-1/T*(E-EOld)):
18        stateVector = stateVectorNew
19
20 S = 1
21 for site in stateVector:
22     if site == 0:
23         S += 1
```

## 3.5 Symmetry optimization

In the case of a single sodium atom inserted in the neon crystal the highly symmetrical structure was exploited to reduce redundant minimization calls in the annealing routine, by identifying symmetrically equivalent structures and caching them. Applying rotation and reflection matrices of the octahedral symmetry group to a single configuration was used to quickly create all equivalent configurations. By sorting them lexicographically and picking e.g. the first, one ensures every redundant set of configurations is always represented by the same configuration.

## 3.6 DFT optical spectra

Listing 3.2: lexicographically sorting point group symmetries

```
1 equivs = np.transpose(np.einsum("klj,ij", oh_group, pos), (1, 0, 2))
2
3 for k in range(equivs.shape[0]):
4     equivs[k] = equivs[k][np.lexsort(equivs[k].T[::-1])].copy()
5     sorting_idx = np.lexsort(
6         np.transpose(equivs, (2, 1, 0)).reshape(pos.size, oh_group.shape[0])
7     )
8     pos = equivs[sorting_idx[0], :, :]
```



## 4 Finding Minima of Neon Replacements

### 4.1 Pair Potential Interaction

#### 4.1.1 Hartree-Fock (HF) equations

We start with the pair potentials of Ne-Ne, Ne-Na and Na-Na to mimic the interaction of long distance van der Waals + short distance repulsion forces. The pair potentials have been calculated using a HF self consistent field cycle with Second Order Møller-Plesset Perturbation Theory (MP2) by varying the atom distance. Now selfconsistently solving the equations:

$$\left[ -\frac{\hbar^2}{2m} \vec{\nabla}^2 - \frac{Ze^2}{4\pi\epsilon_0} - \frac{e^2}{4\epsilon_0} \sum_{\nu,\sigma}^{(\mu\sigma) \neq (\nu\sigma)} \iiint_{\mathbb{R}^3} \frac{\varphi_{\nu\sigma'}(\tilde{r})}{\|\tilde{r} - \tilde{r}'\|} d^3\tilde{r}' - \hat{A}_{\mu\sigma}(\tilde{r}) \right] \varphi_{\mu\sigma}(\tilde{r}) = \varepsilon_{\mu\sigma} \varphi_{\mu\sigma}(\tilde{r}), \quad (4.1)$$

with  $\hat{A}_{\mu\sigma}$  being the exchange correlation term leads to the eigenenergies  $\varepsilon_{\mu\sigma}$ . Since HF minimizes a slater determinant, the energies correspond to the factors of the slater product state, meaning the systems energy is the sum of the energy of the single electrons. From that, to get the actual energy of the whole system, one needs to remove inter-electron repulsion and exchange correlation from the sum of all eigenenergies  $\varepsilon_{\mu\sigma}$ :

$$E_{HF}^0 = \sum_{\mu,\sigma} \varepsilon_{\mu\sigma} - \frac{1}{2} \sum_{\substack{\mu,\nu \\ \sigma,\sigma'}}^{(\mu\sigma) \neq (\nu\sigma)} \left[ C_{\mu\sigma}^{\nu\sigma'} - A_{\mu\sigma}^{\nu\sigma} \delta_{\sigma\sigma'} \right] \quad (4.2)$$

with  $C_{\mu\sigma}^{\nu\sigma'}, A_{\mu\sigma}^{\nu\sigma}$  being defined as:

$$C_{\mu\sigma}^{\nu\sigma'} = \frac{e^2}{4\pi\epsilon_0} \iint \frac{|\varphi_{\mu\sigma}(r)|^2 |\varphi_{\nu\sigma'}(r')|^2}{|r - r'|} d^3r d^3r', \quad (4.3)$$

$$A_{\mu\sigma}^{\nu\sigma} = \frac{e^2}{4\pi\epsilon_0} \iint \frac{\varphi_{\mu\sigma}^*(r) \varphi_{\nu\sigma}^*(r') \varphi_{\mu\sigma}(r') \varphi_{\nu\sigma}(r)}{|r - r'|} d^3r d^3r'. \quad (4.4)$$

Since solving the HF equations in real space is too costly one restricts the variations to coefficients of a superposition of fixed basis vectors. These lead to the Roothaan-Hall equations, which are a discrete Matrix representation of the HF equations:

$$\mathbf{F}\mathbf{C} = \mathbf{S}\mathbf{C}\epsilon, \quad (4.5)$$

with  $\epsilon$  being a diagonal matrix of single particle energies on the diagonal,  $\mathbf{S}$  the basis overlap matrix,  $\mathbf{C}$  the coefficient matrix and finally  $\mathbf{F}$  the Fock operator.

#### 4.1.2 Møller-Plesset Perturbation

Møller-Plesset Perturbation theory extends the idea of HF. We consider the actual true Hamiltonian  $\hat{H}$  in the same basis as 4.5 and define the difference to the Fock operator to be a relatively small perturbation:

$$\hat{H} = \hat{F} + \underbrace{(\hat{H} - \hat{F})}_{\hat{V}_{pert}}. \quad (4.6)$$

The 0-th order Term is just the sum of the eigenvalues  $\varepsilon_i$  of the Fock Problem. Now the 1st order terms of the perturbation are

$$E^{(1)} = \langle \Psi_{HF} | (\hat{H} - \hat{F}) | \Psi_{HF} \rangle = \quad (4.7)$$

$$= -\frac{1}{2} \sum_{\mu\nu} \left[ \langle \varphi_\mu \varphi_\nu | \hat{V}^{(2)}(\hat{r}_\mu, \hat{r}_\nu) | \varphi_\mu \varphi_\nu \rangle - \langle \varphi_\mu \varphi_\nu | \hat{V}^{(2)}(\hat{r}_\mu, \hat{r}_\nu) | \varphi_\nu \varphi_\mu \rangle \right]. \quad (4.8)$$

which is exactly the mean-field and the exchange correlation term in the HF Ansatz, so to go beyond HF we need to consider MP2:

$$E^{(2)} = \sum_{\substack{N < a < b \\ \mu < \nu \leq N}} \frac{\left| \langle \psi_0 | \hat{V}^{(2)}(\hat{r}_\mu, \hat{r}_\nu) | \psi_{\mu\nu}^{ab} \rangle \right|^2}{(\varepsilon_a + \varepsilon_b) - (\varepsilon_\mu + \varepsilon_\nu)}. \quad (4.9)$$

where  $|\psi_{\mu\nu}^{ab}\rangle$  is an excited Slater determinant, where single orbitals  $\mu, \nu$  are removed and replaced by single orbitals  $a, b$  in the slater determinant.  $\varepsilon_i$  are eigenvalues of the slater single particle factors of the HF problem.

### 4.1.3 Basis Set Superposition Error (BSSE)

Now the the superpositions of a fixed basis leads to the BSSE. These errors are being corrected for by using 'ghost calculations', essentially the same calculation with one atom being removed (i.e. the extended two atom centered basis stays the same). This means the basis will have access to the second electron states, but the nucleus and additional electrons are not accounted for. These 'ghost energies' will then subsequently be removed from the HF approximation to correct for the BSSE. This is know as the counterpoise-corrected interaction energy formula or Boys–Bernardi method:

$$E_{\text{int}}^{\text{CP}}(R) = E_{AB}(R) - E_A^{\text{ghost}}(R) - E_B^{\text{ghost}}(R). \quad (4.10)$$

Here  $E_{AB}(R)$  is the energy after MP2 and  $E_A^{\text{ghost}}(R), E_B^{\text{ghost}}(R)$  refer to the same calculation results, just with one atom removed completely.

### 4.1.4 Interpolation

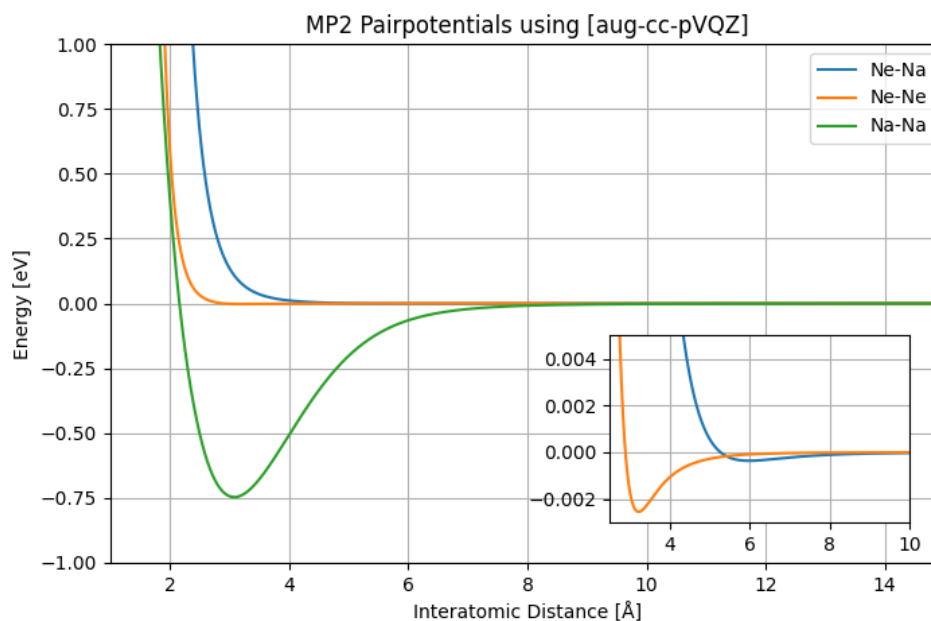


Figure 4.1: Plots of interpolations of pair potentials of Ne-Ne, Na-Na, Ne-Na calculated with Moeller Plesset 2 perturbation theory.

- explain exact interpolation
- paper citation where brute force plot shows up as well
- which basis is being used in the roothaan equations

- explain setup of calculation with ball being carved out
- maybe here explain the nearest neighbours
- lammmps gets linear interpolation
- since plot is in principle linearly interpolated this is the exact pair interactions that lammmps will read

## 4.2 Geometric LAMMPS Setup

The simulation models a finite spherical cluster of atoms extracted from a bulk face-centered cubic lattice. To balance computational efficiency with physical realism, the system is partitioned into two concentric spherical regions defined in units of the lattice constant. The inner spherical region, with radius  $n_{inner}$  contains atoms that are free to move and evolve dynamically. This region represents the core of the cluster where physical process and in this case the energy minimization via a relaxation calculation of LAMMPS takes place.

Surrounding this core is an immobilized shell with thickness  $n_{outer}$ . Atoms within this outer shell are constrained by setting their forces to zero, effectively fixing them in space. This fixed shell acts as a rigid boundary that suppresses surface effects and mimics the presence of an extended bulk lattice, thereby reducing finite size artifact in the dynamics of the inner region.

Consequently the total cluster is confined within a spherical domain of radius  $R_{tot} = (n_{inner} + n_{outer}) \cdot a$  with  $a$  being the lattice constant of the pure neon face-centered-cubic (fcc) lattice.

Technically the simulation is defined as a cubic volume large enough to contain the entire spherical cluster, with half-length at least  $R_{tot}$ . For the relaxation mechanics atomic positions are set according to 4.2 on a perfect fcc lattice no matter their role (sodium defect, fixed neon, dynamic neon) as long as they are inside the cutoff radius  $R_{tot}$ . This approach is commonly employed in molecular dynamics studies to simulate nanoparticles or finite clusters embedded in bulk-like surroundings.

The centered red dots in 4.2 represent sodium atoms. They represent the initial pre-relaxation positions of sodium inside the dynamic inner sphere. These initial positions are the same for every relaxation step of the simulated annealing. Note that the mini-

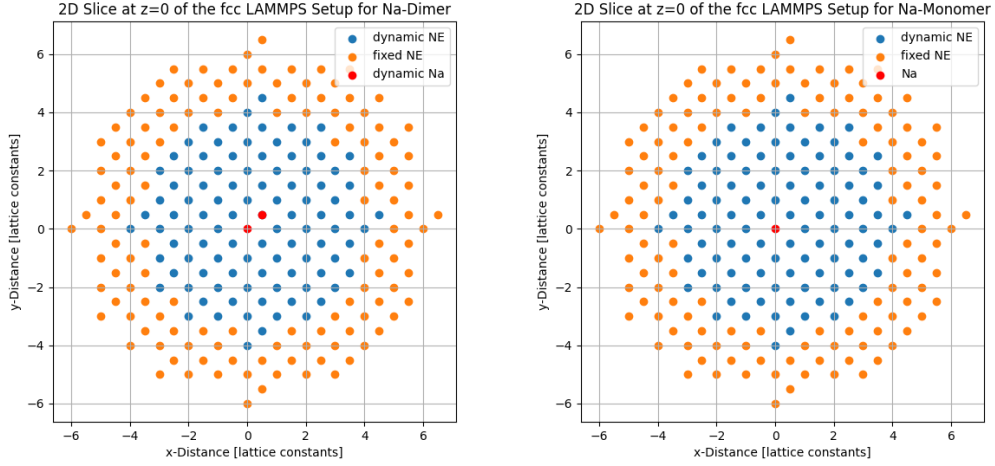


Figure 4.2: Initial setup for the [LAMMPS](#) relaxation. Only the inner blue sphere is allowed to relax, forces on the orange atom positions are overridden to 0.

num energy of the pair-potential in [4.1](#) is roughly at  $3.1\text{\AA}$  which is the nearest neighbor distance in the neon [fcc](#) lattice. With  $a = 4.4637\text{\AA}$ :

$$\vec{a}_1 = \begin{bmatrix} a \\ 0 \end{bmatrix} \quad \vec{a}_2 = \begin{bmatrix} 0 \\ a \end{bmatrix} \quad (4.11)$$

$$\Rightarrow \left\| \frac{1}{2} (\vec{a}_1 + \vec{a}_2) \right\| = \frac{1}{\sqrt{2}} \cdot a = 3.1563\text{\AA}. \quad (4.12)$$

We therefore put the second sodium atom in the dimer calculations at the nearest neighbor lattice site and Set th initial removal  $S$  to  $S = 2$ .

### 4.3 Sodium Monomer

Now the annealing was done as described in the chapters before for a single sodium atom at the center. Since the known energetically minimal configurations for every  $S$  and it's corresponding energy was known due to a brute force calculation, this calculation serves as a benchmark for the annealing algorithm. The brute force minimal energies for the vacancies are shown in [4.3](#) on the right axis. The left axis shows the relative counts (with respect to the total counts) of one annealing sweep consisting of itself 1000 random removals of neon atoms from the state vector. This annealed approach quickly converges to a statistical distribution shown in the same figure [4.3](#) on the left  $y$ -axis. The pattern surfaces quickly after about 50 sweeps, the total amount of sweeps for the plot were 736. The true local minima at  $S = 10$  and  $S = 13$  could

be picked up upon quite sharply. At  $S = 8$  the annealing algorithm seems to pick up on the notch, that is not quite an actual minima but could look like a local minima if approached from many directions in the high dimensional phase space.

- explain how plot is created with sweeps
- more complicated symmetries
- compare figures
- calculation for dimer
- discuss noteworthy structure (e.g. inner shell carved out)
- citation of paper that has the same plot Now first by brute force search we know the minima for a single sodium atom.

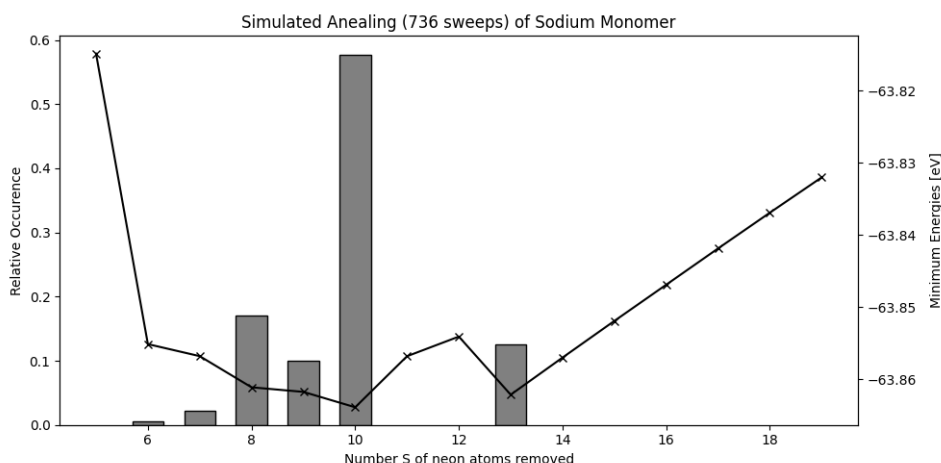


Figure 4.3: Simulated Annealing for single sodium atom inserted, compared to a brute force minima search for a fixed number of removed atoms  $S$ .

- brute force lexiciraphically sorting algorithm here

### 4.4 Sodium Dimer

Since 4.3 shows the simulated annealing algorithm accurately picks up on the location of the minima (albeit with no quantitative measure and misleading local minima being blown out of proportion (See  $S = 8$ )) we try the same approach for the sodium dimer. Here the energetically optimal  $S$  replacement will expectantly going to exceed the number of available sites as in the monomer calculation (i.e. up to second nearest neighbor). The sweeps and their random removal of neon atoms now act upon an extended state vector containing every lattice site up to the 3rd nearest neighbor. This

is also the very reason a brute force approach is unfeasible, since the volume of phase space to be explored grows quickly. Considering the octahedral symmetry  $O_h$  of the host, which the defect cannot possibly exceed and roughly estimating one minimization run to take about 20s, we get  $\frac{2^{42}}{48} \cdot 20s \approx 10^{12}s$  which exceeds realistic resources for a brute force attack. This is the reason the statistical approach was chosen in the first place.

- symmetry
- heuristically take the maxima of these plots
- give structures
- discuss structures (inner shell carved out)

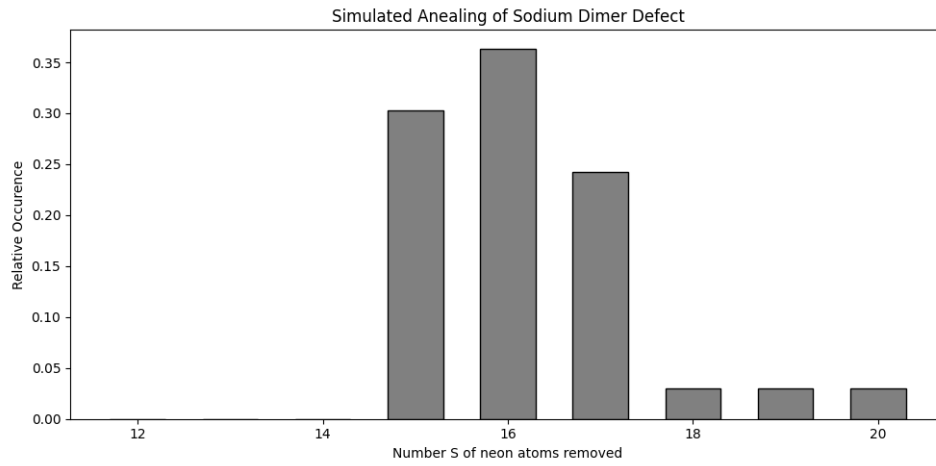


Figure 4.4: Simulated annealing results and their relative occurrence after x sweeps

## **5 DFT optical spectra results**

Zweites Kapitel



## 6 Conclusion

Unfortunately no confidence or error can be estimated since the approach was purely heuristical. Optical spectrum does not depend much on replacements and precise structure.

## **7 Ausblick**

Ausblick

## Bibliography

- [1] P. Bitzek et al. “Structural Relaxation Made Simple”. In: *Physical Review Letters* 97.17 (2006), p. 170201. DOI: [10.1103/PhysRevLett.97.170201](https://doi.org/10.1103/PhysRevLett.97.170201).
- [2] David J. Griffiths. *Introduction to Quantum Mechanics*. 3rd ed. Cambridge University Press, 2018. ISBN: 9781107189638.

# A Anhang

## A.1 Inhalt des beigefügten Datenträgers

 Datenträger



A



a



b



c



B



a



b



A

## Eidesstattliche Erklärung

Ich, Gabriel Sommer, versichere hiermit, dass ich die vorliegende Arbeit selbstständig verfasst und keine anderen als die angegebenen Quellen und Hilfsmittel benutzt habe, wobei ich alle wörtlichen und sinngemäßen Zitate als solche gekennzeichnet habe.

Diese Arbeit wurde bisher in gleicher oder ähnlicher Form keiner anderen Prüfungsbehörde vorgelegt und auch nicht veröffentlicht.

Vienna, den August 28, 2025

---

GABRIEL SOMMER

## Declaration of Authorship

I, Gabriel Sommer, hereby declare that I have written the present thesis independently and have used no sources or aids other than those explicitly stated. All direct quotations and all paraphrased ideas have been clearly marked as such.

This work has not been submitted, either in the same or a substantially similar form, to any other examination board, nor has it been published elsewhere.

Vienna, August 28, 2025

---

GABRIEL SOMMER